A Generalized Levene’s Scale Test for Variance Heterogeneity in the Presence of Sample Correlation and Group Uncertainty

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SUMMARY. We generalize Levene’s test for variance (scale) heterogeneity between \( k \) groups for more complex data, when there are sample correlation and group membership uncertainty. Following a two-stage regression framework, we show that least absolute deviation regression must be used in the stage 1 analysis to ensure a correct asymptotic \( \chi^2_{k-1}/(k-1) \) distribution of the generalized scale (gS) test statistic. We then show that the proposed gS test is independent of the generalized location test, under the joint null hypothesis of no mean and no variance heterogeneity. Consequently, we generalize the recently proposed joint location-scale (gJLS) test, valuable in settings where there is an interaction effect but one interacting variable is not available. We evaluate the proposed method via an extensive simulation study and two genetic association application studies.

KEY WORDS: Association studies; Heteroscedasticity; Joint location-scale test; Scale test.

1. Introduction

Testing for scale (variance) heterogeneity, prior to the main inference of location (mean) parameters, is a common diagnostic method in linear regression to evaluate the assumption of homoscedasticity. In some research areas, such as statistical genetics, testing for heteroscedasticity itself can be of primary interest.

With the goal of detecting a genetic association between a single-nucleotide polymorphism (SNP, \( G \)) and a quantitative outcome (phenotype, \( Y \)), the traditional approach is to conduct a location test, testing mean differences in \( Y \) across the three genotype groups of the SNP (\( G = 0, 1, \) or \( 2 \) copies of the minor allele, the variant with population frequency \(<0.5 \)). However, it has been noted that a number of biologically meaningful scenarios can lead to variance differences in \( Y \) across the genotype groups of a SNP of interest (say \( G_1 \)). For example, an underlying interaction effect, between \( G_1 \) and another SNP \( G_2 \) (\( G_2 \times G_3 \)) or an environmental factor \( E \) (\( G_1 \times E \)), on \( Y \) can lead to heteroscedasticity across \( G_1 \), if the interacting \( G_2 \) or \( E \) variable was not collected and the interaction term may not be directly modeled (Pare et al., 2010). Transformations on a phenotype can also result in variance heterogeneity (Sun et al., 2013). This transformation can occur knowingly for statistical purposes, for example, \( \log(Y) \), or unknowingly, for example, choosing a phenotype measurement that does not directly represent the true underlying biological outcome of a gene. In each of these scenarios, a scale test can be used either alone to indirectly detect associated SNPs (Pare et al., 2010), or combined with a location test to increase testing power (Cao et al., 2014; Soave et al., 2015). Heteroscedasticity due to interaction effects has also been investigated for variable selection via sliced inverse regression methods (Jiang and Liu, 2014).

Genotype uncertainty is inherent in sequenced and imputed SNP data. For such data, the genotype of a SNP for an individual (\( G = 0, 1, \) or \( 2 \)) is represented by three genotype probabilities (\( p_0, p_1, p_2, \) and \( p_0 + p_1 + p_2 = 1 \)). For testing methods that require genotype to be known unambiguously, the probabilistic data are typically transformed into the so-called “best-guess” (most likely or hard-call) genotype, selected as the one with the largest probability. In the context of location-testing, several groups have proposed methods that incorporate the probabilistic data and showed that this improves power (Acar and Sun, 2013; Kutalik et al., 2011). The corresponding development for scale-testing, however, is lacking.

Genetic association studies often involve related individuals, where individuals in a sample are correlated or clustered. The correlation structure may be specified based on the known genealogy information or accurately estimated using the genome-wide genetic SNP data collected (Sun and Dimitrakas, 2012). A number of generalized location tests allowing for family data have been proposed (Horvath et al., 2001; Jakobsdottir and McPeek, 2013), and their power gain over analyzing only a subset of independent individuals is a direct consequence of the increase in sample size. However,
few scale tests deal with correlated data, with the exception of methods proposed specifically for clustered data present in twin studies (Haseman and Elston, 1970; Iachine et al., 2010). Further, these methods have been reported to have type 1 error issues in the presence of non-normal data or small, unequal group sizes (Iachine et al., 2010), and they have not been extended to incorporate group membership uncertainty.

Both classical statistical tests and graphical procedures have been proposed to investigate heteroscedasticity (Bartlett, 1937; Breusch and Pagan, 1979; Cook and Weisberg, 1983; Levene, 1960; White, 1980). In big data settings, such as genome-wide association studies where possibly millions of SNPs are scanned for association with an outcome, graphical and other computationally burdensome approaches are not ideal. Levene’s test (Levene, 1960) is known for its simplicity and robustness to modeling assumptions, and it is perhaps the most popular method for evaluating variance heterogeneity between k groups. Therefore, our development here focuses on Levene’s method.

In this article, we extend Levene’s test for equality of variances across k groups to allow for both group membership uncertainty and sample correlation. When groups are known, we show that the proposed method outperforms existing methods for clustered twin data. In the presence of group uncertainty, we demonstrate that our test continues to be accurate and has improved power over the “best-guess” approach. This generalized scale test can be used alone for heteroscedasticity diagnostic purposes but with wider applicability. Motivated by the complex genetic association studies described above, we also show that the proposed generalized scale test can be combined with existing generalized location tests using the joint location-scale framework, previously developed for population samples without group uncertainty (Soave et al., 2015), to further improve power. We apply our methods to two genetic association studies, one of HbA1c levels in individuals with type 1 diabetes, and the other of lung disease in individuals with cystic fibrosis (CF).

2. Methodology

We first consider a sample of independent observations with no group uncertainty, and we formulate Levene’s test as a regression problem. Using this regression framework, we then extend Levene’s test as the generalized scale (gS hereinafter) test to allow for sample dependency and group uncertainty. For clarity in our method comparisons, we also briefly discuss the Iachine et al. (2010) extension of Levene’s test, specifically designed for twin pairs without group uncertainty. Finally, we generalize the joint location-scale test of Soave et al. (2015) for the complex data structure considered here (gILS).

2.1. Notation and Statistical Model

Let \( y_i, i = 1, \ldots, n \), be a sample of independent observations, where each \( y_i \sim N(\mu_i, \sigma^2_i) \). Suppose the \( y_i \)’s fall into \( k \) distinct treatment groups with group-specific variance \( \sigma^2_j, j = 1, \ldots, k, \) and let \( n_j \) be the sample size for group \( j, n = \sum n_j \).

Our motivation concerns testing the null hypothesis of equal variance across the \( k \) groups:

\[
H_0 : \sigma^2_1 = \sigma^2_2 = \cdots = \sigma^2_k. \tag{1}
\]

Here, we use \( \sigma^2_j \) for group-specific variance, \( j = 1, \ldots, k \), and \( \sigma^2 \) for observation-specific variance, \( i = 1, \ldots, n \).

Let \( x_{ij}, j = 1, \ldots, k-1 \), be the standard dummy variables that define group membership for observation \( i \); group 1 is the reference group. Consider the normal linear model of interest here,

\[
y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \cdots + \beta_{k-1} x_{i(k-1)} + e_i, \tag{2}
\]

where \( e_i \sim N(0, \sigma^2) \), \( \sigma^2 \) corresponds to the variance associated with the group that \( y_i \) belongs to, \( i = 1, \ldots, n \). In other words, \( \sigma^2 = \sigma^2_j \) if \( x_{i(j-1)} = 1 \). In matrix notation,

\[
y = X\beta + \epsilon, \tag{3}
\]

where \( X \) is the design matrix obtained by stacking the \( x_{ij} \)'s = \( (1, x_{i1}, x_{i2}, \ldots, x_{i(k-1)}) \), \( \epsilon \sim N_p(0, \Sigma) \), and \( \Sigma \) is the covariance matrix with diagonal elements \( \sigma^2_j \).

2.2. Formulating Levene’s Test as a Regression F-Test and Modifications

The classical formulation of Levene’s test first centers the \( y_i \)’s by their estimated group means and obtains the corresponding absolute deviation \( d_i \)’s. It then tests for mean differences in the \( d_i \)’s across the \( k \) groups using ANOVA. Let \( I_{ij}, j = 1, \ldots, k \), be the group indicator variables, where \( I_{ij} = 1 \) if individual \( i \) belongs to group \( j \). Now, let \( \bar{\mu}_{(j)} = \sum_{i=1}^n I_{ij} \bar{y}_i/n_j \) be the estimated group means of the \( y_i \)’s, such that an estimate of \( E(y_i) \) is \( \bar{\mu}_i = \sum_{j=1}^k I_{ij} \bar{\mu}_{(j)} \). The corresponding absolute deviation is

\[
d_i = |y_i - \bar{\mu}_i|.
\]

Let \( \bar{\mu}_{(j)} = \sum_{i=1}^n I_{ij} \bar{y}_i/n_j \) be the estimated group means of the \( d_i \)’s, such that an estimate of \( E(d_i) \) is \( \bar{d}_i = \sum_{j=1}^k I_{ij} \bar{\mu}_{(j)} \), and let \( \bar{d} = \sum_{i=1}^n d_i/n \) be the grand mean. Finally, Levene’s test statistic has the following form

\[
F(d) = \frac{\sum_{i=1}^n (\bar{d}_i - \bar{d})^2/(k-1)}{\sum_{i=1}^n (d_i - \bar{d})^2/(n-k)},
\]

where \( F(d) \) follows approximately an \( F(k-1, n-k) \) distribution under the null hypothesis of (1), and a \( \chi^2_{k-1}/(k-1) \) distribution asymptotically as \( n \to \infty \).

For the purpose of developing a unified approach, we re-formulate Levene’s test using the following two-stage regression framework:

Stage 1.1. Obtain the residuals, \( \hat{\epsilon}_i = y_i - \hat{\bar{y}}_i = y_i - x_i^T \hat{\beta} \), from the ordinary least squares (OLS) regression of \( y_i \) on \( x_i \); we refer to this as the stage 1 regression.

Stage 1.2. Take the absolute values of these residuals, \( d_i = |\hat{\epsilon}_i| \).

Stage 2. Test for an association between the \( d_i \)’s and \( x_i \)’s using a regression F-test; we refer to this as the stage 2 regression and test.
The justification for this two-stage regression procedure (Levene’s test) being a test of homoscedasticity (1) is as follows. Stage 1 performs OLS regression using a working covariance matrix $\Sigma_{\text{stage}1} = \sigma^2 I$, where $I$ is the identity matrix. Therefore $y = X(\theta + e)^T y = H_0$, $\hat{y} = y - \tilde{y} \sim N(0, \Sigma(I - H))$, and $\hat{y}_i \sim N(0, \sigma^2(1 - h_{ii}))$, where $h_{ii}$ is the $i$th diagonal element of the hat matrix $H$. Consequently $d_i = |\hat{y}_i|$ follows a folded-normal distribution and its mean is a linear function of $\sigma_i$,

$$E(d_i) = \sigma_i \sqrt{2(1 - h_{ii})/\pi}.$$  

This relationship between $d_i$ and $\sigma_i$ is approximated by the following stage 2 working model,

$$d_i = \alpha + \gamma_1 x_{i1} + \gamma_2 x_{i2} + \cdots + \gamma_{k-1} x_{i(k-1)} + e_i,$$

where $e_i \sim N(0, \sigma^2_d)$. In matrix form,

$$d = X\theta + e,$$

where $\theta = (\alpha, \gamma_1, \ldots, \gamma_{k-1})^T$, and $e \sim N(0, \Sigma_{\text{stage}2})$.

$\Sigma_{\text{stage}1} = \sigma^2 I$. Testing the null hypothesis (1) is now reformulated as testing

$$H_0: \gamma_1 = \gamma_2 = \cdots = \gamma_{k-1} = 0,$$

using the classical OLS regression F-test. Note that although the $d_i$’s are folded normal variables, Levene’s variance test takes advantage of the fact that inference from OLS regression is robust to violations of the normality assumption.

This formulation of Levene’s test has a similar structure to the score test of Glejser (1969) proposed for testing heteroscedasticity associated with continuous covariates. Godfrey (1996) showed that when estimating $\hat{\beta}$ by OLS in stage 1, the Glejser score statistic derived in stage 2 is not asymptotically independent of each other and the covariance matrix $\Sigma$ is no longer diagonal. In the stage 1 regression, because we are only interested in obtaining $\hat{\beta}$ to construct $d_i = |y_i - x_i^T \hat{\beta}|$, we can continue to use OLS or LAD regression with the misspecified working covariance matrix, $\Sigma_{\text{stage}1} = \sigma^2_I$, to obtain consistent and unbiased $\hat{\beta}$ estimates.

Stage 2 involves estimating the variance of $\tilde{y}$ to test the null hypothesis of (6), and not accounting for sample dependency can lead to invalid inference. Let $\Sigma_{\text{stage}2} = \sigma^2_I \Sigma_d$ be the working covariance matrix for $d$, a valid inference can be achieved by using a generalized least squares (GLS) approach when $\Sigma_d$ is known (Aitken, 1936). When $\Sigma_d$ is unknown, feasible GLS (FGLS) can be used, with or without iteration, where an estimate of $\Sigma_d$ is obtained, subject to constraints, and then used in GLS. Alternatively, orthogonal-triangular decomposition methods can be used to obtain a compact representation of the profiled log-likelihood, such that maximum likelihood estimates (MLE’s) of all parameters can be obtained jointly through nonlinear optimization (Pinheiro and Bates, 2000).

In many scientific settings, including genetic association studies, the sample correlation structure is often prespecified with constraints on the $n(n - 1)/2$ correlations, for example, a single serial correlation $\rho$ for time series or family data with a single relationship type (e.g. sibling data), or cluster-specific correlations $\rho$’s for different clusters. In this case, let $\Sigma_{\text{stage}2} = \sigma^2_I \Sigma_d(\rho) = \sigma^2_I C(\rho)C(\rho)^T$ be the Cholesky decomposition, and define

$$d^* = C(\rho)^{-1}d, \ X^* = C(\rho)^{-1}X, \ e^* = C(\rho)^{-1}e.$$  

The GLS or FGLS regression, in essence, deals with the transformed model in stage 2

$$d^* = X^* \theta + e^*,$$

where $\theta = (\alpha, \gamma^T)^T$. For a fixed $\rho$, the conditional MLEs for $\theta$ and $\sigma^2_d$ are

$$\hat{\theta} = [X^*T X^*]^{-1}X^*T d^*, \ \hat{\sigma}^2_d = \frac{1}{n} \left\| d^* - X^* \hat{\theta} \right\|^2.$$  

The MLE of $\rho$ can be obtained by optimizing the profiled log-likelihood,

$$l(\rho) = \text{constant} - n \log \left\| d^*(\rho) - X^*(\rho) \hat{\theta}(\rho) \right\| - \frac{1}{2} \log |C(\rho)|.$$  

Thus, the generalized Levene’s scale (gS) test of the null hypothesis of (6), $H_0: \gamma = 0$, using the regression $F$-test in stage 2, has the following test statistic:

$$F(d^*) = \frac{\sum_{i=1}^n (d_i^* - \bar{d}_*^*)^2 / (k - 1)}{\sum_{i=1}^n (d_i^* - \bar{d}_*)^2 / (n - k)}.$$
where $\hat{d}_i = (x_i')\hat{\theta}$, the predicted values from regression model (7), and $\tilde{d}_i = 1_i\tilde{\alpha}$, the predicted values from the regression of $d_i^*$ on $1_i$. Note that $1_i$ is the first column of the transformed design matrix $X^*$, and may not be a vector of 1's. When the observations are independent of each other and group membership is known unambiguously, it is easy to verify that $\hat{d}_i = \tilde{d}_i$ and $\tilde{d}_i = \tilde{d}_i$, and $F(d^*)$ reduces to the original form of $F(d)$.

Under the normal linear model of (4), the $F$-statistic (8) is asymptotically $\chi^2_{r-1}/(k-1)$ distributed (Arnold, 1980). However, for non-symmetric $e$, we show that this is true only when $d$ is estimated using LAD in the stage 1 regression (Web Appendix A, Theorem 1).

In our simulation and application studies that follow, the examples considered involve known sibling pairs. In that case, we specify a compound symmetric correlation structure with one correlation parameter, $\rho$, that assumes equal correlation among all within-group errors pertaining to the same pair/cluster. We use GLS regression in R with the gls() function in the “nlme” package. The gls() function obtains maximum likelihood estimates of the model parameters by using a hybrid expectation maximization and Newton–Raphson algorithm to optimize the profiled likelihood; see Pinheiro and Bates (2000) for details.

2.4. The Iachine et al. (2010) Scale Test for Twin Pairs and Modifications

Focusing on paired-observations, Iachine et al. (2010) extended Levene’s test to determine if the variance of an outcome differs between monzygotic (MZ) and dizygotic (DZ) twin pairs. The proposed twin (TW) test follows Levene’s two-stage regression procedure but makes use of the Huber–White sandwich estimate (White, 1980) of Var($\gamma_i$) in the stage 2 analysis (here $k = 2$ groups, requiring only one dummy variable) to construct an asymptotically $\chi^2_k$ distributed Wald statistic, operationally an $F$-statistic in finite samples.

Complications with the TW test may arise if the number of clusters is small in either group (MZ or DZ) and can be compounded with imbalance between the groups (Iachine et al., 2010). Unfortunately, there is no clear definition of too few clusters (Cameron and Miller, 2015), and empirical type 1 error rates can be inflated for study designs with less than 20 clusters per group, particularly combined with non-symmetric data (see Iachine et al. (2010) and simulation results in Section 3 below).

The original TW method assumes that if two observations are from the same pair/cluster they also belong to the same group $k$. This may not be satisfied in a more general setting such as the genetic association studies discussed above. For example, two individuals from the same DZ pair or familial cluster often have different genotypes at a SNP of interest, so individuals from the same cluster may not share a common $\sigma^2$. However, the sandwich variance estimator can still be used in this setting. In the presence of group uncertainty, the TW method can be modified by replacing the group indicator covariate with corresponding group probabilities.

2.5. Generalized Joint Location-Scale (gJLS) Testing

The standard location test of mean differences in an (approximately) normally distributed outcome across covariate values (e.g., the three genotype groups of a SNP in a genetic association study) is testing

$$H_{0_{location}} : \beta_1 = \cdots = \beta_{k-1} = 0,$$

based on regression model (2). While the location inference tests the $\beta_j$’s, the scale test discussed here uses only the $p$ estimates from the stage 1 regression of model (2) to obtain $d = |y_i - \tilde{y}_i|$ for the stage 2 regression of model (4), and it performs a hypothesis test on the $\gamma_j$’s, testing

$$H_{0_{scale}} : \gamma_1 = \cdots = \gamma_{k-1} = 0.$$

A joint location-scale (JLS) test is interested in the following global null hypothesis,

$$H_{0_{joint}} : \beta_j = 0, \gamma_j = 0, \forall j = 1, \ldots, k-1. \quad (9)$$

One simple yet powerful JLS method proposed in Soave et al. (2015) uses Fisher’s method to combine $p_L$ and $p_S$, the p-values of the individual location and scale tests. One can consider other aggregation statistics, for example, the minimum p-value (Derkach et al., 2014); for a review of this topic see Owen (2009). Focusing on Fisher’s method, the corresponding test statistic is

$$W_F = -2(\log(p_L) + \log(p_S)).$$

For independent observations with no group uncertainty, Soave et al. (2015) showed that, under $H_{0_{joint}}$ of (9) and a Gaussian model, $p_L$ and $p_S$ are independent. Thus, $W_F$ is distributed as a $\chi^2$ random variable.

In the presence of sample correlation with group uncertainty, we propose to use the same framework but obtain $p_L$ from a generalized location test (e.g., a GLS approach to model (2), where the design matrix $X$ includes the group probabilities, and the covariance matrix, $\Sigma_{stage} = \sigma^2_0 \Sigma_0$, incorporates the sample correlation), and $p_S$ from the gS test proposed here. We show that the assumption of independence between $p_L$ and $p_S$ continues to hold theoretically under $H_{0_{joint}}$ of (9) for normally distributed outcomes (Web Appendix B), as well as empirically for approximately normally distributed outcomes in finite samples (Web Figure 1).

3. Simulations

The validity of the generalized joint location-scale (gJLS) testing procedure relies on the accuracy of the individual generalized location ($gL$) test and generalized scale ($gS$) test components. The performance of the $gL$ test has been established in the literature, therefore, our simulation studies here focused on evaluation of the proposed gS test, and when appropriate compared it with Levene’s original test ($Lev$) and the TW test of Iachine et al. (2010). For completeness, we also numerically demonstrate the power improvement of the proposed gJLS test. We use subscripts $OLS$ and $LAD$ to denote if the stage 1 regression was performed using OLS to obtain group-mean-adjusted residuals or LAD for group-median-adjusted residuals. Implementation details of each of
the six tests ($\text{LevOLS}$, $\text{LevLAD}$, $\text{TWOLS}$, $\text{TWLAD}$, $g\text{SOLS}$, $g\text{SLAD}$) is outlined in Web Appendix C.

We considered two main simulation models. Simulation model 1 followed the exact simulation setup of Iachine et al. (2010) to ensure fair comparison. Simulation model 2 extended model 1 by introducing genotype groups for each individual as well as group membership uncertainty. To apply the original $Lev$ test for comparison, we ignored the inherent sample correlation in the presence of correlated data. In all simulations, empirical type 1 error and power were evaluated at the 5% significance level using 10,000 replicates, unless otherwise stated.

3.1. Simulation Model 1

3.1.1. Model setup. Following the exact simulation study design of Iachine et al. (2010), we simulated outcome values for $n_1$ MZ and $n_2$ DZ twin pairs, $n = 2n_1 + 2n_2$, and we tested if the variance of the outcome differed between the two groups of pairs, that is, $\sigma_1^2 = \sigma_2^2$. To study robustness, we simulated outcomes using Gaussian, Students $t_4$ (heavier tailed), and $\chi^2_4$ (non-symmetric) distributions.

We first generated pairs of observations from independent bivariate normal distributions $BV\mathcal{N}(0,1, \rho_k), k = 1,2$, with $\rho_1$ and $\rho_2$ corresponding to the correlation within the MZ and DZ twin pairs, respectively. Let $w$ be the variable for an observation, we then applied a transformation $g(\cdot)$ to $w$ to obtain the desired marginal distribution, $y = \sigma_0 g(w)$, where the $\sigma_0$’s induced different variances between the two groups. The choice of $g(\cdot)$ depended on the desired distribution for $y$:

$$g(w) = \begin{cases} w, & \text{if } y \sim \mathcal{N}(0,1) \\ F^{-1}_t(\Phi(w)), & \text{if } y \sim t_4 \\ F^{-1}_4(\Phi(w)), & \text{if } y \sim \chi^2_4 \end{cases},$$

where $\Phi$, $F_t$, and $F^*_4$ are the cumulative distribution functions for the standard normal, Students $t_4$ and $\chi^2_4$ distributions, respectively.

We varied the sample size ($n_1, n_2 = 5, 10, \text{ or } 20$ for small samples, and $= 500, 1000, \text{ or } 2000$ for large samples, and $n_1 \text{ may or may not equal } n_2$), and group variances ($\sigma_1^2, \sigma_2^2 = 1, 2, \text{ or } 4$). The level of correlation within the MZ and DZ twin pairs was $\rho_1 = 0.75$ and $\rho_2 = 0.5$, respectively.

3.1.2. Results. We were able to replicate the simulation results of Iachine et al. (2010) that studied $\text{LevOLS}$, $\text{LevLAD}$, $\text{TWOLS}$, and $\text{TWLAD}$ (Table 1 and Web Table 1). However, we noticed that results reported in their article for $\text{LevLAD}$ and $\text{TWLAD}$ presumably using median-adjusted residuals (labeled as $W_{50}$ and $TW_{50}$, columns 9 and 12 of Tables 1–4 in Iachine et al. (2010)) were in fact switched with the $Lev$ and $TW$ results obtained using 10% trimmed mean-adjusted residuals (labeled as $W_{10}$ and $TW_{10}$ in Iachine et al. (2010)). Subsequent conclusions in Iachine et al. (2010) that the $TW$ method using the 10% trimmed mean “performed best”, therefore, are incorrect and should instead refer to $\text{TWLAD}$ using median-adjusted residuals from the stage 1 regression.

### Table 1

| $n_1$ | $n_2$ | $\text{LevOLS}$ | $\text{LevLAD}$ | $\text{TWOLS}$ | $\text{TWLAD}$ | $g\text{SOLS}$ | $g\text{SLAD}$ |
|------|------|-----------------|-----------------|----------------|----------------|----------------|----------------|
|      |      |                  |                  |                |                |                |                |
|      | 20   | 0.102            | 0.087            | 0.055          | 0.044          | 0.058          | 0.046          |
| 5    | 20   | 0.115            | 0.071            | 0.085          | 0.041          | 0.099          | 0.049          |
| 10   | 20   | 0.112            | 0.091            | 0.085          | 0.064          | 0.075          | 0.054          |
| 5    | 10   | 0.114            | 0.079            | 0.118          | 0.079          | 0.092          | 0.054          |

Six different tests were evaluated, including the original Levene’s test, $Lev$, the twin test of Iachine et al. (2010), $TW$, and the proposed generalized scale test, $gS$, with subscripts $\text{OLS}$ and $\text{LAD}$ denoting whether the stage 1 regression was performed using OLS or LAD. Parameter values included $n_1$ and $n_2$ for the number of MZ and DZ twin pairs, respectively, and $\rho_1 = 0.75$ and $\rho_2 = 0.5$ for the corresponding within-pair correlations. Without loss of generality, $\sigma_1^2 = \sigma_2^2 = 1$ for type 1 error rate evaluation. The empirical type 1 error was estimated from 10,000 simulated replicates at the nominal 5% level.

Our results in Table 1 clearly show that

- In the presence of sample correlation, Levene’s original method $Lev$ that ignores the correlation had severely increased type 1 error rate, even with Gaussian data. That is, $TW$ and $gS$ performed better than $Lev$.
- When the error structure was non-symmetric ($\chi^2_4$) or the group sizes were small (e.g., $n_1$ or $n_2$ less than 20), using OLS in the stage 1 regression for either $TW$ or $gS$ led to increased type 1 error. That is, $TW_{LAD}$ and $gS_{LAD}$ performed better than $TW_{OLS}$ and $gS_{OLS}$, respectively.
- When the group sizes were unbalanced and small (e.g., $n_1 = 10, n_2 = 20$), $TW_{LAD}$ had increased type 1 error, even with Gaussian data. That is, $gS_{LAD}$ performed better than $TW_{LAD}$.

In large samples, the original $Lev$ test remained problematic with empirical $\alpha = 0.097$ when $n_1 = n_2 = 2000$ even for Gaussian data (Web Table 1). The accuracy of both $TW_{LAD}$ and $gS_{LAD}$ increased as $n$ increased, with empirical $\alpha = 0.052$ when $n_1 = n_2 = 2000$ even for the non-symmetric $\chi^2_4$ data. The accuracy of both $TW_{OLS}$ and $gS_{OLS}$ also improved as $n$ increased, however, only for symmetric Gaussian or $t_4$ data. For $\chi^2_4$ data, their empirical $\alpha$ level remained as high as 0.103 even for $n_1 = n_2 = 2000$; this empirical result is consistent with Theorem 1 (Web Appendix A).
Because most of the six tests did not have good type 1 error control in the presence of sample correlation, small samples, unbalanced group sizes, or non-symmetric data, we delay the discussion of power until simulation model 2 below where we focus on methods comparison between TWLAD and gSLAD, and in a more general simulation set-up.

3.2. Simulation Model 2

3.2.1. Model setup. The second simulation setup was motivated by genetic association studies as previously discussed. We again considered sibling pairs to introduce sample correlation. However, unlike simulation model 1, here we allowed individuals from the same pair to belong to different groups, where the groups were the three different genotypes of a SNP.

Consider a SNP of interest with minor allele frequency (MAF) of $q$ ($= 0.2$ or $0.1$), we first simulated genotypes for $n/2$ ($= 20, 50, 100, 500, or 1000$) pairs of siblings. To account for the inherent correlation of genotypes between a pair of siblings, we started with drawing the number of alleles shared identical by decent (IBD), $D = 0, 1, or 2$, from a multinomial distribution with parameters $(0.25, 0.5, 0.25)$, independently for each sib-pair. Given the IBD status $D$, we then simulated paired genotypes $(G_1, G_2) = (i, j)$, $i, j \in \{0, 1, 2\}$, following the known conditional distribution of $(G_1, G_2|D)$ (Thompson, 1975; Sun, 2012). The distribution depends on $q$ in a way that smaller $q$ leads to greater imbalance in the genotype group sizes. The probabilities of the numbers of individuals with genotype $G = 0, 1,$ and $2$ are $(1 - q)^2$, $2q(1 - q)$, and $q^2$, respectively.

To introduce group membership uncertainty, we converted the simulated true genotypes $G$‘s to probabilistic data $X$’s using a Dirichlet distribution. We used scale parameters $a$ for the correct genotype category and $(1 - a)/2$ for the other two; this error model was used previously by Acar and Sun (2013) to study location tests under genotype group uncertainty. We varied $a$ from 1 to 0.5, where $a = 1$ corresponds to no genotype uncertainty and $a = 0.5$ implies that, on average, 50% of the “best-guess” genotypes correspond to the true genotypes. Thus, the genotype group uncertainty level ranged from 0 to 50% in our simulations.

We then simulated outcome data for each sib-pair similarly as for model 1 above. For each of the $n/2$ sib-pairs, we first simulated paired data from $BVN(0, 1, \rho)$, where $\rho = 0.5$ was the within-sib-pair correlation. For each simulated value $w$, we then applied the $\sigma_g(w)$ transformation to obtain the desired outcome data $y$ as in simulation model 1 (Gaussian, Student’s $t_4$, and $\chi^2_4$). However, $k$ here refers to the corresponding true underlying genotype group of an individual, and two individuals from the same sib-pair might not have the same genotype. We used $(\sigma^2_g, \sigma^2_k, \sigma^2_\rho) = (1, 1, 1)$ to study type 1 error control, and $(1, 1.5, 2)$ or $(1, 2, 4)$ to study power; other values such as $(2, 1.5, 1)$ and $(2, 2, 1)$ were also investigated.

It is evident from the earlier simulation results of model 1 that the original Lev test is not valid in the presence of sample correlation, and TWLAD and gSLAD are inferior, respectively, to $TWLAD$ and $gSLAD$ when the error structure is non-symmetric or the group sizes are small. Therefore, the results presented below focus on comparison between $TWLAD$ and $gSLAD$. In the presence of genotype group uncertainty, we also considered the “best-guess” approach and used $TWLAD$ and $gSLAD$ to represent the corresponding results.

For completeness, we also compared the power of $gSLAD$ along-side $gL$ and $gJLS$ using sibling data with genotype uncertainty. To do this, after simulating the genotype data for sib-pairs as above ($a = 0.7$ for genotype uncertainty at 70%), we then induced both scale differences $(\sigma^2_g, \sigma^2_k, \sigma^2_\rho) = (1, 1.5, 2)$ and location differences $(\mu_0, \mu_1, \mu_2) = (0, 0.3, 0.6)$ in the outcome between the three genotype groups; other parameter values were also considered (results not shown). As before, the outcome simulation used the true underlying genotype group membership for each individual, while the test statistics were constructed using either “best-guess” genotypes or incorporating the group membership probabilities as proposed. To numerically demonstrate the loss in power when limiting analysis to independent observations, we also analyzed the “best-guess” genotype dataset after randomly discarding one individual from each sibling pair.

3.2.2. Results. In the presence of sample correlation but with no group uncertainty, the results in Table 2 show that both $TWLAD$ and $gSLAD$ were accurate in large samples, for example, when sample size was 2000 ($n/2 = 1000$ sib-pairs). However, $TWLAD$ had increased type 1 error when group sizes were unbalanced and relatively small, even for Gaussian data. For example, when the MAF is $q = 0.2$ and the number of sib-pairs is $n/2 = 100$, the expected number of observations for the three genotype groups are $n*(1-q)^2$, $2q(1-q)$, $q^2$. In that case, the empirical type 1 error of $TWLAD$ was 0.060, 0.072, and 0.078 for Gaussian, $t_4$, and $\chi^2_4$, respectively. The problem was exacerbated by a smaller MAF $q = 0.1$ with empirical type 1 error levels of 0.092, 0.115, and

| $n/2$ | TWLAD | gSLAD | TWLAD | gSLAD | TWLAD | gSLAD |
|-------|-------|-------|-------|-------|-------|-------|
| 20    | 0.110 | 0.040 | 0.109 | 0.042 | 0.113 | 0.044 |
| 50    | 0.117 | 0.043 | 0.110 | 0.046 | 0.160 | 0.044 |
| 100   | 0.092 | 0.048 | 0.115 | 0.049 | 0.118 | 0.047 |
| 500   | 0.056 | 0.048 | 0.068 | 0.047 | 0.070 | 0.052 |
| 1000  | 0.055 | 0.050 | 0.061 | 0.049 | 0.058 | 0.045 |

Parameter values included $n/2$ for the number of sib-pairs, $\rho = 0.5$ for the within-pair correlations, and $q = 0.1$ or 0.2 for the minor allele frequency (MAF) of the SNP of interest; on average the expected sizes of the three genotype groups are $n(1-q)^2$, $n2q(1-q)$ and $nq^2$. Without loss of generality, $\sigma^2_g = \sigma^2_k = \sigma^2_\rho = 1$ for type 1 error rate evaluation. The empirical type 1 error was estimated from 10,000 simulated replicates at the nominal 5% level.
0.118, respectively, for the three types of data. In contrast, the proposed gSLAD test remained accurate in most cases and was slightly conservative in small samples when \( n/2 < 100 \).

Results in Table 3 are characteristically similar to those of Table 2. However, we note that group uncertainty somewhat mitigates the problem of unbalanced group sizes, and consequently the accuracy issue of TWLAD. Nevertheless, it is clear that gSLAD had better type 1 error control than TWLAD across the MAF values and the three outcome distributions. As expected, under the null hypothesis TWBG and gSBG using the “best-guess” genotype group have similar type 1 error rates, which mitigates the problem of unbalanced group sizes, and consequently the accuracy issue of TWLAD. Within each method, the estimated power increases as the analysis uses more information available from the data; this

Superscript \( ^BG \) denotes TWLAD and gSLAD applied to the “best-guess” genotype data. The true genotype data were masked using a Dirichlet distribution for the genotype probabilities with scale parameters \( a \) for the correct genotype and \( (1 - a)/2 \) for the other two. On average, \( a = 0.7 \) corresponds to 30% group uncertainty level. See legend of Table 2 for additional simulation details.

### Table 3

| n/2 | TWBG LAD | TWLAD | gSBG LAD | gSLAD | TWBG LAD | TWLAD | gSBG LAD | gSLAD | TWBG LAD | TWLAD | gSBG LAD | gSLAD |
|-----|----------|-------|----------|-------|----------|-------|----------|-------|----------|-------|----------|-------|
| 20  | 0.067    | 0.074 | 0.036    | 0.037 | 0.083    | 0.079 | 0.044    | 0.046 | 0.088    | 0.090 | 0.047    | 0.050 |
| 50  | 0.066    | 0.062 | 0.045    | 0.047 | 0.076    | 0.062 | 0.046    | 0.046 | 0.084    | 0.076 | 0.049    | 0.053 |
| 100 | 0.058    | 0.057 | 0.045    | 0.046 | 0.064    | 0.059 | 0.047    | 0.046 | 0.072    | 0.069 | 0.051    | 0.049 |
| 500 | 0.057    | 0.054 | 0.054    | 0.052 | 0.056    | 0.053 | 0.052    | 0.048 | 0.055    | 0.052 | 0.050    | 0.048 |
| 1000| 0.051    | 0.055 | 0.052    | 0.054 | 0.053    | 0.050 | 0.052    | 0.049 | 0.054    | 0.052 | 0.049    | 0.047 |

MAF = 0.1

| n/2 | TWBG LAD | TWLAD | gSBG LAD | gSLAD | TWBG LAD | TWLAD | gSBG LAD | gSLAD | TWBG LAD | TWLAD | gSBG LAD | gSLAD |
|-----|----------|-------|----------|-------|----------|-------|----------|-------|----------|-------|----------|-------|
| 20  | 0.061    | 0.062 | 0.040    | 0.039 | 0.065    | 0.063 | 0.037    | 0.044 | 0.075    | 0.073 | 0.047    | 0.050 |
| 50  | 0.053    | 0.053 | 0.046    | 0.045 | 0.063    | 0.059 | 0.046    | 0.050 | 0.069    | 0.070 | 0.051    | 0.053 |
| 100 | 0.049    | 0.051 | 0.046    | 0.045 | 0.057    | 0.053 | 0.047    | 0.049 | 0.059    | 0.058 | 0.049    | 0.047 |
| 500 | 0.051    | 0.049 | 0.049    | 0.051 | 0.051    | 0.052 | 0.048    | 0.050 | 0.053    | 0.052 | 0.048    | 0.051 |
| 1000| 0.049    | 0.046 | 0.047    | 0.047 | 0.052    | 0.049 | 0.047    | 0.049 | 0.055    | 0.053 | 0.050    | 0.052 |

MAF = 0.2

### Table 4

| n/2 | gSBG LAD | gSLAD | gSBG LAD | gSLAD | gSBG LAD | gSLAD |
|-----|----------|-------|----------|-------|----------|-------|
| 20  | 0.064    | 0.066 | 0.067    | 0.077 | 0.050    | 0.064 |
| 50  | 0.079    | 0.087 | 0.077    | 0.081 | 0.089    | 0.089 |
| 100 | 0.124    | 0.152 | 0.087    | 0.112 | 0.101    | 0.117 |
| 500 | 0.495    | 0.613 | 0.314    | 0.420 | 0.376    | 0.442 |
| 1000| 0.795    | 0.885 | 0.533    | 0.671 | 0.634    | 0.759 |

MAF = 0.1

| n/2 | gSBG LAD | gSLAD | gSBG LAD | gSLAD | gSBG LAD | gSLAD |
|-----|----------|-------|----------|-------|----------|-------|
| 20  | 0.050    | 0.066 | 0.062    | 0.058 | 0.063    | 0.074 |
| 50  | 0.089    | 0.120 | 0.084    | 0.089 | 0.091    | 0.104 |
| 100 | 0.166    | 0.196 | 0.114    | 0.129 | 0.129    | 0.160 |
| 500 | 0.668    | 0.784 | 0.471    | 0.582 | 0.499    | 0.608 |
| 1000| 0.939    | 0.985 | 0.739    | 0.846 | 0.810    | 0.896 |

\( gSBG \) denotes gSLAD applied to the “best-guess” genotype data. The true genotypes were masked using a Dirichlet distribution for the genotype probabilities with scale parameters \( a \) for the correct genotype and \( (1 - a)/2 \) for the other two. On average, \( a = 0.7 \) corresponds to 30% group uncertainty level. Besides the parameters shown in the table, other values include \( \rho = 0.5 \) for within-pair correlation, and \( (\sigma_1^2, \sigma_2^2, \sigma_3^2) = (1, 1.5, 2) \). Power was estimated from 1000 simulated replicates at the 5% level.
was already demonstrated for the proposed $gS$ in the earlier simulations (e.g., Table 3) and for $gL$ in the existing literature. A direct comparison between $gL$ and $gS$ can be difficult to interpret, because the individual power depends on two different sets of parameters, $(\mu_0, \mu_1, \mu_2)$ and $(\sigma_0^2, \sigma_1^2, \sigma_2^2)$. However, it is clear that Fisher’s method combines the evidence from $gS$ and $gL$ in a cumulative fashion resulting in $gJLS$ being more powerful than either test alone.

4. Applications
To demonstrate the utility of the proposed generalized scale ($gS$) test ($gS_{LAD}$ using LAD in the stage 1 regression) and subsequent generalized joint location-scale ($gJLS$) test, we revisited the two genetic association studies considered in Soave et al. (2015), and compared our results with those using only a sample of unrelated individuals with no genotype group uncertainties. We also used application data combined with simulation methods to further empirically validate the performance of the proposed methods.

4.1. HbA1c Levels in Subjects with Type 1 Diabetes
We use this application to demonstrate the gain in power by incorporating group uncertainty (probabilistic) data. Details of this dataset were previously reported in Soave et al. (2015). Briefly, the outcome of interest was inverse normal transformed HbA1c levels in $n = 1304$ unrelated subjects with type 1 diabetes, and the SNP of interest was rs1358030 near SORCS1 on chromosome 10 with MAF of 0.36. With no sample correlation or group uncertainty, the original $Lev$ test...
There were 1,313 singletons, 94 sib-pairs, and two sib-trios in the whole sample, resulting in 1409 independent individuals and \( n_{\text{all}} = 1313 + 94 \times 2 + 2 \times 3 = 1507 \) individuals. In addition to the eight genotyped SNPs originally analyzed in Soave et al. (2015), three imputed SNPs (rs11240600, rs11134081, and rs62605921) were also included in the current analysis. Results using the \( n_{\text{indep}} \) sample were from Soave et al. (2015) for the eight genotyped SNPs, or obtained here for the three imputed SNPs (using the “best-guess” genotypes), where the standard regression Location test, Levene’s Scale test, and the JLS joint location-scale test were used. Results for the \( n_{\text{all}} \) sample were obtained using the corresponding generalized tests, \( gL, gS, gJLS \), incorporating all available information.

### Table 5

**Application study of lung function severity in 1,507 patients with cystic fibrosis**

| Chr | Gene | SNP          | bp-Positiona  | MAF  | Location | Scale | JLS  | \( gL \) | \( gS \) | \( gJLS \) |
|-----|------|--------------|--------------|------|----------|-------|------|--------|--------|----------|
| 1   | SLC26A9 | rs7512462   | 204,166,218  | 0.41 | 0.30     | 0.58  | 0.48 | 0.30   | 0.39   | 0.36     |
| 1   | SLC26A9 | rs4077468   | 204,181,380  | 0.42 | 0.53     | 0.61  | 0.69 | 0.45   | 0.59   | 0.62     |
| 1   | SLC26A9 | rs12047830  | 204,183,322  | 0.49 | 0.55     | 0.15  | 0.29 | 0.52   | 0.11   | 0.22     |
| 1   | SLC26A9 | rs7419153   | 204,183,932  | 0.37 | 0.50     | 0.06  | 0.14 | 0.73   | 0.09   | 0.24     |
| 1   | SLC26A9 | rs11240600b | 204,187,369  | 0.33 | 0.14     | 0.62  | 0.30 | 0.11   | 0.65   | 0.27     |
| 5   | SLCA93  | rs17563161b | 550,624      | 0.26 | 0.0004   | 0.02  | 0.0001 | 0.0002 | 0.02   | 5.6 \times 10^{-5} |
| 5   | SLCA93  | rs11134081b | 557,404      | 0.35 | 0.0006   | 0.17  | 0.001 | 0.0006 | 0.05   | 0.0003   |
| X    | SLCA14  | rs12839137  | 115,479,578  | 0.24 | 0.02     | 0.08  | 0.01 | 0.01   | 0.16   | 0.02     |
| X    | SLCA14  | rs9056283   | 115,479,909  | 0.49 | 0.009    | 0.07  | 0.005 | 0.005  | 0.18   | 0.007    |
| X    | SLCA14  | rs3788766   | 115,480,867  | 0.40 | 0.001    | 0.01  | 0.0002 | 0.0004 | 0.02   | 9.5 \times 10^{-5} |
| X    | SLCA14  | rs62605921b | 115,475,499  | 0.24 | 0.02     | 0.14  | 0.02 | 0.01   | 0.22   | 0.01     |

There were 1,313 singletons, 94 sib-pairs, and two sib-trios in the whole sample, resulting in \( n_{\text{indep}} = 1.313 + 94 + 2 = 1409 \) unrelated individuals, and \( n_{\text{all}} = 1313 + 94 \times 2 + 2 \times 3 = 1507 \) individuals. In addition to the eight genotyped SNPs originally analyzed in Soave et al. (2015), three imputed SNPs (rs11240600, rs11134081, and rs62605921) were also included in the current analysis. Results using the \( n_{\text{indep}} \) sample were from Soave et al. (2015) for the eight genotyped SNPs, or obtained here for the three imputed SNPs (using the “best-guess” genotypes), where the standard regression Location test, Levene’s Scale test, and the JLS joint location-scale test were used. Results for the \( n_{\text{all}} \) sample were obtained using the corresponding generalized tests, \( gL, gS, gJLS \), incorporating all available information.

\( \text{a}^{\text{hg18 assembly (March 2006; NCBI36).}} \)

\( \text{b}^{\text{The three imputed SNPs, imputed using the Beagle software version 4.1 (Browning and Browning, 2016). The imputation quality is measured by allelic R}^2 \text{ provided by Beagle, and the linkage disequilibrium is measured by r}^2 \text{ with the adjacent genotype SNP.}} \)

\( n_{\text{indep}} = 1409, n_{\text{all}} = 1507 \)

was applied and resulted in a significant result with \( p = 0.01 \) (Soave et al., 2015). Combined with other evidence reported in Paterson et al. (2010), we assume here that the association is real and smaller p-values imply better performance.

To demonstrate the effect of genotype group uncertainty, we masked the true genotypes of rs1358030 using the same Dirichlet distribution as in the simulation studies above, where the value of \( a \) ranged from 1 to 0.5, corresponding to no group uncertainty to 50% uncertainty. We then applied \( gS^{BG} \) to the “best-guess” data and the proposed \( gS \) incorporating the probabilistic data, and obtained the corresponding p-values, \( p_{gS^{BG}} \), and \( p_{gS} \). For a given uncertainty level, we repeated the masking process independently 1000 times and obtained averaged p-values on the log10 scale (10\( \text{average of log10(p)} \)), \( p_{gS^{BG}} \), and \( p_{gS} \). Between the two methods, it was clear that \( gS \) was more efficient than \( gS^{BG} \). For example, when \( a = 0.75 \) for 25% group uncertainty, the \( gS \) test remains significant with \( p_{gS} = 0.048 \) as compared to \( p_{gS^{BG}} = 0.068 \). However, regardless of the method used, the power of the scale tests decreased sharply as genotype uncertainty increased, consistent with results for location tests reported in Acar and Sun (2013).

In addition to simulating group uncertainty for genotyped rs1358030, we also analyzed some ungenotyped variants that were imputed in the region surrounding rs1358030. Imputation was done using the IMPUTE2 software (Howie et al., 2009, 2011); see Paterson et al. (2010) for imputation details. Variants were chosen to include different levels of imputation quality (based on “info score” provided by IMPUTE2) and correlation/linkage disequilibrium (LD as measured by \( r^2 \)) with rs1358030, the original SNP of interest. Web Table 7 presents the results of \( gL, gS, \) and \( gJLS \) using the “best-guess” genotype or incorporating the genotype probabilities. As expected, variants in high LD/correlation with rs1358030 were imputed with high quality (i.e., with little genotype uncertainty), thus they yielded similar results between the two approaches (e.g., rs5787660). In general, we obtained smaller p-values for analysis using the genotype probabilities as compared to the “best-guess” genotypes, with larger differences for SNP of lower imputation quality. However, as discussed earlier, an alternative SNP starts to behave like a null SNP when the genotype uncertainty is too high leading to no power regardless of the methods. These agree with the earlier observations and simulation results in Figure 1.

### 4.2. Lung Disease Severity in Individuals with Cystic Fibrosis

We used this application to demonstrate the gain in power by incorporating all available information including related subjects and genotype probabilities. We also used this dataset combined with permutation methods to further demonstrate the validity of the proposed methods. Details of this dataset were previously reported in Soave et al. (2015). The outcome of interest was a measure of lung function severity based on forced expiratory volume in 1 second, obtained on a total of \( n_{\text{all}} = 1507 \) individuals with CF (1313 singletons, 188 from 94 sib-pairs, and six from two sib-trios).

Focusing on the \( n_{\text{indep}} = 1313 + 94 + 2 = 1409 \) unrelated individuals and eight genotyped SNPs, Soave et al. (2015) performed an association study using the original Location, Scale, and joint location-scale (JLS) tests. These SNPs were from three genes (SLC26A9, SLC9A3, and SLC6A14), chosen
based on association evidence for other CF-related outcomes as reported in Sun et al. (2012) and Li et al. (2014). Soave et al. (2015) concluded that SNPs from SLC9A3 and SLC6A14 were associated with CF lung disease, and their results are included in Table 5.

Here, we used all $n_{sd} = 1507$ related individuals, and we not only re-analyzed the eight genotyped SNPs, but also investigated three imputed SNPs previously not studied, using the generalized tests, $gL$, $gS$, and $gJLS$ (Table 5). The imputation was previously done using Beagle version 4.1 (Browning and Browning, 2016); see Li et al. (2014) for details. The imputed variants were chosen such that each of the three genes contained a new variant and the imputation quality was reasonably high. Since our analysis involved only (known) siblings and singletons, we used a compound symmetric correlation structure (a single correlation parameter $\rho$) to model within family dependence for each application of the GLS regression for the $gS$ component.

We first note that the conclusions for the presumed null SNPs from SLC26A9 did not change, as desired. The conclusions for the presumed associated SNPs from SLC9A3 and SLC6A14 did not change either, but using all available data led to overall smaller $p$-values for the generalized tests. The apparent lack of a large efficiency gain was somewhat disappointing, but it was also expected given the few number of siblings ($n_{sam} = 94 + 2 \times 2 = 98$) added to the full sample; see Section 5 for additional comments. Nevertheless, the application clearly demonstrates the advantage of using all the available information (all samples and the genotype probabilistic data). For example, for the imputed rs11134081 variant, the $p$-value of the original Levene’s Scale test is $0.17$ analyzing the smaller sample of independent individuals and their “best guess” genotypes, while the $p$-value of the $gS$ test is $0.05$. Lastly, we note that the $JLS$ framework indeed yields increased power when aggregating evidence from the individual tests; see Soave et al. (2015) for detailed discussions of the motivation and merits of the joint-testing framework.

To further examine the accuracy of the proposed $gS$ and $gJLS$ tests (as well as the $gL$ test for completeness), we generated 10,000 permutation replicates of the outcome to assess the empirical type 1 error control. Permutation was performed separately between singletons and between sib-pairs; see Abney (2015) for permutation techniques for more general family data. Without loss of generality, we focused on SNP rs17563161 from SLC9A3 (Web Figure 1). Testing the resulting $p$-values for deviation from the expected Uniform(0,1) distribution using the Kolmogorov–Smirnov test showed that all tests were valid.

5. Discussion

Levene’s scale test is widely used as a model diagnostic tool in linear regression, and more recently it has been employed as an indirect test for interaction effects. Increased data complexity due to sample correlation or group uncertainty, however, limits its applicability. Here, we proposed a generalization of Levene’s scale test, $gS$, that has good type 1 error control in the presence of sample correlation, small samples, unbalanced group sizes, and non-symmetric outcome data. We showed that the least absolute deviation (LAD) regression approach to obtain group-median-adjusted residuals is needed to ensure robust performance of $gS$. Based on our results, we recommend the use of $gSLAD$ over $gSO LS$ (and other existing tests) uniformly for all studies analyzed.

In the presence of group membership uncertainty, $gS$ incorporating the probabilistic data increases power compared to using the “best-guess” group data. However, based on the simulations considered here, we note that when the group uncertainty level is moderate (e.g., 30%), the efficiency gain is also moderate (Table 4 and Figure 1). When the group uncertainty is too high, the relative efficiency gain may diminish because the absolute power decreases considerably and eventually converges to the type 1 error rate.

In the presence of sample correlation, the original Lev test is inadequate due to inflated type 1 error. Using a subset of only unrelated individuals would improve the accuracy of Lev but at a cost to the power. The size of the efficiency loss depends on the proportion omitted from the sample as well as the dependency structure. The TW method of Iachini et al. (2010) extends the Lev test for twin data. Their simulation study as well as ours showed that TW has an increased type 1 error rate when group sizes are unbalanced and relatively small, in contrast to the proposed $gS$. When all group sizes were large, $gS$ and TW were empirically equivalent.

To further study the effect of misspecification of sample correlation structure on type 1 error and power, we conducted an additional simulation study considering three types of misspecifications. Without loss of generality, we revisited simulation model 1 with 20 MZ and 20 DZ twin pairs of Gaussian data, and we focused on the $Lev_{LAD}$, $TW_{LAD}$, and $gS_{LAD}$ tests. Web Table 6 clearly shows that in the absence of sample correlation, methods ($TW_{LAD}$ and $gS_{LAD}$) that model correlation retain correct type 1 error and have negligible power loss. In the presence of correlation, methods ($Lev_{LAD}$) that ignore correlation are not robust as demonstrated before (cf. Tables 2 and 4 in Iachini et al. (2010)). However, if the true correlation structure is misspecified as considered here, $TW_{LAD}$ and $gS_{LAD}$ can have increased type 1 error rates. Although this type of complete misclassification is unlikely in many practical settings such as the genetic association studies, the potential detrimental effects of correlation misspecification merit further investigation.

In the CF application, although $gS$ yielded comparable or less significant results after incorporating siblings in the analysis, we observed that the corresponding $gL$ test results were more significant. We considered the possibility that even though scale differences existed in the data, the addition of only 98 siblings (7% increase from the independent sample) may not yield a noticeable improvement of $gS$. Using the setup of simulation model 1, we examined the effect of incorporating only a small proportion of additional related subjects to an otherwise independent sample (Web Table 4). We found that, compared with using a sample of 1000 singletons, using $n = 900$ singletons along with 100 sib-pairs (10% increase) led to a <5% power increase. In contrast, the addition of siblings to all unrelated subjects provided a substantial increase in power (Web Table 4). These results, and the noticeable power gain from the $gL$ test when applied to the same CF data, are consistent with observations in genetic association studies that, larger samples are needed to detect variance compared to mean differences (Visscher and Posthuma, 2010).
The expression of $E(d) = \sigma_i \sqrt{\frac{\pi}{2} (1 - h_i)}$ in Section 2.2 suggests that the stage 2 regression of (4) could be improved by rescaling the $d_i$'s by $(1 - h_i)^{-1/2}$. This adjustment has been shown to improve the type 1 error control of Levene's original test for small samples with group design imbalance (Keyes and Levy, 1997). Examination of this rescaling for $gS_\text{LAD}$ under simulations involving correlated data, however, led to instances of increased type 1 error (results not shown). Thus, further investigation is required to propose an appropriate adjustment. Another potential improvement to the analysis of regression model (4) is from the recognition that the $d_i$'s are based on residuals, $\hat{\epsilon}_i$'s, and thus correlated even when there is no sample correlation among the true disturbances, $\epsilon_i$'s, as the group sizes increase. For the complex data scenarios considered here, $gS_{\text{LAD}}$ appears robust for even small samples. Nevertheless, the potential for gain in efficiency by accounting for this type of correlation merits additional consideration.

The developments here did not consider additional covariates, $z$, for example, age and sex in genetic association studies. The extension is straightforward if the effects of $z$ on $y$ are strictly on the mean. In that case, including $z$ as part of the design matrix in stage 1 suffices. However, if $z$ also influences the variance of $y$, not including $z$ as part of the design matrix in stage 2 may lead to increased type 1 error of testing the $\gamma_j$'s that are associated with the primary covariates of interest. This is the same phenomenon as observed in location-testing where omitting potential confounders can lead to spurious association.

Joint location-scale testing is becoming a popular method for complex outcome-covariate association data, where the conventional location-only analysis may be underpowered. This scenario has received attention in many fields, including our motivating example of genetic epidemiology (Soave et al., 2015). The proposed $gS$ test allows investigators to combine evidence from scale tests with existing generalized location tests via the $JLS$ testing framework of (Soave et al., 2015). The CF application study showed that individual location or scale tests can provide more significant results when utilizing related individuals and incorporating the available group probabilistic data, which in turn may lead to a more powerful $gJLS$ test.

6. Supplementary Materials

Web Appendix Sections A, B, and C, Web Figure 1, and Web Tables S1–S7 referenced in Sections 2, 3, 4, and 5 are available with this article at the Biometrics website on Wiley Online Library. R code is available at https://github.com/dsoave/gJLS.

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