

A powerful and adaptive association test for rare variants

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ABSTRACT

This article focuses on conducting global testing for association between a binary trait and a set of rare variants (RVs), though its application can be much broader to other types of traits, common variants (CVs) and gene set or pathway analysis. We show that many of the existing tests have deteriorating performance in the presence of many non-associated RVs: their power can dramatically drop as the proportion of non-associated RVs in the group to be tested increases. We propose a class of so-called **sum of powered score (SPU) tests, each of** which is based on the score vector from a general regression model, hence can deal with different types of traits and adjust for covariates, e.g. principal components accounting for population stratification. The SPU tests generalize the Sum test, a representative burden test based on pooling or collapsing genotypes of RVs, and a sum of squared score (SSU) test that is closely related to several other powerful variance component tests; a previous study (Basu and Pan 2011) has demonstrated good performance of one, but not both, of the Sum and SSU tests in many situations. The SPU tests are versatile in the sense that one of them is often powerful, though its identity varies with the unknown true association parameters. We propose an adaptive **SPU (aSPU) test to approximate the most powerful SPU test for a given scenario**, consequently maintaining high power and being highly adaptive across various scenarios. We conducted extensive simulations to show superior performance of the aSPU test over several state-of-the-art association tests in the presence of many non-associated RVs. Finally we applied the SPU and aSPU tests to the GAW17 mini-exome sequence data to compare its practical performance with some existing tests, demonstrating their potential usefulness.

Key words: aSPU test; GWAS; Score statistic; Sequencing data; SPU test; SSU test; Sum test.

1. INTRODUCTION

The recent advances in sequencing technologies have made it feasible to conduct global testing for association between complex traits and rare variants (RVs) (Bansal et al 2010). The most popular approach in genome-wide association studies (GWASs) is to test on each single nucleotide variant (SNV) one by one, then select the SNVs meeting a stringent significance level after adjusting for multiple testing. However, such a strategy may be low powered due to weak signal contained within each individual RV for its extremely low minor allele frequency (MAF). Hence, developing new association tests tailored to RVs has been an active research area in the last few years. Due to low MAFs of RVs, to achieve practically meaningful power, the majority of existing approaches focus on testing on a group of RVs, rather than on each individual RV (Capanu et al 2011); the main idea is to boost power through aggregating information across multiple RVs in an analysis unit, such as a gene (e.g., Morgenthaler and Thilly 2007; Li and Leal 2008; Madsen and Browning 2009; Liu and Leal 2010; Han and Pan 2010; Price et al 2010; Zhang et al 2010; Zhu et al 2010; Luo et al 2011; Neale et al 2011; Ionita-Laza et al 2011; Feng et al 2011; Pan and Shen 2011; Basu and Pan 2011; Gordon et al 2011; Wu et al 2011; Fan et al 2013). As theoretically shown (Cox and Hinkley 1974) and to be demonstrated in our simulations, there is no uniformly most power test for this purpose, which means that, depending on the unknown truth, including specific association effect directions and sizes, a given and fixed test may or may not be powerful. Hence, there has been intensive efforts in developing adaptive tests for RVs (e.g., Pan and Shen 2011; Lin and Tang 2011; Zhang et al 2011; Lee et al 2012; Chen et al 2012; Derkach et al 2012; Sun et al 2013). However, due to their limited extents of adaptivity (e.g. with a pre-determined and fixed set of the weights on RVs), these adaptive tests are still not flexible (or adaptive) enough with loss of power under some situations. A main motivation in this paper is to develop a broader family of association tests such that

at least one of them is powerful for a given situation. We will develop such a family of tests, called the sum of powered score (SPU) tests, which generalize the Sum (of score) test and the sum of squared score (SSU) test (Pan 2009). The Sum test is a representative of the burden tests based on genotype pooling or collapsing (Morgenthaler and Thilly 2007; Li and Leal 2008; Madsen and Browning 2009), whereas the SSU test is closely related to kernel machine regression (and its implementation for RVs, SKAT) (Wu et al 2010, 2011), C-alpha test (Neale et al 2011) and an empirical Bayes test for high-dimensional data (Goeman et al 2006); see Basu and Pan (2011). In many simulation set-ups, one, but not both, of the Sum test and SSU test has been shown to be powerful (Basu and Pan 2011). For example, with different association directions of causal RVs, the Sum test suffers from a loss of power, while the SSU test performs much better. However, we emphasize that, in analysis of multiple RVs, there exist non-associated RVs. For example, in cancer research it has been observed that the vast majority of RVs do not appear to confer risk (Capanu et al 2011). Hence, it is important to assess the performance of a test in the presence of non-associated RVs in the group of the RVs to be tested. In fact, as to be shown, the performance of the Sum test deteriorates rapidly as the number of non-associated RVs increases, whereas the SSU test is more robust but nevertheless may gradually become less competitive. It seems that the performance of various tests has not been fully investigated for the case with many non-associated RVs, including some new adaptive tests, such as a kernel-based adaptive clustering (KBAC) test (Liu and Leal 2010), a p-value weighted sum test (PWST) (Zhang et al 2011), an estimated regression coefficient (EREC) test (Lin and Tang 2011), an adaptive SSU (aSSU) test (Pan and Shen 2011) and an optimized SKAT (SKAT-O) test (Lee et al 2012). As to be shown, it turns out that these tests suffer more from substantial power loss and are no longer competitive in the presence of a high proportion of non-associated RVs. In contrast, regardless of the number of non-associated RVs, at least one of our

proposed SPU tests may remain relatively more powerful. Since the identity of the most powerful SPU test also changes with the unknown true association pattern with causal and non-causal RVs, we propose a simple yet highly adaptive SPU (aSPU) test to maintain high power across a wide range of scenarios. As to be shown, our proposed aSPU test is often much more powerful than many existing adaptive tests in the presence of many non-associated RVs.

We conducted extensive simulation studies to compare our newly proposed tests with several state-of-the-art tests, such as the PWST, EREC, aSSU and SKAT-O tests, which all appeared after the publication of and thus were not compared in Basu and Pan (2011). As an active research topic, quite a few association tests for RVs have been proposed in the last two or three years. However, unfortunately, most of them have not been fully compared to each other, especially in the presence of many non-associated RVs, which is expected to be a norm instead of an exception in analysis of RVs. Hence, as a second aim of this article, we assess the performance of these existing tests along with our newly proposed ones, offering some insights into their potential in practical use. Our study can be regarded as an update and follow-up to Basu and Pan (2011).

2. METHODS

2.1. Data and notation

Our proposed methods are based on general regression models, and thus can be applied to binary, quantitative and survival responses or traits in the framework of generalized linear models and Cox proportional hazards model while adjusting for covariates, such as environmental variables and principal components accounting for population stratification. To be concrete, we consider only the case-control study design with a binary response/trait and no covariates; more general cases can be

similarly approached, as to be shown in our example. Suppose that for subject $i = 1, \dots, n$, $Y_i = 0$ or 1 is a binary response or trait, e.g. an indicator of disease, and $X_i = (X_{i1}, \dots, X_{ik})'$ is a group of predictors of interest, such as k RVs from a candidate gene or region. We use additive coding for each RV; that is, X_{ij} is the count of the minor allele at RV j for subject i . Consider a logistic regression model:

$$\text{Logit}[Pr(Y_i = 1)] = \beta_0 + \sum_{j=1}^k X_{ij}\beta_j. \quad (1)$$

We'd like to test the null hypothesis $H_0: \beta = (\beta_1, \dots, \beta_k)' = 0$; that is, there is no association between any RVs and the trait under H_0 .

Many of the existing tests and our new tests are based on the score vector $U = (U_1, \dots, U_k)'$ for β in the logistic regression model (1) and $V = Cov(U|H_0)$ (Pan 2009; Basu and Pan 2011; Lin and Tang 2011; Wu et al 2011; Lee et al 2012):

$$U = \sum_{i=1}^n (Y_i - \bar{Y})X_i,$$

$$V = Cov(U|H_0) = \bar{Y}(1 - \bar{Y}) \sum_{i=1}^n (X_i - \bar{X})(X_i - \bar{X})',$$

where \bar{Y} and \bar{X} are the sample means of Y_i 's and X_i 's respectively. An advantage of using a score-based test is the closed form of the score vector and only a null model (i.e. the model under H_0) needs to be fitted, thus is computationally much faster, sometimes even only feasible, as compared to the corresponding Wald or likelihood ratio test, for which a more general and complicated model has to be fitted and may not even converge (e.g. when $k > n$). Furthermore, it is noted that the score vector U in the joint model (1) is the same as $U_M = (U_{M,1}, \dots, U_{M,k})'$ with $U_{M,j}$ being the score statistic for $\beta_{M,j}$ in the marginal model:

$$\text{Logit}[Pr(Y_i = 1)] = \beta_{M,0} + X_{ij}\beta_{M,j}, \quad (2)$$

where subscript M denotes parameters from a marginal model. In contrast, for example, in general, the maximum likelihood estimates (MLEs) $\hat{\beta}$ and $\hat{\beta}_M = (\hat{\beta}_{M,1}, \dots, \hat{\beta}_{M,k})'$

differ, and our experience suggests that the Wald test based on the marginal models is more powerful than that based on the joint model. The most popular single variant-based analysis corresponds to a minimum p-value (UminP) test combining univariate score tests for the marginal models:

$$T_{UminP} = \max_{j=1}^k U_j^2 / V_{jj},$$

where $V_{jj} = Var(U_j|H_0)$ is the j th diagonal element of V . To adjust for multiple testing, one could apply the conservative Bonferroni adjustment, or better, as implemented here, based on the asymptotic null distribution of $U \sim N(0, V)$ under H_0 , use numerical integrations (or simulations) to obtain an asymptotically exact p-value for the UminP test (Conneely and Boehnke 2007).

2.2. A brief review of some existing tests

Basu and Pan (2011) compared the performance of many existing association tests for RVs. Their major conclusion is that, if there is (nearly) a common association strength for causal RVs with no or few non-associated RVs, then the burden tests, such as the Sum test (Pan 2009), were most powerful; otherwise, the SSU test (Pan 2009) and its close relatives, kernel machine regression (KMR or SKAT) (Wu et al 2010, 2011) and C-alpha test (Neale et al 2011) performed best. The Sum test is based on a working assumption that in the joint logistic regression model (1), we have a common association parameter between the k RVs and the trait, say $\beta_1 = \dots = \beta_k = \beta_c$. Then we only need to test a null hypothesis with a **single parameter H_0 : $\beta_c = 0$** , corresponding to fitting a simple logistic regression model:

$$\text{Logit}[Pr(Y_i = 1)] = \beta_{c,0} + \sum_{j=1}^k X_{ij}\beta_c. \quad (3)$$

On the other hand, the SSU test is a **variance-component test**: assuming that β_1, \dots, β_k in model (1) are independent random effects with mean 0 and variance τ^2 , it can be derived as a score test on a null hypothesis with a single parameter H_0 : $\tau^2 = 0$.

Specifically, both the Sum test and SSU test are based on the score vector U :

$$T_{Sum} = 1'U = \sum_{j=1}^k U_j, \quad T_{SSU} = U'U = \sum_{j=1}^k U_j^2,$$

from which it is clear that the Sum test, as other burden tests, such as the CMC test (Li and Leal 2008) and the weighted Sum test (Madsen and Browning 2009), will lose its power if the causal RVs have different association directions, leading to different signs of U_j 's and thus a small test statistic T_{Sum} , failing to reject H_0 . In contrast, since the components of U is squared in the SSU test (and KMR and C-alpha test), the SSU test and its close relatives do not lose power with different association directions due to the sum over U_j^2 , instead of over U_j as in the Sum test. A more general score-based statistic can be written as

$$T_G = \zeta'U = \sum_{j=1}^k \zeta_j U_j,$$

where $\zeta = (\zeta_1, \dots, \zeta_k)'$ is a vector of weights for the k RVs (Lin and Tang 2011). For example, if $\zeta_j = -1$ or 1 depending on whether $\hat{\beta}_{M,j} < 0$ and its p-value < 0.1 , then T_G is the adaptive Sum (aSum) test of Han and Pan (2010). Two new tests that were not reviewed in Basu and Pan (2011) are also special cases of the above general test T_G . First, if $\zeta_j = 2(p_j - 0.5)$, where p_j is the p-value for a one-sided Wald test for $H_{j,0}: \beta_{M,j} = 0$ versus $H_{j,1}: \beta_{M,j} < 0$ with a test statistic $\hat{\beta}_{M,j} / \sqrt{V_{jj}}$, then T_G is the p-value weighted Sum test (PWST) of Zhang et al (2011). Second, if $\zeta_j = \hat{\beta}_{M,j} \pm d$, then the T_G test is the estimated regression coefficient (EREC) test of Lin and Tang (2011);



it is noted that, due to the instability of estimating $\beta_{M,j}$ for a RV, Lin and Tang (2011) proposed to shrink $\hat{\beta}_{M,j}$ towards a constant d or $-d$, with $d = 1$ for binary traits. Each of the above three adaptive tests accommodates different association directions by using the signs of $\hat{\beta}_{M,j}$'s, thus overcoming a main shortcoming of the Sum test, retaining high power in the presence of different association directions. Nevertheless, with RVs, for the same reason that motivates pooling or collapsing RVs

weight

in most association tests proposed so far, there is only limited information contained in each RV, implying that all the above weighting schemes may not work well under some situations, as to be elaborated later.

selection

In addition to differing association directions for causal RVs, a more common issue is the existence of many non-associated RVs among the group of RVs to be tested. In particular, with many non-associated RVs, as shown by Basu and Pan (2011), the burden tests, including the Sum test, lose their power quickly, while the SSU test and its close relatives perform much better. On the other hand, intuitively, if we can exclude non-associated RVs in constructing a test statistic, it may help improve the power. Along this line, Pan and Shen (2011) proposed a class of adaptive Neyman-type tests (Neyman 1937), including an adaptive Sum (aSum+) and an adaptive SSU (aSSU) test, which are based on RV selection, instead of weighting. Specifically, first, one orders the components of the score vector U in a descending order based on the magnitudes of U_j and U_j^2 respectively for the aSum+ and aSSU tests; second, suppose that the p-values for the Sum and SSU tests based on the first j components of the ordered U are $P_{Sum,j}$ and $P_{SSU,j}$ respectively, then the test statistics for the two adaptive tests are

$$T_{aSum+} = \min_{1 \leq j \leq k} P_{Sum,j}, \quad T_{aSSU} = \min_{1 \leq j \leq k} P_{SSU,j},$$

and the final p-values are obtained by permutations or simulations. In short, the aSum+ and aSSU tests work by selecting the first few components of a re-ordered score vector U that are most informative (with smallest p-values) while possibly ignoring other components of U for non- or weakly-associated RVs. In particular, the aSum+ test accounts for possibly different association directions by using only those positively associated RVs; however, it may suffer from power loss due to its ignoring those negatively associated RVs. To improve over the aSum+ test, Pan et al (2011) proposed an adaptive Sum test based on two directional searches, denoted as aSum2d, to use both positively and negatively associated SNVs, and found its

improved power in detecting gene-gene interactions for CVs. Specifically, we first re-order the components of U in a descending order of U_j as for the aSum+ test, and suppose that the p-value of the Sum test applied to the *last* j components of the re-ordered U is $P_{Sum,-j}$, then the aSum2d test statistic is

$$T_{aSum2d} = \min\{T_{aSum+}, T_{aSum-}\} \text{ where } T_{aSum-} = \min_{1 \leq j \leq k} P_{Sum,-j}.$$

We can then use permutations or simulations to obtain p-values for T_{aSum2d} (and T_{aSum-} if needed). If desired, one can also just use T_{aSum-} to test for only negatively-associated RVs.

Another adaptive test, called **kernel-based adaptive weighting** (KBAC), was proposed by **Liu and Leal** (2010). A unique feature of the KBAC test is to detect not only the main effects of, but also possible interactions among RVs. For the latter purpose, rather than weighting on each individual RV, it uses a kernel-based weight on each **unique pattern (or combination)** of the genotypes across the k RVs. It up-weights a genotype pattern that appears more **frequently in cases (i.e. with a higher risk of disease)**, and then contrasts the frequencies of genotype patterns between the case and control groups by taking a weighted sum of their frequency differences. As pointed out by Basu and Pan (2011), there are two **potential limitations**. First, since its test statistic is a weighted sum of the frequency differences between the case and control groups, the presence of opposite association directions may contribute to both positive and negative frequency differences, leading to a small test statistic and thus loss of power. Second, as the number of non-associated RVs increases, there will be a larger number of unique genotype patterns and thus a smaller number of subjects with each genotype pattern, leading to loss of power. These two points will be confirmed later.

2.3. A new **class of tests and a data-adaptive** test

Our basic observation is that, depending on the unknown pattern of association

effects of the group of RVs to be tested, different tests may be more powerful; in spite of the generality of the T_G test, its performance *critically* depends on the choice of the weights, and any fixed choice may or may not be most suitable. Hence our primary goal is to construct a class of versatile tests such that for a **given scenario, at least one of the tests is powerful**. Then we combine these tests to obtain a data-adaptive test that will maintain high power across a wide range of scenarios. For this purpose, we would like to choose **weight ζ_j as informative** and as simple as possible. Since most existing association tests use the score vector U , suggesting that most information is already contained in U , we would simply use U to construct weights. In particular, since we have $U \sim N(0, V)$ under H_0 , we know that a large $|U_j|$ offers strong evidence to reject $H_{0,j}$: $\beta_j = 0$. Specifically, we choose $\zeta_j = U_j^{\gamma-1}$ for an integer $\gamma \geq 1$, leading to a *sum of powered score* (SPU) test:

$$T_{SPU(\gamma)} = \sum_{j=1}^k U_j^\gamma. \quad (4)$$

With various values of $\gamma \geq 1$, we obtain a class of the SPU tests. The SPU tests cover the Sum and SSU tests as two special cases with a corresponding $\gamma = 1$ and $\gamma = 2$ respectively. Importantly, as γ increases, the $SPU(\gamma)$ test puts more weights on the larger components of U while gradually ignoring the remaining components. An extreme case is that, as an even number $\gamma \rightarrow \infty$, we have

$$T_{SPU(\gamma)} \propto \|U\|_\gamma = \left(\sum_{j=1}^k |U_j|^\gamma \right)^{1/\gamma} \rightarrow \|U\|_\infty = \max_{j=1}^k |U_j|.$$

As to be shown, since the SPU tests are based on resampling methods to calculate their p-values, they are invariant to any monotone transformation of their test statistics, such as $(\cdot)^{1/\gamma}$. That is, we can equivalently define $T_{SPU(\infty)} = \max_{j=1}^k |U_j|$, which uses only the largest component of $|U|$. More generally, as we increase the value of γ , we put higher and higher weights on the larger components of U , effectively realizing RV selection. On the other hand, an even integer of γ automatically eliminates the

effects of different signs of U_j 's, avoiding power loss of the Sum test in the presence of different association directions. However, an odd integer of γ might be more suitable, as in the SPU(1) or Sum test, when the associations are all in the same direction.

We know that under H_0 , the score vector U has an asymptotic Normal distribution $N(0, V)$. Hence, in theory, we can derive the asymptotic distribution of $T_{SPU(\gamma)}$, which however may not be easy to calculate. As an alternative, we recourse to permutations (Churchill and Doerge 1994). Specifically, we permute the original set of traits Y to obtain a new set of traits $Y^{(b)}$, based on which we calculate the score vector $U^{(b)}$ and the null statistic $T_{SPU}^{(b)} = T_{SPU}(U^{(b)})$; after $b = 1, \dots, B$ permutations, we calculate the p-value as $\sum_{b=1}^B [I(|T_{SPU}^{(b)}| \geq |T_{SPU}|) + 1] / (B + 1)$. We used $B = 200$ in our simulations for a nominal significance level at 5%.

In the presence of covariates, we propose generalizing the above permutation scheme. Specifically, first we regress Y on the covariates to fit a null model under H_0 to obtain $\hat{\mu}_{i,0} = \hat{E}(Y_i|H_0)$ and residual $r_i = Y_i - \hat{\mu}_{i,0}$; second, we permute the set of the residuals $r = \{r_i | i = 1, \dots, n\}$ to obtain a permuted set $r^{(b)}$; third, we calculate the new score vector based on the permuted residuals as $U^{(b)} = \sum_{i=1}^n X_i r_i^{(b)}$ and the corresponding null statistic $T_{SPU}^{(b)} = T_{SPU}(U^{(b)})$; after repeating the above steps for $b = 1, \dots, B$, we calculate the p-value as $\sum_{b=1}^B [I(|T_{SPU}^{(b)}| \geq |T_{SPU}|) + 1] / (B + 1)$.

Alternatively, we also propose using the parametric bootstrap: we will first fit a null model under H_0 to obtain $\hat{\mu}_{i,0} = \hat{E}(Y_i|H_0)$, then simulate a new set of traits $Y_i^{(b)} \sim \text{Bin}(1, \hat{\mu}_{i,0})$ for $b = 1, \dots, B$; we calculate the test statistic $T_{SPU}^{(b)}$ based on each set of simulated $Y^{(b)}$ and calculate the p-value as in the permutation method.

Since the power of a SPU(γ) test depends on the choice of γ while the optimal choice of γ depends on the unknown true association pattern of the RVs to be tested, it would be desirable to data-adaptively choose the value of γ . For this purpose, we propose an adaptive SPU (aSPU) test that simply combines the p-values of multiple SPU tests (with various values of γ), though other combining methods are also possi-

ble (Pan et al 2010; Cheung et al 2012). Suppose that we have some candidate values of γ in Γ , e.g. $\Gamma = \{1, 2, 3, \dots, 8, 15, 16, 31, 32, \infty\}$ as used in our later simulations, and suppose that the p-value of the $SPU(\gamma)$ test is $P_{SPU(\gamma)}$, then our combining procedure is to take the minimum p-value:

$$T_{aSPU} = \min_{\gamma \in \Gamma} P_{SPU(\gamma)}.$$

Of course, T_{aSPU} is no longer a genuine p-value; we use the permutation or parametric bootstrap to estimate its p-value. It may appear that a double permutation or bootstrap procedure is needed, but indeed not necessary. For example, if we use the permutation method, first, we permute the original set of traits to obtain $Y^{(b)}$ and the corresponding score vector $U^{(b)}$ for $b = 1, 2, \dots, B$. We then calculate the corresponding SPU test statistics $T_{SPU(\gamma)}^{(b)}$ and their p-values $p_{\gamma}^{(b)} = \sum_{b_1 \neq b} [I(T_{SPU(\gamma)}^{(b_1)} \geq T_{SPU(\gamma)}^{(b)}) + 1]/B$. Thus, we have $T_{aSPU}^{(b)} = \min_{\gamma \in \Gamma} p_{\gamma}^{(b)}$, and the final p-value of the aSPU test is $P_{aSPU} = \sum_{b=1}^B [I(T_{aSPU}^{(b)} \leq T_{aSPU}) + 1]/(B+1)$.

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We note the practicality of permutation- or other resampling-based methods for p-value calculations. First, due to extremely low MAFs of some RVs, it is always dubious whether asymptotic results are applicable. Second, it is computationally feasible to use permutation-based tests for genome-wide searches. In practice, we can first use a smaller B , say $B = 1000$, to scan a genome, then gradually and repeatedly increase B for a few groups of RVs that pass an initial significance criterion (e.g. p-value $< 5/B$) in the previous step; in this way, contrary to otherwise claimed, it is indeed feasible to apply a permutation-based test to genome scans and obtain highly significant results (if any). We have applied permutation- or simulation-based aSPU test to genome-wide scans to yield p-values $< 10^{-6}$.

Finally we comment on the choice of Γ . The following considerations guide the choice of the integers $\gamma \geq 1$ in Γ . First, in order to cover the burden and variance-component tests, which have been shown empirically to perform well under some situations (Basu and Pan 2011), we would include $\gamma = 1$ and $\gamma = 2$ in Γ . Second,

depending on whether the phenotype-RV association directions vary, we may need to use either even or odd integers γ 's to yield high power; if unsure, then it is suggested to use both odd and even integers γ 's. Third, depending on how sparse are the association signals, one may use smaller or larger γ 's. For example, the more the RVs to be tested and the fewer associated RVs to be expected, then larger γ 's would be desirable. As to be shown, we have found that often $\Gamma = \{1, 2, 3, \dots, 8, \infty\}$ suffices. For demonstration, we have included $\gamma = \infty$, which is not necessary, but may be beneficial when testing on CVs. In the following, we also show the results of the $\text{SPU}(\gamma)$ tests for $\gamma \in \Gamma$; we have two purposes. The first is to show varying operating characteristics of the various SPU tests. For example, we would like to show higher power of $\text{SPU}(3)$ or $\text{SPU}(4)$ than that of $\text{SPU}(1)$ and $\text{SPU}(2)$, demonstrating the power gain of using some $\text{SPU}(\gamma)$ test with $\gamma > 2$. Second, we will show that often $\text{SPU}(8)$ gives the results almost the same as those of $\text{SPU}(\infty)$, suggesting no need to use other larger γ 's. In practice, we suggest using the aSPU test that combines the strengths (and possibly weaknesses) of various SPU tests; the aSPU test can be regarded as a rigorous (and almost exact) means for multiple testing adjustment with the use of several SPU tests, while the results of the SPU tests may shed light on the underlying genetic architecture. For example, if a large γ gives the most significant p-value, it may indicate a high degree of signal sparsity; if some odd γ 's yield more significant results than even γ 's, then most or all of the large associations are in the same direction. More elaborately, as shown below, an analysis of a SPU test can also imply the relative contribution of each SNV to the aggregated association (if any).

2.4. SNV selection

A limitation of most global tests is their inability for variant selection: even if the global null hypothesis is rejected, they may not give any information on which RVs are (or are not) likely to be associated with disease. We note that the aSPU test can be used to rank the importance of the RVs. First, we estimate the optimal value

of $\gamma < \infty$, $\hat{\gamma} = \arg \min_{\gamma \in \Gamma} P_{SPU}(\gamma)$ chosen by the aSPU test. Second, we assess the relative contribution of each RV r to the aSPU test as $C_r = |U_r|^{\hat{\gamma}} / \sum_{j=1}^k |U_j|^{\hat{\gamma}}$. Third, we rank the RVs based on their C_r values, and we can select the top k_1 RVs such that the sum of their relative contributions $\sum_{r=1}^{k_1} C_r \geq \alpha_1$ with $\alpha_1 = 0.8$, say; the choice of α_1 determines the trade-off between increasing true positives and increasing false positives. Generally, we can use C_r to prioritize and generate hypotheses on the selection of causal RVs.

2.5. Further comments and extensions

Below we briefly comment on the advantages of the aSPU test over several other adaptive tests. First, since the power of any univariate test for a single RV may be low (which is exactly the reason why we would like to combine information across multiple RVs, e.g., through pooling or collapsing), the p-value of such a test may not be much informative; the aSum test and PWST based on such p-values may not perform well. Second, we note that the adaptive Neyman-type tests, such as the aSSU test, is based on the idea of variable or RV selection, while the SPU tests are more based on weighted averaging of variables or RVs. As discussed extensively in the model selection literature (Yuan and Yang 2005; Shen and Huang 2006) and in a genetic application (Newton et al 2007), if signals are strong enough, then model selection is expected to perform better; otherwise, model averaging is preferred. In our current context, again due to extremely low MAFs of RVs, no matter how strong is its association strength, there is only quite limited information contained within each individual RV. Thus we expect that the model averaging-based SPU tests to outperform the model selection-based aSSU or other adaptive Neyman-type tests. Third, we note that the EREC test is related to the SPU tests. As shown in Pan (2009), we have

$$\hat{\beta}_M = \text{Diag}(V)^{-1}U + O_p(1/n),$$

where $\text{Diag}(V)$ is a diagonal matrix with its j th diagonal element as V_{jj} . Hence, if

$|\hat{\beta}_M|$ is much larger than d , then $\zeta = \hat{\beta}_M \pm d \approx \hat{\beta}_M$, which is roughly proportional to U (if the MAFs of the RVs are in a close range), suggesting that the EREC test will be similar to the SPU(2), i.e. SSU test. On the other hand, if $|\hat{\beta}_M|$ is small relative to d , then $\zeta = \hat{\beta}_M \pm d \approx \pm d$, implying that the EREC test will behave similarly to the SPU(1) (or Sum) test. Generally, we expect that the EREC test behaves between the SPU(1) and SPU(2) tests.

Furthermore, several approaches (Lee et al 2012; Derkach et al 2012; Sun et al 2013), including SKAT-O, have been proposed to combine a burden test like SPU(1) and a variance-component test like SPU(2). In contrast, our proposed aSPU test is based on combining a broader set of tests including but beyond SPU(1) and SPU(2), hence is more flexible and adaptive. As to be shown, in the presence of many non-associated RVs, the weight $\zeta = 1$ or $\zeta = U$ may not suffice: we may need weights $U^{\gamma-1}$ with $\gamma > 2$. In other words, with many non-associated RVs, the power of the EREC or SKAT-O (or similar tests combining SPU(1) and SPU(2)) can be much lower than SPU(γ) test with a large γ , and lower than the aSPU test. In addition, since the aSum, PWST and EREC tests use the marginal estimates $\hat{\beta}_{M,j}$, which have to be obtained iteratively during each permutation, whereas the score vector U is much easier to obtain, the SPU tests are much faster.

In the presence of external or prior biological information, as for other tests, it may be helpful to incorporate some *external* weights (differing from ζ discussed earlier) into the SPU tests. Given some external weights w_j , we can have a weighted SPU (wSPU) test as

$$T_{wSPU(\gamma)} = \sum_{j=1}^k w_j U_j^\gamma,$$

while all other aspects, including the construction of an adaptive wSPU test, remain the same as before. For example, if it is believed that causal RVs tend to have lower MAFs, as advocated by Madsen and Browning (2009), one can use a w_j inversely proportional to the MAF of the j th RV. In this way, by suitably weighting both CVs

and RVs, it is possible to use the adaptive wSPU test for a joint analysis of CVs and RVs (Ionita-Laza et al 2013). Alternatively, w_j can be a predicted likelihood of the j th RV's being functional or deleterious based on some computational algorithms (Wei et al 2011). As other tests, the performance of the wSPU tests depends on how informative or correct the external weights are, while the choice of the external weights may not always be clear, hence we skip further discussions on the use of such external weights.

We have proposed using permutations (or the parametric bootstrap) to calculate the p-values for the SPU and aSPU tests. If the asymptotic normality of the score vector is expected to approximately hold, e.g. in analysis of CVs, we may replace the permutation or bootstrap with simulation-based methods, which will be much faster (Zou et al 2004; Seaman and Muller-Myhsok 2004).

3. RESULTS

3.1 Simulation set-ups

We conducted extensive simulation studies to evaluate and compare the performance of various tests. We simulated genotypes as in Wang and Elston (2008). First, a latent vector $Z = (Z_1, \dots, Z_k)'$ was generated from a multivariate Normal distribution $N(0, R)$, where R had a first-order auto-regressive (AR1) covariance structure with its (i, j) th element $R_{ij} = \text{Corr}(Z_i, Z_j) = \rho^{|i-j|}$; we used $\rho = 0$ and $\rho = 0.9$ to generate independent and correlated RVs respectively. Second, the latent vector Z was dichotomized to yield a haplotype with some specified MAFs, each of which was randomly selected from a uniform distribution between 0.001 and 0.01 during each simulation. Third, the above two steps were repeated to generate two independent haplotypes, which were then combined to obtain genotype $X_i = (X_{i1}, \dots, X_{ik})'$ for subject i . Fourth, for a non-null case we randomly chose k_1 causal RVs with their corresponding $\beta_j \neq 0$ while all other $\beta_j = 0$; for a null case, all $\beta_j = 0$. Fifth,

the disease status Y_i of subject i was generated from the logistic regression model (1). We used $\beta_0 = -\log(0.05/0.95)$ for a 5% background disease probability; that is, $Pr(Y_i = 1|X_i = 0) = 0.05$. Finally, as in a case-control study, we sampled $n/2$ cases and $n/2$ controls in each dataset.

We considered a few set-ups with combinations of various values of $\rho = 0$ or 0.9 , $k_1 = 8$ or 1 , and $n = 1000$. We varied the number of non-associated RVs $k - k_1$ between 0 and 128, and a range of possible values of $\beta_j \neq 0$ to cover from a common association effect to varying association strengths or directions, and from a single to multiple causal RVs.

Throughout the simulations, the test significance level was fixed at $\alpha = 0.05$. The results were based on 1000 independent replicates for each set-up. We compared the performance of the SPU tests with several state-of-the-art adaptive tests not reviewed in Basu and Pan (2011), including one based on a Bayesian hierarchical GLM (BhGLM) (Yi et al 2011). As a benchmark, we also included the UminP test that tests on each individual RV separately and then combine them by taking their minimum p-value.

3.2 Simulation results

To save space, we focus on a few cases with correlated RVs (i.e. neighboring RVs were in linkage disequilibrium). It is not only more general to consider correlated RVs (or covariates), but also in agreement with real sequence data as generated from the 1000 Genomes Project (Zhang et al 2013). The extensive simulation results with independent RVs and other association parameters were similar to those presented below and thus are relegated to Supplementary Materials.

First, all the tests maintained well controlled Type I error rates (Table 1). Second, we consider the case with non-zero β_j 's randomly drawn from a uniform distribution $U(1, 2)$, representing the association pattern with varying association strengths but all in the same direction (Table 2). Among the SPU tests, the SPU(1), SPU(2), SPU(4)

or $\text{SPU}(6)$ were the respective winners with no, a medium and a large number of non-associated RVs. This is in agreement with our analysis earlier that an increasing proportion of non-associated RVs requires a larger value of γ in the $\text{SPU}(\gamma)$ test to weed off the effects of non-associated RVs. In particular, the quickly deteriorating performance of the $\text{SPU}(1)$ (i.e. Sum) test was striking. Compared to some more powerful SPU tests, the UminP test was low powered because the UminP test used information from only the most significantly associated RV while ignoring other associated RVs. It is noted that a $\text{SPU}(\gamma)$ test with $8 \leq \gamma < \infty$ was only slightly more powerful than $\text{SPU}(\infty)$, suggesting no need to use $\gamma > 8$.

Among the adaptive tests, the aSPU test was the overall winner; its performance in the presence of many non-associated RVs was most impressive: for example, with 126 non-associated RVs, the power of the aSPU test was 0.811, much higher than 0.749 of SKAT (with a linear kernel used throughout), 0.658 of aSSU, 0.567 of EREC, 0.532 of aSum+, 0.380 of PWST, 0.331 of KBAC and 0.248 of BhGLM. It is noted that the power of the aSPU test was always close to that of the most powerful SPU test, whose identity however changed with the set-up. Although *all* other adaptive tests performed well with no or few non-associated RVs, they failed to do so otherwise. In particular, the aSSU test was much less powerful than the SSU test with many non-associated RVs, presumably due to the difficulty in RV selection with relatively weak signals with each individual RV.

Third, for the case with both varying association directions and effect sizes of the causal RVs (Table 3), among the SPU tests, as the number of non-associated RVs increased, $\text{SPU}(2)$, $\text{SPU}(4)$, $\text{SPU}(6)$ and $\text{SPU}(8)$ became most powerful respectively, and as expected, the $\text{SPU}(1)$ (i.e. Sum) test was the least powerful. For example, with 128 null RVs, the power of $\text{SPU}(1)$ and that of $\text{SPU}(2)$ were only 0.070 and 0.261 respectively, much lower than 0.370 of $\text{SPU}(7)$ and $\text{SPU}(8)$; accordingly, the power of aSPU was 0.329, much higher than 0.235 of SKAT and 0.195 of SKAT-O. It was also

confirmed that there was no need to use a $\text{SPU}(\gamma)$ test with $\gamma > 8$ to gain power. Among the adaptive tests, the aSPU test was the winner, though the PWST and SKAT were most powerful with no or only few non-associated RVs, but quickly lost their edge as the number of non-associated RVs increased. The aSSU test performed second best after the aSPU test, presumably due to easier RV selection with larger effect sizes of some causal RVs. The BhGLM, aSum and KBAC tests did not perform well in this case.

Similar results with higher significance levels α and with covariates were obtained as shown in Supplementary Materials.

3.3 Data example

We applied the methods to the mini-exome sequence data provided by the Genetic Analysis Workshop (GAW) 17 (Almasy et al 2011). The exome sequence data contain 24,487 SNVs in 3205 genes from 697 unrelated subjects. Our analyses focused on RVs with MAFs no larger than 1%; after removing those more frequent SNVs, we had 2476 genes containing at least 1 RV, with a total of 18,131 RVs. We conducted gene-based analyses.

The phenotypes were generated by GAW17 organizers based on some disease liability models with covariates. In particular, biological knowledge on pathways, especially the vascular endothelial growth factor (VEGF) pathway, and on predicted deleterious coding variants, was utilized to design a realistic simulation model. Fixing the sequence data for the 697 subjects, 200 independent sets of a binary phenotype were generated. In addition to some causal SNVs, three risk factors, age, gender and smoking status, were possibly associated with the binary phenotype. An advantage of using the GAW17 data is the opportunity to assess statistical power of any given method due to the known causal SNVs and the availability of replicated phenotypes. Hence, in addition to conducting a genome-wide scan, we also applied the methods to each causal gene with all 200 sets of the binary phenotype to estimate the power.

3.3.1 A genome-wide scan

To demonstrate the practical use of the proposed methods, we first conducted a genome-wide scan on all the 2476 genes with the first set of the binary phenotype. A logistic regression model was fitted to each gene with the three covariates:

$$\text{Logit}[Pr(Y_i = 1)] = \beta_0 + \sum_{j=1}^k X_{ij}\beta_j + \text{Age}_i \cdot \alpha_1 + \text{Gender}_i \cdot \alpha_2 + \text{Smoke}_i \cdot \alpha_3,$$

for $i = 1, \dots, 697$, where X_{i1}, \dots, X_{ik} are the k RVs in the gene to be tested, and Y_i is the binary phenotype. In the presence of the covariates, we used the parametric bootstrap to calculate the p-values for the SPU, aSPU, SKAT and SKAT-O tests.

Throughout the data analysis, we used the following “step-up” procedure to determine the number of bootstraps (or permutations), B . We started with $B = 10^3$, then gradually increased B : if an estimated p-value was less than $5/B$, we increased B to ten times of its current value to re-estimate the p-value, and the process was repeated until no estimated p-value was less than $5/B$. For the genome-scan on the GWA17 data, we tried $B = 10^3$, 10^4 and up to 10^5 to obtain p-values. On a multi-core computer with 100 cores, it took about 0.21 hours to test the 2476 genes based on $B = 10^3$, 0.05 hours to test the 50 genes (with their p-values < 0.005 in the previous round) with $B = 10^4$, and 0.12 hours for testing the 5 genes based on $B = 10^5$. If a single-core computer was used, a conservative estimate of the time to be taken would be the above time multiplied by 100, which would take less than two days. If we doubled the sample size, it took about three times of the original time. Note that our code was completely in R and not yet optimized; implementing the core part in C or another compiled language is planned and expected for at least a 10-fold speedup. Hence, to be more accurate, we can replace the used threshold $5/B$ with a larger value such as $50/B$.

Figure 1 shows the Manhattan plots for the tests. Since we were testing on 2476 genes, at the usual family-wide significance level of 0.05 and with a Bonferroni

adjustment, we would use a gene-wise significance level of $0.05/2476 = 2.02 \times 10^{-5}$, which would suggest using a bootstrap replication number up to 10^6 to 10^7 .

None of the tests detected any significant genes. Nonetheless, we highlighted the top five most significant genes based on each test to show their differing operating characteristics. In Figure 1, it is clear that the three representative SPU tests gave overlapping but different sets of the top genes. It is interesting to note that the aSPU test combined the results of the SPU tests.

As a comparison, Figure 1 also shows the results for the parametric bootstrap- and asymptotics-based SKAT and SKAT-O tests. First, we note the differences among the top genes between the resampling- and asymptotics-based tests, though their overall trends were similar, implying that one has to be cautious in using asymptotics-based tests for analysis of RVs. Second, we note the difference between the results of SKAT and SKAT-O; the latter had some similarity to that of the SPU(1) test, as expected. Most importantly, although similar to some extent, the top ranked genes by the aSPU, SKAT and SKAT-O tests were still different, suggesting the potential usefulness of the aSPU test as a complement to SKAT and SKAT-O. We also note that, most of the genes contained no more than 30 RVs; otherwise, the difference between the aSPU and SKAT or SKAT-O could be larger, as suggested in our simulations.

3.3.2 Power comparison

With the 200 replicated sets of the binary phenotype, the GAW17 data offer a unique opportunity to estimate the power of any test when applied to real sequence data. Due to the small sample size and relatively small effect sizes of the causal RVs, there was low power to detect any causal RVs in the GAW17 data. Accordingly, we used a less stringent gene-wise significance level of 0.05 (i.e. without multiple testing adjustment) and estimated the power of a test as the sample proportion of its rejecting the null hypothesis among the 200 sets of the replicated phenotypes. We considered all 35 causal genes (with at least one causal RV) and tested on each separately. The

main results are shown in Table 4.

We excluded the genes for which the maximum power of all the tests was lower than 10%, and partitioned the remaining causal genes into three groups based on whether the aSPU test was more powerful than both SKAT and SKAT-O, or between them, or less than them. It is clear that for some genes, the aSPU test was more powerful than SKAT and SKAT-O. For gene PIK3C2B, due to the connections between SPU(1) and burden tests and between SPU(2) and SKAT, given that the SPU(1) was more powerful than SPU(2), it is not surprising to see that SKAT-O (which combines SKAT and a burden test) was more powerful than SKAT; furthermore, perhaps due to the relatively high signal sparsity, the SPU(3) turned out to be the most powerful among the SPU tests, leading to that the aSPU test was more powerful than both SKAT and SKAT-O. A similar but different situation was with gene VNN1. That the SPU(1) test was less powerful than SPU(2) might explain why SKAT was more powerful than SKAT-O; however, interestingly, since the SPU(5) test was most powerful among the SPU tests while SPU(4)-SPU(8) were also relatively high-powered, we had the aSPU test more powerful than both SKAT and SKAT-O. On the other hand, when SPU(1) or SPU(2) was (nearly) more powerful than other SPU tests, the aSPU test combining all the SPU tests (with only one or two high-powered but more low-powered ones) lost edge to either SKAT or SKAT-O, but not both, as for gene RRAS. In some situations, as for gene KDR, since both SPU(1) and SPU(2) were far more superior than other SPU tests, the aSPU test ended up as being less powerful than both SKAT and SKAT-O. In summary, we found that, as expected, there was no uniform winner among the aSPU, SKAT and SKAT-O tests; under some situations, the aSPU test could be more powerful than both SKAT and SKAT-O.

Since all the causal genes contained only a relatively small number of non-causal RVs, and all causal RVs were deleterious (Almasy et al 2011), most often the SPU(3)

test was most powerful among the SPU tests. In contrast, perhaps as expected, the $\text{SPU}(\infty)$ was almost always least powerful. Importantly, we note that the power of the $\text{SPU}(3)$ test could be much larger than $\text{SPU}(1)$ and $\text{SPU}(2)$, representatives of the burden tests and variance-component tests. For example, for gene PIK3C2B with 23 causal RVs among a total of 60 RVs, the power of $\text{SPU}(3)$ was 0.650, much larger than 0.565 and 0.445 for $\text{SPU}(1)$ and $\text{SPU}(2)$ respectively. It is also confirmed that, even with the same association direction but also with non-associated RVs, the $\text{SPU}(1)$ test might lose power, as for gene VNN1 and BCHE. These points reinforce what was observed in simulation studies: we may need to use SPU tests beyond $\text{SPU}(1)$ and $\text{SPU}(2)$ to yield high power, supporting the use of the aSPU test in some applications.

We briefly mention two limitations of the GAW17 data. First, the number of RVs in any gene was often relatively small in this mini-exome sequence dataset, hence the clear advantage of $\text{SPU}(\gamma)$ tests with $\gamma > 3$ did not show up, which in turn limited the potential advantage of the aSPU test. However, with the increasing availability of whole-genome sequence data, we expect a much larger number of RVs in or near a gene, for which we may see a more dramatic advantage of the aSPU test. Second, since all the causal SNVs were deleterious, it favored SKAT-O, which might not perform so well otherwise (as shown in simulations; see Table 3).

We also did a simulation study to confirm that our methods could control Type I error rates satisfactorily with real sequence data at a higher significance level. We randomly selected three genes, RRAS, HIF3A and PIK3C2B, with 5, 15 and 60 RVs respectively. To mimic the GAW17 phenotype data, we randomly generated a binary phenotype Y_i with $\Pr(Y_i = 1) = 0.3$ under H_0 for each of the 697 subjects. We then tested H_0 for possible association between the phenotype and each gene with 10^5 simulation replicates so that we could obtain more accurate Type I error estimates for a higher significance level α . As shown in Table 5, our methods could satisfactorily control the Type I error rate. We note that, for gene RRAS with only 5 RVs, many

tests could be conservative with Type I errors lower than the nominal level α , due to the highly discrete null distributions of the test statistics. Furthermore, due to its higher-level discreteness and extreme-value-type test statistic, the tail distribution of the $\text{SPU}(\infty)$ statistic had larger variability, which in turn could perturb that of the aSPU test. As shown in Table 4, since the $\text{SPU}(\infty)$ test was almost always lowest-powered, we suggest excluding it when testing on a few RVs with a binary phenotype. Here, for the purpose of demonstration, we included the $\text{SPU}(\infty)$ test.

3.3.3 SNV selection

The aSPU test can be used **for SNV selection**. For comparison, we also included two methods, one was the **UminP test as used** in GWAS, and the **other was Lasso penalized logistic regression** (Tibshirani 1996; Zhou et al 2010). For UminP, we ranked the RVs based on their corresponding p-values. We used R package **glmnet** to fit Lasso penalized logistic regression. There is a tuning parameter $\lambda \geq 0$ in Lasso; as one reduces the value of λ , there will be more non-zero coefficient estimates $\tilde{\beta}_j$, thus selecting more RVs to be included. In this way, we counted the number of non-zero $\tilde{\beta}_j$ and the corresponding number of true positives (i.e. the corresponding causal RVs). One problem with Lasso penalized regression with RVs was that it was difficult to control the number of the non-zero coefficient estimates; as shown in Figure 2 for gene PIK3C2B, we could not obtain 40 or so non-zero coefficients, even with some labor-intensive fine-tuning of λ .

The methods were compared based on the number of true positives among a given number of their top ranked RVs for a causal gene. We found that aSPU and Lasso performed similarly, but much better than UminP. Among the 15 causal genes, if we looked at the top six ranked RVs, the frequencies of selecting (0, 1, 2, 3, 4) true positives were the following: (10, 5, 0, 0, 0) by UminP, (3, 6, 4, 1, 1) by Lasso, and (4, 3, 6, 1, 1) by aSPU. Similar results were obtained for other numbers of top ranked RVs. The bad performance of the UminP test could be due to the unstable

variance estimate for a RV, which was too close to 0 (with a too small MAF) and thus dramatically inflated the corresponding test statistic. Figure 2 shows a few more examples in detail. It is confirmed that aSPU and Lasso performed similarly. Given the simplicity of the aSPU test and that Lasso may not yield some given numbers of none-zero coefficient estimates, the use of aSPU for ranking and selecting RVs seems to be promising.

4. DISCUSSION

We have proposed and studied a class of SPU tests, which are versatile in the sense that, in many scenarios at least one of the SPU tests has high power, though the identity of a more powerful SPU test may change with the varying scenario. The SPU tests are based on the score vector of a regression model, rendering it both computationally efficient and general to cover a wide range of applications with binary, quantitative, ordinal and survival traits and possible covariates. In particular, compared to several other adaptive tests based on estimated (marginal) regression coefficients (e.g. Feng et al 2011; Zhang et al 2011; Lin and Tang 2011), the SPU tests, as any score-based test, only requires fitting a simplified model under the null hypothesis. In addition to its computational simplicity, a score-based test is more stable with RVs: for example, with a binary trait, the MLE of a regression coefficient for a RV does not exist if the minor allele appears in only cases or controls (but not both), leading to no convergence with an iterative algorithm to obtain the MLE. On the other hand, as shown earlier, the MLE of a (marginal) regression coefficient is approximately proportional to its score component, implying that, as expected, the score vector is as informative as a vector of the estimated regression coefficients while being computationally much simpler. Our major contribution is that, by recognizing the limitation of the existing adaptive tests with a fixed choice of weights, we allow many possible choices of weights indexed by a single parameter $\gamma \geq 1$. There is a

simple interpretation, and thus a guidance, on parameter γ : as the value of γ increases, we up-weight the larger components of the absolute value of the score vector, $|U|$; that is, with a decreasing proportion of the causal RVs in the group to be tested, we expect a $\text{SPU}(\gamma)$ test with a larger value of γ to be more powerful because its up-weighting of the larger components of $|U|$ essentially reduces or even eliminates noisy perturbations from many non-associated RVs to the test statistic, thus maintaining high power in the presence of many non-associated RVs. In addition, in the presence of both positive and negative association directions for causal RVs, an even number of γ will eliminate the cancelling effect of positive and negative components of U . In particular, as compared with some new adaptive tests, such as KBAC, PWST, aSSU, EREC and SKAT-O, our proposed aSPU test was more adaptive and performed much better in simulations when there were a large number of non-associated RVs.

For its highly adaptive and versatile performance, the aSPU test has a wide spectrum of applications with other types of traits and/or other genetic variants. For example, our preliminary results showed promising performance of the aSPU test for polygenic testing on association between a binary trait and thousands of CVs (The International Schizophrenia Consortium 2009). In principle, the aSPU test can be also applied to gene set or pathway analysis (Liu et al 2008; Wang et al 2010).

The relatively good performance of the SSU test and its close relatives, KMR or SKAT (Wu et al 2010, 2011) and C-alpha test (Neale et al 2011), was attributed to its treating the regression coefficients β as random effects and then testing on the variance component of the random effects (Basu and Pan 2011). Here, based on the formulation of the SPU tests, more general than the SSU test, we feel that it can be viewed from another angle: the good performance of the SSU test, or more generally, any $\text{SPU}(\gamma)$ test, is due to the weighting of each score component U_j by itself or its power $U_j^{\gamma-1}$; since U_j contains association information about RV j , such weights are both simple and informative: specifically, since U has a null distribution

$N(0, V)$, a larger component of $|U_j|$ corresponds to stronger evidence of association between the j th RV and the trait, and thus assigning a higher weight U_j or $U_j^{\gamma-1}$ will help boost power by reducing the noises introduced to the test statistic with non-associated RVs. However, depending on the unknown truth, such as the proportion of associated RVs and their specific association effects, a $\text{SPU}(\gamma)$ test with a suitable γ will be more powerful than others. For example, in the presence of many non-associated RVs, we would expect a larger γ to be more effective: a non-associated RV is expected to have a smaller $|U_j|$, and thus almost a zero weight with $U_j^{\gamma-1}$, in which way we may successfully eliminate the negative effects of many non-associated RVs on testing. In particular, when the group of RVs to be tested contains a large proportion of non-associated RVs, we found that a $\text{SPU}(\gamma)$ test with $\gamma > 2$ could be much more powerful than the Sum and SSU tests, explaining when and why the aSPU test could outperform SKAT-O and other tests combining a burden test like $\text{SPU}(1)$ and a variance component test like $\text{SPU}(2)$ (Lee et al 2012; Derkach et al 2012; Sun et al 2013). We also found that a $\text{SPU}(\gamma)$ test with a large $\gamma > 8$ gave results similar to that of the $\text{SPU}(\infty)$ test in our simulations, suggesting that searching γ over 1 to 8 perhaps suffices in many applications.

Since the SPU tests use a mechanism of weighting to minimize the effects of non-associated RVs, they are analogous to model averaging in the literature of model selection, in contrast to the adaptive Neyman-type tests, such as the aSSU test (Pan and Shen 2011), which are more in the line of model or RV selection. In the current context of analysis of RVs, due to limited information contained within each individual RV, model averaging is expected to outperform model selection, as supported by our empirical results when comparing between the aSPU and aSSU tests. Since in general neither model selection nor model averaging can dominate the other (Yuan and Yang 2005; Shen and Huang 2006), there may be a merit in combining the two approaches in other applications. In addition, a few modifications or extensions are

possible. First, in our current implementation of the aSPU test, we simply take the minimum p-value (minP) to combine multiple SPU tests; other combining methods may be equally applied and may be preferred under certain situations (Pan et al 2010). Second, it appears straightforward to extend the SPU tests to the case with variable thresholds (Price et al 2010), which is related to adaptive Neyman-type tests (Pan and Shen 2011). Third, we have not evaluated the performance of the SPU tests in the presence of interactions among RVs; in particular, it would be interesting to compare their performance with the KBAC test that was partly designed to detect interactions (Liu and Leal 2010). Finally, although we have pointed out some possible extensions of the aSPU test for 1) analysis of CVs, 2) joint analysis of both CVs and RVs, and 3) pathway analysis of CVs and/or RVs, these topics warrant further investigation in future research.

R code will be posted on our web site at
<http://www.biostat.umn.edu/~weip/prog.html>.

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Table 1: Empirical Type I error rates of various tests for the cases with a group of 8 non-associated RVs and another group of non-associated RVs; the RVs within each group were correlated, but there was no between-group correlation; all results were based on 1000 simulation replicates.

Test	# non-associated RVs						
	0	8	16	32	64	96	128
UminP	.025	.020	.017	.018	.015	.007	.011
SPU(1)	.051	.055	.044	.049	.048	.041	.051
SPU(2)	.048	.055	.034	.029	.039	.037	.035
SPU(3)	.049	.051	.040	.041	.032	.038	.037
SPU(4)	.051	.046	.033	.040	.025	.037	.029
SPU(5)	.053	.048	.039	.050	.026	.043	.030
SPU(6)	.056	.046	.042	.048	.024	.039	.022
SPU(7)	.057	.044	.040	.046	.028	.038	.023
SPU(8)	.054	.042	.040	.044	.022	.036	.023
SPU(16)	.055	.041	.041	.047	.029	.037	.025
SPU(32)	.053	.042	.043	.047	.030	.039	.025
SPU(∞)	.053	.042	.042	.047	.031	.039	.025
aSPU	.055	.044	.041	.048	.038	.047	.038
aSum+	.045	.058	.045	.053	.048	.051	.046
aSum2d	.062	.054	.042	.049	.052	.043	.045
aSSU	.066	.047	.045	.045	.046	.048	.052
KBAC	.049	.061	.035	.049	.047	.057	.051
aSum	.066	.056	.033	.048	.048	.055	.042
PWST	.061	.047	.031	.049	.040	.040	.042
EREC	.056	.058	.044	.046	.050	.045	.046
BhGLM	.044	.061	.042	.042	.043	.039	.048
SKAT	.057	.064	.042	.046	.050	.046	.049
SKAT-O	.058	.065	.047	.040	.051	.047	.052

Table 2: Empirical power of various tests for the cases with a group of 8 causal RVs with ORs randomly drawn from $U(1,2)$, and another group of non-associated RVs; all results were based on 1000 simulation replicates. The highest powered non-adaptive and adaptive tests are bold-faced.

Test	# non-associated RVs						
	0	8	16	32	64	96	128
UminP	.874	.812	.768	.733	.659	.619	.586
SPU(1)	.939	.852	.746	.577	.411	.300	.244
SPU(2)	.926	.908	.904	.872	.832	.801	.769
SPU(3)	.917	.903	.893	.870	.829	.802	.786
SPU(4)	.909	.896	.890	.882	.854	.840	.835
SPU(5)	.902	.894	.879	.875	.843	.834	.834
SPU(6)	.901	.882	.872	.873	.843	.835	.835
SPU(7)	.899	.881	.868	.869	.836	.830	.828
SPU(8)	.898	.876	.863	.864	.833	.834	.826
SPU(16)	.885	.860	.848	.852	.821	.814	.814
SPU(32)	.878	.855	.844	.850	.817	.807	.810
SPU(∞)	.877	.852	.844	.846	.814	.801	.806
aSPU	.923	.898	.894	.869	.842	.829	.811
aSum+	.940	.893	.866	.801	.677	.607	.532
aSum2d	.921	.857	.805	.719	.618	.504	.453
aSSU	.900	.867	.847	.819	.767	.695	.658
KBAC	.903	.785	.722	.613	.466	.331	.283
aSum	.948	.892	.855	.756	.636	.480	.407
PWST	.823	.729	.698	.613	.508	.400	.380
EREC	.943	.901	.887	.833	.738	.656	.579
BhGLM	.934	.863	.779	.619	.434	.299	.248
SKAT	.927	.914	.906	.870	.823	.800	.749
SKAT-O	.940	.915	.899	.858	.799	.767	.696

Table 3: Empirical power of various tests for the cases with a group of 8 causal RVs with ORs=(3, 1/3, 2, 2, 2, 1/2, 1/2, 1/2) and another group of non-associated RVs; all results were based on 1000 simulation replicates. The highest powered non-adaptive and adaptive tests are bold-faced.

Test	# non-associated RVs						
	0	8	16	32	64	96	128
UminP	.507	.379	.324	.288	.208	.197	.157
SPU(1)	.341	.227	.159	.111	.074	.070	.070
SPU(2)	.631	.542	.485	.435	.332	.279	.261
SPU(3)	.563	.493	.451	.413	.303	.323	.274
SPU(4)	.625	.540	.508	.478	.402	.386	.351
SPU(5)	.601	.514	.480	.465	.374	.383	.358
SPU(6)	.619	.529	.504	.490	.398	.401	.367
SPU(7)	.600	.521	.485	.480	.386	.399	.370
SPU(8)	.610	.530	.494	.485	.402	.404	.370
SPU(16)	.595	.524	.488	.470	.390	.395	.356
SPU(32)	.595	.523	.487	.469	.388	.389	.353
SPU(∞)	.592	.520	.484	.467	.386	.387	.355
aSPU	.589	.511	.467	.461	.366	.361	.329
aSum+	.596	.497	.419	.370	.268	.245	.214
aSum2d	.560	.450	.391	.326	.220	.193	.169
aSSU	.598	.511	.464	.424	.337	.311	.296
KBAC	.525	.392	.327	.254	.181	.135	.129
aSum	.549	.415	.299	.232	.138	.145	.127
PWST	.675	.554	.460	.354	.267	.197	.196
EREC	.545	.448	.365	.298	.189	.176	.154
BhGLM	.434	.290	.186	.123	.082	.072	.071
SKAT	.650	.557	.474	.425	.318	.272	.235
SKAT-O	.622	.495	.406	.378	.260	.226	.195

Table 4: Estimated power for some causal genes with the GAW17 data.

Chr	Gene	#RVs	#Causal	SPU(1)	SPU(2)	SPU(3)	SPU(4)	SPU(5)	SPU(6)	SPU(7)	SPU(8)	SPU(∞)	aSPU	SKAT	SKAT-O
1	PIK3C2B	60	23	0.565	0.445	0.650	0.400	0.395	0.355	0.360	0.350	0.340	0.600	0.435	0.560
6	VNN1	6	1	0.185	0.230	0.315	0.235	0.380	0.320	0.350	0.325	0.140	0.270	0.215	0.185
3	BCHE	28	13	0.110	0.190	0.215	0.185	0.175	0.165	0.165	0.165	0.160	0.210	0.195	0.175
8	LPL	15	2	0.090	0.130	0.135	0.110	0.115	0.110	0.115	0.115	0.110	0.135	0.125	0.125
10	SIRT1	23	9	0.095	0.105	0.110	0.065	0.070	0.060	0.055	0.055	0.015	0.105	0.090	0.100
14	SOS2	7	2	0.100	0.270	0.285	0.265	0.275	0.265	0.275	0.265	0.245	0.220	0.255	0.200
19	RRAS	5	2	0.235	0.140	0.155	0.145	0.155	0.150	0.155	0.150	0.100	0.180	0.135	0.200
8	PLAT	25	8	0.225	0.135	0.145	0.110	0.105	0.100	0.100	0.100	0.070	0.155	0.130	0.195
9	VLDLR	23	8	0.080	0.120	0.125	0.110	0.120	0.105	0.115	0.105	0.075	0.090	0.125	0.090
17	SREBF1	21	10	0.050	0.085	0.090	0.105	0.100	0.105	0.100	0.105	0.100	0.085	0.090	0.070
4	KDR	14	8	0.365	0.350	0.160	0.105	0.105	0.100	0.100	0.100	0.020	0.280	0.365	0.390
13	FLT1	25	8	0.125	0.160	0.170	0.150	0.160	0.155	0.155	0.160	0.065	0.125	0.150	0.165
14	HSP90AA1	20	3	0.050	0.275	0.180	0.195	0.170	0.140	0.110	0.120	0.030	0.155	0.335	0.250

Table 5: Empirical Type I error rates at the nominal significance level α based on 10^5 simulation replicates.

Gene	α	SPU(1)	SPU(2)	SPU(3)	SPU(4)	SPU(5)	SPU(6)	SPU(7)	SPU(8)	SPU(∞)	aSPU	SKAT	SKAT-O
RRAS	0.01	0.00689	0.00974	0.00968	0.00963	0.00967	0.00965	0.00965	0.00968	0.00600	0.00791	0.00940	0.00908
	0.001	0.00041	0.00099	0.00102	0.00100	0.00097	0.00100	0.00100	0.00100	0.00032	0.00063	0.00085	0.00072
HIF3A	0.01	0.00902	0.00905	0.00922	0.00927	0.00924	0.00938	0.00931	0.00936	0.01019	0.00958	0.00980	0.00808
	0.001	0.00106	0.00086	0.00099	0.00095	0.00101	0.00098	0.00102	0.00097	0.00126	0.00101	0.00088	0.00074
PIK3C2B	0.01	0.00943	0.00901	0.00903	0.00894	0.00898	0.00915	0.00902	0.00917	0.00977	0.00933	0.00980	0.00995
	0.001	0.00094	0.00081	0.00097	0.00095	0.00090	0.00093	0.00089	0.00095	0.00093	0.00116	0.00086	0.00106

Figure 1: Manhattan plots for the GAW17 data.

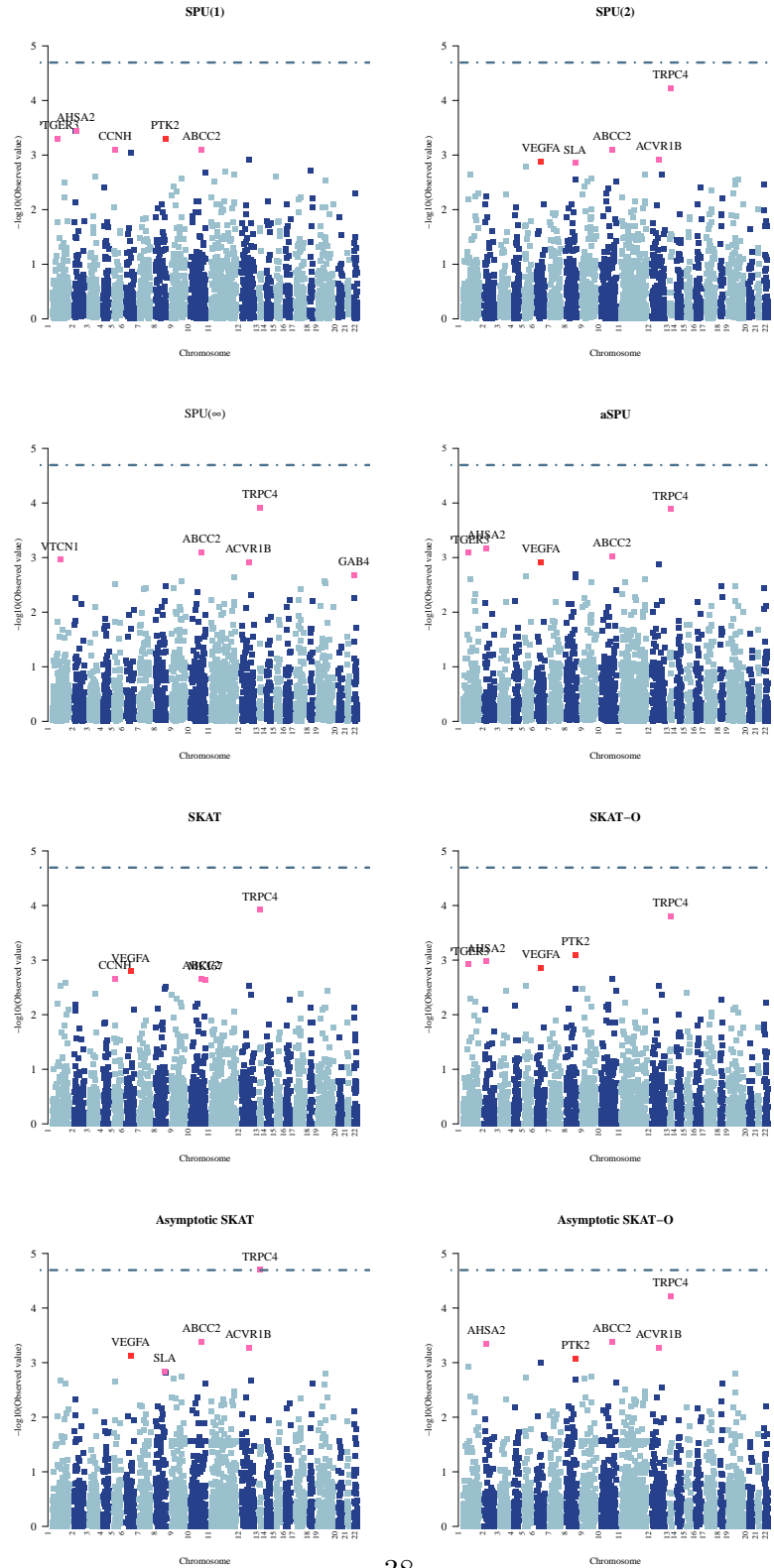


Figure 2: RV selection by aSPU and Lasso for a few causal genes with the GAW17 data.

