Modeling and Testing for Joint Association Using a Genetic Random Field Model

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SUMMARY. Substantial progress has been made in identifying single genetic variants predisposing to common complex diseases. Nonetheless, the genetic etiology of human diseases remains largely unknown. Human complex diseases are likely influenced by the joint effect of a large number of genetic variants instead of a single variant. The joint analysis of multiple genetic variants considering linkage disequilibrium (LD) and potential interactions can further enhance the discovery process, leading to the identification of new disease-susceptibility genetic variants. Motivated by development in spatial statistics, we propose a new statistical model based on the random field theory, referred to as a genetic random field model (GenRF), for joint association analysis with the consideration of possible gene–gene interactions and LD. Using a pseudo-likelihood approach, a GenRF test for the joint association of multiple genetic variants is developed, which has the following advantages: (1) accommodating complex interactions for improved performance; (2) natural dimension reduction; (3) boosting power in the presence of LD; and (4) computationally efficient. Simulation studies are conducted under various scenarios. The development has been focused on quantitative traits and robustness of the GenRF test to other traits, for example, binary traits, is also discussed. Compared with a commonly adopted kernel machine approach, SKAT, as well as other more standard methods, GenRF shows overall comparable performance and better performance in the presence of complex interactions. The method is further illustrated by an application to the Dallas Heart Study.

KEY WORDS: Complex interaction; Genetic association; Linkage disequilibrium; Multi-marker test; Pseudo-likelihood; Random field.

1. Introduction

With the advance of high-throughput technologies, high-dimensional genetic data have been widely used in association studies for the identification of genetic variants contributing to common complex diseases. While a large number of genetic variants have been revealed today to be individually associated with complex diseases, they only explain a small proportion of heritability (Manolio et al., 2009). Complex diseases are likely influenced by the joint effect of genetic variants through complex biology pathways, given the fact that genes are the functional sets. However, the multiple testing problem occurs when one considers a set of single locus analyses, which dramatically diminishes power. Therefore, the joint analysis of a functional set of genetic variants simultaneously can further enhance the discovery process, leading to the identification of new genetic variants associated with complex diseases (Chatterjee et al., 2006). While the conventional linear or logistic regression models can easily be used for joint association analyses, they are subject to several issues, such as multiple-collinearity, when dealing with a large ensemble of dense genetic markers. The exponentially increased number of parameters also makes them impractical to model two-way or high-order interactions among a large number of genetic variants (Ritchie et al., 2001).

Several new statistical methods have been recently developed for joint association analysis, including the kernel machine-based method (well known as SKAT) (Wu et al., 2010, 2011) and the similarity regression (SIMreg) (Tzeng et al., 2009). Both methods significantly reduce the number of regression parameters, making it feasible and computationally efficient to handle high-dimensional variants. In addition, they account for linkage disequilibrium (LD) and potential interactions, which further improve performance. Both SKAT and SIMreg can be thought of as being developed from the general idea that, if genetic association exists, then genetic similarity leads to trait similarity, which is also the intuition behind our method.

In this article, we propose a random field framework for modeling and testing for the joint association of multiple genetic variants. We view outcomes as stochastic realizations of random vectors and propose to use a random vector field model, referred to as a genetic random field model (GenRF), to model the joint association. This approach is motivated by development in spatial statistics where outcomes are viewed as stochastic realizations of a random field on a Euclidean space (Cressie, 1993). This perspective leads to a very distinctive model from the aforementioned methods; specifically, GenRF regresses the response of one subject on responses of all other subjects. GenRF can be understood from the intuition that genetic similarity leads to trait similarity if variants are associated with the trait. Under the GenRF model, testing for the joint association reduces to a test involving a scalar parameter. Using the pseudo-likelihood method, a test for the joint association is developed, which
enjoys many appealing features as SKAT and can achieve comparable or better performance than existing methods, as demonstrated by simulation studies in Section 3 and a real data application in Section 4. Much of the development is focused on quantitative traits and robustness of the test to other traits, for example, binary traits, is also discussed.

There is a long history of applying spatial statistical methods to the analysis of genetic data (e.g., Molitor, Marjoram, and Thomas, 2003a, 2003b; Thomas, et al., 2003; De Iorio and Verzilli, 2007). For example, De Iorio and Verzilli (2007) used a spatial probit model to account for the local spatial correlation between variants physically close for fine-scale mapping of disease genes. Molitor et al. (2003b) used spatial clustering techniques for fine-scale gene mapping. Probably, the most closely related work to ours is Molitor et al. (2003a), which used a spatial auto-regressive model for analysis of haplotype effects and gene mapping. It differs from our work in two ways. First, it is haplotype-based whereas ours is genotype-based. Second, it is developed from the Bayesian framework where the trait is related to haplotypes through a linear model and the spatial model is used to model the prior distribution of haplotype effects, whereas our method directly models traits using a spatial model via a frequentist approach. In this article, we focus on multi-marker association testing and the direct spatial modeling of traits using a frequentist approach leads to a test that is analytically tractable and easy to implement.

2. Method

2.1. Genetic Random Field Model

Consider a study where $n$ subjects are sequenced in a region of interest. For subject $i$, $i = 1, \ldots, n$, let $G_i$ denote the genotype for the $p$ variants within the region, $Y_i$ the trait or phenotype, and $X_i$ the other covariates including, for example, demographic and environmental factors. We are interested in studying the joint association between variants $G_i$ and trait $Y_i$, possibly adjusted for the effect of $X_i$.

As SKAT and SIMreg, our method is also motivated by the general idea that, if the genetic variants are jointly associated with a trait, then the genetic similarity across subjects will contribute to the trait similarity. To put it in another way, if variants are jointly associated with the trait, then the response of a subject would be close to the response of other subjects who share similar genetic and possibly other variables. Based on this key idea, we propose to directly model the response of each subject as a function of all other responses and the contribution of other responses to $Y_i$ is weighted by their genetic similarity.

For simplicity, we temporarily assume $Y_i$’s are centered (have mean zero) and there are no other adjustment covariates. Specifically, based on the idea discussed above, we model the conditional distribution of $Y_i$ given all other responses as

$$Y_i | Y_{-i} \sim \gamma \sum_{j \neq i} s(G_i, G_j) Y_j + \varepsilon_i,$$  \hspace{1cm} (1)

where $Y_{-i}$ denotes responses for all other subjects except $Y_i$; $s(G_i, G_j)$ is known weights, weighting the contribution of $Y_j$ on approximating (or predicting) $Y_i$ via their genetic similarity; $\gamma$ is a non-negative coefficient measuring the magnitude of the overall contribution, further discussed below; and $\varepsilon_i$’s are random errors. A proper weight function $s(G_i, G_j)$ gives higher value when the two subjects are more similar in terms of genetic variants and, as discussed below, can be viewed as a measure for proximity of two subjects in a genetic space. The random errors $\varepsilon_i$’s are assumed to be independent and identically distributed with normal $(0, \sigma^2)$; extension to distributions other than normal is discussed in Section 2.2.

A main distinction between model (1) and the usual regression is that (1) models the conditional distribution of $Y_i$ given traits of other subjects, whereas in the usual regression one models the conditional distribution of a subject’s traits given his/her genetic variants. Intuitively, model (1) states that the trait of a subject can be approximated by traits of other subjects who are similar in genetic variants, if variants are associated with the trait. The coefficient $\gamma$ indicates the magnitude of the trait similarity as a result of genetic similarity. Thus, $\gamma$ can also be interpreted as a measure for the magnitude of the joint association of $G_i$ with $Y_i$. Specifically, if $G_i$ is not associated with $Y_i$, then regardless of how similar subject $i$ is to other subjects in terms of their genetic variants, the trait $Y_i$ is independent of all other $Y_j$’s for $j \neq i$; that is, $\gamma = 0$. On the contrary, if $G_i$ is strongly associated with $Y_i$, then one may expect $Y_i$ can largely be predicted by traits of subjects having the same or similar genetic variants and a large $\gamma$ indicates a strong joint association. Therefore, we can test the joint association of genetic variants with the trait by testing a null hypothesis involving a single parameter, that is, $H_0 : \gamma = 0$.

Models like (1), where responses are regressed on responses themselves, are referred as auto-regressive models and are commonly used in spatial statistics. In this article, we view the trait as a random field on a genetic space, and from this perspective, model (1) is formally a conditional auto-regressive (CAR) model (Cressie, 1993). A random field is a generalization of the notation of a stochastic process (Adler and Taylor, 2007). Informally, a stochastic process is a set of random variables indexed by integers or real numbers. A random field can be defined in more general spaces with the index set being an Euclidean space of dimension greater than one or other spaces. For example, in spatial statistics, crop yields of regions can be viewed as a random field defined in a two-dimensional space. Regions that are closer in location have more similar crop yields if spatial correlation exists. For our problem, we may view observed traits as realizations of a random field defined in a $p$-dimensional space of the $p$ genetic variants; that is, corresponding to each “location” in the $p$-dimensional genetic space, there is a random response variable associated with it. Similarly, responses from locations that are “closer” in the genetic space are expected to be more similar if the genetic association exists. In this sense, our model is a generalization of the auto-regressive model in spatial statistics. Models like (1) were firstly studied in the seminar work of Besag (1974) for random fields and we will term our model (1) as a genetic random field (GenRF) model. As a matter of fact, the GenRF model is closely related to the CAR model in spatial statistics; that is $s(G_i, G_j)$ analogously defines the proximity of neighbor $G_i$ to $G_j$ and $\gamma$ is the counterpart of a spatial dependence parameter. However, we note that the
usual tests of spatial dependence, for example, the Cliff-Ord test (Cliff and Ord, 1972) and the Lagrange multiplier test (Burridge, 1980), do not apply in our setting to test for the joint association of variants. The reason is that the matrix S, defined below, in our GenRF model does not satisfy the regularity condition usually assumed in spatial statistics for deriving the asymptotic distribution, as each subject has infinite neighbors in the genetic space.

We have yet to define a measure for “closeness” in the genetic space. Suppose each component of G records the number of minor alleles in a single locus and takes on values 0, 1, 2, respectively, corresponding to AA, Aa, aa. Then a sensible measure for closeness or similarity is the so called identity-by-state (IBS) (Wu et al., 2010), defined as

\[ s(G_i, G_j) = \sum_{k=1}^{p} (2 - |G_{ik} - G_{jk}|). \]

That is, the IBS measures the number of alleles in the region of interest shared by two individuals; for example, for p = 1, s(AA, AA) = 2, s(Aa, aa) = 1, s(AA, aa) = 0. Other measures for closeness in the genetic space rather than IBS are also possible, for example, the other kernel functions discussed in Wu et al. (2010), providing flexibility in our GenRF model. Similar to SKAT, our GenRF model can also incorporate weights for the joint association of variants. The reason is that the matrix \( L \) is known, one may estimate \( \gamma \) by the maximum pseudo-likelihood method. The estimator for \( \gamma \) can be obtained by minimizing \( \sum_{i=1}^{n} (Y_i - \beta^T X_i - \gamma \sum_{j \neq i} s(G_i, G_j)(Y_j - \beta^T X_j))^2 \), which in matrix notation is equal to

\[ \{(I - \gamma S)(Y - X\beta)\}^T(I - \gamma S)(Y - X\beta). \]

The minimization leads to an estimator for \( \gamma \) given by

\[ \hat{\gamma} = \frac{(Y - X\beta)^T S(Y - X\beta)}{(Y - X\beta)^T S^2(Y - X\beta)}. \]

Intuitively, one expects that a large value of \( \hat{\gamma} \) would give us evidence to reject the null hypothesis that \( \gamma = 0 \). In practice, \( \beta \) in unknown. We propose to replace \( \beta \) by its least square estimator \( \hat{\beta} \) under the null hypothesis, that is, \( \hat{\beta} = (X^T X)^{-1} X^T Y \), which is unbiased for \( \beta \). Substitute \( \hat{\beta} \) into the expression for \( \hat{\gamma} \) and straightforward algebra leads to the final test statistic:

\[ \hat{\gamma} = \frac{Y^T BS BY}{Y^T BS^2 BY}. \]

where \( B = I - X(X^T X)^{-1} X^T \). Again a large value of \( \hat{\gamma} \) supports the rejection of the null hypothesis.

We next show how the p-value for testing \( \gamma = 0 \) can be obtained based on the test statistic \( \hat{\gamma} \); that is, we would like to calculate the probability of \( \hat{\gamma} \) greater than the observed value of the statistic under the null hypothesis. Suppose \( \eta \) is the observed value of the test statistic \( \hat{\gamma} \). Since \( BS^2 B \) is positive-definite, we have

\[ P_{\eta \mid 0}(\frac{Y^T BS BY}{Y^T BS^2 BY} > \eta) = P_{\eta \mid 0}(BY^T (S - \eta S^2) BY > 0) \]
As it is assumed that \( \varepsilon_i \sim N(0, \varepsilon^2) \), i.i.d. across \( i \), it follows that \( BY \sim N(0, \varepsilon^2 B^2) \) under the null hypothesis. On the other hand, the statistic \( \tilde{\gamma} \) in (6) is ancillary to \( \varepsilon^2 \) because \( \varepsilon^2 \) in the numerator and denominator cancels out. Therefore, the above equation becomes

\[
P_{\theta_0}\left((BY)^T(S - \eta S^2)BY > 0\right) = P\left(Z^T(S - \eta S^2)Z > 0\right),
\]

where \( Z \) is an \( n \times 1 \) random vector following \( N(0, B^2) \). Applying standard results on the distribution of quadratic form in normal random variables, we have

\[
Z^T(S - \eta S^2)Z \sim \sum_i \lambda_i \Phi_i,
\]

where \( \Phi_i \)’s are i.i.d random variables with \( \chi^2 \) distribution, and \( \lambda_i \) are the eigenvalues of \( B(S - \eta S^2)B \). The final \( p \)-value can be obtained by Davies’ exact method (1980) for the weighted summation of independent Chi-square variables.

The proposed test has several appealing properties. First, due to the analytical form of the test statistic, the computational burden is well controlled. Second, as \( \tilde{\gamma} \) in (6) is ancillary to \( \varepsilon^2 \), unlike SKAT, there is no need to plug in a consistent estimator for \( \varepsilon^2 \). Third, the proposed method improves power by exploiting LD and allowing for possible complex interactions among variants. LD can cause correlations between variants, especially when we consider nearby loci. Considering similarity in variants can naturally reduce the degree of freedom. In the extreme case where components of \( G \) are “perfectly correlated,” the similarity argument will consider the whole set as a single variable. In addition, genetic variants involved in the disease pathway are more likely to interact with each other than to contribute to risk individually, known as the epistatic variants effect. Specifying two-way interactions in a set of loci is a challenging high-dimensional problem and the situation gets even worse in modeling higher order interactions. Since GenRF does not directly model the relationship of \( G_i \) with \( Y_i \), the difficulty of modeling complex interactions are circumvented and the interaction effect is naturally incorporated through measuring genetic similarity. Finally, as SKAT, the GenRF test can boost power of testing rare variants by increasing their weights by specifying \( w_k \) appropriately for variant \( k \).

2.3. Robustness to Other Distributions

The derivation of the GenRF test given above is built on the normal distribution assumption. Asymptotically, the proposed test is robust to distributions other than normal with slight modification. Consider \( P_{\theta_0}((BY)^T(S - \eta S^2)BY > 0) \), where it is now assumed \( Y \) follows an arbitrary distribution with mean zero and possibly heteroscedastic variances. The random quantity \( (BY)^T(S - \eta S^2)BY \) is a quadratic form in \( BY \) (with mean 0) with matrix \( A = S - \eta S^2 \). Rotar (1974) proved that under sufficiently weak conditions on matrix \( A \) and for large \( n \), \( P_{\theta_0}((BY)^T(S - \eta S^2)BY > 0) \) is close to \( P_{\theta_0}(\tilde{Z}^T(S - \eta S^2)\tilde{Z} > 0) \), where \( \tilde{Z} \) follows \( N(0, \Sigma) \) with \( \Sigma \) being the covariance matrix of \( BY \). In addition, Gotze and Tikhomirov (1999) gave an upper bound on \( \sup_x \left| P_{\theta_0}(B^2 ABY < x) - P_{\theta_0}(\tilde{Z}^T\tilde{A}\tilde{Z} < x) \right| \). These properties lead to the robustness of the GenRF test, with minor modification, as long as \( BY \) has expectation zero under the null hypothesis, which is true since the least squares estimator \( X(X^TX)^{-1}X^TY \) is unbiased for the mean of \( Y \) regardless of the distribution of \( Y \). For example, for binary traits, \( \Sigma = BWB \), where \( W = \text{diag}(\mu_1(1 - \mu_1), \ldots, \mu_n(1 - \mu_n)) \) and \( \mu_i = \beta^T X_i \). Then

\[
\tilde{Z}^T(S - \eta S^2)\tilde{Z} \sim \sum_i \lambda_i \Phi_i,
\]

where \( \Phi_i \)’s are i.i.d random variables with \( \chi^2 \) distribution, and \( \lambda_i \) are the eigenvalues of \( W^{1/2}B(S - \eta S^2)BW^{-1/2} \). The final \( p \)-value can be also obtained by Davies’ exact method (1980). We comment that, as the score test in SKAT is of similar quadratic form, one would expect that SKAT may share this property as well.

Therefore, one can directly use the test statistic in (6) for binary traits or traits that have distributions other than normal and the test, with a minor modification on the null distribution considering heteroscedastic variances, would be asymptotically valid. Note, this test corresponds to a model where the trait mean is related to a linear predictor through an identity link and may seem unnatural for binary traits. However, we argue that the model is mostly viewed as a mean leading to a sensible test. We also note that the commonly used trend test for testing genetic associations in an additive genetic model can be developed from a linear model for the mean of a binary trait (Laird and Lange, 2011), and a linear model is used for testing genetic associations for a binary trait in Ballard, Cho, and Zhao, (2010) as well. We note that a possible practical issue for binary traits may arise in practice, that is, the estimated means \( \mu_i \) may be outside of \([0,1]\) and consequently \( \hat{W}^{1/2} \) is not well defined. In this case, a remedy is to truncate the predictions \( \hat{\mu}_i \) at 0 or 1. The practical issue may arise when covariates have a wide support and a very strong effect and is less of a concern otherwise, for example, when covariates are categorical. Certainly, studying other link functions, for example, the logit link, to avoid this practical problem is important in the future. The validity of the test, corresponding to an identity link, is further studied by simulations shown in Sections 1 and 2 of the supplemental materials.

3. Simulation Studies

We report results of several simulations, each based on 1000 Monte Carlo (MC) replicates, to evaluate the performance of the GenRF test, relative to existing methods including SKAT. Four sets of simulations are conducted to evaluate: (1) type-1 error rates under different minor allele frequencies (MAF) and sample sizes, (2) power for common variant analysis under different LD, interaction effect, and proportions of causal SNPs, (3) power under scenarios where the causal SNPs include rare variants, and (4) robustness of the GenRF test to different distributions of the response variable.

In the first set of simulations, we evaluated type-1 error rates using sample size \( n = 50, 100, 200, 500 \). Genotypes
for $p = 20$ loci without LD were simulated, with MAF for each locus 0.005, 0.01, 0.1, or 0.2. Responses were generated according to

$$Y_i = \varepsilon_i, \text{ where } \varepsilon_i \sim N(0, 1),$$

so that no genetic variant is associated with the trait.

In the second set of simulations, we evaluated power under scenarios varying in LD, interaction effects, or the proportions of causal SNPs, setting $p = 20$ and $n = 500$. To simulate LD, the 20 loci were evenly divided into two regions. For each region, the haplotype allele was simulated one by one with MAF 0.2 and correlation coefficient ($\rho$) between adjacent pairs of alleles equal to 0, 0.2, 0.4, and 0.8, respectively, for each scenario. Genotypes were then generated by summing up two haplotype vectors. This way, all the loci are positively correlated with others in the same region. Responses were generated according to

$$Y_i = 0.2G_{i,5} + 0.2G_{i,15} + \varepsilon_i, \text{ where } \varepsilon_i \sim N(0, 1).$$

That is, variants 5 and 15, belonging to different LD regions, are associated with the trait.

To generate data with complex interactions, we set MAF 0.2, and the LD parameter $\rho = 0.4$. Data were generated such that two-way interactions exist between $K$ ($K = 1, 2, 3$, or 4) pairs of alleles, with alleles in each pair belonging to the two different LD regions as described above. Responses were then generated according to the following model,

$$Y_i = 0.2 \sum_{k=1}^{K} G_{i,4k+4} G_{i,14k+4} + \varepsilon_i, \text{ where } \varepsilon_i \sim N(0, 1).$$

We see that these models contain only interactions but no main effect of each locus.

To examine the effect of causal proportion, we set MAF 0.2 and $\rho = 0.4$. For each MC data set, $K$ causal SNPs were randomly selected with $K = 1, 2, 3$, or 4, each corresponding to 5%, 10%, 15%, and 20% causal SNPs. Responses were then generated according to

$$Y_i = 0.15 \sum_{k=1}^{K} G_{i,k_0} + \varepsilon_i, \text{ where } \varepsilon_i \sim N(0, 1),$$

where $(G_{B_1}, \ldots, G_{B_K})$ are the selected causal SNPs.

Simulation set 2 has focused on common variants. The third set of simulations considered scenarios involving rare variants and the scenarios vary in proportions of causal variants. We set $p = 20$, $n = 500$, and $\rho = 0$. The 20 SNPs were divided into two regions, one with 16 rare variants (MAF 0.008) and one with 4 common variants (MAF 0.1). Note, the proportion of rare variants is chosen according to the Dallas Heart Study. Two scenarios were considered where traits were associated with: (1) rare variants only or (2) both common and rare variants. For each scenario, $K$ rare SNPs were causal with $K = 1, 2, 4, 6, 8, 10, 12, \text{ or } 14$, that is, $K \times 6.25\%$ SNPs in the rare region are causal. In the scenario that both rare and common variants are causal, we set one of the common SNP as causal additionally. The effect size $\beta$ was set to be a decreasing function of MAF with $\beta = 0.2 \times |\log_{10} \text{MAF}|$ as in Wu et al. (2011). Responses were generated according to the following model,

$$Y_i = \beta_1 \sum_{k=1}^{K} G_{i,4k} + \beta_2 G_{i,20} + \varepsilon_i, \text{ where } \varepsilon_i \sim N(0, 1),$$

where $\beta_1 = 0.2 \times |\log_{10} 0.008|$, $\beta_2 = 0$ for scenario 1 and $\beta_1 = 0.2 \times |\log_{10} 0.008|$, $\beta_2 = 0.2 \times |\log_{10} 0.1|$ for scenario 2.

We considered one additional scenario where the 500 subjects’ genotypes were simulated based on data from the Dallas Heart Study. For each MC data set, we randomly selected one gene, then we randomly choose 10%, 20%, ..., 80% causal variants from those rare variants with true MAF less than 1%. Traits were simulated by

$$Y_i = \sum_{k=1}^{K} \beta_{B_i} G_{i,B_k} + \varepsilon_i, \text{ where } \varepsilon_i \sim N(0, 1),$$

where $(G_{B_1}, \ldots, G_{B_K})$ are the selected causal variants and $\beta_{B_i} = 0.2 \times |\log_{10} \text{MAF}_{B_i}|$.

In the fourth set of simulations, we further evaluated the robustness of the GenRF test to distributions other than normal, specifically, exponential, binary and mixture normal distributions. Details on the simulation setup is described in the supplemental materials.

In terms of type-I error rates, we only evaluated the proposed GenRF test and SKAT. In both GenRF and SKAT, we adopted the IBS kernel and considered both weighted and unweighted (i.e., $w_i = 1$) versions; in the weighted version, Beta (1, 25) weight as in Wu et al. was used (2011). In addition to SKAT, we compared GenRF test to other more standard methods. For common variant scenarios, we included the principle component regression test (PCR) (Guadernan et al., 2007); the MinSNP test (Ballard et al., 2010), and the $F$-test in linear regression model including only main effects. For scenarios involving rare variants, the variable-threshold (VT) test (Price et al., 2010) was included.

Table 1 shows results for the first set of simulations with different MAF and sample sizes. The GenRF test achieves the type I error rate close to the nominal level. However, SKAT is conservative in some scenarios due to the estimation of nuisance parameters, especially when the sample size is small. Since the GenRF test is an exact test without asymptotic approximation under normal assumption, the type I error rate is better controlled.

Table 2 shows the power of various methods under common variant scenarios. The first part shows the effect of LD on power. When LD does not exist or is low, for example, $\rho < 0.4$, the three linear regression-based tests, PCR, MinSNP, and $F$-test, are more powerful as expected because the data were generated exactly from a linear model. Among them, the PCR and MinSNP can exploit LD and have increasing power when LD is higher. When LD is moderate or high, both the GenRF test and SKAT have higher or even substantially higher power than the other tests by borrowing information from other loci. The power of the GenRF test is comparable to that of SKAT.
Table 1
Type I error rate simulation results under different levels of MAF and sample size (1000 replicates)

| Methods   | MAF n | 50  | 100 | 200 | 500  | 50  | 100 | 200 | 500  |
|-----------|-------|-----|-----|-----|------|-----|-----|-----|------|
| GenRF     | 0.005 | 0.043| 0.049| 0.050| 0.045| 0.040| 0.043| 0.061| 0.048|
|           |       |     |     |     |      |     |     |     |      |
| GenRF.w   | 0.01  | 0.048| 0.056| 0.051| 0.045| 0.043| 0.046| 0.060| 0.046|
| SKAT      |       | 0.035| 0.051| 0.057| 0.057| 0.034| 0.046| 0.050| 0.039|
| SKAT.w    |       | 0.034| 0.050| 0.053| 0.059| 0.029| 0.042| 0.046| 0.035|
| GenRF     | 0.1   | 0.051| 0.049| 0.055| 0.044| 0.050| 0.058| 0.055| 0.049|
|           |       |     |     |     |      |     |     |     |      |
| GenRF.w   | 0.2   | 0.052| 0.052| 0.053| 0.048| 0.047| 0.051| 0.052| 0.046|
| SKAT      |       | 0.022| 0.039| 0.041| 0.041| 0.016| 0.030| 0.041| 0.043|
| SKAT.w    |       | 0.041| 0.035| 0.043| 0.054| 0.045| 0.043| 0.046| 0.044|

Each cell contains the type I error rate, that is, rejection rate when data are generated under the null model. GenRF, the unweighted genetic random field test; SKAT: the unweighted sequential kernel association test of Wu et al. (2011); GenRF.w, GenRF with Beta (1,25) weight as in Wu et al. (2011); SKAT.w, SKAT with Beta (1,25) weight.

The second part shows results when there are complex interactions between variants but no main effects. Note the LD structure is the same as that in part 1 with $\rho = 0.4$ in which the five methods have comparable power. Therefore, the power difference is mainly due to the complex interactions. In these scenarios, the linear regression-based methods has low power in detecting the joint association. Both GenRF test and SKAT attain much larger power. Moreover, the proposed GenRF test has larger power than SKAT in detecting the joint association effect when complex interactions exist.

The third part shows results when the causal proportion varies. Similarly, the LD parameter $\rho$ is set to be 0.4 to eliminate the impact of factors other than the causal proportion. Because MinSNP is based on single SNP analysis, the test is less powerful especially when causal proportion is high, that is 15% or 20%. GenRF and SKAT show comparable power in general, but GenRF performs better as causal proportion gets higher.

Table 2
Power simulation results for common variant analysis under different levels of linkage disequilibrium (LD), interaction effects, and causal proportion (1000 replicates)

| Method | Different level of LD | Number of two-way interactions | Different causal proportion |
|--------|-----------------------|--------------------------------|----------------------------|
|        | 0  | 0.2 | 0.4 | 0.8 | 1   | 2   | 3   | 4   | 5%  | 10% | 15% | 20% |
| GenRF  | 0.462 | 0.472 | 0.566 | 0.816 | 0.119 | 0.364 | 0.652 | 0.862 | 0.124 | 0.321 | 0.539 | 0.776 |
| SKAT   | 0.491 | 0.487 | 0.545 | 0.764 | 0.100 | 0.299 | 0.546 | 0.746 | 0.150 | 0.324 | 0.506 | 0.727 |
| PCR    | 0.495 | 0.467 | 0.518 | 0.676 | 0.119 | 0.268 | 0.470 | 0.657 | 0.159 | 0.308 | 0.473 | 0.679 |
| MinSNP | 0.570 | 0.507 | 0.543 | 0.656 | 0.098 | 0.252 | 0.408 | 0.576 | 0.180 | 0.342 | 0.463 | 0.624 |
| F-test | 0.545 | 0.514 | 0.524 | 0.538 | 0.112 | 0.231 | 0.394 | 0.562 | 0.145 | 0.278 | 0.471 | 0.665 |

GenRF, the unweighted genetic random field test; SKAT, the unweighted sequential kernel association test of Wu et al. (2011); PCR, the principle component regression test of Guaderman et al. (2007); MinSNP, the MinSNP test considered by Ballard et al. (2010); F-test, the F-test in linear regression.
Table 3
Power simulation results under scenarios involving rare variants with different proportion of causal variants (1000 replicates)

| Method     | 6.25% | 12.5% | 25%  | 37.5% | 50%  | 62.5% | 75%  | 87.5% |
|------------|-------|-------|------|-------|------|-------|------|-------|
| GenRF      | 0.045 | 0.073 | 0.139| 0.209 | 0.305| 0.466 | 0.593| 0.739 |
| SKAT       | 0.048 | 0.052 | 0.066| 0.073 | 0.086| 0.100 | 0.111| 0.126 |
| GenRF.w    | 0.062 | 0.087 | 0.212| 0.429 | 0.660| 0.848 | 0.950| 0.980 |
| SKAT.w     | 0.083 | 0.125 | 0.252| 0.368 | 0.515| 0.654 | 0.736| 0.814 |
| VT         | 0.065 | 0.082 | 0.128| 0.209 | 0.314| 0.487 | 0.680| 0.852 |
| F-test     | 0.080 | 0.113 | 0.190| 0.302 | 0.449| 0.556 | 0.670| 0.765 |
| GenRF      | 0.191 | 0.259 | 0.380| 0.501 | 0.625| 0.761 | 0.861| 0.927 |
| SKAT       | 0.274 | 0.281 | 0.287| 0.313 | 0.331| 0.359 | 0.387| 0.416 |
| GenRF.w    | 0.061 | 0.097 | 0.232| 0.434 | 0.646| 0.853 | 0.939| 0.981 |
| SKAT.w     | 0.078 | 0.155 | 0.277| 0.386 | 0.523| 0.631 | 0.732| 0.818 |
| VT         | 0.217 | 0.306 | 0.418| 0.504 | 0.603| 0.720 | 0.845| 0.930 |
| F-test     | 0.163 | 0.270 | 0.354| 0.477 | 0.618| 0.701 | 0.779| 0.843 |

Common and rare causal variants

| Method     | 10%  | 20%  | 30%  | 40%  | 50%  | 60%  | 70%  | 80%  |
|------------|------|------|------|------|------|------|------|------|
| GenRF      | 0.080| 0.140| 0.169| 0.247| 0.329| 0.414| 0.507| 0.600|
| SKAT       | 0.071| 0.089| 0.114| 0.117| 0.153| 0.191| 0.205| 0.271|
| GenRF.w    | 0.100| 0.204| 0.321| 0.434| 0.588| 0.696| 0.796| 0.875|
| SKAT.w     | 0.118| 0.196| 0.294| 0.330| 0.433| 0.544| 0.600| 0.688|
| VT         | 0.095| 0.159| 0.254| 0.359| 0.498| 0.612| 0.721| 0.827|
| F-test     | 0.147| 0.239| 0.355| 0.423| 0.528| 0.653| 0.721| 0.795|

DHS

| Method     | 10%  | 20%  | 30%  | 40%  | 50%  | 60%  | 70%  | 80%  |
|------------|------|------|------|------|------|------|------|------|

Rare causal variants: causal variants are rare only; common and rare causal variants: causal variants are both rare and common; DHS: scenario based on the Dallas Heart Study. GenRF.w, the genetic random field test with Beta (1, 25) weight as in Wu et al. (2011); SKAT.w, the sequential kernel association test with Beta (1, 25) weight; VT, the variable-threshold test of Price et al. (2010); other entries as in Table 2.

and the number of common variants is fixed at one, the results change dramatically. When the rare causal proportion is higher than 37.5%, the weighted GenRF and SKAT show higher power than the unweighted counterpart. Overall, for scenarios considered here, the GenRF test has very good performance relative to others.

Supplementary Tables 1 and 2 show the robustness of the GenRF test to distributions other than normal. In implementing GenRF and SKAT, the identity link is used in modeling the responses. The GenRF test achieves the type I error rate close to the nominal level even when the distribution of the response is not normal; the same holds for SKAT. Particularly, for binary traits, we evaluated the robustness of the GenRF test to heteroscedastic variances in Section 2 of the Supplementary Material. Valid type I error rates and reasonably good power are achieved. In addition, the remedy by truncation when predictions fall outside of the range of [0,1] works well in practice, even under extreme and possibly unrealistic scenarios.

4. Application
We applied our method to the Dallas Heart Study (Browning et al., 2004), a population-based, multi-ethnic study on 3551 subjects whose lipids and glucose metabolism were measured. In this study, 348 sequence variations in the coding regions of the four genes, ANGPTL3, ANGPTL4, ANGPTL5, and ANGPTL6 were discovered. Most of these variants (86%) are rare with MAF less than 1%. More information regarding the number of rare variants is shown in the Supplementary Material. Individuals who have diabetes mellitus, alcohol dependency, or have taken lipids lowering drugs were excluded as these factors may confound the interpretation of associations. Our final analysis was based on data on 2812 subjects after quality control steps.

We assessed the association between ANGPTL gene families and two traits, specifically high-density lipoprotein (HDL) and triglyceride, using the proposed GenRF test and SKAT, both with and without weighting. As in the simulation studies, the IBS kernel and the Beta (1, 25) weight were applied.
Analyses were also carried out using the more traditional methods including PCR, MinSNP, VT, and F-test. Our analysis were done for the non-synonymous variants, adjusted for gender and ethnicity.

The association between ANGPTL4 gene and the level of HDL and triglyceride was previously discovered by Romeo et al. (2007). In our analysis, both weighted GenRF and SKAT gave evidence for the ANGPTL4 and triglyceride association ($p$-values: 0.019 and 0.006). Among all the methods considered, only weighted SKAT showed marginal evidence for the association between ANGPTL4 and HDL ($p$-value: 0.040). One possible explanation is that the causal proportion of ANGPTL4 is low and SKAT performs better in this case as shown in simulation studies. Note that the weighted GenRF and SKAT uncovered these associations while the unweighted tests did not, possibly indicating the causal variants in ANGPTL4 might be rare (MAF $< 5\%$), or the effect size is negatively correlated with allele frequency. As for ANGPTL5, our analysis using GenRF provided evidence to support the association with HDL ($p$-value: 0.009 and 0.036 for weighted and unweighted analyses) while SKAT provided marginal evidence ($p$-value: 0.035 and 0.050). Note the unweighted tests gave larger $p$ values. Since all variants in ANGPTL5 are rare (MAF $< 5\%$), the result suggests that the causal variants might be the rare variants with relatively higher allele frequency. This finding was supported by standard approaches like MinSNP ($p$-value: 0.033), F-test ($p$-value: 0.051), and VT test ($p$-value: 0.051). More results are shown in Table 4.

Overall, for this study, GenRF performs comparably to SKAT and seems to perform better than the other more standard methods.

5. Discussion

We have proposed a novel framework for modeling and testing for the joint association of genetic variants with a trait from the perspective of viewing traits as a random field on a genetic space. The development has been focused on quantitative traits with a normal distribution. Based on the GenRF model, a test for genetic associations was developed and this test enjoys many appealing features. The GenRF test is based on testing a null hypothesis involving a single parameter, allowing it to exploit LD to improve power. When LD is moderate or high, our simulations showed that the GenRF test achieves much higher power than the more traditional regression-based methods. The GenRF model is flexible to allow for complex interaction effects and, as demonstrated by simulations, the GenRF test is even much more powerful than SKAT in the presence of complex interaction effects. Moreover, as SKAT, prespecified variant-specific weights can be incorporated to boost power for rare variants. Unlike SKAT, the GenRF test is an exact test under the normal assumption and thus not overly conservative in finite samples. Finally, the test is computationally easy to implement since an analytical form is available. In summary, the GenRF test is an appealing alternative to SKAT and other existing methods for testing the joint association of variants with a trait. It can achieve overall comparable performance and sometimes even much better
performance relative to SKAT as well as other methods.

Although we focus on quantitative traits, we note that the GenRF test is robust to distributions other than normal as discussed previously and demonstrated by simulation studies. Specifically for binary traits, although the GenRF model with an identity link function may seem a bit unnatural, the resulting test with a minor modification is still valid and can achieve good power. However, due to the conceptual difficulty associated with modeling binary traits using a linear model and the possible practical issue that can arise, it would be interesting to study, within the framework of random field model, other link functions for binary traits as well as other distributions in the future.

6. Supplementary Materials
Web Appendices referenced in Sections 2 and 3, and the R code implementing the method are available with this paper at the Biometrics website on Wiley Online Library.

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