A unified quantile framework for nonlinear heterogeneous transcriptome-wide associations

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Abstract

Transcriptome-wide association studies (TWAS) are powerful tools for identifying gene-level associations by integrating genome-wide association studies and gene expression data. However, most TWAS methods focus on linear associations between genes and traits, ignoring the complex nonlinear relationships that may be present in biological systems. To address this limitation, we propose a novel framework, QTWAS, which integrates a quantile-based gene expression model into the TWAS model, allowing for the discovery of nonlinear and heterogeneous gene-trait associations. Via comprehensive simulations and applications to both continuous and binary traits, we demonstrate that the proposed model is more powerful than conventional TWAS in identifying gene-trait associations.

Key Words: Regression quantile process; Nonlinear association test; Transcriptome-Wide Association Studies.
1 Introduction

Over the past twenty years, genome-wide association studies (GWAS) have collectively identified tens of thousands of genetic variants associated with various complex traits and diseases. However, most of these variants are located in non-coding regions of the genome, making it difficult to interpret their functional roles. Since it is assumed that most functional genetic variants exert their effects on traits through their influence on gene expression, directly linking gene expression levels to phenotypes can provide a better understanding of the underlying biological mechanisms and identify potential therapeutic targets more effectively (Tang et al., 2021; Li et al., 2021). The main challenge to such transcriptomic studies is that gene expression levels are not easily available in large-scale disease studies. To overcome this, several approaches have been developed to impute or predict gene expression levels based on DNA sequence data. One of the most widely adopted approaches is the transcriptome-wide association studies (TWAS) (Gusev et al., 2016; Gamazon et al., 2015; Zhao et al., 2021; Wainberg et al., 2019). TWAS leverages large-scale data on both genotype and gene expression across various human tissues as available in projects such as the Genotype-Tissue Expression (GTEx) project (GTEx Consortium, 2020) to learn the relationship between gene expression and genetic variation (Gusev et al., 2016; Gamazon et al., 2015; Zhao et al., 2021; Wainberg et al., 2019). By integrating this model with GWAS data, TWAS links predicted gene expression levels to traits of interest, providing insights into gene-trait associations.

As illustrated in Figure 1, TWAS combines two distinct models – a gene expression model (Model A) that models gene expression as a function of expression quantitative trait loci (eQTLs, i.e., genetic variants that are associated with gene expression), and a GWAS model (Model B ) that captures the associations between a trait and individual genetic variants.
The two models are estimated separately and then combined to infer associations between genetically regulated gene expression levels and phenotypes (Model C). For example, the widely-used PrediXcan (Gamazon et al., 2015) first uses a sparse linear model such as elastic net or lasso to estimate the cis-eQTL effects on gene expression and then imputes gene expression levels. Then, in the second step, it formally tests the association between imputed gene expression (genetically regulated gene expression levels) and the trait of interest. However access to individual-level GWAS data is often restricted due to privacy concerns and data-sharing limitations. Instead, GWAS summary statistics (estimated effect sizes and their standard errors) are more readily available. These summary statistics not only facilitate easy access to GWAS results but also allow researchers to integrate data from multiple GWAS, resulting in more powerful analyses. Along this direction, S-PrediXcan (Barbeira et al., 2018) extends PrediXcan to situations where only GWAS summary statistics are available. More recently, Gusev et al. (2016) and Nagpal et al. (2019) employed Bayesian gene expression models, while UTMOST (Hu et al., 2019) introduced a multi-task learning approach to jointly model gene expression across tissues. Given the practicality of working with summary statistics, our proposed framework also focuses on scenarios where only GWAS summary statistics are accessible.

Current TWAS methodologies rely on the assumption that both the gene expression model (Model A, SNP-gene expression association) and the GWAS model (Model B, SNP-trait association) follow linear relationships. However, multiple lines of evidence suggest substantial heterogeneity in gene expression patterns, driven by genetic variation, cellular and molecular diversity, and environmental, demographic, and technical factors (Leek and Storey, 2007; Somel et al., 2006; Budinska et al., 2013). Moreover, how eQTLs regulate gene expression can be highly context-dependent, influenced by factors such as gene-gene
interactions and gene-environment interactions (GEI) (Umans et al., 2021), leading to heterogeneous eQTL effects. Therefore, linear and mean-based gene-expression models can be inadequate for capturing the complexity of SNP-gene expression relations. Recent work by Lin et al. (2022) considered a quadratic gene expression model in TWAS, and observed improved power in identifying gene-trait association.

Our strategy is based on quantile regression (Koenker and Bassett, 1978) which models conditional quantiles across multiple quantile levels, and is an effective approach to capture heterogeneous genetic associations (Song et al., 2017; Wang et al., 2022). The flexibility of quantile regression allows it to more effectively capture the variability in gene expression patterns driven by genetic and environmental factors (Wang et al., 2024). In our proposed Quantile TWAS (QTWAS) framework, we use quantile regression along with a quantile-specific variable screening scheme to model the entire conditional distribution of gene expression given the underlying genotype profile, which helps capture the heterogeneity in eQTL-regulated gene expressions.

As we demonstrate in Appendix Section 3.1, when SNP-gene expression associations are non-linear, heteroskedastic, or exhibit heavy tails, the resulting gene-trait associations
(Model C) can become non-linear even if the GWAS associations themselves are linear. This type of non-linearity has not been adequately explored in existing TWAS literature. Allowing potentially nonlinear gene-trait associations could enhance the detection power, provide more nuanced insights into gene-trait associations and help refine targeted interventions. Our proposed QTWAS framework addresses this gap, by designing a new integration strategy that combines the conditional quantiles of genetically regulated gene expression with GWAS summary statistics to effectively capture non-linear TWAS associations (Model C in Figure 1). By doing so, we observe significantly improved power in detecting gene-trait associations.

Our approach differs from existing TWAS methods in two fundamental ways. First, we employ quantile regression with quantile-specific screening to model the full conditional distribution of gene expression. Second, our unique integration of conditional gene expression quantiles with GWAS summary statistics accommodates non-linear TWAS associations. We also establish a theoretical framework and develop rigorous statistical inference tools. Our numerical studies indicate that by relaxing the assumption of linear gene-trait associations, QTWAS outperforms the traditional linear model-based TWAS in terms of statistical power. We applied the proposed QTWAS framework to summary statistics from several GWAS datasets, including continuous traits (low-density lipoprotein) from the UK Biobank and binary traits (schizophrenia) where we confirm the increased power of QTWAS over traditional TWAS approaches. In particular, we show that the unique genes identified by QTWAS have functional enrichments among gene sets relevant for the corresponding traits/diseases, and furthermore these genes are more likely to exhibit heterogeneity in the eQTL model (Model A in Figure 1), further highlighting the advantages of the QTWAS approach.
2 Methodology

2.1 Notations and background

We denote by $X$ the gene expression level of a target gene, by $Y$ the trait/phenotype, and by $Z = (Z_1, \cdots, Z_p)^\top$ a vector of $p$ genetic variants. For now we assume that $Y$ is continuous but will relax this assumption later. SNPs $Z_1, \cdots, Z_p$ are discrete random variables with possible values of 0, 1, 2, signifying the number of reference alleles at a locus. As with other TWAS methods we focus on cis-QTLs, i.e. SNPs located within $\pm 1$Mb from the target gene, as such regions encompass most of the identified eQTLs for a gene (GTEx Consortium, 2020). Possible confounders in genetic association studies including race and ethnicity, principal components (PCs) of genotype data, and probabilistic estimation of expression residuals (PEER) factors are normally used as covariates in linear regression models to remove their effects (Stegle et al., 2012; Hu et al., 2019; Stegle et al., 2010). Without loss of generality, we assume that these confounding effects on phenotypes $Y$ and gene expression $X$ have been removed. Furthermore, we use $C$ to represent other covariates that are uncorrelated to $Z$, such as age, gender, and other postnatal non-genetic factors.

As we introduced in Section 1, TWAS aims to identify the gene ($X$) - trait ($Y$) associations by integrating two separate models – a GWAS model for SNP-trait association, and a gene expression model capturing SNP-gene expression relationships. Since we rely solely on GWAS summary statistics, we inherit the GWAS linear model from which the summary statistics are derived. Specifically, a typical GWAS study assumes the following model:

\[
Y = \alpha_0 + Z_j \beta_{GWAS,j} + C^T \eta + e, \tag{1}
\]

where $\alpha_0$ is the intercept, $\beta_{GWAS,j}$ and $\eta$ are the coefficients regarding the $j$th SNP and covariates, and $e$ is the random error. We use the GWAS summary statistics, including the
estimated $\beta_{\text{GWAS},j}$ and their standard errors derived from this model.

To fully capture the heterogeneity in genotype - gene expression relationships, we model the conditional quantile of gene expression $X$, denoted as $Q_X(\tau \mid Z, C)$, as:

$$Q_X(\tau \mid Z, C) = \alpha_0(\tau) + C^T \alpha(\tau) + Z^T \beta(\tau) \quad \text{for all } \tau \in (0, 1),$$

where $\alpha_0(\tau), \alpha(\tau), \text{ and } \beta(\tau)$ are quantile-specific intercepts and slopes for covariates and genotypes, respectively. Using the GTEx data for a specific tissue, $\{X_i, Z_i, C_i\}_{i=1}^n$ with sample size $n$, we can estimate Model (2) by solving

$$(\hat{\alpha}_0, \hat{\alpha}, \hat{\beta}) = \arg \min_{\alpha_0, \alpha, \beta} \sum_{i=1}^n \rho(\tau)(X_i - \alpha_0 - C_i \alpha - Z_i \beta),$$

where $\rho(\tau)(u) = |u|\{(1 - \tau)I(u < 0) + \tau I(u > 0)\}$ is the quantile regression check function with $u \in \mathbb{R}$, and $I(\cdot)$ is an indicator function. In the later section 2.4.2, we introduce a quantile-specific screening procedure to further improve the prediction accuracy.

## 2.2 Generalized gene-trait association model

Observed gene expression is influenced by both genetic and environmental factors. In TWAS, the focus is on the association between a trait and genetically regulated gene expression, which is denoted as $X_Z$ in this paper. By isolating the genetically regulated component, we reduce the noise introduced by environmental and other non-genetic influences. Consequently, we increase the power to detect gene-trait associations that are directly mediated by genetic variation. In a linear gene expression model, $X_Z$ can be explicitly written as $E(X_Z|Z) = \sum_{j=1}^p Z_j \beta_j$. With the assumption that $Z$ and $C$ are independent, which will be discussed in detail in Section 2.2.1, our proposed quantile gene expression model below (3)
implicitly assumes that the conditional distribution of $X_Z$ given $Z$ is $Z^T \beta(\tau)$, and $\beta(\tau)$ is unspecified. As mentioned in the Introduction, when the relationship between gene expression and eQTLs is heterogeneous, the gene-trait associations could be complex and non-linear. To account for this complexity, we propose a generalized and additive gene-trait association model:

$$Y = g_1(X_Z) + g_2(C) + \epsilon_y,$$  \hspace{1cm} (3)

where $g_1(\cdot)$ and $g_2(\cdot)$ are unknown functions. The primary function of interest is $g_1(\cdot)$ which allows for a nonlinear association between $Y$ and $X_Z$. Note that $g_1(\cdot) = 0$ corresponds to no gene-trait association between $X$ and $Y$. When both $g_1$ and $g_2$ are linear functions eq, (3) degenerates to the linear model in traditional TWAS, e.g. S-PrediXcan; see Appendix Section 1 for a review of traditional linear model based TWAS approaches.

### 2.2.1 Model Assumptions

There are several key assumptions underlying conventional TWAS. These assumptions are also assumed in the proposed QTWAS. The first assumption is that the SNP set $(Z_1, \cdots, Z_p)$ is independent of the covariates $C$. This assumption is naturally guaranteed as most covariates are postnatal and related to environmental (non-genetic) factors. It enables the separation of the genotype-contributed gene expression, denoted as $X_Z$, from the other factors, making it possible to focus on testing the association between the trait $Y$ and $X_Z$ in the gene-trait models.

The second assumption is that the SNP set $(Z_1, \cdots, Z_p)$ only affects $Y$ through $X$. This assumption implies the conditional independence between $Y$ and $X_Z$ given $Z$, which leads to valid TWAS inference (Barbeira et al., 2018; Hu et al., 2019). However, this assumption can be invalidated in practice by horizontal pleiotropy (van der Graaf et al., 2024; Barfield
et al., 2018), the scenario where a genetic variant may have an independent effect on multiple traits. Horizontal pleiotropy can lead to false positive discoveries in TWAS associations if the genetic variant has an independent effect on both the gene expression and the trait through different mechanisms. Recently, TWAS approaches that remove this assumption have been proposed (Dong et al., 2020; Zhao et al., 2024; Deng and Pan, 2021). In Section 5, we outline strategies to remove this assumption for QTWAS as well.

The last assumption is on model transferability, in the sense that the eQTL effects on gene expression are the same in both reference (GTEx) population and GWAS population. This assumption may not hold perfectly in reality due to differences in populations and environmental effects.

## 2.2.2 Inference on Gene-Trait association

For each gene-trait pair, we aim to test a global hypothesis \( H_0 : g_1(\cdot) = 0 \), indicating no gene-trait association. Although \( g_1(\cdot) \) is unknown and cannot be fully estimated without having individual-level GWAS data, we can approximate it using a piece-wise linear function over quantile regions of \( X_Z \). We can show that the local slopes can be estimated and inferred by combining the conditional quantile function of gene expression and GWAS summary statistics. Specifically, let \( A_k = \{Q_{X_Z}(\tau_k), Q_{X_Z}(\tau_{k+1})\} \) represent the interval between the \( \tau_k \)th and \( \tau_{k+1} \)th quantiles of \( X_Z \), with \( \bigcup_k A_k \) covering the full range of \( X_Z \). The approximation of \( g_1(\cdot) \) is given by

\[
g_1(X_Z) \approx \sum_{k=1}^{K} \gamma_k X_Z I\{X_Z \in A_k\}, \tag{4}
\]

where \( I(\cdot) \) is an indicator function. Then testing the gene-trait association is equivalent to testing the null hypothesis:

\[
H_0 : \gamma_k = 0 \text{ for } k = 1, ..., K; \quad H_a : \text{at least one } \gamma_k \neq 0.
\]
One can view $\gamma_k$ as a population minimizer such that $\gamma_k = \arg \min_{\gamma} \mathbb{E}_Y (Y - \gamma_k X_Z)^2 I\{X_Z \in A_k\}$. Thus, the slope coefficient $\gamma_k$ summarizes the local gene-trait association within a quantile sub-region of $X_Z$, which can be written as

$$\gamma_k = \frac{\text{cov}(X_Z, Y \mid X_Z \in A_k)}{\text{var}(X_Z \mid X_Z \in A_k)}, \quad \text{for } k = 1, \ldots, K. \tag{5}$$

The above eq. (5) implies that $A_k$ cannot be too small; otherwise, the estimation of $\text{var}(X_Z \mid X_Z \in A_k)$ will be highly unstable. Meanwhile, $A_k$ cannot be too large as that would lead to a poor estimation of the unknown function $g_1(\cdot)$ according to eq. (4). Our recommendation for choosing $A_k$ empirically is discussed in Section 2.4.

**Estimating $\gamma_k$ through model integration** $\gamma_k$ can be estimated by leveraging a conditional quantile process model of the gene expression and GWAS summary statistics. We first decompose the covariance $\text{cov}(X_Z, Y \mid X_Z \in A_k)$ in eq. (5) by the law of total variance:

$$\text{cov}(X_Z, Y \mid X_Z \in A_k) = \mathbb{E}\{\text{cov}(X_Z, Y \mid Z, X_Z \in A_k)\}$$

$$+ \text{cov}\{\mathbb{E}(X_Z \mid Z, X_Z \in A_k), \mathbb{E}(Y \mid Z, X_Z \in A_k)\}. \tag{6}$$

The second model assumption outlined in Section 2.2.1 implies the conditional independence $Y \perp \perp X_Z \mid Z$, which further implies $\mathbb{E}\{\text{cov}(X_Z, Y \mid Z, X_Z \in A_k)\} = 0$, and $\mathbb{E}(Y \mid Z, X_Z \in A_k) = Z^\top \beta_{\text{GWAS}}$, where $\beta_{\text{GWAS}}$ is the SNP-level effect size.

For any continuous random variable $X$, its quantile function denoted as $Q_X$ has the property that $Q_X(U) \overset{\text{dist.}}{=} X$, where $U$ is Uniform (0, 1) random variable, and $\overset{\text{dist.}}{=} \overset{\text{dist.}}{=}$ represents equality in distribution. Therefore, under Model (3), the distribution of gene expression $X$ can be expressed as a convolution of a genotype-related random variable $Z^\top \beta(U)$ and the remainder term, $R(U) = \alpha_0(U) + C^\top \alpha(U) + Q_\epsilon(U)$. Together with the assumption of
independence between $X$ and the covariates $C$, we conclude that

$$X_Z \overset{\text{dist.}}{=} Z^\top \beta(U),$$

i.e. the genotype-related gene expression $X_Z$ has the same distribution as $Z^\top \beta(U)$. Therefore, it is easy to derive that $\mathbb{E}(X_Z \mid Z, X_Z \in A_k) = \int_{\tau_k}^{\tau_{k+1}} Z^\top \beta(u)du = Z^\top \beta_{A_k}$, where $\beta_{A_k} := \int_{\tau_k}^{\tau_{k+1}} \beta(u)du$. Accordingly, the estimator for $\beta_{A_k}$ is given by $\hat{\beta}_{A_k} = \int_{\tau_k}^{\tau_{k+1}} \hat{\beta}(\tau)d\tau$, where $\tau_k$ and $\tau_{k+1}$ define the range of $A_k$. Together with the estimated $\hat{\beta}_{\text{GWAS}}$ from the GWAS models, we can estimate $\gamma_k$ by

$$\hat{\gamma}_k = \frac{\widehat{\text{cov}}(Z^\top \beta_{A_k}, Z^\top \beta_{\text{GWAS}})}{\text{var}(X_Z \mid X_Z \in A_k)} = \frac{\hat{\beta}_{A_k}^\top \hat{\Sigma}_Z \hat{\beta}_{\text{GWAS}}}{\hat{\sigma}_{X_Z \in A_k}^2}, \quad (7)$$

where $\hat{\sigma}_{X_Z \in A_k}^2$ is the variance of imputed gene expression in the region $A_k$ and acts as an estimate of $\sigma_{X_Z \in A_k}^2 := \text{var}(X_Z \mid X_Z \in A_k)$. Additionally, $\hat{\Sigma}_Z$ is the sample estimate of the covariance matrix of $Z$, with its true value given by

$$\Sigma_Z := \text{var}(Z) = \text{diag}\{\sigma_1, \ldots, \sigma_p\}D_Z \text{diag}\{\sigma_1, \ldots, \sigma_p\},$$

where $\sigma_j$ stands for the standard deviation of the $j$th SNP and $D_Z$ is the linkage disequilibrium (LD), which describes the correlation structure of the genotypes $Z$. The LD matrix, $D_Z$, can be estimated using data from the GTEx project or other external reference datasets. The standard deviation $\sigma_j$ for each SNP can be estimated based on its minor allele frequency.

**Constructing test statistics for the quantile-stratified gene-trait association.** The standard errors of $\hat{\gamma}_k$ can be estimated by

$$\text{se}(\hat{\gamma}_k) = \sqrt{\frac{\text{var}(\epsilon_y)}{N_{\text{GWAS}} \text{var}(X_Z \mid X_Z \in A_k)}} \approx \frac{\delta_Y}{\sqrt{N_{\text{GWAS}} \hat{\sigma}_{X_Z \in A_k}}},$$

where $\delta_Y$ is the estimated standard deviation of trait $Y$, and $N_{\text{GWAS}}$ is the sample size of GWAS data. In the formula above, we approximate $\text{var}(\epsilon_y)$ using $\delta_Y$, since the variance of
the trait explained by a single gene is typically minimal (Hu et al., 2019; O’Connor et al., 2017).

The ratio \( Z_k = \frac{\hat{\gamma}_k}{se(\hat{\gamma}_k)} \) naturally forms a Wald-type test statistic for testing \( H_0 : \gamma_k = 0 \) as below

\[
Z_k = \frac{\hat{\gamma}_k}{se(\hat{\gamma}_k)} \approx \frac{\sqrt{N_{\text{GWAS}}}}{\hat{\sigma}_Y \hat{\sigma}_{X} \in A_k} \hat{\beta}_{A_k}^T \hat{\Sigma}_Z \hat{\beta}_{\text{GWAS}},
\]

where \( \hat{\beta}_{\text{GWAS}}, j \)'s are estimated from separate marginal linear regression models (eq. (1)), and thus, they are correlated due to the linkage disequilibrium (LD) structure among the SNPs \( Z \). Denote \( \Delta = \frac{1}{\hat{\sigma}_{X} \in A_k} \hat{\beta}_{A_k}^T \hat{\Sigma}_Z \hat{\beta}_{A_k} \), then we have, under \( H_0 : \gamma_k = 0 \),

\[
\Delta^{-1/2} Z_k \approx N(0, I),
\]

because \( \frac{\sqrt{N_{\text{GWAS}}}}{\hat{\sigma}_Y \Delta^{1/2}} \hat{\beta}_{\text{GWAS}} \approx N(0, I) \) under the null hypothesis of no SNP-trait associations (Hu et al., 2019).

The \( p \)-value for testing \( \gamma_k = 0 \) in each region \( A_k \) can be computed as \( p_k = 2\Phi(-|\Delta^{-1/2} Z_k|) \), where \( \Phi(\cdot) \) is the standard normal CDF. Finally, we combine all \( p_k \)'s from \( K \) regions by the Cauchy combination method (Liu and Xie, 2020), which offers a convenient analytical solution to combine \( p \)-values from correlated tests. Alternative methods for combining \( p \)-values could also be implemented, such as Fisher’s combination method or the minimum \( p \)-value (Dudoit et al., 2003). However, these methods often rely on the assumption of independence among \( p \)-values (which is unrealistic in our setting because of correlations across quantile regions) or require computationally intensive approximations. In Figure 2, we summarize the flowchart of QTWAS. We provide further discussion on selecting \( K \) and other implementation details in Section 2.4.
2.3 From linear models to generalized linear models

Although the derivation of the QTWAS test statistics above assumes a linear GWAS model with a continuous $Y$, the same derivation applies to generalized linear models with any link function. We denote the GWAS data as $\{Y_i, Z_i, C_i\}_{i=1}^{N_{\text{GWAS}}}$, where $Z_i = (Z_{i1}, \ldots, Z_{ip})$ represents genotype data while $C_i = (C_{i1}, \ldots, C_{iq})$ represents covariates. Assume there is an arbitrary link function $h(\cdot)$, such that $\mathbb{E}(Y_i | Z_{ij}, C_i) = h(\alpha_0 + Z_{ij} \beta_{\text{GWAS,}j} + C_i \eta)$ and $\mathbb{E}(Y_i | X_{Z,i}, C_i) = h_1(X_{Z,i} + g_2(C_i))$. Common link functions include $h(x) = \exp(x)/(1+\exp(x))$ for logistic regression, $h(x) = x$ for linear regression, $h(x) = \exp(x)$ for loglinear models.
We can treat the GWAS summary statistics from generalized linear models as if they were estimated from linear regression with pseudo-response $Y_i^*$. Specifically, we can define two models

**Model B:** \[ Y_i^* = h^{-1}\{E(Y_i | Z_{ij}, C_i)\} + \epsilon_i^* = \alpha_0 + Z_{ij}\beta_{GWAS,j} + C_i\eta + \epsilon_i^* , \]

**Model C:** \[ Y_i^* = h^{-1}\{E(Y_i | X_{Z,i}, C_i)\} + \epsilon_i^* = g_1(X_{Z,i}) + g_2(C_i) + \epsilon_i^* , \]

where $E(\epsilon_i^*) = E(\epsilon_i^*) = 0$ and $var(\epsilon_i^*) \approx var(\epsilon_i^*) \approx \sigma_i^{*2}$. A reasonable estimation for $\sigma_i^{*2}$ is $\sqrt{N_{GWAS}}\hat{\sigma}_jse(\hat{\beta}_{GWAS,j})$, where $\hat{\sigma}_j$ is the estimated standard deviation of the $j$-th SNP. This estimate is based on the derivation of the standard error $se(\hat{\beta}_{GWAS,j})$. Therefore, the GWAS summary statistics $\{\hat{\beta}_{GWAS,j} , se(\hat{\beta}_{GWAS,j})\}$, which were obtained from generalized linear models using $\{Y_i, Z_{ij}, C_i\}_{i=1}^{N_{GWAS}}$, can be equivalently viewed as if they were estimated from linear models with the pseudo-response data $\{Y_i^*, Z_{ij}, C_i\}_{i=1}^{N_{GWAS}}$. As a result, the derivation of the QTWAS statistics remains valid, meaning that the GWAS summary statistics can be used in eq. (8) regardless of the specific models from which they were originally generated.

### 2.4 Model implementation

#### 2.4.1 Implementation details in the GTEx data

We trained the gene expression prediction model for 49 tissues from the GTEx project (v8), as described below. Gene expression levels were normalized and adjusted for covariates and confounders, including sex, sequencing platform, and the top five principal components of genotype data, as well as the top 15 probabilistic estimation of expression residuals (PEER) factors (Hu et al., 2019; Stegle et al., 2010). We considered protein-coding genes, removed ambiguously stranded SNPs, and only considered ref/alt pairs A/T, C/G, T/A, and G/C. SNPs with minor allele frequency less than 0.01 were excluded from the analyses. For each
gene, we used SNPs between 1Mb upstream and downstream of the transcription start site. The LD matrix $D_Z$ is estimated from the genotype data in the GTEx data.

### 2.4.2 Variant screening procedure

Based on empirical evidence, $X_Z$ often depends on a sparse set of SNPs (Barbeira et al., 2018; Gamazon et al., 2015). Most existing TWAS approaches use penalized linear regression to select significant SNPs associated with the mean of $X_Z$ (Barbeira et al., 2018; Gamazon et al., 2015; Hu et al., 2019), which may not be optimal at identifying more local (quantile-stratified) associations. We introduce a new variant screening procedure based on the quantile rank score test to identify important SNPs separately for each region $A_k$. Specifically, we aggregate multiple quantile rank score tests (Gutenbrunner et al., 1993) at selected quantile levels within $A_{X,k}$ to select region-specific SNPs while controlling the false discovery rate at the 5% level using the method of Benjamini and Hochberg (1995). The new screening procedure is more effective at identifying heterogeneous distributional associations and non-gaussian errors. We outline the detailed algorithm and a flowchart of the screening procedure in Appendix Section 1.1. Note that the variant screening step is a data pre-processing procedure for training the Genotype-GeneExpression model by GTEx data, which is independent of the subsequent steps for constructing test statistics. Thus, it does not affect the multiple testing burden at the gene-level QTWAS $p$-values.

### 2.4.3 Selection of $K$

The length of $A_k$ and the number of regions ($K$) can be set depending on applications. Though the piecewise linear approximation eq. (4) assumes $\bigcup_{k=1}^{K} A_k = (0, 1)$ for estimation, this assumption can be relaxed in the context of hypothesis testing. To test the overall associations between $X_Z$ and $Y$, integrating $\beta(\tau)$’s over a larger region $A_k$ is especially
helpful in detecting weak genetic associations. Based on our empirical experience, $K = 3$ or $4$ should be sufficient to detect homogeneous associations such as location shift, and a relatively larger $K$ (e.g., $K = 9$) facilitates detecting local associations, whereas a very large $K$ is not recommended because of the risk of power loss. As the underlying association patterns are unknown in real applications, one can further consider using partially overlapped regions and multiple choices of $K$ to improve the power. In practice, we use the Cauchy combination method to combine results from $K \in \{3, 4, 5, 9\}$ with slightly overlapped regions to obtain robust and powerful results (see the region segments in Appendix Section 1.2). This approach is data-driven, insensitive to the underlying association patterns, and avoids selecting $K$ as a tuning parameter. Furthermore, for a specified $K$, we recommend considering the partition such that $\bigcup_{k=1}^{K} A_k$ covers the 5% percentile to the 95% percentile of the value of $X_Z$. We do not recommend considering $\tau < 0.05$ and $\tau > 0.95$, as coefficients of extreme quantiles are more challenging to estimate. Note that excluding the extreme regions $\tau \in (0, 0.05)$ and $\tau \in (0.95, 1)$ may lead to loss of power if local association only manifest at these extreme tails. Investigators can also choose the regions based on specific applications.

3 Simulation studies

3.1 Simulation settings

The simulation studies are based on the data in whole blood tissue from GTEx v8 ($n = 670$). We generate gene expression based on the genotype data on 670 individuals from GTEx (see details below on the genotype-gene expression models). Then, we resample $n = 1,000$ subjects and generate their trait values based on their genotypes. For each gene, $Z$ includes all SNPs within $\pm 1$ Mb from its TSS. Gene expression $X$ is normalized before analysis as common practice in genetic association tests. The set of covariates $C$ includes the top five
principal components, top 15 PEER factors, platform, and sex. Similar to Hu et al. (2019), we randomly select 500 genes and generate the gene expression data and traits independently for each gene, as described below.

To evaluate Type I error, we generate the gene expression \(X\) from the model: 
\[ X = Z^\top \beta + C^\top \alpha_x + \epsilon_x, \]
in which \(\beta\) is estimated based on true GTEx data via the elastic net with the tuning parameter set as 0.5. The trait \(Y\) is generated by \(Y = C^\top \eta + e\). Both error terms \(e\) and \(\epsilon_x\) follow a standard normal distribution; \(\alpha_x\) and \(\eta\) are vectors with each element randomly drawn from \(\text{Unif}(0, 1)\). This null model preserves the associations between gene expression and SNPs from GTEx data but assumes no gene-trait association. A similar setting has been simulated in Hu et al. (2019).

For power analyses, we consider three different Genotype-GeneExpression models, and we assume a simple linear Genotype-Trait model to mimic the setting of GWAS summary statistics from linear models.

**Genotype-GeneExpression models.** We consider the following three models: 

- **(a) Location shift:** 
  \[ X = Z^\top \beta + C^\top \alpha_x + \epsilon_x; \]
- **(b) Location-scale:** 
  \[ X = Z^\top \beta + C^\top \alpha_x + (1 + 0.5Z^\top \beta)\epsilon_x; \]
- **(c) Local signal:** 
  \[ Q_X(\tau > 0.7 \mid Z, C) = 5\tau^{-0.7}Z^\top \beta + C^\top \alpha_x + F^{-1}_{\epsilon_x}(\tau). \]

In the location shift model (a), genetic variants \(Z\) only affect the mean of \(X\), while in the location-scale model (b), genetic variants \(Z\) affect both the mean and variance of \(X\). In the local signal model (c), variants \(Z\) only affect part of the distribution of \(X\), i.e., \(Z\) only affects the upper quantile (\(> 0.7\)th quantile) of \(X\). In each of the three scenarios, we consider two error distributions for \(\epsilon_x\): standard normal and Cauchy distributions, where the Cauchy distribution is commonly considered as a challenging case of heavy-tailed distribution in association studies (Song et al., 2017; Wang et al., 2022). Under models (b) and (c), when
the quantile specific coefficients are different across quantiles, the Genotype-GeneExpression
association is heterogeneous, and the transcriptome-wide association is nonlinear (Appendix
Section 2.1).

**Genotype-Trait model.** We consider a simple linear model
\[ Y = Z^\top \beta_{\text{GWAS}} + C^\top \eta + \epsilon, \]
where \( \epsilon \) follows a standard normal distribution.

To illustrate the performance in different scenarios, we randomly select 1% of SNPs from
the 2Mb region around TSS to be causal (i.e., with non-zero effect sizes \( \beta \) and \( \beta_{\text{GWAS}} \)). We
set \( \beta_{\text{GWAS}} = 1_p \) and \( \beta = 2 \cdot 1_p \) for local signal model, \( \beta_{\text{GWAS}} = 0.2 \cdot 1_p \) and \( \beta = 0.4 \cdot 1_p \) for
location-scale model, and \( \beta_{\text{GWAS}} = 0.1 \cdot 1_p \) and \( \beta = 0.2 \cdot 1_p \) for location shift model, where
\( 1_p \) represents a column vector with all elements being 1. \( \alpha_x \) and \( \eta \) are vectors with each
element randomly drawn from Unif(0,1).

For power analyses, we repeat the data generation procedure two times per gene and
report the statistical power based on 1,000 replicates at the significance threshold \( \alpha = 2.5e^{-6} \)
(corresponding to the usual Bonferroni threshold when testing 20,000 protein-coding genes).
For type I error analysis, we repeat the procedure for each gene 20,000 times and report
the results based on \( 10^7 \) replicates at different significance thresholds ranging from 0.05 to
\( 2.5e^{-6} \). In addition, we compare the proposed framework with S-PrediXcan (note that
we have re-implemented S-PrediXcan as it needs to be trained based on simulated data,
and we denote it as “S-PrediXcan*”). For our method, we report results based on a single
region partition (\( K = 3/4/5/9 \)) and the unified results combining all partitions. The detailed
partitions are described in Appendix Section 1.2. For one gene, we generate random \( p \)-values
\( p \sim \text{Unif}(0,1) \) if the elastic net model in S-PrediXcan* does not select any variables, or if
none of the four regions \( A_1-A_4 \) in QTWAS has valid \( p \)-value (e.g., no variant is selected).
3.2 Simulation results

The type I error for QTWAS, either with a single choice of $K$ or a unified result based on four choices of $K$, is controlled at all significance levels (Table 1). Regarding power performance, QTWAS, combining different quantile intervals, has improved power in most scenarios compared to S-PrediXcan* (Figure 3). When the error is Gaussian, both methods have comparable power for the location shift and location-scale models. However, QTWAS has substantially improved power over S-PrediXcan* when the association is local and only at upper quantiles. When the error follows the Cauchy distribution, QTWAS performs well compared to S-PrediXcan. Additionally, we observe that QTWAS is not very sensitive to the choice of $K$, and the unified approach performs best.

| $\alpha$ | S-PrediXcan* | Unified | $K = 3$ | $K = 4$ | $K = 5$ | $K = 9$ |
|----------|--------------|---------|---------|---------|---------|---------|
| 0.05     | 5.024E-02    | 5.154E-02 | 4.993E-02 | 4.910E-02 | 4.924E-02 | 4.937E-02 |
| 1e-2     | 1.075E-02    | 9.784E-03 | 9.908E-03 | 1.017E-02 | 9.841E-03 | 9.843E-03 |
| 1e-3     | 1.248E-03    | 6.536E-04 | 9.267E-04 | 8.847E-04 | 1.004E-03 | 8.584E-04 |
| 1e-4     | 9.770E-05    | 3.900E-05 | 2.380E-05 | 6.140E-05 | 6.490E-05 | 5.910E-05 |
| 1e-5     | 4.600E-06    | 1.800E-06 | 1.700E-06 | 1.200E-06 | 1.300E-06 | 1.500E-06 |
| 2.5e-6   | 1.400E-06    | 1.200E-06 | 2.000E-07 | 3.000E-07 | 3.000E-07 | 1.200E-06 |

Table 1: Type I error results for S-PrediXcan* and QTWAS ($n_{GTEx} = 670$), as well as for quantile region stratified QTWAS based on $10^7$ replicates. “Unified” combines the $p$-value of $K = 3/4/5/9$ via the Cauchy combination method.

We further illustrate the power improvement of QTWAS using the partition $K = 4$ as an example. We provide the figure of region-specific power of the QTWAS test statistics under three models in Appendix Section 2.1. The power of QTWAS in each region reveals the true underlying signal. For example, the location shift model with normal errors has equally high power in each region, and the location-scale model with normal errors shows increasing power from lower quantile to upper quantile, corresponding to the assumptions of our model. For
local signal models, we observed a dominant power boost for QTWAS, owing to the power of QTWAS test statistics in the upper quantile region ($A_4$), corresponding to the true signals being simulated at upper quantiles (i.e., $\tau > 0.7$). Therefore, the region-specific quantile test statistics can reveal more complex and detailed association patterns.

Under alternative models, we assess the robustness of the QTWAS approach. That is, we report how sensitive the QTWAS is to the choice of $K$. Among the significant results of QTWAS in Figure 3, we report the proportion identified by at least two partitions with the significance threshold $2.5e^{-6}$ (see Appendix Section 2.2). We observe that most of the QTWAS discoveries are identified by at least two partitions for all models, with the proportion slightly decreasing for the models with a higher level of heterogeneity. Overall, these results suggest that QTWAS results are robust.
3.3 Model evaluation

3.3.1 Imputation accuracy

To evaluate the accuracy of the gene expression imputation model (2), we consider the goodness of fit criterion $R^Q(\tau) = 1 - \hat{V}(\tau)/\tilde{V}(\tau)$ (Koenker and Machado, 1999), a measure of explained deviance by the quantile model associated to genetic effects at a fixed quantile level, where $\hat{V}(\tau) = \min \sum_i^n \rho_\tau(X_i - C_i^T \alpha_\tau - Z_i^T \beta_{\tau} - \alpha_{0, \tau})$ and $\tilde{V}(\tau) = \min \sum_i^n \rho_\tau(X_i - C_i^T \alpha_\tau - \alpha_{0, \tau})$ are optimized quantile loss under the alternative and null model, respectively. It is a natural analog to $R^2$ in linear models. We use $K = 4$ as an example and consider the largest $R^Q(\tau)$ over the four intervals as the explained deviance by QTWAS, which coincides with the fact that the Cauchy combination is practically driven by the smallest $p$-value in the combination.

To compare the imputation accuracy for QTWAS and S-PrediXcan*, we plotted $R^Q$ against $R^2$ (Figure 4). Except for the location shift model with normal error, QTWAS generally explained more deviance than S-PrediXcan*. Specifically, in the location-scale and local signal models, S-PrediXcan* explains a low proportion of the total deviance, indicating a relatively poor goodness of fit compared to quantile models.

3.3.2 Evaluation of variant screening procedure

To evaluate the quantile variant screening, we measure the canonical correlation between selected sets and the causal set in the three alternative models with normal or Cauchy errors. For ease of presentation, we again consider $K = 4$ as an example and use “QTWAS$_{Ak}$” to denote the results for region $k = 1, 2, 3, 4$. The proportion of replicates with a correlation greater than 0.95 is reported in Table 2 based on 1,000 replicates. For the location shift model, both QTWAS and S-PrediXcan* select SNP sets highly correlated with the true causal set. In the location-scale model, the proportion of replicates selecting highly correlated
SNPs is increasing with the quantile levels and is comparable to S-PrediXcan* for interval $A_4$, consistent with the power results. In the local shift model, QTWAS selected a set of SNPs with high correlation with the true causal set in the upper quantile interval $A_4$ more often than S-PrediXcan*, as expected. The ability of QTWAS to select variants that are more correlated to the underlying causal variants in heterogeneous cases is likely due to the specific quantile-oriented screening procedure we use here.

| Model               | QTWAS$_{A_1}$ | QTWAS$_{A_2}$ | QTWAS$_{A_3}$ | QTWAS$_{A_4}$ | S-PrediXcan* |
|---------------------|---------------|---------------|---------------|---------------|--------------|
| location shift      | 97.9%         | 99.1%         | 99.1%         | 99.0%         | 99.8%        |
| location-scale      | 31.3%         | 54.2%         | 70.4%         | 83.7%         | 85.4%        |
| local shift         | 4.5%          | 0.5%          | 0.5%          | 39.1%         | 15.4%        |

Table 2: The proportion of replicates (out of 1,000) with canonical correlation values between the selected variable set and the true causal set greater than 0.95.

### 3.4 Comparisons to sMiST

We also compare the performance of S-PrediXcan and QTWAS to sMiST (Dong et al., 2020), a method based on mixed effect models to test the total effect of genetic variants, including
their direct effects and indirect effects through gene expression. Though sMiST can also be performed based on summary statistics, the goal of sMiST is to test not only the effect of imputed gene expression (e.g., similar to TWAS) but also the direct effect of genetic variants, which violates the second assumption outlined in Section 2.2.1 for classic TWAS. By adding the genotype information into the Genotype-Trait model with random effects, their model can be written in our notations as below:

$$g\{E(Y \mid X, Z, C)\} = C^\top \eta + X\gamma_x + Z^\top \delta,$$

where $\delta_j$ for $j = 1, \cdots, p$ are random effects with mean zero and variance $\sigma^2_\delta$. The null hypothesis that sMist tests is $H_0 : \gamma_x = 0$ and $\sigma^2_\delta = 0$. Though S-PrediXcan and sMiST are both based on GWAS summary statistics, their performance are not directly comparable because sMiST needs S-PrediXcan estimated coefficients as an input, and it tests both fixed effects and random effects. Thus, it is more powerful than S-PrediXcan when the $\sigma^2_\delta \neq 0$. Nevertheless, we applied sMiST to the three models we considered in our simulations. The R package is obtained from the author’s website (https://research.fredhutch.org/hsu/en/software.html).

We report results based on three $p$-value combination procedures offered by their R package. From Table 3, we can see that sMiST is more powerful than S-PrediXcan in the local model, as expected. Compared to QTWAS, sMiST is comparable in the location and location-scale models but less powerful in the local model, which suggests that it is promising to develop a similar mixed effects model under the quantile framework to further improve the power of the association test.

### 3.5 Additional simulations

The previously presented local signal model is in favor of quantile regression, as the association only appears at upper quantiles. In the Appendix (Section 3.4), we consider
Table 3: Comparisons with sMiST. Power results based on 1000 Monte Carlo replicates.

| Setting          | S-PrediXcan* | QTWAS | sMiST     |
|------------------|--------------|-------|-----------|
|                  |              |       | p.oMiST   | p.aMiST   | p.fMiST   |
| location shift   | 0.999        | 0.996 | 0.997     | 0.996     | 0.997     |
| location-scale   | 0.901        | 0.905 | 0.911     | 0.909     | 0.911     |
| local shift      | 0.132        | 0.440 | 0.244     | 0.237     | 0.236     |

Two additional models: \( Q_X(\tau \mid Z, C) = Z^\top \beta(\tau) + C^\top \alpha_x + F^{-1}_\epsilon(\tau) \) with \( \beta(\tau) = \sqrt{\tau} \) and \( \beta(\tau) = \sin(2\pi\tau) \), respectively. Different from the previous local model, in which \( \beta(\tau) \neq 0 \) only for \( \tau \in (0.7, 1) \), \( \beta(\tau) \) here changes smoothly for \( \tau \in (0, 1) \). For these two heterogeneous models, QTWAS also outperformed S-PrediXcan*. Specifically, QTWAS is powerful for the model with \( \beta(\tau) = \sin(2\pi\tau) \) while S-PrediXcan* has almost no power because there is no association when \( \tau = 0.5 \). We further consider the location model with unobserved gene-environment interaction for the Genotype-GeneExpression model (Appendix Section 3.3). Results suggest that when there exists gene-environment interaction, but the environmental factor is unobserved, QTWAS is equivalently powerful or more powerful than S-PrediXcan in detecting gene-trait associations.

We further conduct simulations based on other tissues from GTEx data with smaller sample sizes. In particular, we considered the breast mammary tissue (\( n = 396 \)) and the brain cerebellum tissue (\( n = 209 \)). The Genotype-GeneExpression model and the Genotype-Trait model remain the same as in Section 3.1. With a smaller sample size of the genotype-gene expression data, we still observe significant power improvement of QTWAS over S-PrediXcan*. The detailed results are shown in Appendix Section 3.5.

Additionally, we consider another data-generating mechanism under the same Genotype-GeneExpression model and the Genotype-Trait model, but with cis-heritability controlled at 10% and 30% levels, respectively. At both levels, QTWAS exhibits higher power than
S-PrediXcan. Results are shown in Appendix Section 3.6. At the cis-heritability levels 10% and 30%, we also consider the scenario with horizontal pleiotropy. That is, we add a direct genetic effect in the Genotype-Trait model in addition to the classic homogeneous model (Appendix Section 3.7). The Genotype-GeneExpression model is the same as before. Results suggest that QTWAS is more powerful than S-PrediXcan at 10% and 30% cis-heritability levels. However, we acknowledge that with horizontal pleiotropy, both S-PrediXcan and QTWAS cannot infer causality and only detect associations (see Appendix Figure 8); a detailed discussion of robust methods that detect causality against horizontal pleiotropy is provided in Section 5.

4 Applications

We apply S-PrediXcan and QTWAS to publicly available GWAS summary statistics from UK Biobank (UKBB) from Pan-UKB team (2020). Specifically, we focus on two traits, one continuous (low-density lipoprotein (LDL)) and a binary trait (schizophrenia (SCZ)) (Pardiñas et al., 2018). For LDL (“LDL direct, adjusted by medication”), \( N_{GWAS} = 398,414 \) and we leverage data on whole-blood tissue from GTEx with \( n = 670 \). For SCZ, we leverage summary statistics on \( N_{GWAS} = 35,802 \) individuals and gene expression data on 13 brain tissues from GTEx. For QTWAS, we consider different quantile partitions with \( K \in \{3, 4, 5, 9\} \) and use the Cauchy combination to combine all \( p \)-values. Note that we only keep genes with \( R^2 > 0.1 \) for QTWAS, and, similarly, we consider \( R^2 > 0.1 \) for S-PrediXcan results. We focus on protein-coding genes and further restrict to the set of 6,560 genes with valid pre-trained S-PrediXcan models available from the PredictDB website (PredictDB Team, 2021). Further, we use genomic control (Devlin and Roeder, 1999) to adjust for possible inflation induced by polygenic effects, although future model developments based on mixed effect
models will be implemented in the QTWAS framework (see Section 5). The significance threshold we used is $2.5e^{-6}$.

### 4.1 Results for LDL

QTWAS identified 136 genes while S-PrediXcan identified 39 genes, with 29 genes identified by both methods (Figure 5).

![Manhattan plot for LDL with significance threshold $2.5e^{-6}$](image)

Figure 5: Manhattan plot for LDL with significance threshold $2.5e^{-6}$.

Further, we explore the reproducibility of our findings using another GWAS of lipid levels Graham et al. (2021), which contains 1.65 million individuals of mixed ancestries (although the European ancestry is dominant at 79.8%). Among the 107 genes that were uniquely identified by QTWAS in UKBB, 73 genes were also significantly identified in the new study at significance level $2.5e^{-6}$. In contrast, for the 10 genes that were uniquely identified by S-PrediXcan, none of them were successfully reproduced in the second study. These results suggest that QTWAS may be more powerful.

### 4.2 Assessing nonlinearity and functional enrichment analysis

As the two-step TWAS framework combines both eQTL and GWAS information from two datasets, the genes only identified by QTWAS but not by S-PrediXcan are highly likely to be
detected due to nonlinear associations. To further assess the heterogeneity and nonlinearity of gene expression for the genes uniquely identified by QTWAS, we first obtain their residuals from an elastic net model that fits gene expression to genotypes and covariates. We then fit another elastic net model with the squared residuals against the genotypes. The explained deviations in this model, similar to $R^2$ in linear models, measure the correlation between the squared residuals and genotypes. A higher value indicates stronger heteroskedasticity, as the squared residuals represent the variance in gene expression. We compare the nonlinearity between the group of genes identified by both QTWAS and S-PrediXcan and those identified only by QTWAS. Our results show that the genes identified exclusively by QTWAS exhibit a higher degree of nonlinearity (Table 4).

We further performed enrichment analysis to explore the function of the 107 genes uniquely identified by QTWAS but not by S-PrediXcan. We used the ToppGene database (https://toppgene.cchmc.org/enrichment.jsp) and focused on traits enriched in this group of genes. As presented in Table 5, the most enriched trait is LDL measurement, which is our primary response trait. Following LDL measurement, the traits with the highest number of gene hits are total cholesterol, apolipoprotein B, and triglycerides, all of which are closely associated with LDL levels. The fifth trait, cholesteryl ester, is a form of cholesterol in which a fatty acid is attached to the cholesterol molecule. In total, 61 out of the 107 genes are enriched across these five traits.

| variance of gene expression explained by genotypes | number of genes identified by QTWAS | only identified by QTWAS | identified by both QTWAS and S-PrediXcan |
|--------------------------------------------------|------------------------------------|-------------------------|----------------------------------------|
| less than 5%                                      | 96                                 | 78%                     | 22%                                    |
| between 5% and 10%                                | 24                                 | 71%                     | 29%                                    |
| greater than 10%                                  | 16                                 | 94%                     | 6%                                     |

Table 4: Analysis of heterogeneity and nonlinearity of genes identified by QTWAS using GTEx data.
| Trait                                      | q-value (Bonferroni) | Number of Gene Hits |
|-------------------------------------------|---------------------|---------------------|
| low density lipoprotein cholesterol       | 5.932e-32           | 52                  |
| measurement                               |                     |                     |
| total cholesterol measurement             | 1.709e-24           | 43                  |
| apolipoprotein B measurement              | 2.034e-19           | 28                  |
| triglycerides measurement                 | 5.628e-14           | 36                  |
| cholesteryl ester measurement             | 1.758e-12           | 16                  |

Table 5: Genes uniquely identified by QTWAS are enriched in LDL-related traits in ToppGene.

### 4.3 Application to schizophrenia

We applied both QTWAS and S-PrediXcan to summary statistics from a GWAS on schizophrenia Pardiñas et al. (2018), which included 11,260 cases and 24,542 controls. We use 13 brain tissues from GTEx data, with sample sizes ranging from 114 to 205. The p-values from the 13 brain tissues are combined by the Cauchy method for the final p-value. QTWAS identified 76 genes, and S-PrediXcan identified 33 genes, with 18 genes overlapping between the two methods (Figure 6). We performed a similar enrichment analysis as for LDL, identifying 17 out of 58 genes as enriched in the top five traits, including autism and schizophrenia and complement C4, which is genetically and neurobiologically related to schizophrenia (Table 6). We also include the analysis for the other 7 binary traits in Appendix Section 4.

![Manhattan plot for SCZ with significance threshold 2.5e − 6.](image)

Figure 6: Manhattan plot for SCZ with significance threshold $2.5 \times 10^{-6}$. 
| Trait                                           | q-value (Bonferroni) | Number of Gene Hits |
|------------------------------------------------|----------------------|---------------------|
| autism spectrum disorder, schizophrenia        | 9.507e-10            | 12                  |
| complement C4 measurement                     | 8.927e-09            | 7                   |
| Takayasu arteritis                            | 4.392e-07            | 8                   |
| ubiquitin carboxyl-terminal hydrolase 25 measurement | 6.755e-07            | 5                   |
| Epstein Barr virus nuclear antigen 1 IgG measurement | 1.938e-06            | 4                   |

Table 6: Enrichment analysis of the top 5 traits associated with genes uniquely identified by QTWAS for schizophrenia.

5 Discussion

We have proposed a novel quantile-based TWAS approach that utilizes a conditional quantile process to model heterogeneous gene expressions and combines it with GWAS summary statistics to infer nonlinear gene-trait associations. Our framework is particularly useful in scenarios where gene expression patterns and underlying genetic regulation exhibit significant heterogeneity, allowing for more flexible and precise modeling of gene-trait associations. As demonstrated in both simulations and applications, our quantile-based TWAS is able to identify more gene-trait associations than traditional methods and provide deeper insights regarding how gene expression levels regulate phenotypes, especially when their relationships vary across different quantiles of gene expression. These findings underscore the potential of our approach to enhance the discovery of genetic contributions to complex traits and diseases.

Through validation analyses, we show that the novel genes identified by QTWAS are likely to be functional and relevant to the trait under study. There are, however, several sources of confounding that lead to false positive associations in TWAS analyses. First, LD confounding and co-regulation (Wainberg et al., 2019) can lead to false positive associations and fine-mapping methods can further prioritize relevant genes at each locus (Ma et al., 2023, 2021; Mancuso et al., 2019). Second, QTWAS, as described here, focuses on esti-
mating the association between genetically predicted gene expression and traits, but there are no guarantees of causal inference. Colocalization methods can identify genetic variants that are causal for two phenotypes, including gene expression and a trait of interest (Zuber et al., 2022). Examples include Bayesian approaches and likelihood-based methods (Giambartolomei et al., 2014; Plagnol et al., 2009). Similarly, Mendelian Randomization (MR) methods have been proposed to test for causal relationships between gene expression and traits, by treating eQTLs as the instrumental valuables (Shi et al., 2020). A thorough review of the connections between TWAS and MR can be found in Zhu and Zhou (2021). Future research will be needed to adapt MR to quantile association tests. Furthermore, violations of assumptions, such as no horizontal pleiotropy, can also lead to false positives for both QTWAS and MR-based approaches. Increasing efforts have been made to adjust TWAS for horizontal pleiotropy (Deng and Pan, 2021; Zhao et al., 2024; Gleason et al., 2021). We provide additional details on how horizontal pleiotropy could be included in QTWAS in future work in Appendix Section 6.

In addition, several emerging topics are worth exploring for future work. It is desirable to develop approaches that combine data on multiple tissues to increase the total sample size of eQTL studies for the estimation of the conditional distribution of gene expression. Such approaches have been developed before, e.g., UTMOST (Hu et al., 2019), and have been shown to effectively increase imputation accuracy and power. Multi-tissue quantile modeling may allow investigations of more comprehensive nonlinear associations across tissues. Furthermore, the current QTWAS framework can be better developed when individual GWAS data are available, which would allow nonparametric approaches to estimate higher-resolution nonlinear gene-trait associations and multiplicative errors instead of additive errors, depending on the original transformation applied to the trait.
Software

QTWAS has been implemented in publicly available software. We have posted a comprehensive demonstration on Github: https://github.com/tianyingw/QTWAS, which contains the tutorial for downloading and using the pre-trained QTWAS models to conduct analysis.

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