#### Method-

# Identifying *cis*-mediators for *trans*-eQTLs across many human tissues using genomic mediation analysis

Fan Yang,<sup>1,5</sup> Jiebiao Wang,<sup>1</sup> The GTEx Consortium,<sup>1,4</sup> Brandon L. Pierce,<sup>1,2,3</sup> and Lin S. Chen<sup>1</sup>

<sup>1</sup> Department of Public Health Sciences, <sup>2</sup> Department of Human Genetics, The University of Chicago, Chicago, Illinois 60637, USA; <sup>3</sup> Comprehensive Cancer Center, The University of Chicago, Chicago, Illinois 60637, USA

The impact of inherited genetic variation on gene expression in humans is well-established. The majority of known expression quantitative trait loci (eQTLs) impact expression of local genes (*cis*-eQTLs). More research is needed to identify effects of genetic variation on distant genes (*trans*-eQTLs) and understand their biological mechanisms. One common *trans*-eQTLs mechanism is "mediation" by a local (*cis*) transcript. Thus, mediation analysis can be applied to genome-wide SNP and expression data in order to identify transcripts that are "*cis*-mediators" of *trans*-eQTLs, including those "*cis*-hubs" involved in regulation of many *trans*-genes. Identifying such mediators helps us understand regulatory networks and suggests biological mechanisms underlying *trans*-eQTLs, both of which are relevant for understanding susceptibility to complex diseases. The multitissue expression data from the Genotype-Tissue Expression (GTEx) program provides a unique opportunity to study *cis*-mediation analyses challenging and prone to confounding bias, particularly when conducted among diverse samples. To address this problem, we propose a new method: Genomic Mediation analysis with Adaptive Confounding adjust-ment (GMAC). It enables the search of a very large pool of variables, and adaptively selects potential confounding variables for each mediation test. Analyses of simulated data and GTEx data demonstrate that the adaptive selection of confounders by GMAC improves the power and precision of mediation analysis. Application of GMAC to GTEx data provides new insights into the observed patterns of *cis*-hubs and *trans*-eQTL regulation across tissue types.

[Supplemental material is available for this article.]

Recent studies have demonstrated that many expression quantitative trait loci (eQTLs) that affect expression of local transcripts (ciseQTLs) also affect the expression of distant genes (trans-eQTLs) (Battle et al. 2014; Pierce et al. 2014). This observation suggests the effects of trans-eQTLs are "mediated" by the local (cis-) gene transcripts near the eQTLs (Fehrmann et al. 2011; Pierce et al. 2014). In other words, some cis-eQTLs are also trans-eQTLs because the variation in the expression of the cis-gene affects the expression of a trans-gene or genes. In the simplest scenario, a cis-eQTL affects expression of a nearby gene that is a transcription factor, which then regulates the transcription of a distant gene; thus, the transcription factor "mediates" the effect of the eQTL on the distant gene. By studying eQTLs that have both the cis- and trans-effects, one may identify the cis-genes that mediate the effects of transeQTLs on expression of distant genes, including "cis-hub" genes that regulate the expression of many trans-genes (Chen et al. 2007; Stranger et al. 2012). Studying mediation (causation) moves beyond the analysis of cis- and trans- associations (correlation). Prior studies have applied mediation tests to genome-wide SNPs and expression data (from blood cells) to identify transcripts that

<sup>5</sup>Present address: Department of Biostatistics and Informatics, Colorado School of Public Health, University of Colorado Denver, Aurora, CO 80045, USA

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are *cis*-mediators of the effects of *trans*-eQTLs (Chen et al. 2007; Battle et al. 2014; Pierce et al. 2014). Characterizing these regulatory relationships will allow us to better understand regulatory networks and their roles in complex diseases (Veyrieras et al. 2008), as it is well known that SNPs influencing human traits tend to be eQTLs (Nicolae et al. 2010). Analyses of *cis*-mediation will also provide us with a better understanding of the biological mechanisms underlying *trans*-eQTLs (Westra et al. 2013).

The expression levels of a given gene can vary substantially across human cell types, and the regulatory relationships between SNPs and gene expression levels may also depend on cell type (Torres et al. 2014; Wang et al. 2016). To date, most large-scale eQTL studies have been conducted using RNA extracted from peripheral blood cells, which are mixtures of different cell types and may not be informative for gene regulation in other human tissues. In order to study gene expression and regulation in a variety of human tissues, the National Institutes of Health commonfund GTEx (Genotype-Tissue Expression) project has collected expression data on 44 tissue types from hundreds of post-mortem donors (Lonsdale et al. 2013; Ardlie et al. 2015). This rich transcriptome data, coupled with data on inherited genetic variation, provides an unprecedented opportunity to study gene expression and regulation patterns from both cross-tissue and tissue-specific perspectives.

<sup>&</sup>lt;sup>4</sup>A full list of Consortium members and their affiliations is available at the end of the text.

Corresponding authors: Ichen@health.bsd.uchicago.edu, brandonpierce@uchicago.edu

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One major challenge in mediation analysis is the presence of unmeasured or unknown variables that affect both the mediator (i.e., cis-gene) and outcome (i.e., trans-gene) variables. The presence of such a variable is known as "mediator-outcome confounding," and in such a scenario, estimates obtained from mediation analysis can be biased (Robins and Greenland 1992; Pearl 2001; Cole and Hernan 2002). In other words, in the presence of an unmeasured confounding variable(s), the association between the two cis- and trans-genes will be a biased estimate of the causal relationship between the two genes, and estimates obtained from mediation analysis will be biased. It is well recognized that measures of transcriptional variation can be affected by genetic, environmental, demographic, technical, and biological factors. The presence of unmeasured or unknown confounding effects may induce inflated rates of false detection of mediation relationships or jeopardize the power to detect real mediation, if those confounding effects are not well accounted for. Given that eQTL analyses are conducted in the context of complex biological systems, there is a wide array of biological variables that could potentially confound the mediator-outcome association and bias mediation estimates, a problem that may be exacerbated by the diversity of GTEx participants, with respect to ethnicity, age, and cause of death. Given these challenges, it is desirable to have methods that consider a large pool of potential confounding variables.

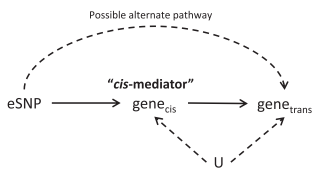
To adjust for unmeasured or unknown confounding effects in genomics studies, existing literature focused on the construction of sets of "hidden" variables that capture a substantial amount of the variation in a large set of variables (Price et al. 2006; Leek and Storey 2007; Stegle et al. 2012). A commonality of those approaches is that they model the effects of hidden confounding factors and summarize those effects into a set of constructed variables, sort those variables decreasingly by their estimated impacts, and adjust the top ones as a set of covariates to eliminate major confounding effects in the subsequent analysis. For example, in GTEx eQTL analyses (Ardlie et al. 2015; The GTEx Consortium 2017), the top Probabilistic Estimation of Expression Residuals (PEER) factors were estimated for each tissue type, and up to 35 factors were adjusted. One aspect that is largely ignored is that not all potential gene pairs (or pairs of regulator and regulated genes) are affected by the same set of hidden confounders. There are likely thousands of cis-mediated trans-eQTLs in the human genome, i.e., trios consisting of a genetic variant, a *cis*-gene transcript, and a trans-gene transcript in a specific tissue type. However, for each trio, mediator-outcome confounding will be present only when a hidden variable is causally related to the regulator and regulated genes. By this criterion, the potential confounder set varies by different trios. Adjusting a universal set of variables for all mediation trios is not only inefficient but also may limit our ability to consider a larger pool of potential confounding variables in genomic mediation analyses.

We propose to adaptively select the variables to adjust for each trio given a large set of constructed or directly measured potential confounding variables. This strategy supplements existing confounding adjustment approaches that focus on the construction of variables for capturing confounding effects and enlarges the pool of variables to be considered. Additionally, by leveraging the *cis* genetic variant as an "instrumental variable," we are able to select the variables capturing confounding effects rather than variables only correlated with *cis*- and *trans*-genes. We further propose a mediation test with nonparametric *P*-value calculation, adjusting for the adaptively selected sets of confounders. We term the proposed algorithm Genomic Mediation analysis with Adaptive Confounding adjustment (GMAC). The GMAC algorithm improves the efficiency and precision of confounding adjustment and the subsequent genomic mediation analyses. We applied GMAC to each of the 44 tissue types of GTEx data in order to study the *trans*-regulatory mechanism in human tissues. Our algorithm identifies genes that mediate *trans*-eQTLs in multiple tissues, as well as "*cis*-hubs" that mediate the effects of a *trans*-eQTL on multiple genes.

#### Results

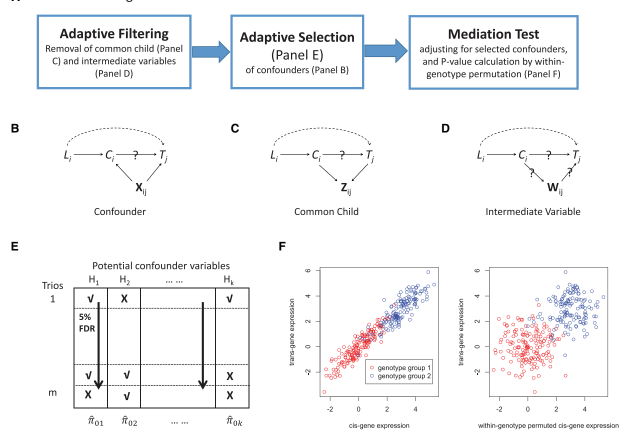
#### GMAC improves power and precision of analysis of GTEx data

We performed genomic mediation analysis with data from each tissue type in GTEx. Taking the tissue, adipose subcutaneous, as an example, there are 298 samples for this tissue type, and genelevel expression measures for 27,182 unique transcripts are available after quality control. Consider a candidate mediation trio consisting of a gene transcript  $i(C_i)$ , its *cis*-associated genetic locus  $(L_i)$ , and another gene transcript  $j(T_i)$  in *trans*-association with the locus. The goal is to test for mediation of the effect of the genetic locus on the trans-gene by the cis-gene (see Fig. 1). We focused on only the trios  $(L_i, C_i, T_i)$  in the genome showing both *cis*- and *trans*-eQTL associations, i.e.,  $L_i \rightarrow C_i$  and  $L_i \rightarrow T_j$ . Because associations are necessary but not sufficient conditions for inferring mediation, by considering only the trios with cis- and transassociations, we effectively reduced the search space to a promising pool of candidate mediation trios and alleviated the multiple testing burdens. We detected and selected a lead cis-eQTL for 8500 of these transcripts, corresponding to 8216 unique ciseSNPs for subsequent analysis (see Methods). We applied Matrix eQTL (Shabalin 2012) to the 8216 SNPs and the 27,182 gene expression levels to calculate the pair-wise trans-associations. At the P-value cutoff of 10<sup>-5</sup>, there were 3169 significant pairs of SNP and trans-gene transcripts. Since some cis-eSNPs were the lead cis-eSNPs for multiple local gene transcripts, those significant SNP and trans-gene pairs entailed a total of 3332 trios (i.e., SNPcis-trans) for this tissue type. We applied GMAC (see Methods; see Fig. 2 for a graphical illustration of the main steps of the GMAC) to the 3332 trios in this tissue type to test for mediation



Possible mediator-outcome confounding

**Figure 1.** Causal diagram demonstrating mediation and "mediator-outcome confounding." Here, the variable set "**U**" represents a set of unmeasured or unknown variables that may show confounding effects in the mediation analysis. Mediation analysis can detect mediation of the effect of the eSNP on the *trans*-gene by the *cis*-gene, assuming mediator-outcome confounding is absent or adjusted for in the analysis. Mediation will not be detected if the effect of the eSNP on the *trans*-gene is through some alternative pathway that does not involve the *cis*-gene.



#### A The GMAC Algorithm

**Figure 2.** Graphical illustrations of GMAC and its main ideas. (*A*) A summary of the GMAC algorithm; (*B*) a mediation relationship among an eQTL,  $L_{ii}$  its *cis*-gene transcript,  $C_{ii}$  and a *trans*-gene transcript,  $T_{j}$ , with confounders, **X**<sub>ij</sub>, allowing  $L_i$  to affect  $T_j$  via a pathway independent of  $C_{ii}$  (*C*) a mediation trio where  $C_i$  and  $T_j$  have common child variable(s), **Z**<sub>ij</sub>; (*D*) a mediation trio where  $C_i$  affects  $T_j$  through intermediate variable(s), **W**<sub>ij</sub>. (*E*) The adaptive confounder selection procedure: Based on the *P*-value matrix for the association of each potential confounder variable to at least one of the *cis*- or the *trans*-gene transcripts, we apply a stratified FDR approach by considering the *P*-values for each potential confounder (each column) as a stratum, with the significant ones indicated by a check mark ( $\sqrt{}$ ). When conducting the mediation test for each trio, we only adjust for the significant confounding variables (the ones with  $\sqrt{}$  in each row). (*F*) A mediation trio  $L_i \rightarrow C_i \rightarrow T_j$  (*left*) and a trio under the null with both *cis*-linkage and *trans*-linkage but no mediation (*right*). Withingenotype permutation of the *cis*-gene expression levels maintains the *cis*- and *trans*-linkage (different mean levels) while breaking the potential correlation between the *cis*- and *trans*-expression levels within each genotype group. Note that **X**<sub>ij</sub>, **X**<sub>ij</sub>, **W**<sub>ij</sub> may vary by trios and are all subsets of **H**. We assume that either **X**<sub>ii</sub> or a combination of variables in **X**<sub>ij</sub> would capture the variation of the unmeasured confounder **U** in Figure 1.

and obtained the mediation *P*-values for those trios. Since different tissue types have different sample sizes in GTEx and in addition to cross-tissue confounders, there are many tissue-specific confounding effects, we constructed Principal Components (PCs) from the expression data of each tissue type as potential confounders (Fig. 2B). The number of PCs for each tissue type is equal to the tissue sample size minus 1. We analyzed trios for mediation in a similar fashion for all other GTEx tissue types.

At the 5% false discovery rate (FDR) (Storey and Tibshirani 2003) level, we identified 6145 instances of significant mediation out of 64,824 trios tested in the 44 tissue types. These trios represent potential examples of *cis*-mediation of *trans*-eQTLs within a specific tissue. Table 1 lists the number of significant mediation trios at 5% FDR and the number of trios with suggestive mediation (*P*-value < 0.05), as well as the total number of trios with significant *cis*- and *trans*-associations for all tissue types. The number of confounders selected for each mediation test ranged from 0 to 22 across all tissue types, with a mean of 7.695 and a median of 8. The median number of confounders selected for each tissue type ranged from 3 to 12, while the pool of variables (PCs) from

which we selected confounders ranged from 69 to 360. Supplemental Table S1 presents the descriptive statistics for the number of selected confounders for all the trios in each tissue type. It is clear that with GMAC, on average, we adjust an efficient number while considering a large pool of confounding variables in the mediation tests, and that may improve the power and accuracy of the analyses.

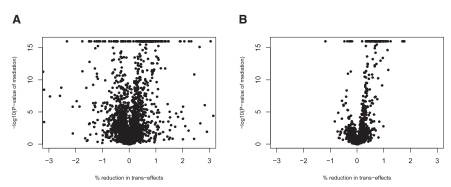
Again taking the tissue, adipose subcutaneous, as an illustration, in Figure 3, we plotted the negative log base 10 of the mediation *P*-values versus the percentages of reduction in *trans*effects after adjusting for a potential *cis*-mediator, based on mediation tests without adjusting for hidden confounders (Fig. 3A) and mediation tests by GMAC considering all PCs as potential confounders (Fig. 3B). The percentage of reduction in *trans*-effects is calculated by  $(\beta_2^m - \beta_2)/\beta_2^m \times 100\%$ , where  $\beta_2^m$  is the marginal *trans*-effect of the eQTL on the *trans*-gene expression levels, and  $\beta_2$  is the *trans*-effect after adjusting for *cis*-mediation. For trios representing true *cis*-mediation, we expect the *trans*-effects to be substantially reduced after adjusting for the mediator; that is, we expect the trios with very significant mediation *P*-values

Tissue name	Tissue sample size	# Trios tested	<pre># Trios with suggestive mediation</pre>	# Trios significant at 5% FDR	
Muscle skeletal	361	2387	496	264	
Whole blood	338	2274	508	281	
Skin sun-exposed lower leg	302	3273	629	330	
Adipose subcutaneous	298	3332	640	325	
Artery tibial	285	2699	527	281	
Lung	278	2762	543	323	
Thyroid	278	3894	696	376	
Cells transformed fibroblasts	272	3000	642	340	
Nerve tibial	256	3812	677	326	
Esophagus mucosa	241	2640	465	242	
Esophagus muscularis	218	2431	447	230	
Artery aorta	197	2009	368	186	
Skin not sun-exposed suprapubic	196	1961	365	177	
Heart left ventricle	190	1290	242	115	
Adipose visceral omentum	190	1410	242	125	
Breast mammary tissue	183	1410	257	125	
Stomach	170	1153	234	128	
	169	1585	309	161	
Colon transverse	159				
Heart atrial appendage		1221	243	103	
Testis	157	3896	607	267	
Pancreas	149	1270	208	102	
Esophagus gastroesophageal junction	127	857	148	74	
Adrenal gland	126	981	185	94	
Colon sigmoid	124	968	220	108	
Artery coronary	118	874	191	95	
Cells EBV-transformed lymphocytes	114	856	152	78	
Brain cerebellum	103	1295	187	84	
Brain caudate basal ganglia	100	763	139	61	
Liver	97	496	87	41	
Brain cortex	96	754	134	45	
Brain nucleus accumbens basal ganglia	93	592	97	43	
Brain frontal cortex BA9	92	595	102	52	
Brain cerebellar hemisphere	89	1072	222	116	
Spleen	89	825	157	68	
Pituitary	87	732	132	61	
Prostate	87	474	101	54	
Ovary	85	469	95	43	
Brain putamen basal ganglia	82	481	94	35	
Brain hippocampus	81	343	93	47	
Brain hypothalamus	81	342	74	41	
Vagina	79	248	58	25	
Small intestine terminal ileum	77	434	82	39	
Brain anterior cingulate cortex BA24	72	365	81	29	
Uterus	70	287	62	25	

to have positive percent reduction in the *trans*-effect. In Figure 3A, we observed many trios with significant mediation *P*-values, but for a substantial number of these trios, the percentages of reduction in *trans*-effects are negative. At the mediation *P*-value threshold of 0.05, 1577 out of 3332 trios were significant; however, 712 trios (712/3332 = 21.3%) have negative percent reduction in *trans*-effects. This contradicting result is expected in the presence of unadjusted confounders, and many of these trios may be false positives. Thus, mediation analyses of GTEx data without adjusting for hidden confounding effects will lead to many spurious findings.

In addition to our main analysis based on GMAC (adaptively selecting confounders from all expression PCs), we also conducted mediation tests adjusting for only the 35 PEER factors used in the GTEx eQTL analyses (The GTEx Consortium 2017). At the 5% FDR level, 3356 out of 64,824 trios from all tissue types were significant. Using GMAC adjusting for adaptively selected PEER factors, 5131 trios were significant at the 5% FDR level. The comparison of adjusting for all (up to 35) PEER factors versus GMAC (considering a larger pool of potential confounders with up to 360 PCs) demon-

strates that adaptive selection enables more efficient adjustment of confounding effects with much fewer selected confounding variables (Supplemental Table S1) and improves power to detect mediation. Furthermore, using GMAC to adaptively select confounders from all PCs identifies 6145 significant trios, suggesting an increase in power. It can also be seen that all three methods-(1) GMAC with adaptively-selected PCs, (2) GMAC with adaptively-selected PEER factors, and (3) adjusting for all PEER factorswould yield reasonable mediation estimates (i.e., percentages of reduction in trans-effects versus mediation P-values), compared to no confounder adjustment (see Supplemental Fig. S1). In conclusion, motivated by the fact that the potential confounder set may vary by different trios, GMAC adaptively adjusts for only the variables that are causally related to both cis- and trans-genes and may show confounding effects in the mediation analysis of each trio (Fig. 2B). Compared with adjusting for a universal set of (top) variables for all mediation trios, GMAC considers a larger pool of potential confounding variables in genomic mediation analyses and enjoys improved power while controlling for false positives.



**Figure 3.** Plots of negative log base 10 of mediation *P*-values versus the percentages of reduction in *trans*-effects after accounting for *cis*-mediation. The *P*-values are calculated based on (*A*) mediation tests without adjusting for hidden confounders, and (*B*) mediation tests by GMAC considering all PCs as potential confounders. *P*-values are truncated at  $10^{-16}$ . The plots are based on the results from the adipose subcutaneous tissue. The percentage of reduction in *trans*-effects is calculated by  $(\beta_2^m - \beta_2)/\beta_2^m \times 100^{\circ}$ , where  $\beta_2^m$  is the marginal *trans*-effect of the eQTL on the *trans*-gene expression levels, and  $\beta_2$  is the *trans*-effect after adjusting for a potential *cis*-mediator and other covariates. For trios with true *cis*-mediations, the marginal *trans*-effects are nonzero, and after adjusting for the true *cis*-mediators, we expect the adjusted *trans*-effects  $\beta_2$  to be substantially reduced; that is, we expect the trios with very significant mediation *P*-values to have positive percent reduction in *trans*-effects. For results based on no adjustment of hidden confounders (*A*), we observed many trios with significant mediation *P*-values but the percentages of reduction in *trans*-effects below the threshold and percent reduction in *trans*-effects being negative in *A* and *B*, respectively.

The majority of the cis-mediators and trans target genes observed among our trios showing mediation have high mappability scores (Supplemental Fig. S2). However, nonuniquely mapping reads can result in false positive eQTLs, so we consider the mappability of each gene as a quality control filter for studying specific examples of cis-mediation (see Methods). Examining the mappability for genes involved in cis-mediation, we observed that cis-genes showing evidence of cis-mediation for multiple trans-genes were enriched for cis-genes with low mappability scores (Supplemental Fig. S2). Similarly, genes showing evidence of cis-mediation across many different tissue types were also enriched for genes showing low mappability scores (Supplemental Fig. S2). This finding demonstrates that transcripts that do not uniquely map to the genome are an important source of false positives when conducting genomic mediation analysis. More specifically, we find that analyzing low-mappability genes can lead to the identification of spurious cis-hubs and cross-tissue cis-mediators.

We attempted to identify "cis-hubs" with high mappability in the GTEx data, defined as a transcript that appears to mediate the effect of a nearby eSNP on expression of multiple distant (i.e., trans) gene transcripts. Restricting our analysis to cis- and transgenes with mappability > 0.95, we observed 685 *cis*-genes with at least two trans targets (considering all tissues), representing 21% of the 3168 cis-genes observed among the trios with a mediation P-value < 0.05 (Table 2). In addition, we attempted to identify cis-genes that have at least one trans target in multiple tissues. Restricting to high-mappability genes, we observed 531 cis-genes with trans targets in more than one tissue, representing 17% of the 3168 cis-genes observed among the trios with a mediation Pvalue < 0.05 (Table 2). We observed only six examples of cis-genes that had the same trans targets in multiple tissues. In other words, the vast majority of cis-hubs observed were of two distinct types: (1) those that mediated the effect of a trans-eQTL on multiple trans-genes within a single tissue type; and (2) those that were mediators in multiple tissues but with unique trans targets in each tissue type. All instances of *cis*-mediation of *trans*-eQTLs with a mediation *P*-value < 0.1 (16,648 trios) are listed in Supplemental Table S2, including trios containing transcripts with low mappability.

#### Examples of mediation across tissues

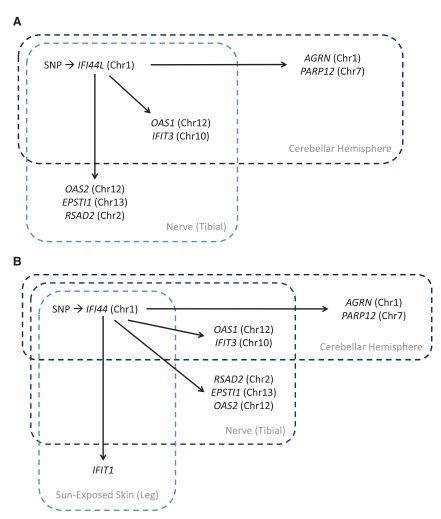
In analyses restricted to cis- and transgenes with mappability scores > 0.95, one biologically interpretable example of a *cis*-gene that appears to mediate the effects of trans-eSNPs in multiple tissues is the IFI44L gene on Chromosome 1 (Fig. 4A). IFI44L is a cis-eGene in two GTEx tissues (cerebellar hemisphere and tibial nerve), and the cis-eSNPs associated with IFI44L expression are also associated with expression of multiple genes in trans in both cerebellar and tibial nerve tissue. OAS1 is a trans target of these SNPs in both tissues, while other trans targets are observed in only cerebellar (AGRN and PARP12) or tibial nerve (RSAD2, OAS2, and EPSTI1). Below the

mappability threshold of 0.95, we observe an additional potential *trans* target of *IFI44L*, present in both cerebellar and tibial nerve tissue, *IFIT3* (mappability = 0.87). These relationships are depicted in Figure 4A.

Interestingly, if we expand our analysis to include cis- and trans-genes with mappability > 0.90, we detect IFI44 (mappability of 0.93) as a cis-mediator regulating a nearly identical set of trans genes across three tissues: cerebellar hemisphere (OAS1, IFIT2, AGRN, and PARP12), tibial nerve (OAS1, IFIT3, RSAD2, and EPSTI1), and sun-exposed skin (IFIT1) (Fig. 4B). IFI44 and IFI44L are paralogs and reside adjacent to each other on 1p31.1. These genes are regulated by the same SNP in each tissue. It is highly unlikely that sequence similarity between these two genes causes our RNAseq-based expression measurements for IFI44 (and/or IFI44L) to reflect the expression variation of both genes. The two regions of similarity shared between these two transcripts are 753 and 202 bp in length, and these regions share 69% and 67% similarity, respectively (Supplemental Fig. S3). These two regions contain no identical sequences longer than ~10 bp, making it impossible for an RNA-seq read (76 bp) to be ambiguous in terms of its mapping to IFI44 vs. IFI44L. Furthermore, a prior study of array-based

**Table 2.** Frequency of *cis*-genes that mediate the effect of a *trans*eQTL on multiple *trans*-genes or in multiple tissue types

Observed number of for each <i>cis</i> -gene	trans targets	Number of tissues for which each <i>cis</i> -gene is a mediator			
Number of <i>trans</i> targets	Cis-gene count	Number of tissues	Cis-gene count		
1	2510	1	2637		
2	482	2	420		
3	123	3	91		
4	33	4	16		
5–6	12	5	3		
7–14	8	6	1		



**Figure 4.** A biologically interpretable example of a *cis*-eGene (*IFI44L*) that appears to mediate the effects of *trans*-eSNPs in multiple tissues. The gene *IFI44L* (*A*) resides <5 kb away from *IFI44* (*B*), and expression of these genes is associated with a common *cis*-eQTL that also impacts the expression of multiple genes in *trans* in multiple tissues. Both *IFI44* and *IFI44L* show statistical evidence of mediation for a similar set of interferon-related genes. Thus, based on this evidence, we infer that at least one of these genes is a *cis*-mediator, although we cannot know which is (or if both are) the true mediator.

expression measures in endometrial cancer tissue reported genetic coregulation of *IF144* and *IF144L* (in *trans*) (Kompass and Witte 2011).

Based on these observations, it is unclear which of these two genes is truly a *cis*-mediator of the observed *trans*-eQTLs (or if both are mediators). The causal *cis*-eSNP for *IFI44* (and/or *IFI44L*) appears to be different in different tissues, as the LD between the lead *cis*-eSNPs in cerebellar (rs12129932) and the lead eSNP in tibial nerve (rs74998911) is quite low, with  $r^2 < 0.01$  in EUR 1000 Genomes data (The 1000 Genomes Project Consortium 2015).

Regardless of the uncertainty whether *IFI44L* or *IFI44* (or both) is the true *cis*-hub of this *trans*-eQTL, nearly all of the genes involved in the putative regulatory pathways identified here are interferon-regulated/inducible genes, namely *OAS1*, *OAS2*, *IFIT1*, *IFIT3*, *IFI44*, *IFI44L*, *RSAD2*, and *AGRN* (Cheon and Stark 2009; Kyogoku et al. 2013). These genes have been previously reported to be co-expressed and/or coregulated in various human cell types, including interferon-exposed fibroblasts and mammary epithelial

cell lines (Cheon and Stark 2009), virusinfected airway epithelial cell cultures (Ioannidis et al. 2012), and peripheral blood of individuals with acute respiratory infections (Zaas et al. 2009), as well as in both normal and cancerous human tissue (Cancer Cell Metabolism Gene DB, https://bioinfo.uth.edu/ccmGDB/). These previously reported co-expression findings also extend to EPSTI1 (Cheon and Stark 2009), the one gene we find to be a trans target of IFI44L (and/or IFI44) that does not have a well-established function in immune response, providing additional evidence of an immune-related function for this gene.

Variation in the *IFI44L* gene is associated with risk for MMR (measles, mumps, and rubella) vaccination-related febrile seizures, with a missense variant in *IFI44L* showing the strongest association (Feenstra et al. 2014). Variation in *IFI44L* has also been implicated in schizophrenia risk (Ruderfer et al. 2014) as well as bipolar disorder (Chen et al. 2013). These findings suggest that the putative cross-tissue *cis*-hub identified here may be relevant to multiple neurological and psychological disorders, particularly those with etiologies related to immune function.

### Comparison of GMAC with other methods using simulated data

We evaluate the performance of the proposed GMAC in various simulated data scenarios. For each scenario described below, we simulated 1000 mediation trios  $(L_i, C_i, T_j)$  for a sample size of n = 350, similar to the sample size of the GTEx data. Each mediation trio consists of a gene transcript *i*  $(C_i)$ , its *cis*-associated genetic locus  $(L_i)$ , and a gene transcript *j* 

 $(T_j)$  in *trans*-association with the locus. Note that, in the mediation analysis in this work (simulations and real data analysis), we consider only the trios with evidence of *cis*- and *trans*-associations,  $L_i \rightarrow C_i$  and  $L_i \rightarrow T_j$ . We are interested in testing whether an observed *trans*-eQTL association is mediated by the *cis*-gene transcript, i.e.,  $L_i \rightarrow C_i \rightarrow T_j$ . We compared GMAC with other methods in different scenarios, including in the presence of confounders, common child variables, and intermediate variables. A common child variable is a variable that is affected by both  $C_i$ and  $T_j$  (Fig. 2C). An intermediate variable is a variable that is affected by  $C_i$  and affecting  $T_j$ , that is, at least partially mediating the effects from  $C_i$  to  $T_j$  (Fig. 2D).

#### Scenario 1: Under the null in the presence of common child variables

This is a scenario in which there is one common child variable for each pair of *cis*- and *trans*-gene transcripts (Fig. 2C). In this scenario, adjusting for common child variables in mediation analyses would "marry"  $C_i$  and  $T_j$  and make  $C_i$  appear to be regulating  $T_j$ 

	rate in the presen		i child variable		rate in the presen		
Significance level	Oracle adjustment	GMAC algorithm	Child adjustment	Significance level	Oracle adjustment	GMAC algorithm	No adjustment
0.01	0.011	0.010	0.287	0.01	0.007	0.008	0.459
0.05	0.049	0.050	0.413	0.05	0.045	0.048	0.585
(C) Power in th	e presence of an ir	ntermediate var	iable	(D) Power in the	presence of conf	ounders	
Significance level	Oracle adjustment	GMAC algorithm	Adjusting intermediate variable	Significance level	Oracle adjustment	GMAC algorithm	Adjusting all
0.01	0.999	0.868	0.006	0.01	0.258	0.257	0.161
0.05	0.999	0.871	0.041	0.05	0.473	0.468	0.364

**Table 3.** Comparison of the type I error rate and power of GMAC compared to other methods for mediation analysis under the null (*A* and *B*) and the alternative (*C* and *D*) hypotheses, based on simulated data

even if there is no such effect (i.e., "collider bias") (Greenland 2003), increasing the false positive rate for detecting mediation. Therefore, we consider it as "improper" to adjust for common child variables. We simulated a pool of independent and normally distributed variables H, with dimensionality being the same as the sample size of 350. For each of the 1000 mediation trios, we simulated the genetic locus L<sub>i</sub> under Hardy-Weinberg Equilibrium with a minor allele frequency of 0.1. Given  $L_i$ , the *cis*-gene transcript  $C_i$ and the *trans*-gene transcript  $T_i$  are generated according to the models:  $C_i = \beta_{i0c} + \beta_{i1c} L_i + \varepsilon_{ic}$  and  $T_i = \beta_{i0t} + \beta_{i1t} L_i + \varepsilon_{it}$ . In this scenario, the trans-effect is not mediated by the cis-gene transcript. We let the parameters in the above models vary across the 1000 trios, with  $\beta_{i1c}$  sampled uniformly from 0.5 to 1.5, and the rest sampled uniformly from 0.5 to 1.0. The error terms  $\epsilon_{ic}$  and  $\epsilon_{it}$  are normally distributed. For each mediation trio, one candidate variable in **H** is randomly chosen to be the common child variable,  $Z_i$ , and the effects of *cis*- and *trans*-gene transcripts on  $Z_i$  are sampled uniformly from 1 to 1.5.

#### Scenario 2: Under the null in the presence of confounders

Scenario 2 is generated under the null in the presence of confounders (Fig. 2B). Each candidate confounding variable has a 5% probability of being a true confounder of the cis-trans relationship for a randomly chosen proportion of trios where the proportion follows a uniform distribution from 0 to 0.2. This specification results in, on average, 1.85 confounders for each trio in our simulated data. Suppose for the  $i^{th}$  trio there are  $n_i$ number of variables in H selected to be confounders; we denote the confounders as  $X_{i1}, \ldots, X_{in_i}$ . The *cis*-gene transcript  $C_i$  and trans-gene transcript  $T_i$  are generated according to the regression models  $C_i = \beta_{i0c} + \beta_{i1c}L_i + \alpha_{i1}X_{i1} + \ldots + \alpha_{in_i}X_{in_i} + \epsilon_{ic}$  and  $T_j = \beta_{i0t} + \beta_{i1t}L_i + \gamma_{i1}X_{i1} + \ldots + \gamma_{in_i}X_{in_i} + \epsilon_{it}$ . We let the parameters in the above models vary across the 1000 trios with similar parameter specification as before. In this scenario, there are no cis- to trans-gene mediation effects. Failure to adjust for confounders may induce false positive results, and it is improper to not adjust for confounders.

#### Scenario 3: Under the alternative in the presence of intermediate variables

We consider another scenario in which there is one intermediate variable for each *cis-trans* relationship (Fig. 2D). For each mediation trio, we simulated the genetic locus and the *cis*-gene transcript as before and further simulated a child variable,  $W_i$  of the *cis*-gene transcript. The *trans*-gene transcript,  $T_j$  is then simulated to be affected by  $W_i$ , according to  $T_j = \beta_{i0t} + \beta_{i1t}L_i + \gamma_i W_i + \varepsilon_{it}$ . Note that the

*trans*-gene  $T_i$  is simulated to be affected by the *cis*-gene  $C_i$  only via the intermediate variable  $W_i$ . The mediation effects from *cis*- to *trans*-gene transcript (via  $W_i$ ) is nonzero in this scenario. Because  $W_i$  is on the causal pathway from  $C_i$  to  $T_j$ , it is improper to adjust for  $W_i$ , and the adjustment will reduce or eliminate power to detect true mediation.

#### Scenario 4: Under the alternative in the presence of confounders

To compare with the existing approach that adjusts for a universal set of variables, we consider a scenario in which the dimensionality of potential confounding variables **H** is 100. For each trio, up to five variables in **H** are randomly selected to confound the *cis-trans* gene relationship. The absolute effects of confounders on *cis* transcripts are sampled uniformly from 0.15 to 0.5 with a 50% probability to be negative; the effects of confounders on *trans* transcripts are sampled uniformly from 0.15 to 0.5, with all to be positive. We set the effect of *cis* transcript on *trans* transcript to be 0.1, i.e., non-zero mediation effects. When the number of potential confound-ing variables is large, although one may still adjust them all for the simulated sample size, this adjustment is inefficient and may hurt the power.

#### Simulation results

For each scenario, we compared the results based on the following methods: (1) Oracle adjustment, which correctly adjusts for the true confounders but no child or intermediate variables in the mediation test; (2) the GMAC algorithm; and (3) improper (or inefficient) adjustment, which corresponds to incorrectly adjusting for the common child variables in scenario 1, failure to adjust for confounders in scenario 2, incorrectly adjusting for the intermediate variables in scenario 3, and universally adjusting for all variables in H in scenario 4, including variables which are not true confounders. Table 3, A and B, shows the true type I error rates at the significance levels of 0.01 and 0.05 in scenarios 1 and 2, respectively. As expected, adjusting for child variables "marries" the cisand trans-genes in the mediation test, resulting in inflated rates of false positive findings. Failure to adjust for confounding also leads to inflated type I error rates. In contrast, both the Oracle and GMAC adjustment control the type I error rates. Table 3C shows that, when the power to detect mediation is high (by Oracle and GMAC), incorrectly adjusting for an intermediate variable in this setting greatly reduces the power to detect mediation. In comparison, GMAC correctly filters out most of the true intermediate variables in the adjustment for mediation tests and maintains power comparable to Oracle adjustment. Table 3D shows that GMAC has

comparable power to Oracle adjustment in scenario 4. In our simulation, 2962 out of 3023 generated confounders across the 1000 mediation trios are correctly selected. In comparison, adjusting for all variables in the pool of confounders is inefficient and reduces power to detect mediation.

#### Discussion

In this work, we have developed the GMAC algorithm for conducting mediation analysis to identify cis transcripts that mediate the effects of trans-eQTLs on distant genes. We address a central problem in mediation analysis, "mediator-outcome confounding," by developing an algorithm that can (1) search a very large pool of variables (surrogate and/or measured) for variables likely to have confounding effects and (2) adaptively adjust for such variables in each mediation test conducted. We acknowledge that we cannot make definitive causal claims regarding any of the mediation/regulatory relationships for which we detect evidence. Instead, the focus of our work is to strengthen the evidence that mediation analysis can provide and to identify candidate cis-hub genes likely to mediate the effects of eQTLs on many trans-genes within or across human tissue types. Cis-hub genes are likely to be key players in the regulatory networks relevant to human disease, thus it is important that we understand their patterns of regulation. By applying GMAC to 44 human tissues from the GTEx project, we are able to characterize cis-hubs with potential disease relevance by aggregating information across many different tissue types. Analyses of simulated data show that the GMAC algorithm improves the power to detect true mediation compared with existing methods, while controlling the true false positive rate.

In analyses of GTEx data, over 20% of *cis*-mediators we observe appear to mediate the effects of a *trans*-eQTL on multiple genes, but the vast majority of these *cis*-hubs are either tissue-specific (i.e., mediating multiple *trans*-genes in a single tissue type) or have unique *trans* targets in each tissue type. We provided one example of a biologically plausible multitissue *cis*-hub, whereby a *cis*mediator of *trans*-eQTLs appears to have common *trans* targets across multiple tissue types. The *cis*-hub identified (*IFI44L*) has potential relevance for neurological and psychological disorders, particularly those with etiologies related to immune function, demonstrating the potential value of our approach for understanding disease-relevant pathways.

One innovative aspect of this work is our algorithm that rigorously addresses the problem of "mediator-outcome confounding" in the context of genomic mediation analysis. In eQTLbased mediation analysis, potential confounders of the *cis-trans* association include demographic and environmental factors, as well as a wide array of biological phenomena, such as expression of specific genes or other biological processes that may be represented by the expression of sets of genes. Neglecting to control for such confounding variables can lead to substantial bias in estimates of mediation, resulting in spurious findings, as we have described previously (Pierce et al. 2014). Considering the complexity of the biological systems under study, as well as the diversity of the GTEx donors, a careful control for such confounding variables is critically important.

Most existing methods control for confounding variables by constructing a set of variables that represent the largest components of variation in the transcriptome and adjusting for the selected set for all tests conducted. However, only when a variable is causally related to both the *cis*- and *trans*-genes (as shown in Fig. 2B) will the variable potentially show confounding effects in mediation analysis. GMAC adaptively selects a set of confounding variables for each trio undergoing mediation analysis, enabling large-scale genomic mediation analyses adjusting only for the confounding variables that could potentially bias a specific mediation estimate. As opposed to adjusting for all known covariates, our strategy of selecting only potential confounders for adjustment purposes is important for three reasons: (1) Adjusting for fewer variables increases power (i.e., fewer degrees of freedom) (see Supplemental Table S1); (2) the number of variables from which one selects covariates could be extremely large (e.g., all expressed genes), making adjustment for all covariates impossible; and (3) inadvertently adjusting for "common child" or intermediate variables can result in substantial biases. In this work, we select potential confounders from all expression PCs, but one could also select from among transcripts that are not well-represented by PCs. By efficiently selecting confounders from a very large pool of potential variables, GMAC improves both power and precision in mediation analyses.

There are several limitations of our approach and its application to GTEx data. First, when working with real genomic data, we can never be sure that we have measured and accounted for all possible mediator-outcome confounding. Potential confounders include participant characteristics, environmental factors, and tissue micro-environmental factors, as well as a wide array of biological factors which may or may not be captured by the expression data being analyzed. Second, in the analysis presented here, we only consider the trios with both strong cis- and trans-eQTL effects. For any given tissue type we are analyzing, our sample size is too small for robust genome-wide detection/analysis of transeQTLs. As such, the mediation trios we considered are only a subset of the true mediation trios in the genome, and the small sample sizes may also result in underpowered mediation tests. As the sample size of GTEx increases, future studies will have increased power to identify cis-mediators using GMAC. Third, we did not consider the full complexity of gene isoforms and splice variants in this work; future studies should consider the possibility of mediation relationships that are isoform-specific. Lastly, some trans-eQTLs may not be mediated by variation in the expression of a cis-gene. Other potential mediating mechanisms could include variation in coding sequence, physical inter-chromosomal interaction, or variation in noncoding RNA. Our work is not intended to identify and analyze such trans-eQTLs, as we perform trans-eQTL analyses using only SNPs known to be cis-eSNPs.

It is important to note that our expectation is that most *trans*eQTLs are fully mediated by a transcript that is regulated in *cis* by the causal *trans*-eQTL variant. We did not observe "complete mediation" (i.e., % mediation = 100%) for the majority of the significant mediation *P*-values we observed. However, as we have explained and demonstrated previously (Pierce et al. 2014), full mediation will be observed as partial mediation in the presence of mediator measurement error and/or imperfect LD between the causal variant and the variant used for analysis purposes. Thus, considering RNA quantification is not error-free and causal variants are often unknown, we expect to often observe partial mediation when full mediation is present.

We also demonstrate that it is critical to consider mappability for both *cis*- and *trans*-genes involved in mediation analysis. For genes containing sequences that do not uniquely map to the human transcriptome, it is possible that gene expression measures may be comprised of signals coming from multiple genes, which can produce false positives in mediation analysis, including spurious detection of *cis*-hubs and cross-tissue *cis*-mediators.

Our application of the GMAC algorithm to the multitissue expression data from GTEx provides a unique cross-tissue perspective on cis-mediation of trans-regulatory relationships across human tissues. This multitissue perspective is important because observing mediation relationships that are consistent across multiple tissues provides confidence that a significant mediation P-value reflects a true instance of mediation. For the "cis-hub" genes and genes that appear to be cis-mediators in multiple tissues, further investigation is warranted, as these genes may have many regulatory relationships that we are not powered to detect in this work. Thus, a multitissue mediation analysis approach has the potential to increase power to identify true mediators while controlling for false positives. In future work, attempts at joint analyses of multiple tissue types may provide a more complete picture of the cross-tissue and tissue-specific trans-regulatory mechanisms. The GMAC approach described here will be a valuable tool for such studies as well as any future studies that aim to understand the relationships among cis- and trans-eQTLs and characterize the biological mechanisms and networks involved in human disease biology.

We have developed an R GMAC package to perform the proposed genomic mediation analysis with adaptive selection of confounding variables. The package is currently available through R CRAN.

#### **Methods**

#### Biospecimen collection and processing of GTEx data

A total of 7051 tissues samples were obtained from 44 distinct tissue types from 449 post-mortem tissue donors (with 65.6% male). Those donors were from multiple ethnicity groups, spanned a wide age range, and have various causes of death (see GTEx portal for descriptive statistics). Donor enrollment, consent processes, and biospecimen collection and processing have been described previously (Lonsdale et al. 2013; Ardlie et al. 2015). Briefly, each tissue sample was preserved in a PAXgene tissue kit and stored as both frozen and paraffin-embedded tissue. Total RNA was isolated from PAXgene fixed tissue samples using the PAXgene Tissue mRNA kit. For whole blood, total RNA was isolated from samples collected and preserved in PAXgene blood RNA tubes.

Blood samples were used as the primary source of DNA. Genotyping was conducted using the Illumina Human Omni5-Quad and Infinium ExomeChip arrays. Standard QC procedures were performed using PLINK software (Purcell et al. 2007), and genotype imputation was performed using IMPUTE2 software (Howie et al. 2009) and reference haplotypes from the 1000 Genomes Project (The 1000 Genomes Project Consortium 2015). The first three PCs representing ancestry (Price et al. 2006) were included as covariates in all analyses.

RNA-seq data were generated for RNA samples with an RIN value of six or greater. Nonstrand-specific RNA sequencing was performed using an automated version of the Illumina TruSeq RNA sample preparation protocol. Sequencing was done on an Illumina HiSeq 2000, to a median depth of 78M 76-bp pairedend reads per sample. RNA-seq data were aligned to the human genome using TopHat (Trapnell et al. 2009). Gene-level expression was estimated in RPKM units using RNA-SeQC (DeLuca et al. 2012). RNA-seq expression samples that passed various quality control measures (as previously described) were included in the final analysis data set.

#### Mappability of transcripts

Because nonuniquely mapping reads can result in false positive eQTLs, we use the mappability of each gene as a quality control

filter, as described by The GTEx Consortium (2017). The mappability was calculated as follows: Mappability of all *k*-mers in the reference human genome (hg19) computed by ENCODE (The ENCODE Project Consortium 2012) was downloaded from the UCSC Genome Browser (accession: wgEncodeEH000318, wgEncodeEH00032) (Rosenbloom et al. 2013). The exon- and UTR-mappability of a gene were computed as the average mappability of all *k*-mers in exons and UTRs, respectively. We used k =75 for exonic regions, as it is the closest to GTEx read length among all possible *k*'s. UTRs are generally quite small, so k = 36 was used, the smallest among all possible *k*'s. Mappability of a gene was computed as the weighted average of its exonmappability and UTR-mappability, with the weights being proportional to the total length of exonic regions and UTRs, respectively.

#### The selection of trios for mediation tests

In the mediation analysis presented in this work, we consider only the trios with evidence of *cis* and *trans* associations,  $L_i \rightarrow C_i$  and  $L_i \rightarrow T_j$ . The identification of *cis*-eQTLs is described elsewhere (Ardlie et al. 2015; The GTEx Consortium 2017). For genes with multiple *cis*-eSNPs as eQTLs, only one *cis*-eSNP for each gene (i.e., the high-quality SNP with the smallest *P*-value) was selected and was included in the subsequent *trans*-eQTL and mediation analyses. The complete *cis*-eQTL list is available through the GTEx portal (https://gtexportal.org/), and all data can be obtained through dbGaP (phs000424.v6.p1).

Furthermore, using Matrix eQTL (Shabalin 2012), we conducted genome-wide *trans*-eQTL analyses, restricted to the *cis*eSNPs described above and examining association for all genes located at least 1 Mb away from the *cis*-eSNPs. Up to 35 PEER factors and other covariates were adjusted. In each tissue type, when a *trans*-association *P*-value is less than  $10^{-5}$ , the eSNP, its corresponding *cis* transcript, and the *trans* transcript were treated as a candidate trio and were retained for mediation analysis.

#### The GMAC algorithm

In order to identify *cis*-mediators of *trans*-eQTLs across the genome, we propose the GMAC algorithm (Fig. 2A). Here, we present a brief description of each step. A detailed description and justification for each can be found in the Supplemental Methods. Specifically,

- Step 0. We focus on only the trios (*L<sub>i</sub>*, *C<sub>i</sub>*, *T<sub>j</sub>*) in the genome showing both *cis* and *trans*-eQTL associations, i.e., *L<sub>i</sub>* → *C<sub>i</sub>* and *L<sub>i</sub>* → *T<sub>j</sub>*. Consider a pool of candidate variables **H** consisting of either real covariates, constructed surrogate variables, or both.
- Step 1. Filter out common child and intermediate variables from the pool of potential confounders. For each trio  $(L_i, C_i, T_j)$ , we calculate the marginal associations of variables in **H** to  $L_i$  and filter the ones with significant associations at the 10% FDR level. As shown in Figure 2B–D, common child and intermediate variables are directly associated with  $L_i$ , while confounders are assumed to be unassociated with  $L_i$ . Note that since genetic loci are "Mendelian randomized" (Smith and Ebrahim 2003), without loss of generality we assume the confounders are not associated with  $L_i$ . Let **H**<sub>ij</sub> denote the retained pool of candidate variables specific to the trio  $(L_i, C_i, T_j)$ .
- Step 2. Adaptively select confounders. For each trio and each of its potential confounding variables in **H**<sub>ij</sub>, we calculate the *P*-value of the overall *F*-test to assess the association of the variable to at least one of the *cis* and *trans* transcripts. Considering the *P*-values for one potential confounding variable to all trios

as one stratum, we apply a 10% FDR significance threshold to each stratum (each column in Fig. 2E)-a stratified FDR approach (Sun et al. 2006). The significant variables corresponding to a trio (each row in Fig. 2E) will be selected in the mediation analyses as the adaptively selected confounders specific to that trio (see Supplemental Methods for details). Let X<sub>ii</sub> denote the list of adaptively selected confounding variables for the trio,  $(L_i, C_i, T_i)$ .

• Step 3. Test for mediation. For each trio and its adaptively selected confounder set, we calculate the mediation statistic as the Wald statistic for testing the indirect mediation effect  $H_0$ :  $\beta_1 =$ 0 based on the following regression regressing the trans-gene expression levels on the cis-gene expression levels adjusting for the cis-eQTL, other covariates, and selected confounders:

$$T_i = \beta_0 + \beta_1 C_i + \beta_2 L_i + \Gamma X_{ij} + \varepsilon.$$

We perform within-genotype group permutation on the cisgene transcript at least 10,000 times and recalculate each null mediation statistic based on the locus, a permuted cis-gene transcript, and the trans-gene transcript,  $(L_i, C_{i0}, T_j)$ . Figure 2F shows the expression variation patterns of a hypothetical mediation relationship  $L_i \rightarrow C_i \rightarrow T_i$  on the left panel, and a null relationship entailed by  $(L_i, C_{i0}, T_j)$  with  $L_i \rightarrow C_{i0}$  and  $L_i \rightarrow T_j$  but no mediation. It justifies that by permuting the cis-gene expression levels within each genotype group, one maintains the cisassociations while breaking the potential mediation effects from the cis- to the trans-gene transcript (i.e., conditional correlations of *cis* and *trans* transcript conditioning on L<sub>i</sub>, or correlation within each genotype group). We calculate the *P*-value of mediation for the trio  $(L_i, C_i, T_i)$  by comparing the observed mediation statistic with the null statistics.

The proposed algorithm is superior to existing approaches for mediation analysis that adjust a universal set of variables for all trios. GMAC avoids the adjustment of common child variables, intermediate variables, and unrelated variables in genomic mediation analysis, and it is able to search a much larger pool of variables for potential confounders, not just those captured by the top few surrogate variables or PCs.

#### Software availability

The R software package GMAC is available in the Supplemental Materials and also online through R CRAN https://cran.r-project. org/.

#### **GTEx** Consortium

#### Laboratory, Data Analysis & Coordinating Center (LDACC)-Analysis Working Group

François Aguet,<sup>6</sup> Kristin G. Ardlie,<sup>6</sup> Beryl B. Cummings,<sup>6,7</sup> Ellen T. Gelfand,<sup>6</sup> Gad Getz,<sup>6,8</sup> Kane Hadley,<sup>6</sup> Robert E. Handsaker,<sup>6,9</sup> Katherine H. Huang,<sup>6</sup> Seva Kashin,<sup>6,9</sup> Konrad J. Karczewski,<sup>6,7</sup> Monkol Lek,<sup>6,7</sup> Xiao Li,<sup>6</sup> Daniel G. MacArthur,<sup>6,7</sup> Jared L. Nedzel,<sup>6</sup> Duyen T. Nguyen,<sup>6</sup> Michael S. Noble,<sup>6</sup> Ayellet V. Segrè,<sup>6</sup> Casandra A. Trowbridge,<sup>6</sup> and Taru Tukiainen<sup>6,7</sup>

Nathan S. Abell,<sup>10,11</sup> Brunilda Balliu,<sup>11</sup> Ruth Barshir,<sup>12</sup> Omer Basha,<sup>12</sup> Alexis Battle,<sup>13</sup> Gireesh K. Bogu,<sup>14,15</sup> Andrew Brown,<sup>16,17,18</sup> Christopher D. Brown,<sup>19</sup> Stephane E. Castel,<sup>20,21</sup> Lin S. Chen,<sup>22</sup> Colby Chiang,<sup>23</sup> Donald F. Conrad,<sup>24,25</sup> Nancy J. Cox,<sup>26</sup> Farhan N. Damani,<sup>13</sup> Joe R. Davis,<sup>10,11</sup> Olivier Delaneau,<sup>16,17,18</sup> Emmanouil T. Dermitzakis,<sup>16,17,18</sup> Barbara E. Engelhardt,<sup>27</sup> Eleazar Eskin,<sup>28,29</sup> Pedro G. Ferreira,<sup>30,31</sup> Laure Frésard,<sup>10,11</sup> Eric R. Gamazon,<sup>26,32,33</sup> Diego Garrido-Martín,<sup>14,15</sup> Ariel D.H. Gewirtz,<sup>34</sup> Genna Gliner,<sup>35</sup> Michael Gloudemans,<sup>10,11,36</sup> Roderic Guigo,<sup>14,15,37</sup> Ira M. Hall,<sup>23,24,38</sup> Buhm Han,<sup>39</sup> Yuan He,<sup>40</sup> Farhad Hormozdiari,<sup>28</sup> Cedric Howald,<sup>16,17,18</sup> Hae Kyung Im,<sup>41</sup> Brian Jo,<sup>34</sup> Eun Yong Kang,<sup>28</sup> Yungil Kim,<sup>13</sup> Sarah Kim-Hellmuth,<sup>20,21</sup> Tuuli Lappalainen,<sup>20,21</sup>

Learning, Princeton University, Princeton, NJ 08540, USA

<sup>28</sup>Department of Computer Science, University of California, Los Angeles, CA 90095, USA <sup>29</sup>Department of Human Genetics, University of California, Los Angeles, CA

90095, USA <sup>30</sup>Instituto de Investigação e Inovação em Saúde (i3S), Universidade do Porto,

4200-135 Porto, Portugal

<sup>31</sup>Institute of Molecular Pathology and Immunology (IPATIMUP), University of Porto, 4200-625 Porto, Portugal

<sup>32</sup>Department of Clinical Epidemiology, Biostatistics and Bioinformatics, Academic Medical Center, University of Amsterdam, 1105 AZ Amsterdam, The Netherlands

<sup>33</sup>Department of Psychiatry, Academic Medical Center, University of Amsterdam, 1105 AZ Amsterdam, The Netherlands

<sup>34</sup>Lewis Sigler Institute, Princeton University, Princeton, NJ 08540, USA

<sup>35</sup>Department of Operations Research and Financial Engineering, Princeton University, Princeton, NJ 08540, USA <sup>36</sup>Biomedical Informatics Program, Stanford University, Stanford, CA 94305,

USA <sup>37</sup>Institut Hospital del Mar d'Investigacions Mèdiques (IMIM), 08003 Barcelona,

Spain <sup>38</sup>Department of Medicine, Washington University School of Medicine, St. Louis, MO 63108, USA. <sup>39</sup>Department of Convergence Medicine, University of Ulsan College of

<sup>&</sup>lt;sup>6</sup>The Broad Institute of Massachusetts Institute of Technology and Harvard University, Cambridge, MA 02142, USA

<sup>&</sup>lt;sup>7</sup>Analytic and Translational Genetics Unit, Massachusetts General Hospital, Boston, MA 02114, USA

<sup>&</sup>lt;sup>8</sup>Massachusetts General Hospital Cancer Center and Dept. of Pathology, Massachusetts General Hospital, Boston, MA 02114, USA

<sup>&</sup>lt;sup>9</sup>Department of Genetics, Harvard Medical School, Boston, MA 02114, USA

<sup>&</sup>lt;sup>10</sup>Department of Genetics, Stanford University, Stanford, CA 94305, USA <sup>11</sup>Department of Pathology, Stanford University, Stanford, CA 94305, USA

<sup>&</sup>lt;sup>12</sup>Department of Clinical Biochemistry and Pharmacology, Faculty of Health Sciences, Ben-Gurion University of the Negev, Beer-Sheva 84105, Israel <sup>13</sup>Department of Computer Science, Johns Hopkins University, Baltimore, MD

<sup>21218,</sup> USA <sup>14</sup>Centre for Genomic Regulation (CRG), The Barcelona Institute for Science and

Technology, 08003 Barcelona, Spain

<sup>&</sup>lt;sup>15</sup>Universitat Pompeu Fabra (UPF), 08003 Barcelona, Spain

<sup>&</sup>lt;sup>16</sup>Department of Genetic Medicine and Development, University of Geneva Medical School, 1211 Geneva, Switzerland

<sup>&</sup>lt;sup>7</sup>Institute for Genetics and Genomics in Geneva (iG3), University of Geneva, 1211 Geneva, Switzerland

<sup>&</sup>lt;sup>18</sup>Swiss Institute of Bioinformatics, 1211 Geneva, Switzerland

<sup>&</sup>lt;sup>19</sup>Department of Genetics, Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA 19104, USA

<sup>&</sup>lt;sup>20</sup>New York Genome Center, New York, NY 10013, USA

<sup>&</sup>lt;sup>21</sup>Department of Systems Biology, Columbia University Medical Center, New York, NY 10032, USA <sup>22</sup>Department of Public Health Sciences, The University of Chicago, Chicago, IL

<sup>60637,</sup> USA <sup>23</sup>McDonnell Genome Institute, Washington University School of Medicine, St.

Louis, MO 63108, USA <sup>24</sup>Department of Genetics, Washington University School of Medicine, St. Louis,

MO 63108, USA <sup>25</sup>Department of Pathology & Immunology, Washington University School of

Medicine, St. Louis, MO 63108, USA <sup>26</sup>Division of Genetic Medicine, Department of Medicine, Vanderbilt University

Medical Center, Nashville, TN 37232, USA <sup>27</sup>Department of Computer Science, Center for Statistics and Machine

Medicine, Asan Medical Center, Seoul 138-736, South Korea <sup>40</sup>Department of Biomedical Engineering, Johns Hopkins University, Baltimore,

MD 21218, USA <sup>41</sup>Section of Genetic Medicine, Department of Medicine, The University of

Chicago, Chicago, IL 60637, USA

Gen Li,<sup>42</sup> Xin Li,<sup>11</sup> Boxiang Liu,<sup>10,11,43</sup> Serghei Mangul,<sup>28</sup> Mark I. Gen Ll, <sup>12</sup> Ain Ll, <sup>14</sup> Boxlang Llu, <sup>14</sup> Sergirer Mangui, <sup>14</sup> Marxi, <sup>15</sup> McCarthy, <sup>44,45,46</sup> Ian C. McDowell, <sup>47</sup> Pejman Mohammadi, <sup>20,21</sup> Jean Monlong, <sup>14,15,48</sup> Stephen B. Montgomery, <sup>10,11</sup> Manuel Muñoz-Aguirre, <sup>14,15,49</sup> Anne W. Ndungu, <sup>44</sup> Dan L. Nicolae,<sup>41,50,51</sup> Andrew B. Nobel,<sup>52,53</sup> Meritxell Oliva,<sup>41,54</sup> Halit Nicolae,<sup>41,30,31</sup> Andrew B. Nobel,<sup>22,33</sup> Meritxell Oliva,<sup>41,34</sup> Halit Ongen,<sup>16,17,18</sup> John J. Palowitch,<sup>52</sup> Nikolaos Panousis,<sup>16,17,18</sup> Panagiotis Papasaikas,<sup>14,15</sup> YoSon Park,<sup>19</sup> Princy Parsana,<sup>13</sup> Anthony J. Payne,<sup>44</sup> Christine B. Peterson,<sup>55</sup> Jie Quan,<sup>56</sup> Ferran Reverter,<sup>14,15,57</sup> Chiara Sabatti,<sup>58,59</sup> Ashis Saha,<sup>13</sup> Michael Sammeth,<sup>60</sup> Alexandra J. Scott,<sup>23</sup> Andrey A. Shabalin,<sup>61</sup> Reza Sodaei,<sup>14,15</sup> Matthew Stephens,<sup>50,51</sup> Barbara E. Stranger,<sup>41,54,62</sup> Benjamin J. Strober,<sup>40</sup> Jae Hoon Sul,<sup>63</sup> Emily K. Tsang,<sup>11,36</sup> Sarah Urbut,<sup>51</sup> Martijn van de Bunt,<sup>44,45</sup> Gao Wang,<sup>51</sup> Xiaoquan Wen,<sup>64</sup> Fred A. Wright,<sup>65</sup> Hualin S. Xi,<sup>56</sup> Esti Yeger-Lotem,<sup>12,66</sup> Zachary Zappala,<sup>10,11</sup> Judith B. Zaugg,<sup>67</sup> and Yi-Hui Zhou<sup>65</sup>

#### Enhancing GTEx (eGTEx) groups

Joshua M. Akey, <sup>34,68</sup> Daniel Bates, <sup>69</sup> Joanne Chan, <sup>10</sup> Lin S. Chen, <sup>22</sup> Melina Claussnitzer,<sup>6,70,71</sup> Kathryn Demanelis,<sup>22</sup> Morgan Diegel,<sup>69</sup> Jennifer A. Doherty,<sup>72</sup> Andrew P. Feinberg,<sup>40,73,74,75</sup>

<sup>42</sup>Department of Biostatistics, Mailman School of Public Health, Columbia University, New York, NY 10032, USA

<sup>3</sup>Department of Biology, Stanford University, Stanford, CA 94305, USA

<sup>44</sup>Wellcome Trust Centre for Human Genetics, Nuffield Department of Medicine, University of Oxford, OX3 7BN, UK
 <sup>45</sup>Oxford Centre for Diabetes, Endocrinology and Metabolism, University of Oxford, OX3 7LE, UK
 <sup>46</sup>Oxford NIHR Biomedical Research Centre, Churchill Hospital, Oxford, OX3

7LJ, UK <sup>47</sup>Computational Biology & Bioinformatics Graduate Program, Duke University, Durham, NC 27708, USA <sup>48</sup>Human Genetics Department, McGill University, Montreal, Quebec H3A 0G1,

<sup>49</sup>Departament d'Estadística i Investigació Operativa, Universitat Politècnica de Catalunya, 08034 Barcelona, Spain <sup>50</sup>Department of Statistics, The University of Chicago, Chicago, IL 60637, USA

<sup>51</sup>Department of Human Genetics, The University of Chicago, Chicago, IL

50637, USA <sup>52</sup>Department of Statistics and Operations Research, University of North Carolina, Chapel Hill, NC 27599, USA <sup>53</sup>Department of Biostatistics, University of North Carolina, Chapel Hill, NC

27599, USA <sup>54</sup>Institute for Genomics and Systems Biology, The University of Chicago, <sup>55</sup>Department of Biostatistics, The University of Texas MD Anderson Cancer

<sup>56</sup>Computational Sciences, Pfizer Inc, Cambridge, MA 02139, USA

<sup>58</sup>Diversitat de Barcelona, Os028 Barcelona, Catalonia, Spain <sup>58</sup>Department of Biomedical Data Science, Stanford University, Stanford, CA <sup>59</sup>Department of Statistics, Stanford University, Stanford, CA 94305, USA

- <sup>60</sup>Institute of Biophysics Carlos Chagas Filho (IBCCF), Federal University of Rio

de Janeiro (UFRJ), 21941902 Rio de Janeiro, Brazil <sup>61</sup>Department of Psychiatry, University of Utah, Salt Lake City, UT 84108, USA <sup>62</sup>Center for Data Intensive Science, The University of Chicago, Chicago, IL

60637, USA <sup>63</sup>Department of Psychiatry and Biobehavioral Sciences, University of California,

Los Angeles, CA 90095, USA <sup>64</sup>Department of Biostatistics, University of Michigan, Ann Arbor, MI 48109,

USA <sup>65</sup>Bioinformatics Research Center and Departments of Statistics and Biological Sciences, North Carolina State University, Raleigh, NC 27695, USA

<sup>6</sup>National Institute for Biotechnology in the Negev, Beer-Sheva, 84105 Israel

<sup>67</sup>European Molecular Biology Laboratory, 69117 Heidelberg, Germany <sup>68</sup>Department of Ecology and Evolutionary Biology, Princeton University,

Princeton, NJ 08540, USA <sup>69</sup>Altius Institute for Biomedical Sciences, Seattle, WA 98121, USA

<sup>70</sup>Beth Israel Deaconess Medical Center, Harvard Medical School, Boston, MA

02215, USA <sup>71</sup>University of Hohenheim, 70599 Stuttgart, Germany

Marian S. Fernando,<sup>41,54</sup> Jessica Halow,<sup>69</sup> Kasper D. Hansen,<sup>73,76,77</sup> Eric Haugen,<sup>69</sup> Peter F. Hickey,<sup>77</sup> Lei Hou,<sup>6,78</sup> Farzana Jasmine,<sup>22</sup> Ruiqi Jian,<sup>10</sup> Lihua Jiang,<sup>10</sup> Audra Johnson,<sup>69</sup> Rajinder Kaul,<sup>69</sup> Manolis Kellis,<sup>6,78</sup> Muhammad G. Kibriya,<sup>22</sup> Kristen Lee,<sup>69</sup> Jin Billy Li,<sup>10</sup> Qin Li,<sup>10</sup> Xiao Li,<sup>10</sup> Jessica Lin,<sup>10,79</sup> Shin Lin,<sup>10,80</sup> Sandra Linder,<sup>10,11</sup> Caroline Linke,<sup>41,54</sup> Yaping Liu,<sup>6,78</sup> Matthew T. Maurano,<sup>81</sup> Benoit Molinie,<sup>6</sup> Stephen B. Montgomery,<sup>10,11</sup> Jemma Nelson,<sup>69</sup> Fidencio J. Neri,<sup>69</sup> Meritxell Oliva,<sup>41,54</sup> Yongjin Park,<sup>6,78</sup> Brandon L. Pierce,<sup>22</sup> Nicola J. Rinaldi,<sup>6,78</sup> Lindsay F. Rizzardi,<sup>73</sup> Richard Sandstrom,<sup>69</sup> Andrew Skol,<sup>41,54,62</sup> Kevin S. Smith,<sup>10,11</sup> Michael P. Snyder,<sup>10</sup> John Stamatoyannopoulos,<sup>69,79,82</sup> Barbara E. Stranger,<sup>41,54,62</sup> Hua Stamatoyannopoulos,<sup>69,79,82</sup> Barbara E. Stranger,<sup>41,54,62</sup> Hua Tang,<sup>10</sup> Emily K. Tsang,<sup>11,36</sup> Li Wang,<sup>6</sup> Meng Wang,<sup>10</sup> Nicholas Van Wittenberghe,<sup>6</sup> Fan Wu,<sup>41,54</sup> and Rui Zhang<sup>10</sup>

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Philip A. Branton,<sup>84</sup> Latarsha J. Carithers,<sup>84,85</sup> Ping Guan,<sup>84</sup> Helen M. Moore,<sup>84</sup> Abhi Rao,<sup>84</sup> and Jimmie B. Vaught<sup>84</sup>

#### NIH/NHGRI

Sarah E. Gould,<sup>86</sup> Nicole C. Lockart,<sup>86</sup> Casey Martin,<sup>86</sup> Jeffery P. Struewing,86 and Simona Volpi86

#### NIH/NIMH

Anjene M. Addington<sup>87</sup> and Susan E. Koester<sup>87</sup>

#### NIH/NIDA

A. Roger Little<sup>88</sup>

Baltimore, MD 21205, USA

<sup>74</sup>Department of Medicine, Johns Hopkins University School of Medicine, Baltimore, MD 21205, USA

<sup>75</sup>Department of Mental Health, Johns Hopkins University School of Public Health, Baltimore, MD 21205, USA

<sup>76</sup>McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins School of Medicine, Baltimore, MD 21205, USA

<sup>77</sup>Department of Biostatistics, Johns Hopkins University, Baltimore, MD 21205,

USA 78 Computer Science and Artificial Intelligence Laboratory, Massachusetts Institute of Technology, Cambridge, MA 02139, USA

<sup>79</sup>Department of Medicine, University of Washington, Seattle, WA 98195, USA <sup>80</sup>Division of Cardiology, University of Washington, Seattle, WA 98195, USA

<sup>81</sup>Institute for Systems Genetics, New York University Langone Medical Center, New York, NY 10016, USA

<sup>82</sup>Department of Genome Sciences, University of Washington, Seattle, WA 98195, USA

<sup>83</sup>Office of Strategic Coordination, Division of Program Coordination, Planning and Strategic Initiatives, Office of the Director, NIH, Rockville, MD 20852, USA <sup>84</sup>Biorepositories and Biospecimen Research Branch, Division of Cancer

Treatment and Diagnosis, National Cancer Institute, Bethesda, MD 20892, USA <sup>85</sup>National Institute of Dental and Craniofacial Research, Bethesda, MD 20892,

USA <sup>86</sup>Division of Genomic Medicine, National Human Genome Research Institute, Rockville, MD 20852, USA <sup>87</sup>Division of Neuroscience and Basic Behavioral Science, National Institute of

Mental Health, NIH, Bethesda, MD 20892, USA <sup>88</sup>Division of Neuroscience and Behavior, National Institute on Drug Abuse,

NIH, Bethesda, MD 20892, USA

<sup>&</sup>lt;sup>72</sup>Huntsman Cancer Institute, Department of Population Health Sciences, University of Utah, Salt Lake City, UT 84112, USA <sup>73</sup>Center for Epigenetics, Johns Hopkins University School of Medicine,

#### Biospecimen Collection Source Site-NDRI

Lori E. Brigham,<sup>89</sup> Richard Hasz,<sup>90</sup> Marcus Hunter,<sup>91</sup> Christopher Johns,<sup>92</sup> Mark Johnson,<sup>93</sup> Gene Kopen,<sup>94</sup> William F. Leinweber,<sup>94</sup> John T. Lonsdale,<sup>94</sup> Alisa McDonald,<sup>94</sup> Bernadette Mestichelli,<sup>94</sup> Kevin Myer,<sup>91</sup> Brian Roe,<sup>91</sup> Michael Salvatore,<sup>94</sup> Saboor Shad,<sup>94</sup> Jeffrey A. Thomas,<sup>94</sup> Gary Walters,<sup>93</sup> Michael Washington,<sup>93</sup> and Joseph Wheeler<sup>92</sup>

#### Biospecimen Collection Source Site—RPCI

Jason Bridge,<sup>95</sup> Barbara A. Foster,<sup>96</sup> Bryan M. Gillard,<sup>96</sup> Ellen Karasik,<sup>96</sup> Rachna Kumar,<sup>96</sup> Mark Miklos,<sup>95</sup> and Michael T. Moser<sup>96</sup>

#### Biospecimen Core Resource—VARI

Scott D. Jewell,<sup>97</sup> Robert G. Montroy,<sup>97</sup> Daniel C. Rohrer,<sup>97</sup> and Dana R. Valley<sup>97</sup>

#### Brain Bank Repository—University of Miami Brain Endowment Bank

David A. Davis<sup>98</sup> and Deborah C. Mash<sup>98</sup>

#### Leidos Biomedical—Project Management

Anita H. Undale,<sup>99</sup> Anna M. Smith,<sup>100</sup> David E. Tabor,<sup>100</sup> Nancy V. Roche,<sup>100</sup> Jeffrey A. McLean,<sup>100</sup> Negin Vatanian,<sup>100</sup> Karna L. Robinson,<sup>100</sup> Leslie Sobin,<sup>100</sup> Mary E. Barcus,<sup>101</sup> Kimberly M. Valentino,<sup>100</sup> Liqun Qi,<sup>100</sup> Steven Hunter,<sup>100</sup> Pushpa Hariharan,<sup>100</sup> Shilpi Singh,<sup>100</sup> Ki Sung Um,<sup>100</sup> Takunda Matose,<sup>100</sup> and Maria M. Tomaszewski<sup>100</sup>

#### **ELSI Study**

Laura K. Barker,  $^{102}$  Maghboeba Mosavel,  $^{103}$  Laura A. Siminoff,  $^{102}$  and Heather M. Traino  $^{102}$ 

#### Genome Browser Data Integration & Visualization-EBI

Paul Flicek,<sup>104</sup> Thomas Juettemann,<sup>104</sup> Magali Ruffier,<sup>104</sup> Dan Sheppard,<sup>104</sup> Kieron Taylor,<sup>104</sup> Stephen J. Trevanion,<sup>104</sup> and Daniel R. Zerbino<sup>104</sup>

<sup>89</sup>Washington Regional Transplant Community, Falls Church, VA 22003, USA <sup>90</sup>Gift of Life Donor Program, Philadelphia, PA 19103, USA <sup>91</sup>LifeGift, Houston, TX 77055, USA

- <sup>92</sup>Center for Organ Recovery and Education, Pittsburgh, PA 15238, USA
- <sup>93</sup>LifeNet Health, Virginia Beach, VA 23453, USA
   <sup>94</sup>National Disease Research Interchange, Philadelphia, PA 19103, USA
- <sup>95</sup>Unyts, Buffalo, NY 14203, USA
- <sup>96</sup>Pharmacology and Therapeutics, Roswell Park Cancer Institute, Buffalo, NY 14263. USA
- <sup>97</sup>Van Andel Research Institute, Grand Rapids, MI 49503, USA
- <sup>98</sup>Brain Endowment Bank, Miller School of Medicine, University of Miami, Miami, FL 33136, USA
- <sup>99</sup>National Institute of Allergy and Infectious Diseases, NIH, Rockville, MD

20852, USA <sup>100</sup>Biospecimen Research Group, Clinical Research Directorate, Leidos Biomedical Research, Inc., Rockville, MD 20852, USA <sup>101</sup>Leidos Biomedical Research, Inc., Frederick, MD 21701, USA

- <sup>102</sup>Temple University, Philadelphia, PA 19122, USA
- <sup>103</sup>Department of Health Behavior and Policy, School of Medicine, Virginia Commonwealth University, Richmond, VA 23298, USA <sup>104</sup>European Molecular Biology Laboratory, European Bioinformatics Institute,

Hinxton CB10 1SD, UK

#### Genome Browser Data Integration & Visualization—UCSC Genomics Institute, University of California Santa Cruz

Brian Craft,<sup>105</sup> Mary Goldman,<sup>105</sup> Maximilian Haeussler,<sup>105</sup> W. James Kent,<sup>105</sup> Christopher M. Lee,<sup>105</sup> Benedict Paten,<sup>105</sup> Kate R. Rosenbloom,<sup>105</sup> John Vivian,<sup>105</sup> and Jingchun Zhu<sup>105</sup>

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<sup>105</sup>UCSC Genomics Institute, University of California Santa Cruz, Santa Cruz, CA 95064, USA

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## Identifying *cis*-mediators for *trans*-eQTLs across many human tissues using genomic mediation analysis

Fan Yang, Jiebiao Wang, The GTEx Consortium, et al.

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