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STATISTICAL INFERENCE FOR GENETIC RELATEDNESS BASED ON HIGH-DIMENSIONAL LOGISTIC REGRESSION

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Abstract: We examine statistical inference for genetic relatedness between binary traits, based on individual-level genome-wide association data. Specifically, for high-dimensional logistic regression models, we define parameters characterizing the cross-trait genetic correlation, genetic covariance, and trait-specific genetic variance. We develop a novel weighted debiasing method for the logistic Lasso estimator and propose computationally efficient debiased estimators. Furthermore, we study the rates of convergence for these estimators and establish their
asymptotic normality under mild conditions. Moreover, we construct confidence intervals and statistical tests for these parameters, and provide theoretical justifications for the methods, including the coverage probability and expected length of the confidence intervals, and the size and power of the proposed tests. Numerical studies under both model-generated data and simulated genetic data show the superiority of the proposed methods. By analyzing a real data set on autoimmune diseases, we demonstrate their ability to obtain novel insights about the shared genetic architecture between 10 pediatric autoimmune diseases.

*Key words and phrases:* Confidence interval; debiasing methods; functional estimation; genetic correlation; hypothesis testing.

1. Introduction

Genome-wide association studies (GWAS) have identified thousands of genetic variants or single nucleotide polymorphisms (SNPs) associated with various complex phenotypes. Among them, many variants were found to be associated with multiple complex traits, reflecting the pleiotropic action of genes or the correlation between causal loci in two traits. Understanding the shared genetic architecture among different traits can potentially lead to further insights into the biological etiology of diseases and inform therapeutic interventions (Van Rheenen et al., 2019).

Various definitions of genetic relatedness or correlation have been pro-
posed in different contexts to characterize quantitatively the shared genetic associations between complex traits, based on GWAS data. Understanding the genetic relatedness between complex traits helps to identify new trait-associated variants (Turley et al., 2018), improve genetic risk prediction (Maier et al., 2015), and assist inference on causality (O’Connor and Price, 2018). Compared with methods of traditional approaches from family studies, where measurements of both traits are required for the same individuals, those based on GWAS enjoy the advantages of increased sample sizes and a reduced risk of confounding or ascertainment biases, and thus have greater potential for large-scale analyses involving multiple traits (Zhang et al., 2020).

Bivariate linear mixed-effects models have been widely applied to estimate the genetic covariance and genetic correlation between two traits from individual-level GWAS data (Lee et al., 2011, 2012; Vattikuti et al., 2012; Lee et al., 2013). These models decompose the phenotypic variance into genetic and residual variance components, and define the genetic correlation as that between the two trait-specific random generic effects. However, the mixed-effect model approach requires knowledge about the genetic relationship matrix, which is commonly approximated by the genetic relationship across the set of all genotyped variants (Yang et al., 2010). Computationally
efficient methods have been developed based on the cross-trait linkage disequilibrium (LD) score regression (Bulik-Sullivan et al., 2015; Ning et al., 2020) to estimate a genetic correlation using GWAS summary statistics over a large set of SNPs. This approach relies on the classical asymptotics, which do not consider the high dimensionality of the SNPs relative to the sample sizes, resulting in possibly inaccurate inference results (Zhao and Zhu, 2019a). Other approaches, such as those of Shi et al. (2017), Lu et al. (2017), and Guo et al. (2021), explore differences in local genetic correlations between traits using genome partitioning based on genomic annotations. Weissbrod et al. (2018) notes that many existing methods are geared primarily toward quantitative traits. Thus, applying them directly to data sets with binary outcomes may suffer from reduced statistical power. They propose a mixed-effects model for estimating the genetic correlation between binary traits.

In this study, we take a high-dimensional regression approach, with fixed genetic effects to identify trait-associated genetic variants and quantify the genetic relatedness between two traits. An important advantage of a multiple regression over the simple univariate regression is its potential to identify more trait-associated variants (Wu et al., 2009). Existing studies on heritability or co-heritability in a high-dimensional regression framework in-
Inference on Genetic Relatedness

Inclusion, for example, those of Bonnet et al. (2015), Janson et al. (2017), Verzelten and Gassiat (2018), Guo et al. (2019), Zhao and Zhu (2019a), Zhao and Zhu (2019b), and Guo et al. (2021). Under the linear regression model, Guo et al. (2019) propose bias-corrected estimators for the genetic covariance and correlation parameters, based on individual-level GWAS data, and Zhao and Zhu (2019a) propose consistent estimators for a polygenic risk score and a genetic correlation, based on GWAS summary statistics. However, these works focus on the genetic relatedness between continuous traits, and do not provide inference procedures such as statistical tests.

We address the following two questions concerning binary traits. How can we define and study the genetic relatedness between two binary traits in a high-dimensional regression framework? How can we perform a valid statistical inference, such as testing hypotheses or constructing confidence intervals (CIs) for the genetic-relatedness parameters? We address these questions in a principled way with rigorous statistical justifications.

To that end, for a pair of binary traits \((y, w) \in \{0, 1\}^2\), we consider the following high-dimensional logistic regression models:

\[
y \mid X \sim \text{Bernoulli}(\pi_y(X)), \quad \log \left\{ \frac{\pi_y(X)}{1 - \pi_y(X)} \right\} = \alpha + X^\top \beta, \quad (1.1)
\]

\[
w \mid X \sim \text{Bernoulli}(\pi_w(X)), \quad \log \left\{ \frac{\pi_w(X)}{1 - \pi_w(X)} \right\} = \zeta + X^\top \gamma, \quad (1.2)
\]
where $\pi_y(X) = P(y = 1|X)$, $\pi_w(X) = P(w = 1|X)$, $X \in \mathbb{R}^p$ is a random vector of $p$ genetic variants with population covariance matrix $\Sigma \in \mathbb{R}^{p \times p}$, $\beta, \gamma \in \mathbb{R}^p$ are the corresponding trait-specific regression coefficients, which are assumed to be sparse vectors, and $\alpha, \zeta \in \mathbb{R}$ are the trait-specific intercepts. The genetic covariance between two traits is defined as the covariance between the log-odds ratios associated with the two traits, that is, 

$$\text{genetic covariance}(y, w) = \text{Cov}\left(\log\left\{\frac{\pi_y(X)}{1-\pi_y(X)}\right\}, \log\left\{\frac{\pi_w(X)}{1-\pi_w(X)}\right\}\right),$$

which, by definition, admits the following expressions:

$$\text{Cov}\left(\log\left\{\frac{\pi_y(X)}{1-\pi_y(X)}\right\}, \log\left\{\frac{\pi_w(X)}{1-\pi_w(X)}\right\}\right) = \text{Cov}(X^\top \beta, X^\top \gamma) = \beta^\top \Sigma \gamma.$$ 

Similarly, we define the genetic variance of the binary trait $y$ as the variance of its associated log-odds ratio, that is, 

$$\text{genetic variance}(y) = \text{Var}\left(\log\left\{\frac{\pi_y(X)}{1-\pi_y(X)}\right\}\right),$$

which satisfies 

$$\text{Var}(X^\top \beta) = \beta^\top \Sigma \beta.$$ 

We define the genetic variance of the trait $w$ as 

$$\text{Var}\left(\log\left\{\frac{\pi_w(X)}{1-\pi_w(X)}\right\}\right) = \text{Var}(X^\top \gamma) = \gamma^\top \Sigma \gamma.$$ 

Whenever the genetic variances of $y$ and $w$ are both nonzero, we can define the genetic correlation $R(y, w)$ between the two traits as the correlation between the associated log-odds ratios, that is, 

$$\text{Corr}\left(\log\left\{\frac{\pi_y(X)}{1-\pi_y(X)}\right\}, \log\left\{\frac{\pi_w(X)}{1-\pi_w(X)}\right\}\right) = \frac{\beta^\top \Sigma \gamma}{\sqrt{\beta^\top \Sigma \beta \cdot \gamma^\top \Sigma \gamma}},$$

and set $R(y, w) = 0$ whenever $\beta^\top \Sigma \beta \cdot \gamma^\top \Sigma \gamma = 0$.

The concept of covariance or correlation between two log-odds ratios is both statistically and empirically meaningful. It is used by Wei and Higgins (2013) to account for correlated outcomes in meta-analysis, and by Bagos
when the data take the form of contingency tables. In our context, as parameters or functionals quantifying the conditional co-occurrence risk of two traits, the genetic covariance and correlation defined above characterize the shared effect size of the genetic variants by considering the true covariance structure of the variants.

We examine the problem of statistical inference for these genetic relatedness functionals, based on individual-level GWAS data with binary outcomes. By carefully analyzing the logistic Lasso estimator, we develop a novel weighted debiasing method and propose computationally efficient debiased estimators for these functionals. We further study their rates of convergence and obtain their asymptotic normality under mild theoretical conditions. Moreover, confidence intervals and statistical tests for these functionals are constructed. We provide theoretical justifications for the methods, including the coverage probability and expected length of the CIs, and the size and power of the proposed tests. Our results provide a rigorous statistical inference framework for studying the genetic relatedness between binary traits.

Throughout, for a symmetric matrix $A \in \mathbb{R}^{p \times p}$, $\lambda_i(A)$ denotes its $i$th largest eigenvalue and $\lambda_{\max}(A) = \lambda_1(A)$ and $\lambda_{\min}(A) = \lambda_p(A)$. For a smooth function $f(x)$ defined on $\mathbb{R}$, we denote $\dot{f}(x) = df(x)/dx$ and
\[ \ddot{f}(x) = \frac{d^2 f(x)}{dx^2}. \]

For sequences \( \{a_n\} \) and \( \{b_n\} \), we write \( a_n = o(b_n) \), \( a_n \ll b_n \) or \( b_n \gg a_n \) if \( \lim_n a_n/b_n = 0 \), and write \( a_n = O(b_n) \), \( a_n \lesssim b_n \) or \( b_n \gtrsim a_n \) if there exists a constant \( C \) such that \( a_n \leq C b_n \) for all \( n \). We write \( a_n \asymp b_n \) if \( a_n \lesssim b_n \) and \( a_n \gtrsim b_n \).

2. Estimation of Genetic Relatedness

2.1 Genetic Relatedness under Various Settings of Data Availability

We consider two types of data collection scenarios commonly used to study the genetic relatedness between two traits based on individual-level GWAS data. Data sets obtained from these two scenarios are widely available in current genetic research. In the first scenario, measurements of two traits, along with the subject genotypes, are obtained from different groups of unrelated individuals. In other words, there are two independent data sets, each containing measurements of a single trait and genotypes for a group of unrelated individuals. This scenario is common in cross-trait analyses based on multiple independent GWAS data. In the second scenario, measurements of multiple traits of interest, along with the subject genotypes, may be obtained from the same group of unrelated individuals. This type of data set is also widely available by virtue of many large-scale studies, such as...
UK Biobank (Sudlow et al., 2015). The above two scenarios are formally defined as follows.

**Scenario (I): Data from independent samples.** The observations are \( \{(y_i, X_i)\}_{i=1}^{n_1} \) and \( \{(w_i, Z_i)\}_{i=1}^{n_2} \), where \( X_i \) and \( Z_i \) are drawn independently from some probability measure \( P_\theta \) on \( \mathbb{R}^p \) with covariance matrix \( \Sigma \), and \( y_i \) and \( w_i \) are generated based on (1.1) and (1.2), respectively.

**Scenario (II): Data from overlapped samples.** The observations are \( \{(y_i, X_i)\}_{i=1}^{n_1} \) and \( \{(w_i, Z_i)\}_{i=1}^{n_2} \), where \( Z_i = X_i \), for \( i \in \{1, 2, ..., m\} \), \( 1 \leq m \leq \min\{n_1, n_2\} \). The samples in \( \{Z_i\}_{i=1}^{m}, \{X_i\}_{i=m+1}^{n_1} \) and \( \{Z_i\}_{i=m+1}^{n_2} \) are drawn independently from some probability measure \( P_\theta \) on \( \mathbb{R}^p \) with covariance matrix \( \Sigma \), and \( y_i \) and \( w_i \) are generated from (1.1) and (1.2), respectively.

Note that Scenario (I) corresponds to Scenario (II) with \( m = 0 \). In what follows, we introduce our main results by focusing on Scenario (I) to avoid unnecessary complications in the notation. A discussion of Scenario II is provided in Section S5 of the Supplementary Material (Ma et al., 2021), because our methods and results in this case are very similar.
2.2 Weighted Bias Correction and the Proposed Estimators

Estimating the genetic correlation $R$ can be reduced to estimating the genetic covariance functional $\beta^T \Sigma \gamma$ and the genetic variance functionals $\beta^T \Sigma \beta$ and $\gamma^T \Sigma \gamma$. The novel bias-correction method proposed here yields nearly unbiased estimators of these functionals of interest. We construct the estimators using the following two-step procedure. In the first step, we obtain an initial plug-in estimator of the functional based on the pooled sample covariance matrix $\hat{\Sigma} = \frac{1}{n_1 + n_2} \left[ \sum_{i=1}^{n_1} X_i X_i^T + \sum_{i=1}^{n_2} Z_i Z_i^T \right]$, and the logistic Lasso estimators

$$
(\hat{\alpha}, \hat{\beta}) = \arg \min_{\alpha, \beta} \left\{ \frac{1}{n_1} \sum_{i=1}^{n_1} \left\{ -y_i (\alpha + \beta^T X_i) + \log(1 + e^{\alpha + \beta^T X_i}) \right\} + \lambda(\|\beta\|_1 + |\alpha|) \right\},
$$

$$
(\hat{\zeta}, \hat{\gamma}) = \arg \min_{\zeta, \gamma} \left\{ \frac{1}{n_2} \sum_{i=1}^{n_2} \left\{ -w_i (\zeta + \gamma^T Z_i) + \log(1 + e^{\zeta + \gamma^T Z_i}) \right\} + \lambda(\|\gamma\|_1 + |\zeta|) \right\},
$$

(2.1)

with $\lambda = C \sqrt{\log p / n}$ for some constant $C > 0$. In the second step, we obtain the final estimator by modifying the initial estimator using a carefully designed bias-correction term.

We begin with the genetic covariance functional $\beta^T \Sigma \gamma$. With the logistic Lasso estimators (2.1) and $\hat{\Sigma}$, the corresponding plug-in estimator is defined as $\hat{\beta}^T \hat{\Sigma} \hat{\gamma}$, the error of which can be decomposed as $\hat{\beta}^T \hat{\Sigma} \hat{\gamma} - \beta^T \Sigma \gamma = \hat{\gamma}^T \Sigma (\hat{\beta} - \beta) + \hat{\beta}^T \Sigma (\hat{\gamma} - \gamma) - (\hat{\beta} - \beta)^T \Sigma (\hat{\gamma} - \gamma) + \hat{\beta}^T (\hat{\Sigma} - \Sigma) \hat{\gamma}$. It turns out that the term $\hat{\beta}^T (\hat{\Sigma} - \Sigma) \hat{\gamma}$ contributes only to the variance of the
plug-in estimator, the terms $\gamma^T \Sigma(\hat{\beta} - \beta)$ and $\hat{\beta}^T \Sigma(\hat{\gamma} - \gamma)$ contribute to the leading-order bias of the plug-in estimator, and the contribution from $(\hat{\beta} - \beta)^T \Sigma(\hat{\gamma} - \gamma)$ is negligible. Therefore, the bias of the plug-in estimator can be further reduced by estimating $\gamma^T \Sigma(\hat{\beta} - \beta)$ and $\hat{\beta}^T \Sigma(\hat{\gamma} - \gamma)$ directly. To accomplish this, set $h(u) = \frac{e^u}{1+e^u}$. Then by Taylor’s expansion, $h(\hat{\alpha} + X_i^T \hat{\beta}) - h(\alpha + X_i^T \beta) = \frac{e^{\hat{\alpha} + X_i^T \hat{\beta} \cdot X_i^T (\hat{\beta} - \beta)}}{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2} + \frac{e^{\hat{\alpha} + X_i^T \hat{\beta} \cdot (\hat{\alpha} - \alpha)}}{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2} + \Delta_i$, where $\Delta_i = \hat{h}[X_i^T \{t\beta' + (1 - t)\beta''\}]\{X_i^T (\beta' - \beta'')\}^2$, for some $t \in (0, 1)$, $\beta' = (\alpha, \beta^T)^T$, $\beta'' = (\hat{\alpha}, \hat{\beta}^T)^T$, and $X_i' = (1, X_i^T)^T$. Furthermore, if we define $\epsilon_i = y_i - h(\alpha + X_i^T \beta)$, then

$$
\{h(\hat{\alpha} + X_i^T \hat{\beta}) - y_i\} X_i
= \left\{ \frac{e^{\hat{\alpha} + X_i^T \hat{\beta}}}{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2} X_i^T (\hat{\beta} - \beta) + \frac{e^{\hat{\alpha} + X_i^T \hat{\beta}}}{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2} (\hat{\alpha} - \alpha) + \Delta_i - \epsilon_i \right\} X_i
= \frac{e^{\hat{\alpha} + X_i^T \hat{\beta}}}{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2} X_i X_i^T (\hat{\beta} - \beta) + (\Delta_i - \epsilon_i) X_i + \frac{e^{\hat{\alpha} + X_i^T \hat{\beta}}}{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2} (\hat{\alpha} - \alpha) X_i.
$$

In order to construct a good estimator of $\Sigma(\hat{\beta} - \beta)$, we rescale each item $\{h(\hat{\alpha} + X_i^T \hat{\beta}) - y_i\} X_i$ by a sample-specific weight $\frac{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2}{e^{\hat{\alpha} + X_i^T \hat{\beta}}}$ so that

$$
\sum_{i=1}^{n_1} \frac{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2}{e^{\hat{\alpha} + X_i^T \hat{\beta}}} \{h(\hat{\alpha} + X_i^T \hat{\beta}) - y_i\} X_i
= \left( \sum_{i=1}^{n_1} X_i X_i^T \right) (\hat{\beta} - \beta) + \sum_{i=1}^{n_1} \frac{(1 + e^{\hat{\alpha} + X_i^T \hat{\beta}})^2}{e^{\hat{\alpha} + X_i^T \hat{\beta}}} (\Delta_i - \epsilon_i) X_i + (\hat{\alpha} - \alpha) \sum_{i=1}^{n_1} X_i.
$$

Consequently, as long as the last two terms in the above equation are negligible relative to the leading term $\left( \sum_{i=1}^{n_1} X_i X_i^T \right) (\hat{\beta} - \beta)$, we can construct
an estimator of $\hat{\gamma}^\top \Sigma (\hat{\beta} - \beta)$ as

$$
\hat{\gamma}^\top \frac{1}{n_1} \sum_{i=1}^{n_1} \frac{(1 + e^{\hat{\alpha} + X_i^\top \hat{\beta}})^2}{e^{\hat{\alpha} + X_i^\top \hat{\beta}}} \{ h(\hat{\alpha} + X_i^\top \hat{\beta}) - y_i \} X_i. \tag{2.2}
$$

Similarly, we can estimate the error term $\hat{\beta}^\top \Sigma (\hat{\gamma} - \gamma)$ using

$$
\hat{\beta}^\top \frac{1}{n_2} \sum_{i=1}^{n_2} \frac{(1 + e^{\hat{\zeta} + Z_i^\top \hat{\gamma}})^2}{e^{\hat{\zeta} + Z_i^\top \hat{\gamma}}} \{ h(\hat{\zeta} + Z_i^\top \hat{\gamma}) - w_i \} Z_i. \tag{2.3}
$$

As a result, in light of the error decomposition, a bias-corrected estimator for $\beta^\top \Sigma \gamma$ is defined as

$$
\tilde{\beta}^\top \Sigma \gamma = \hat{\beta}^\top \Sigma \hat{\gamma} - \hat{\gamma}^\top \frac{1}{n_1} \sum_{i=1}^{n_1} \frac{(1 + e^{\hat{\alpha} + X_i^\top \hat{\beta}})^2}{e^{\hat{\alpha} + X_i^\top \hat{\beta}}} \{ h(\hat{\alpha} + X_i^\top \hat{\beta}) - y_i \} X_i,
\tilde{\gamma}^\top \frac{1}{n_2} \sum_{i=1}^{n_2} \frac{(1 + e^{\hat{\zeta} + Z_i^\top \hat{\gamma}})^2}{e^{\hat{\zeta} + Z_i^\top \hat{\gamma}}} \{ h(\hat{\zeta} + Z_i^\top \hat{\gamma}) - w_i \} Z_i. \tag{2.4}
$$

The above estimator modifies the simple plug-in estimator by adding a carefully constructed bias-correction term that accounts for the leading-order bias of the plug-in estimator. The bias-correction terms (2.2) and (2.3) are weighted averages, where the weights, from the nonlinearity of the link function, reflect each sample’s contribution to the overall bias.

In the same vein of our construction of the estimator $\tilde{\beta}^\top \Sigma \gamma$, bias-corrected estimators for the genetic variances can be defined similarly as

$$
\tilde{\beta}^\top \Sigma \beta = \hat{\beta}^\top \Sigma \hat{\beta} - 2 \hat{\beta}^\top \frac{1}{n_1} \sum_{i=1}^{n_1} \frac{(1 + e^{\hat{\alpha} + X_i^\top \hat{\beta}})^2}{e^{\hat{\alpha} + X_i^\top \hat{\beta}}} \{ h(\hat{\alpha} + X_i^\top \hat{\beta}) - y_i \} X_i, \tag{2.5}
$$

$$
\tilde{\gamma}^\top \Sigma \gamma = \hat{\gamma}^\top \Sigma \hat{\gamma} - 2 \hat{\gamma}^\top \frac{1}{n_2} \sum_{i=1}^{n_2} \frac{(1 + e^{\hat{\zeta} + Z_i^\top \hat{\gamma}})^2}{e^{\hat{\zeta} + Z_i^\top \hat{\gamma}}} \{ h(\hat{\zeta} + Z_i^\top \hat{\gamma}) - w_i \} Z_i. \tag{2.6}
$$
Based on the above genetic variance and covariance estimators, a natural estimator of the genetic correlation is \( \hat{R} = \frac{\hat{\beta}^T \Sigma \gamma}{\sqrt{\hat{\beta}^T \Sigma \beta \hat{\gamma}^T \Sigma \gamma}} \). Taking into account the actual range of \( R \), we propose its final estimator as

\[
\hat{R} = \begin{cases} 
\hat{R}, & \text{if } (\hat{\beta}^T \Sigma \gamma)^2 < \hat{\beta}^T \Sigma \beta \hat{\gamma}^T \Sigma \gamma \\
0, & \text{if } \hat{\beta}^T \Sigma \beta \hat{\gamma}^T \Sigma \gamma = 0 \\
\text{sign}(\hat{R}), & \text{otherwise}
\end{cases}
\]  

\[(2.7)\]

Compared with existing methods for constructing debiased estimators in high-dimensional regression (Zhang and Zhang, 2014; Javanmard and Montanari, 2014a,b; van de Geer et al., 2014; Cai and Guo, 2017; Guo et al., 2019; Ma et al., 2020; Cai and Guo, 2020; Cai et al., 2021; Guo et al., 2021), our proposed method has two distinct advantages. First, the proposed estimators can be obtained directly from their explicit expressions, as in (2.4) to (2.7), which rely only on the initial logistic Lasso estimator, and simple plug-in procedures. Its main computational task is to solve for the initial Lasso estimator, which can be achieved efficiently using a standard tuning process (Section 5), and therefore is more scalable to the large data sets in genetic studies. In contrast, existing methods involve solving other high-dimensional optimization problems, in addition to the initial estimator, for bias correction. These additional problems are computationally challenging, time-consuming, and subject to difficult tuning processes. Second, by
using our carefully constructed weighted bias-correction method, we can avoid many commonly used, but stringent technical conditions. This significantly expands the range of applicability of our proposed methods; see also the discussions after Theorems 1 and 5.

3. CIs and Statistical Tests

As an important consequence, it can be shown that each of the above proposed estimators is asymptotically normally distributed. This can be used to construct CIs and statistical tests for the functionals.

Specifically, it can be shown that the estimator $\hat{\beta}^\top \Sigma \hat{\gamma}$ has variance

$$v^2 = \frac{n_1 + n_2}{n_1} E\{\eta_i(X) (\hat{\gamma}^\top X_i)^2\} + \frac{n_1 + n_2}{n_2} E\{\eta_i(Z) (\hat{\beta}^\top Z_i)^2\} + E\{\hat{\beta}^\top (X_i X_i^\top - \Sigma) \hat{\gamma}\}^2,$$

where $\eta_i(X) = \frac{(1+e^\hat{\alpha}+X_i^\top \hat{\beta})^4 e^\hat{\alpha}+X_i^\top \hat{\beta}}{(1+e^\hat{\alpha}+X_i^\top \hat{\beta})^2 e^2+2X_i^\top \hat{\beta}}$ and $\eta_i(Z) = \frac{(1+e^\hat{\zeta}+Z_i^\top \hat{\gamma})^4 e^\hat{\zeta}+Z_i^\top \hat{\gamma}}{(1+e^\hat{\zeta}+Z_i^\top \hat{\gamma})^2 e^2+2Z_i^\top \hat{\gamma}}$. Intuitively, the parameters $\beta$ and $\gamma$ in the above expressions can be estimated using their initial Lasso estimators. Thus, we can define a moment estimator of the asymptotic variance as

$$\hat{v}^2 = \frac{n_1 + n_2}{n_1} \sum_{i=1}^{n_1} \frac{(1+e^\hat{\alpha}+X_i^\top \hat{\beta})^2 (\hat{\gamma}^\top X_i)^2}{e^\hat{\alpha}+X_i^\top \hat{\beta}} + \frac{n_1 + n_2}{n_2} \sum_{i=1}^{n_2} \frac{(1+e^\hat{\zeta}+Z_i^\top \hat{\gamma})^2 (\hat{\beta}^\top Z_i)^2}{e^\hat{\zeta}+Z_i^\top \hat{\gamma}} + \sum_{i=1}^{n_1} (\hat{\beta} X_i X_i^\top \hat{\gamma} - \hat{\beta} \Sigma \hat{\gamma})^2 + \sum_{i=1}^{n_2} (\hat{\beta} Z_i Z_i^\top \hat{\gamma} - \hat{\beta} \Sigma \hat{\gamma})^2.$$

Hence, a $(1 - \alpha)$-level CI for the genetic covariance is $\text{CI}_\alpha(\beta^\top \Sigma \gamma, \mathcal{D}) = [\hat{\beta}^\top \Sigma \gamma - \hat{\rho}, \hat{\beta}^\top \Sigma \gamma + \hat{\rho}]$, where $\hat{\rho} = \frac{z_{\alpha/2} \hat{v}}{\sqrt{n_1 + n_2}}$, and $z_{\alpha/2} = \Phi^{-1}(1 - \alpha/2)$ is the upper $\alpha/2$-quantile of the standard normal distribution. Similarly, the asymptotic variance of the genetic variance estimator $\hat{\beta}^\top \Sigma \hat{\beta}$ can be derived as $v^2_{\hat{\beta}} = \frac{n_1 + n_2}{n_1} E\{\eta_i(X) (\hat{\gamma}^\top X_i)^2\}$.
The CI for the genetic correlation $R$ is a direct consequence of Slutsky’s theorem. Specifically, for the estimator $\hat{R}$ defined in (2.7), whenever $\hat{\beta}^\top \Sigma \hat{\beta} \neq 0$, we can estimate its asymptotic variance by $\hat{v}_R^2 = \frac{\hat{v}^2}{\beta^\top \Sigma \beta}$, and define the corresponding $(1 - \alpha)$-level CI as $\text{CI}_\alpha(\beta^\top \Sigma \beta, \mathcal{D}) = \left[ \hat{\beta}^\top \Sigma \beta - \hat{\rho}_\beta, \hat{\beta}^\top \Sigma \beta + \hat{\rho}_\beta \right]$, where $\hat{\rho}_\beta = \frac{z_{\alpha/2} \hat{v}_R}{\sqrt{n_1 + n_2}}$; $\text{CI}_\alpha(\gamma^\top \Sigma \gamma, \mathcal{D})$ can be obtained by symmetry.

Converting the above CIs, we obtain statistical tests for each of the null hypotheses, $H_{0,1} : \beta^\top \Sigma \gamma = B_0$, $H_{0,2} : \beta^\top \Sigma \beta = Q_0$, and $H_{0,3} : R = R_0$, for some $B_0 \in \mathbb{R}$, $Q_0 \geq 0$ and $R_0 \in [-1, 1]$. Specifically, we define test statistics $T_1 = \frac{\sqrt{n_1 + n_2} (\hat{\beta}^\top \Sigma \gamma - B_0)}{\hat{v}_\beta}$, $T_2 = \frac{\sqrt{n_1 + n_2} (\hat{\beta}^\top \Sigma \beta - Q_0)}{\hat{v}_\beta}$, and $T_3 = \frac{\sqrt{n_1 + n_2} (\hat{R} - R_0)}{\hat{v}_R}$, so that for each $\ell \in \{1, 2, 3\}$, to obtain an $\alpha$-level test, we reject the null hypothesis $H_{0,\ell}$ whenever $|T_\ell| > z_{\alpha/2}$.

4. Theoretical Properties

4.1 Rates of Convergence and Optimality

The random covariates are characterized by the following conditions.
(A1) For each $1 \leq i \leq n_1$ and $1 \leq j \leq n_2$, $X_i$ and $Z_j$ are centered independent and identically distributed i.i.d. sub-Gaussian random vectors, where $\Sigma = E(X_iX_i^\top) \in \mathbb{R}^{p \times p}$ satisfies $M^{-1} \leq \lambda_{\min}(\Sigma) \leq \lambda_{\max}(\Sigma) \leq M$, for some constant $M > 1$.

(A2) There exists a positive constant $c_0$ such that $E\left(\frac{\beta^\top X_iX_i^\top \gamma}{\beta^\top \Sigma \gamma} - 1\right)^2 > c_0$.

For the regression coefficients, we denote $k = \max\{\|\beta\|_0, \|\gamma\|_0\}$, $U(\beta, \gamma) = \max\{\|\beta\|_2, \|\gamma\|_2\}$, and $L(\beta, \gamma) = \min\{\|\beta\|_2, \|\gamma\|_2\}$. We assume

(A3) $\max\{|\alpha|, |\zeta|\} \leq C$ and $U(\beta, \gamma) \leq C$, for some constant $C > 0$.

Intuitively, assumptions (A1) and (A3) imply that the marginal case probabilities $P(y_i = 1)$ and $P(w_i = 1)$ are balanced, or bounded away from zero and one, whereas (A2) ensures that the asymptotic variances do not diminish.

For technical reasons, for each trait, we split the corresponding samples into halves, so that the initial Lasso estimation step and the other steps, such as the covariance estimation and bias-correction, are conducted on independent data sets. Without loss of generality, we assume under Scenario I that there are $2(n_1 + n_2)$ samples in $D$, divided into two disjoint subsets, $D_1$ and $D_2$, each containing $n_1 + n_2$ independent samples, with $n_1$ samples corresponding to trait $y_i$, and $n_2$ samples corresponding to trait $w_i$. The initial Lasso estimators are obtained from $D_1$; the sample covariance, bias-
correction terms, and asymptotic variance estimators are based on $\mathcal{D}_2$ and the initial Lasso estimators. Note that the sample-splitting procedure is only used to facilitate the theoretical analysis, and is not needed in practice. We demonstrate this point numerically in Section 5; see also Section 7.

The following theorem concerns the rate of convergence of the bias-corrected estimators $\hat{\beta}^\top \Sigma \gamma$ and $\hat{\beta}^\top \Sigma \beta$; the results for $\hat{\gamma}^\top \Sigma \gamma$ are similar.

**Theorem 1** (Rates of Convergence). Suppose (A1) and (A3) hold, $n_1 \asymp n_2 \asymp n$, and $k \lesssim \frac{n \log \log n}{\log p}$. Then, for sufficiently large $(n, p)$ and any $t > 0$,

\begin{align*}
|\hat{\beta}^\top \Sigma \gamma - \beta^\top \Sigma \gamma| &\lesssim \frac{tU(\beta, \gamma)}{\sqrt{n}} + \left\{1 + U(\beta, \gamma)\sqrt{\log n}\right\}\frac{k \log p}{n}, \\
|\hat{\beta}^\top \Sigma \beta - \beta^\top \Sigma \beta| &\lesssim \frac{t\|\beta\|_2}{\sqrt{n}} + (1 + \|\beta\|_2\sqrt{\log n})\frac{k \log p}{n},
\end{align*}

with probability at least $1 - p^{-c} - n^{-c} - t^{-2}$, for some constant $c > 0$.

In Theorem 1, in addition to the mild sparsity condition, the consistency of the proposed estimators requires only the balanced marginal case probabilities from (A1) and (A3), and the general sub-Gaussian design with a regular covariance matrix, which includes many important cases, such as Gaussian, bounded, and binary designs, or any combinations of them. Thus the proposed methods are widely applicable to various practical settings.

To establish the optimality of the proposed genetic covariance estimator, our next result concerns the minimax lower bound for estimating $\beta^\top \Sigma \gamma$. \[\text{Statistica Sinica: Preprint} \quad \text{doi:10.5705/ss.202021.0386}\]
To this end, we define the parameter space for $\theta = (\beta, \gamma, \Sigma)$ as

$$\Theta(k, L_n) = \left\{ (\beta, \gamma, \Sigma) : \max\{\|\beta\|_0, \|\gamma\|_0\} \leq k, U(\beta, \gamma) \leq L_n, M^{-1} \leq \lambda_{\min}(\Sigma) \leq \lambda_{\max}(\Sigma) \leq M \right\},$$

for some constant $M > 1$, and denote $\xi = \beta^\top \Sigma \gamma$.

**Theorem 2** (Minimax Lower Bound). Suppose $X_i$ and $Z_i \overset{i.i.d.}{\sim} N(0, \Sigma)$, for $i = 1, \ldots, n$, and $k \lessapprox \min\{p^{\nu}, \frac{n}{\log p}\}$, for some $0 < \nu < 1/2$. Then,

$$\inf_{\xi} \sup_{\theta \in \Theta(k, L_n)} P_\theta \left( |\hat{\xi} - \xi| \gtrsim \frac{L_n^2}{\sqrt{n}} + \min \left\{ \frac{L_n}{\sqrt{n}}, \frac{k \log p}{n}, L_n^2 \right\} \right) \geq c,$$

for some constant $c > 0$.

By Theorem 1, a uniform upper bound over the parameter space $\Theta(k, L_n)$ can be obtained as $\sup_{\theta \in \Theta(k, L_n)} P_\theta \left( |\hat{\beta}^\top \Sigma \gamma - \beta^\top \Sigma \gamma| \lessapprox \frac{U_n}{\sqrt{n}} + (1 + L_n \sqrt{\log n}) \frac{k \log p}{n} \right) \geq 1 - p^{-c} - n^{-c} - t^{-2}$. Combining this with the lower bound from Theorem 2, we conclude that, for all $k \lessapprox \min\{\frac{n}{\log p \log n}, p^{\nu}\}$, with any $\nu \in (0, 1/2)$, and $\sqrt{\frac{k \log p}{n}} \lessapprox L_n \lessapprox 1$, our genetic covariance estimator $\hat{\beta}^\top \Sigma \gamma$ is minimax rate-optimal over $\Theta(k, L_n)$, up to a $\sqrt{\log n}$ factor. In particular, in this case, the exact rate optimality of $\hat{\beta}^\top \Sigma \gamma$ is guaranteed over the ultra-sparse region $k \lessapprox \frac{\sqrt{n}}{\log p \log n}$, or the weak signal regime $L_n \lessapprox (\log n)^{-1/2}$, over which the minimax rate is $\frac{L_n}{\sqrt{n}} + \frac{k \log p}{n}$. Moreover, this suggests that the uncertainty due to the covariance estimation $\hat{\beta}^\top (\hat{\Sigma} - \Sigma) \hat{\gamma}$ in the plug-in estimator is fundamental and may not be removed, as for the leading-order biases.
Theorem 3 (Rate of Convergence). Suppose (A1), (A2), and (A3) hold, \( n_1 \asymp n_2 \asymp n \), \( k \ll \frac{n}{\log p \log n} \), and \( L(\beta, \gamma) \gg \sqrt{k \log p / n} \). Then, \( |\hat{R} - R| \rightarrow 0 \) in probability. In particular, for sufficiently large \((n, p)\) and any constant \( t > \sqrt{2} \), with probability at least \( 1 - 2t^{-2} \), it holds that

\[
|\hat{R} - R| \lesssim t \left\{ \frac{U(\beta, \gamma)}{L^2(\beta, \gamma) \sqrt{n}} + \frac{1 + U(\beta, \gamma) \sqrt{\log n}}{L^2(\beta, \gamma)} \cdot \frac{k \log p}{n} \right\}.
\]

(4.4)

Compared with Theorem 1, the consistency of \( \hat{R} \) requires an additional condition (A2) and a lower bound on the minimal effect size. These conditions are necessary to ensure that the true genetic variances are bounded away from zero and the genetic correlation is well defined.

4.2 Theoretical Properties of the Inference Procedures

We establish the asymptotic normality of the proposed bias-corrected estimators and provide theoretical justifications for the CIs and statistical tests. We start with a theorem that provides a refined analysis of the estimation errors, and consequently, the asymptotic normality of the estimators.

Theorem 4 (Asymptotic Normality). Suppose (A1), (A2), and (A3) hold, \( n_1 \asymp n_2 \asymp n \), \( k \lesssim \frac{n}{\log p \log n} \), and \( L(\beta, \gamma) \gg \sqrt{k \log p / n} \). Then, we have the following:

1. It holds that \( \hat{\beta}^T \Sigma \gamma - \beta^T \Sigma \gamma = A_n + B_n \), where \( P\{A_n \lesssim \{U(\beta, \gamma) \sqrt{\log n} + 1\} \frac{k \log p}{n}\} \geq 1 - p^{-c} - n^{-c} \), and \( \frac{\sqrt{n_1 + n_2} \beta_n}{v} \rightarrow_d N(0, 1) \) as \((n, p) \rightarrow \ldots\)
Additionally, if $k \ll \frac{U(\beta, \gamma) \sqrt{n}}{1 + U(\beta, \gamma) \sqrt{\log n} \log p}$, we establish the asymptotic normality $\sqrt{n_1 + n_2} \left( \frac{\beta^\top \Sigma \beta - \beta^\top \Sigma \beta}{v} \right) \mid \mathcal{D}_1 \rightarrow_d N(0, 1)$.

2. It holds that $\beta^\top \Sigma \beta - \beta^\top \Sigma \beta = A_n' + B_n'$, where $P \{ A_n' \lesssim (\|\beta\|_2 \sqrt{\log n} + 1) \frac{k \log p}{n} \} \geq 1 - p^{-c} - n^{-c}$, and $\frac{\sqrt{n_1 + n_2} B_n'}{v_{\beta}} \mid \mathcal{D}_1 \rightarrow_d N(0, 1)$ as $(n, p) \rightarrow \infty$. Additionally, if $k \ll \frac{\||\beta\|_2 \sqrt{n}}{1 + \||\beta\|_2 \sqrt{\log n} \log p}$, we establish the asymptotic normality $\sqrt{n_1 + n_2} \left( \frac{\beta^\top \Sigma \beta - \beta^\top \Sigma \beta}{v_{\beta}} \right) \mid \mathcal{D}_1 \rightarrow_d N(0, 1)$.

The second part of the theorem applies to the estimator $\gamma^\top \Sigma \gamma$, by symmetry. A direct consequence of Theorems 1 and 4, in combination with Slutsky’s theorem, is the following theorem concerning the asymptotic normality of the genetic correlation estimator $\bar{R}$ in Section 2.2.

**Theorem 5** (Asymptotic Normality). Under the conditions of Theorem 4, if $k \ll \min \left\{ \frac{n}{\log p \log n + \{1 + U(\beta, \gamma) \sqrt{\log n} \log p \}} \right\}$, we have $\frac{\sqrt{n_1 + n_2} (\bar{R} - R)}{v_{\bar{R}}} \mid \mathcal{D}_1 \rightarrow_d N(0, 1)$ as $(n, p) \rightarrow \infty$.

Some remarks about the technical innovations leading to the above theorems are in order. First, in contrast to existing works on statistical inference in high-dimensional logistic regression, the proposed methods do not require several commonly assumed, but stringent theoretical conditions, such as the bounded individual probability condition (van de Geer, 2008; van de Geer et al., 2014; Ning and Liu, 2017; Ma et al., 2020; Guo et al.,
(2021), where \( P(y_i = 1|X_i) \in (\delta, 1 - \delta) \), for all \( 1 \leq i \leq n \) and some \( \delta \in (0, 1/2) \), the sparse inverse population Hessian condition (van de Geer et al., 2014; Belloni et al., 2016; Ning and Liu, 2017; Janková and van de Geer, 2018), and the sparse precision condition (Ma et al., 2020). Second, from a practical viewpoint, removing these technical assumptions significantly expands the range of applicability of the proposed methods. For example, as argued by Cai et al. (2021) and Xia et al. (2020), in practice, the bounded individual probability and the sparse inverse population Hessian conditions are seldom satisfied or verifiable from the data. In contrast, the balanced marginal case probability condition holds easily, and can be checked based on the observed outcomes.

Using Theorems 4 and 5, we obtain theoretical justifications, such as the asymptotic coverage probability and the expected length of the proposed CIs, namely, \( \text{CI}_\alpha(\beta^T \Sigma \gamma, \mathcal{D}) \), \( \text{CI}_\alpha(\beta^T \Sigma \beta, \mathcal{D}) \), and \( \text{CI}_\alpha(R, \mathcal{D}) \).

**Theorem 6 (CIs).** Under the conditions of Theorem 4, for any constant \( 0 < \alpha < 1 \), if \( k \ll \min\{n \log p \log n, \frac{U(\delta, \gamma) \sqrt{n}}{1 + U(\delta, \gamma) \sqrt{\log n}} \} \), then, we have the following:

1. *(Coverage)* \( \lim_{n,p \to \infty} P_\theta\{\beta^T \Sigma \gamma \in \text{CI}_\alpha(\beta^T \Sigma \gamma, \mathcal{D})\} \geq 1 - \alpha \), \( \lim_{n,p \to \infty} P_\theta\{\beta^T \Sigma \beta \in \text{CI}_\alpha(\beta^T \Sigma \beta, \mathcal{D})\} \geq 1 - \alpha \), and \( \lim_{n,p \to \infty} P_\theta\{R \in \text{CI}_\alpha(R, \mathcal{D})\} \geq 1 - \alpha \);
2. (Length) if we denote \( L\{\text{CI}_\alpha(\cdot, \mathcal{D})\} \) as the length of \( \text{CI}_\alpha(\cdot, \mathcal{D}) \), then with probability at least \( 1 - p^{-c} \), we have \( L\{\text{CI}_\alpha(\beta^T\Sigma\gamma, \mathcal{D})\} \asymp \frac{U(\beta, \gamma)}{\sqrt{n}} \), \( L\{\text{CI}_\alpha(\beta^T\Sigma\beta, \mathcal{D})\} \asymp \|\beta\|_2 \sqrt{n} \), and \( L\{\text{CI}_\alpha(R, \mathcal{D})\} \asymp \frac{1}{L(\beta, \gamma)} \sqrt{n} \).

This theorem implies that the statistical tests proposed in Section 3 have the following theoretical properties related to their size and power under certain local alternatives.

**Corollary 1** (Hypothesis Testing). Under the conditions of Theorem 6, we have the following:

1. (Size) for each \( \ell \in \{1, 2, 3\} \), for any constant \( 0 < \alpha < 1 \), under the null hypothesis \( H_{0,\ell} \), we have \( \lim_{n,p \to \infty} P_\theta(|T_\ell| > z_{\alpha/2}) \leq \alpha \);

2. (Power) for any \( 0 < \delta < 1 \), there exists some \( c > 0 \) such that, for any \( |\beta^T\Sigma\gamma - B_0| \geq cU(\beta, \gamma)n^{-1/2} \), \( \lim_{n,p \to \infty} P_\theta(|T_1| > z_{\alpha/2}) \geq 1 - \delta \); for any \( |\beta^T\Sigma\beta - Q_0| \geq c\|\beta\|_2n^{-1/2} \), \( \lim_{n,p \to \infty} P_\theta(|T_2| > z_{\alpha/2}) \geq 1 - \delta \); and for any \( |R - R_0| \geq cL^{-1}(\beta, \gamma)n^{-1/2} \), \( \lim_{n,p \to \infty} P_\theta(|T_3| > z_{\alpha/2}) \geq 1 - \delta \).

5. **Simulations**

5.1 Evaluations Based on Simulated Genetic Data

To justify our proposed methods for analyzing real genetic data sets, we carried out numerical experiments under settings in which the covariates
are simulated genotypes with possible LD structures that resemble those of the human genome, and inferences are made at a chromosomal basis. Specifically, focusing on Scenario I with $n_1 = n_2 = n$, for given choices of $p$ and $n$, using the R package sim1000G (Dimitromanolakis et al., 2019), we generate genotypes of $2n$ unrelated individuals containing $p$ SNPs, based on the sequencing data over a region (GrCH37: bp 40,900 to bp 2,000,000) on chromosome 9 of 503 European samples from the 1000 Genomes Project Phase 3 (1000 Genomes Project Consortium, 2015). We also generate a comprehensive haplotype map integrated over 1,184 reference individuals (International HapMap 3 Consortium, 2010); see Section S4 of the Supplementary Material for the resulting correlation matrix among the generated SNPs. The true effect sizes for the two binary traits were generated such that for each trait there are 25 associated SNPs, with 12 of them shared by both traits. The effect sizes of the associated SNPs are drawn uniformly from $[-1, 1]$. For reasons of practical interest, we focus mainly on the estimation, CIs and hypothesis testing about the genetic correlation parameter. The results for the genetic covariance and variance can be found in Section 5.2 below and Section S4 of the Supplementary Material.

For the parameter estimation, in addition to our proposed estimators ("pro"), we also considered (i) the simple plug-in ("plg") estimators $\hat{\beta}^T \hat{\Sigma} \hat{\gamma}$,
\[ \tilde{\beta}^\top \hat{\Sigma} \tilde{\beta}, \text{ and } \hat{R}_{plg} = \frac{\beta^\top \Sigma \gamma}{\sqrt{\beta^\top \Sigma \beta \gamma^\top \Sigma \gamma}}; \]

(iii) the component-wise projected Lasso ("lpj") estimators \( \tilde{\beta}^\top \hat{\Sigma} \tilde{\gamma}, \tilde{\beta}^\top \hat{\Sigma} \tilde{\beta}, \) and \( \hat{R}_{lpj} = \frac{\hat{\beta}^\top \hat{\Sigma} \hat{\gamma}}{\sqrt{\beta^\top \Sigma \beta \gamma^\top \Sigma \gamma}}, \) where each component of \( \tilde{\beta} \) and \( \tilde{\gamma} \) is the debiased Lasso estimator implemented by the function \texttt{lasso.proj} in the R package \texttt{hdi} using the default settings; and

(iii) the component-wise projected Ridge ("rpj") estimators \( \tilde{\beta}^\top \hat{\Sigma} \tilde{\gamma}, \tilde{\beta}^\top \hat{\Sigma} \tilde{\beta}, \) and \( \hat{R}_{rpj} = \frac{\hat{\beta}^\top \hat{\Sigma} \hat{\gamma}}{\sqrt{\beta^\top \Sigma \beta \gamma^\top \Sigma \gamma}}, \) where each component of \( \tilde{\beta} \) and \( \tilde{\gamma} \) is obtained from the function \texttt{ridge.proj} in the R package \texttt{hdi} using the default settings.

For the proposed method, we use cross-validation to determine the tuning parameter (see Section S4.1 for details). Table 1 contains the empirical estimation errors (square roots of the empirical mean-squared errors) for the genetic correlation estimators, which demonstrate the superior performance of the proposed method.

For the CIs, we compare our proposed CIs ("pro") with alternative bootstrap CIs. Specifically, the bootstrap CIs are based on the plg estimators calculated from 100 observations sampled from the original data set, so that the final CIs are constructed based on the empirical distributions of 500 bootstrap estimators. Table 2 contains the averaged coverage probabilities and lengths of the proposed and the plg-based bootstrap CIs, denoted as "boot," with 500 rounds of simulation for each setting. Our results suggest the desirable coverage and shorter length of the proposed
CIs. Finally, for hypothesis testing, we evaluate the empirical type-I errors and statistical power of the proposed tests and the plg-based bootstrap tests in a setup in which the effect sizes are generated using an additional constraint $|\beta^T \Sigma \gamma| > 3$. Table 3 shows the empirical type-I errors and statistical power of the proposed tests over different settings, each based on 500 rounds of simulations. Our results suggest that the proposed test is empirically valid and has advantages over the bootstrap tests. In Tables 2 and 3, the proposed method becomes a little conservative when $n$ increases from 200 to 400, likely because of the limitation of our empirically determined tuning parameter. Nevertheless, we still observe greater power for the test and shorter lengths for the CIs with larger $n$, and in both cases, the advantage over the alternative methods. For additional simulations under a slightly different setting of the association structure, see Section S4.5 of the Supplementary Material (Table S8).

5.2 Evaluation Based on Model-Generated Data

We consider the high-dimensional setting $p > n$, and set the sparsity level as $k = 25$. For the true regression coefficients, given the support $S$ such that $|S| = k$, we generate $\beta_j$ and $\gamma_j$ uniformly from $[-1, 1]$, for all $j \in S$. For the design covariates, we focus on Scenario I, where $n_1 = n_2 = n$. The
Table 1: Estimation errors for the genetic correlation under simulated genetic data with $k = 25$. pro: proposed estimators; plg: simple plug-in estimators; lpj: component-wise projected Lasso estimators; rpj: the component-wise projected Ridge estimators.

| $p$  | $n = 200$ |           |           |           |           |           |           |           |           |           |           |           |           |           |           |
|------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
|      | pro | plg | lpj | rpj | pro | plg | lpj | rpj | pro | plg | lpj | rpj | pro | plg | lpj | rpj | pro | plg | lpj | rpj | pro | plg | lpj | rpj | pro | plg | lpj | rpj |
| 700  | 0.09 | 0.12 | 0.15 | 0.16 | 0.09 | 0.11 | 0.14 | 0.13 | 0.08 | 0.11 | 0.13 | 0.12 |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 800  | 0.08 | 0.10 | 0.15 | 0.14 | 0.08 | 0.11 | 0.15 | 0.11 | 0.09 | 0.11 | 0.15 | 0.12 |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 900  | 0.09 | 0.13 | 0.16 | 0.15 | 0.11 | 0.12 | 0.15 | 0.13 | 0.07 | 0.11 | 0.14 | 0.11 |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 1000 | 0.10 | 0.12 | 0.14 | 0.15 | 0.09 | 0.11 | 0.14 | 0.12 | 0.08 | 0.09 | 0.14 | 0.09 |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |

Table 2: Coverage and length of the CIs for the genetic correlation under simulated genetic data with $\alpha = 0.05$.

| $p$  | $n = 200$ |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |           |
|------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
|      | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length | coverage | length |
| 700  | 96.4     | 82.4     | 0.30     | 0.37     | 97.6     | 85.8     | 0.26     | 0.39     | 97.0     | 82.6     | 0.27     | 0.41     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 800  | 97.0     | 85.4     | 0.29     | 0.37     | 98.0     | 82.5     | 0.27     | 0.39     | 98.2     | 85.2     | 0.26     | 0.39     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 900  | 96.6     | 84.2     | 0.31     | 0.36     | 96.8     | 86.2     | 0.26     | 0.38     | 97.6     | 84.0     | 0.25     | 0.39     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 1000 | 97.5     | 86.0     | 0.30     | 0.34     | 97.6     | 80.0     | 0.26     | 0.36     | 97.8     | 84.9     | 0.26     | 0.41     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
Table 3: Type-I errors and power when testing the genetic correlation under simulated genetic data with $\alpha = 0.05$.

| $p$ | $n = 200$ |  |  |  | $n = 300$ |  |  |  | $n = 400$ |  |  |  |
|-----|-----------|---|---|---|-----------|---|---|---|-----------|---|---|---|
|     | type I error | power | type I error | power | type I error | power | type I error | power |
|     | pro | boot | pro | boot | pro | boot | pro | boot | pro | boot | pro | boot |
| 700 | 0.04 | 0.41 | 0.47 | 0.72 | 0.04 | 0.35 | 0.63 | 0.68 | 0.02 | 0.34 | 0.69 | 0.65 |
| 800 | 0.04 | 0.42 | 0.46 | 0.74 | 0.03 | 0.37 | 0.59 | 0.71 | 0.03 | 0.34 | 0.70 | 0.66 |
| 900 | 0.04 | 0.42 | 0.45 | 0.70 | 0.03 | 0.35 | 0.64 | 0.66 | 0.02 | 0.32 | 0.69 | 0.73 |
| 1000| 0.06 | 0.41 | 0.42 | 0.71 | 0.02 | 0.36 | 0.63 | 0.70 | 0.02 | 0.36 | 0.68 | 0.70 |

covariates are generated from a multivariate Gaussian distribution with covariance matrix as either $\Sigma = \Sigma_B$, where $\Sigma_B$ is a $p \times p$ blockwise-diagonal matrix of 10 identical unit-diagonal Toeplitz matrices, with off-diagonal entries that descend from 0.3 to 0 (see Section S4.1 of the Supplementary Materia for its explicit form), or $\Sigma = \Sigma_E$, where $\Sigma_E$ is an exchangeable covariance matrix with unit diagonals and off-diagonals equal to 0.2. The numerical result for each setting is based on 500 rounds of simulations.

For the parameter estimation, we evaluate the proposed method and the three alternative methods defined in the previous section. The results are provided in Section S4.2 of the Supplementary Material (Tables S1, S2),
which show the superiority of each of the proposed estimators over the alternatives. For the same simulation setups, we evaluate and compare various methods for constructing 95% CIs for the parameters. Specifically, we compare our proposed CIs (“pro”) with two alternative bootstrap CIs, based on 500 plg estimators or rpj estimators, calculated from 100 observations sampled from the original data set. Table 4 contains the averaged coverage probabilities and lengths of the proposed and the plg-based bootstrap CIs (“boot”) under the blockwise-diagonal covariant matrix. In general, the coverage of the rpj-based bootstrap CIs is poorer than that of the plg-based CIs for $\beta^\top \Sigma \gamma$ and $\beta^\top \Sigma \beta$, and only slightly better than that of the plg-based CIs for $R$; these results and those under the exchangeable covariance are provided in Supplementary Material Section S4.3 (Tables S3 - S5). In general, our proposed CIs achieve the 95% nominal confidence levels, whereas the bootstrap CIs are off target or biased. In particular, for the genetic correlation $R$, the proposed CI has better coverage and a smaller length. In addition, our proposed methods are computationally more efficient than the bootstrap CIs, because the averaged running time (MacBook Pro, with 2.2 GHz 6-Core Intel Core i7) for the proposed CIs is only about 1 second, whereas the bootstrap CIs take more than 1.6 minutes for the plg-based CIs, and 1 hour for the rpj-based CIs. When the sample size increases from 300
to 500, the empirical coverage of the proposed CIs for $\beta^T \Sigma \gamma$ and $R$ seems to inflate slightly, which is again likely due to our empirically determined tuning parameter. Nevertheless, the proposed CIs have a shorter length for larger $n$, and the advantage of the proposed method over the alternative methods is evident.

For the hypothesis testing, we also compare the empirical Type-I errors and statistical power of our proposed tests and the plg-based bootstrap tests, demonstrating the empirical superiority of the proposed method; these results are provided in Section S4.4 (Tables S6 and S7) of the Supplementary Material.

6. Analysis of 10 Pediatric Autoimmune Diseases

We investigate the genetic correlations between each pair of 10 pediatric autoimmune diseases, including autoimmune thyroiditis (THY), psoriasis (PSOR), juvenile idiopathic arthritis (JIA), ankylosing spondylitis (AS), common variable immunodeficiency (CVID), celiac disease (CEL), Crohn’s disease (CD), ulcerative colitis (UC), type 1 diabetes (T1D) and systemic lupus erythematosus (SLE). We identified the subjects with a disease and controls either directly from previous studies, or from de-identified samples and associated electronic medical records in the genomics biorepository at
Table 4: Coverage and length of the CIs with $\Sigma = \Sigma_B$, $\alpha = 0.05$, and sparsity $k = 25$.

| $p$ | $\beta^T \Sigma \gamma$ | $\beta^T \Sigma \beta$ | $R$ |
|-----|--------------------------|--------------------------|-----|
|     | pro | boot | pro | boot | pro | boot |
| cov | len | cov | len | cov | len | cov | len |
|-----|-----|-----|-----|-----|-----|-----|-----|
|   |     |     |     |     |     |     |     |
| $n = 300$ | | | | | | | |
| 700 | 94.8 | 6.24 | 46.4 | 2.05 | 94.4 | 7.61 | 13.5 | 2.42 | 96.6 | 0.35 | 76.0 | 0.37 |
| 800 | 97.4 | 7.72 | 47.8 | 1.91 | 92.4 | 7.89 | 13.2 | 2.30 | 95.0 | 0.37 | 76.4 | 0.36 |
| 900 | 93.6 | 5.59 | 50.2 | 1.85 | 93.8 | 6.71 | 14.6 | 2.27 | 96.4 | 0.34 | 73.6 | 0.35 |
| 1000 | 93.2 | 5.85 | 42.6 | 1.93 | 92.6 | 7.88 | 7.2 | 2.39 | 93.0 | 0.32 | 76.4 | 0.36 |
| $n = 400$ | | | | | | | | |
| 700 | 96.0 | 6.11 | 56.6 | 2.30 | 92.0 | 7.85 | 30.0 | 2.96 | 96.6 | 0.32 | 76.6 | 0.37 |
| 800 | 97.4 | 5.91 | 55.4 | 2.20 | 92.4 | 7.47 | 22.8 | 2.63 | 96.2 | 0.32 | 74.4 | 0.37 |
| 900 | 96.6 | 5.81 | 51.0 | 2.19 | 90.6 | 7.32 | 21.6 | 2.69 | 96.6 | 0.31 | 73.0 | 0.37 |
| 1000 | 93.8 | 5.65 | 47.8 | 2.07 | 90.4 | 7.11 | 19.8 | 2.58 | 93.4 | 0.31 | 72.6 | 0.36 |
| $n = 500$ | | | | | | | | |
| 700 | 99.0 | 5.71 | 61.0 | 2.40 | 95.2 | 6.93 | 43.2 | 2.92 | 98.6 | 0.30 | 73.4 | 0.37 |
| 800 | 98.6 | 5.70 | 60.6 | 2.38 | 93.4 | 7.07 | 41.2 | 2.83 | 97.2 | 0.29 | 78.0 | 0.37 |
| 900 | 99.2 | 5.92 | 58.0 | 2.32 | 92.6 | 7.36 | 31.2 | 2.88 | 98.4 | 0.30 | 76.6 | 0.36 |
| 1000 | 98.6 | 5.44 | 57.8 | 2.18 | 90.4 | 6.70 | 30.0 | 2.73 | 98.2 | 0.29 | 76.6 | 0.36 |
The data set includes 10,718 normal controls, 97 THY cases, 107 AS cases, 100 PSOR cases, 173 CEL cases, 254 SLE cases, 308 CVID cases, 865 UC cases, 1086 T1D cases, 1123 JIA cases, and 1922 CD cases. Specifically, for each pair of the 10 diseases, we evaluate their chromosome-specific genetic relatedness by estimating and performing hypothesis testing on the genetic correlation parameter for each of the 22 autosomes. By focusing on the chromosome-specific genetic correlations, we can make a better inference from a limited sample size for many diseases, and obtain insights on the genomic regions that relate the two diseases of interest.

For each subject, after removing the SNPs with a minor allele frequency less than 0.05, we have a total of 475,324 SNPs were obtained across 22 autosomes (see Supplementary Material for details). To apply our proposed methods, for each pair of diseases, we randomly split the controls into two groups of equal size, combine them with each of the cases, and fit two high-dimensional logistic regressions between the disease outcomes and the SNPs to obtain the initial logistic Lasso estimators for each disease. Then we obtain the bias-corrected estimators, where the sample covariance matrix is calculated based on all samples. Moreover, using our proposed method, we test the individual null hypothesis that the chromosome-specific genetic
correlation is zero between each pair of diseases in order to identify i) dis-
eases that are genetically associated; and ii) specific chromosomes in which
diseases have a shared genetic architecture.

The results are summarized in Figure 1. The top panel shows the
estimated chromosome-specific genetic correlations between each pair of
diseases, where the disease pairs with larger absolute values are annotated.
The bottom panel shows the negative log-transformed p-values for each
pair of diseases. Our tests suggest strong genetic sharing between UC and
CD on chromosomes 1, 12, 17, 20, and 21, between CVID and JIA on
chromosome 8, and between CD and PSOR on chromosome 13. Many
pairs of these diseases showed genetic relatedness at the nominal p-value
of 0.05. However, however, because of the small sample sizes, they do not
reach the statistical significance after the Bonferroni adjustment of multiple
comparisons. Note that the pairs UC and CD, and CVID and JIA were
also found to be statistically significant by Li et al. (2015) using different
measures of genetic sharing. However, our proposed methods also locate
genetic sharing with specific chromosomes and provide theoretically valid
uncertainty quantifications.
Figure 1: Analysis of genetic sharing between 10 autoimmune diseases. Top panel: estimated genetic correlations between each pair of diseases on each autosome. Bottom panel: negative log-transformed p-values for each pair of diseases, based on the proposed method. The red and blue dashed lines represent the original and Bonferroni-adjusted significance levels, respectively, at 0.05.
7. Discussion

In this paper, we propose a statistical inference framework for studying the genetic relatedness between two binary traits in high-dimensional logistic regression models. Our model allows the number of SNPs to far exceed the sample size while producing efficient and valid statistical inferences under mild conditions on the sparsity and the effect size of the true associations, and on the covariance structure or linkage disequilibrium of the variants. Many works have tried to improve the speed of optimization and operation for genome-scale and ultrahigh-dimensional data sets. For example, Qian et al. (2019) propose a new computational framework in which scalable Lasso solutions can be obtained for a very large Biobank data set involving about 300,000 individuals and 800,000 genetic variants. We expect that these new computational methods will increase the utility of the proposed methods in genetic correlation analysis at a whole-genome sequencing scale.

Supplementary Materials

The Supplementary Material includes proofs of the main theorems and the technical lemmas, additional simulation results, supplementary notes, figures and tables.
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