SIMULTANEOUS SELECTION OF MULTIPLE IMPORTANT SINGLE NUCLEOTIDE POLYMORPHISMS IN FAMILIAL GENOME WIDE ASSOCIATION STUDIES DATA

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We propose a resampling-based fast variable selection technique for selecting important Single Nucleotide Polymorphisms (SNP) in multi-marker mixed effect models used in twin studies. Due to computational complexity, current practice includes testing the effect of one SNP at a time, commonly termed as 'single SNP association analysis'. Joint modeling of genetic variants within a gene or pathway may have better power to detect the relevant genetic variants, hence we adapt our recently proposed framework of e-values to address this. To our knowledge, this is the first computationally efficient method of SNP detection in twin studies that uses multi-marker models. We achieve this through improvements in two aspects. First, unlike other model selection techniques, our method only requires training a model with all possible predictors. Second, we utilize a fast and scalable bootstrap procedure that only requires Monte-Carlo sampling to obtain bootstrapped copies of the estimated vector of coefficients. Using this bootstrap sample, we obtain the e-value for each SNP, and select SNPs having e-values below a threshold. Numerical studies reveal our method to be more effective in detecting causal SNPs than either single-marker analysis on mixed models or model selection methods that ignore the familial dependency structure. We also use the evalues to perform gene-level analysis in a familial GWAS dataset and detect several SNPs that have potential effect on alcohol consumption in individuals.

1. Introduction. Genome Wide Association Studies (GWAS) have identified a large number of genetic variants associated with complex diseases [3, 39]. The advent of economical high-throughput genotyping technology enables researchers to scan the genome with millions of SNPs, and improvements in computational efficiency in analysis techniques has facilitated parsing throught this huge amount of data to detect significant associations [35]. Detection of the small effects of individual SNPs requires large sample sizes [26], which is a major challenge of these studies. For quantitative behavioral traits such as alcohol consumption, drug abuse, anorexia and depression,

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variation due to the environment the subject grew up in brings in additional noise, further amplifying the issue. This is one of the motivations of performing GWAS on families instead of unrelated individuals, through which the environmental variation can be reduced; so as to require smaller samples to detect the same magnitude of SNP effect. Resolving the Minnesota Twin Family Study [29] where data were observed on identical twins, non-identical twins, biological offsprings and adoptees, serve as the motivation for our methodology development in this paper.

Single-marker analysis, i.e. analyzing the effect of SNPs individually on the phenotype of interest and reporting the top SNPs by setting a suitable threshold on the resulting p-values is perhaps the most commonly used method to detect SNPs. The GRAMMAR method of [1] and the association test of [6] are examples of such techniques that are applied in familial data. While they are able to efficiently analyze GWAS data, they assume that phenotypic similarity within families is entirely due to their genetic similarity and ignore the effect of shared environment. As a result, they tend to lose power when analyzing data where shared environmental effects explain a substantial proportion of the total phenotypic variation (see [28] and [10] for example). In contrast, the RFGLS method proposed by [23] takes into account genetic and environmental sources of familial similarity and still provides fast inference through a rapid approximation of the single-SNP mixed effect model.

Single-marker methods are prone to be less effective in detecting SNPs with weak signals [26]. This includes cases where multiple SNPs are jointly associated with the phenotype [18, 40] and SNP-SNP interactions-[33]. Several methods of multi-marker analysis have been proposed as alternatives. The kernel based association tests of [5, 15, 32, 33] are prominent among such techniques. However, all of them test for whether a group of SNPs is associated with the phenotype being analyzed, and are not able prioritize within the group and detect the exact causal SNPs which affect the quantitative phenotype.

One way to solve this problem is to perform model selection. The methods of [11] and [41] take this approach, and are able to perform SNP seletion from a multi-SNP model on GWAS data from *unrelated individuals*. However, they rely on fitting models corresponding to multiple predictor sets, hence are computationally very intensive to implement in a linear mixed-effect framework that is necessary to model familial data.

In this paper we propose a fast and scalable model selection technique thats fits a single model, and identifies important genetic variants with weak signals through modelling multiple variants jointly. We achieve this by extending the recently proposed framework of e-values [25]. For any estimation method that provides consistent estimates (at a certain rate relative to the sample size) of the vector of parameters, e-values quantify the proximity of the sampling distribution for a restricted parameter estimate to that of the full model estimate in a regression-like setup. A variable selection algorithm using the e-values has the following simple and generic steps:

- 1. Fit the full model, i.e. where all predictor effects are being estimated from the data, and use resampling to estimate its *e*-value;
- 2. Set an element of the full model coefficient estimate to 0 and get *e*-value for that predictor, repeat this for all predictors;
- 3. Select predictors that have e-values below a pre-determined threshold.

The above algorithm offers multiple important benefits in the SNP selection scenario. Unlike other model selection methods, only the full model needs to be computed here. It thus offers the user more flexibility in utilizing a suitable method of estimation. Our method allows for fitting multi-SNP models, thereby accommodating cases of modelling multiple correlated SNPs or closelt located multiple causal SNPs simultaneously. Finally, we use the Generalized Bootstrap [4] as our chosen resampling technique. Instead of fitting a separate model for each bootstrap sample, it computes bootstrap estimates using Monte-Carlo samples from the resampling distribution and reusing model objects obtained while fitting the full model. Consequently, the resampling step becomes very fast and parallelizable.

The rest of the paper is organized as follows. Section 2 provides background information on the GWAS Family dataset we use in our case study, as well as introduces the statistical framework we use to model this data. We start Section 3 by providing a technical introduction to the e-values framework, then elaborate on the necessary modifications for adapting it to our modelling scenario, and present details of the bootstrap procedure. We illustrate the performance of this method on synthetic datasets in 4. In Section 5 we analyze our GWAS dataset using the e-values technique to select SNPs from multiple genes that have been reported to influence alcohol consumption in individuals. Finally in Section 6 we outline directions of future research. We include the proofs of all new results stated, specifically, theorems 3.2 and 3.3, in the supplementary material.

2. Data and model.

2.1. The MCTFR data. The familial GWAS dataset collected and studied by Minnesota Center for Twin and Family Research (MCTFR)[23, 28, 29] consists of samples from three longitudinal studies conducted by the

MCTFR: (1) the Minnesota Twin Family Study (MTFS: [14]) that covers twins and their parent, (2) the Sibling Interaction and Behavior Study (SIBS: [27]) that includes adopted and biological sibling pairs and their parents, and (3) the enrichment study (ES: [19]) that extended the MTFS by oversampling 11 year old twins who are highly likely to develop substance abuse. While 9827 individuals completed the initial assessments for participation in the study, after several steps of screening the final sample consisted of 7188 Caucasian individuals clustered in 2300 nuclear families.

DNA samples collected from the subjects are analyzed using Illumina's Human660W-Quad Array, and after standard quality control steps [29], 527,829 SNPs are retained. Covariates for each sample included age, sex, birth year, generation (parent or offspring), as well as two-way interactions between generation and other three covariates each. Five quantitative phenotypes were studied in this GWAS: (1) Nicotine dependence, (2) Alcohol consumption, (3) Alcohol dependence, (4) Illegal drug usage, and (5) Behavioral disinhibition. The response variables corresponding to these phenotypes are derived from questionnaires using a hierarchical approach based on factor analysis [12].

A detailed description of the data is available in [29]. Several studies have been performed that focus on different aspects of this dataset. The study in [23] used RFGLS to single out causal SNPs behind the height of participants, while [28] used the same method to study SNPs behind the development of all five indicators of behavioral disinhibition mentioned above. [16] focused on the effect of several factors affecting alcohol use in the study population, namely the effects of polymorphisms in the ALDH2 gene and the GABA system genes, as well as the effect of early exposure to alcohols as adolescents to adult outcomes. Finally [7] used a bootstrap-based combination test and a sequential score test to evaluate gene-environment interactions behind phenotypic outcomes in the data.

2.2. Statistical model. We use a Linear Mixed Model (LMM) with three variance components accounting for several potential sources of variation to model effect of SNPs behind a quantitative phenotype. This is known as ACE model in the literature [20]. While the-state-of-the-art focuses on using a single SNP and other covariates as fixed effects and trains multiple models, we shall incorporate all SNPs that are genotyped within a gene (or group of genes in some cases) into the set of fixed effects in a single model.

Our model fitting process is invariant across pedigree sizes. In the present context we adopt the standard protocol of assuming nuclear pedigrees, that is, the families are unrelated, as implemented by [6, 23, 28]. Suppose there

are m families in total, with the i^{th} pedigree containing n_i individuals. Denote by $y_i = (y_{i1}, \ldots, y_{in_i})^T$ the quantitative trait values for individuals in that pedigree, while the matrix $\mathbf{G}_i \in \mathbb{R}^{n_i \times p_g}$ contains their genotypes for a number of SNPs. Let $\mathbf{C}_i \in \mathbb{R}^{n_i \times p}$ denote the data on p covariates for individuals in the pedigree i. Given these, we consider the following model.

(2.1)
$$\mathbf{Y}_i = \alpha + \mathbf{G}_i \beta_q + \mathbf{C}_i \beta_c + \epsilon_i$$

with α the intercept term, β_g and β_c fixed coefficient terms corresponding to the multiple SNPs and covariates, respectively, and $\epsilon_i \sim \mathcal{N}_{n_i}(\mathbf{0}, \mathbf{V}_i)$ the random error term. To account for the within-family dependency structure, we break up the random error variance into three independent components:

(2.2)
$$\mathbf{V}_i = \sigma_a^2 \mathbf{\Phi}_i + \sigma_c^2 \mathbf{1} \mathbf{1}^T + \sigma_e^2 \mathbf{I}_{n_i}$$

The first component above represents a within-family random effect term to account for effects of other SNPs. The matrix Φ_i is the relationship matrix within the i^{th} pedigree. Its $(s,t)^{\text{th}}$ element represents two times the kinship coefficient, which is the probability that two alleles, one randomly chosen from individual s in pedigree i and the other from individual t, are 'identical by descent', i.e. come from same common ancestor. The second variance component accounts for shared environmental effect within the family, while the third term finally quantifies other sources of variation unique to an individual.

Following basic probability, the kinship coefficient of a parent-child pair is 1/4, a full sibling pair or non-identical (or dizygous = DZ) twins is 1/4, and for identical (or monozygous = MZ) twins is 1/2 in a nuclear pedigree. Following this, we can construct the Φ_i matrices for different types of families:

$$\mathbf{\Phi}_{MZ} = \begin{bmatrix} 1 & 0 & 1/2 & 1/2 \\ 0 & 1 & 1/2 & 1/2 \\ 1/2 & 1/2 & 1 & 1 \\ 1/2 & 1/2 & 1 & 1 \end{bmatrix}, \mathbf{\Phi}_{DZ} = \begin{bmatrix} 1 & 0 & 1/2 & 1/2 \\ 0 & 1 & 1/2 & 1/2 \\ 1/2 & 1/2 & 1 & 1/2 \\ 1/2 & 1/2 & 1/2 & 1 \end{bmatrix}, \mathbf{\Phi}_{Adopted} = \mathbf{I}_4$$

for families with parents (indices 1 and 2) and MZ twins, DZ twins, or two adapted children (indices 3 and 4), respectively.

3. Variable selection using *e***-values.** We present the details of our methodology in this section. Sections 3.1 and 3.2 summarize the existing method of *e*-values that performs best subset variable selection in a wide range of statistical models [25]. We build on this framework in Section 3.3, where we present new results for better detection of weak SNP signals. Section 3.4 elaborates on the bootstrap implementation of this methodology using the model in (2.1).

3.1. Models and evaluation maps. In a general modelling situation where one needs to estimate a set of parameters $\theta \in \mathbb{R}^p$ from a triangular array of samples $\mathcal{B}_n = \{B_{n1}, \ldots, B_{nk_n}\}$ at stage n, any hypothesis or statistical model corresponds to a subset of the full parameter space. Here we consider the model spaces $\Theta_{mn} \subseteq \mathbb{R}^p$ in which some elements of the parameter vector have fixed values, while others are estimated from the data. Formally, a generic parameter vector $\theta_{mn} \in \Theta_{mn}$ consists of entries

$$\theta_{mnj} = \begin{cases} & \text{Unknown } \theta_{mnj} & \text{for } j \in \mathcal{S}_n; \\ & \text{Known } c_{nj} & \text{for } j \notin \mathcal{S}_n. \end{cases}$$

for some $S_n \subseteq \{1, \ldots, p\}$. Thus the estimable index set S_n and fixed elements $c_n = (c_{nj} : j \notin S_n)$ fully specify any model in this setup.

At this point, the original framework in [25] introduces a few concepts in order to provide a detailed treatment considering a general scenario. For our specific problem, i.e. variable selection, we only require vastly simplified versions of them. To start with, we consider sample size $k_n = n$ for all n, and assume constant sequences of candidate models: $\mathcal{M}_n = \mathcal{M}$ for all n. Consequently we also drop the subscripts in \mathcal{S}_n and c_n . Thus the 'full model', i.e. the model with all covariates is denoted by $\mathcal{M}_* = (\{1, \ldots, p\}, \emptyset)$.

Given data of size n, we obtain the full model estimates as minimizers of an estimating equation:

(3.1)
$$\hat{\theta} = \underset{\theta}{\operatorname{arg\,min}} \Psi(\theta) = \underset{\theta}{\operatorname{arg\,min}} \sum_{i=1}^{n} \Psi_i(\theta, B_i)$$

The only condition we impose on these generic estimating functionals $\Psi_i(.)$ are:

(P1) The population version of (3.1) has a unique minimizer θ_0 , i.e.

$$\theta_0 = \operatorname*{arg\,min}_{\theta} \mathbb{E} \sum_{i=1}^n \Psi_i(\theta, B_i)$$

(P2) There exist a sequence of positive numbers $a_n \uparrow \infty$ and a p-dimensional probability distribution \mathbb{T}_0 such that $a_n(\hat{\theta} - \theta_0) \leadsto \mathbb{T}_0$.

We designate θ_0 as the true parameter vector, some elements of which are potentially set to 0. We can now classify any candidate model \mathcal{M} into one of the two classes: those satisfying $\theta_0 \in \Theta_m$, and those not satisfying the condition. We denote these two types of models by adequate and inadequate models, respectively. Given the data and unknown θ_0 , we want to determine if a candidate model belongs to which of the above two categories.

For this we need coefficient estimates $\hat{\theta}_m$ corresponding to a model. We do so by just replacing elements of $\hat{\theta}$ not in \mathcal{S} by corresponding elements of c. This means that for the j^{th} element, $j = 1, \ldots, p$, we have

$$\hat{\theta}_{mj} = \begin{cases} & \text{Unknown } \hat{\theta}_j & \text{ for } j \in \mathcal{S}; \\ & \text{Known } c_j & \text{ for } j \notin \mathcal{S} \end{cases}$$

We denote the probability distribution of a random variable **T** by [**T**]. With this notation, we want to compare the above model estimate distributions with the full model distribution, i.e. $[\hat{\theta}_m]$ with $[\hat{\theta}]$. For this we define an evaluation map function $E: \mathbb{R}^p \times \tilde{\mathbb{R}}^p \to [0, \infty)$ that measures the relative position of $\hat{\theta}_m$ with respect to $[\hat{\theta}]$. Here $\tilde{\mathbb{R}}^p$ is the set of probability measures on \mathbb{R}^p . We assume that this function satisfies the following conditions:

(E1) For any probability distribution $\mathbb{G} \in \mathbb{R}^p$ and $x \in \mathbb{R}^p$, E is invariant under location and scale transformations:

$$E(x, \mathbb{G}) = E(ax + b, [a\mathbf{G} + b]); \quad a \in \mathbb{R} \neq 0, b \in \mathbb{R}^p$$

where the random variable G has distribution G.

(E2) The evaluation map E is lipschitz continuous under the first argument:

$$|E(x,\mathbb{G}) - E(y,\mathbb{G})| < ||x - y||^{\alpha}; \quad x, y \in \mathbb{R}^p, \alpha > 0$$

- **(E3)** Suppose $\{\mathbb{Y}_n\}$ is a tight sequence of probability measures in \mathbb{R}^p with weak limit \mathbb{Y}_{∞} . Then $E(x, \mathbb{Y}_n)$ converges uniformly to $E(x, \mathbb{Y}_{\infty})$.
- **(E4)** Suppose \mathbf{Z}_n is a sequence of random variables such that $\|\mathbf{Z}_n\| \stackrel{P}{\to} \infty$. Then $E(\mathbf{Z}_n, \mathbb{Y}_n) \stackrel{P}{\to} 0$.

For any $x \in \mathbb{R}^p$ and $[\mathbf{X}] \in \mathbb{R}^p$ such that $\mathbb{V}\mathbf{X}$ is positive definite, following are examples of the evaluations functions covered by the above set of conditions:

(3.2)

$$E_1(x, [\mathbf{X}]) = \left[1 + \left\| \frac{x - \mathbb{E}\mathbf{X}}{\sqrt{\operatorname{diag}(\mathbb{V}\mathbf{X})}} \right\|^2 \right]^{-1}; \quad E_2(x, [\mathbf{X}]) = \exp\left[-\left\| \frac{x - \mathbb{E}\mathbf{X}}{\sqrt{\operatorname{diag}(\mathbb{V}\mathbf{X})}} \right\| \right]$$

Data depths [34, 42, 43] also constitute a very broad class of point-to-distribution proximity functions that satisfy the above regularity conditions for evaluation maps. Indeed, [25] used halfspace depth [34] as evaluation function to perform model selection. However, the conditions (E1) and (E4) are weaker than those imposed on a traditional depth function [43]. Conditions (E2) and (E3) are not required of depth functions in general, but they

arise implicitly in several implementations of data depth [30]. The theoretical results we state here are based on a general evaluation map and not depth functions per se. To emphasize this point, in the numerical sections that follow we use the non-depth evaluation functions E_1 and E_2 given in (3.2) above.

3.2. Model selection using e-values. Depending on the choice of the data sequence \mathcal{B}_n , $E(\hat{\theta}_m, [\hat{\theta}])$ can take different values. For any candidate model \mathcal{M} , we denote the distribution of the corresponding random evaluation map by \mathbb{E}_{mn} . For simplicity we drop the n in the subscripts for such sampling distributions, i.e. $\mathbb{E}_{mn} \equiv \mathbb{E}_m$. These sampling distributions are informative of how model estimates behave, and we use them as a tool to distinguish between inadequate and adequate models. Given a single set of samples, we use resampling schemes that satisfy standard regularity conditions [25] to get consistent approximations of \mathbb{E}_m .

We now define a quantity called the e-value to compare the different model estimates and eventually perform selection of important SNPs from a multi-SNP model. Loosely construed, any functional of the evaluation map distribution \mathbb{E}_m that can act as model evidence is an e-value. For example, [25] took the mean functional of \mathbb{E}_m (say $\mu(\mathbb{E}_m)$) as e-value, and proved a result that, when adapted to our setting, states as:

THEOREM 3.1. Consider estimators satisfying conditions (P1) and (P2), and an evaluation map E satisfying the conditions (E1), (E2) and (E4). Also suppose that

$$\lim_{n\to\infty}\mu(\mathbb{Y}_n)=\mu(\mathbb{Y}_\infty)<\infty$$

for any tight sequence of probability measures $\{\mathbb{Y}_n\}$ in \mathbb{R}^p with weak limit \mathbb{Y}_{∞} . Then as $n \to \infty$,

- 1. For the full model, $\mu(\mathbb{E}_*) \to \mu_{\infty}$ for some $0 < \mu_{\infty} < \infty$;
- 2. For any adequate model, $|\mu(\mathbb{E}_m) \mu(\mathbb{E}_*)| \to 0$;
- 3. For any inadequate model, $\mu(\mathbb{E}_m) \to 0$.

Taking data depths as evaluation functions leads to a further result that $\mu(\mathbb{E}_*) < \mu(\mathbb{E}_m)$ for any adequate model \mathcal{M} and large enough n. Following this, non-zero indices of θ_0 (say \mathcal{S}_0) can be recovered through a fast algorithm that has these generic steps:

1. Estimate the e-value of the full model, i.e. $\hat{\mu}(\mathbb{E}_*)$, through bootstrap approximation of \mathbb{E}_* ;

- 2. For the j^{th} predictor, j = 1, ..., p, consider the model with the j^{th} coefficient of $\hat{\theta}$ replaced by 0, and get its e-value. Suppose this is $\hat{\mu}(\mathbb{E}_{-j})$;
- 3. Collect the predictors for which $\hat{\mu}(\mathbb{E}_{-j}) < \hat{\mu}(\mathbb{E}_*)$. Name this index set $\hat{\mathcal{S}}_0$: this is the estimated set of non-zero coefficients in $\hat{\theta}$.

As $n \to \infty$, the above algorithm provides consistent model selection, i.e. $\mathbb{P}(\hat{S}_0 = S_0) \to 1$, with the underlying resampling distribution having mean 1 and variance τ_n^2 such that $\tau_n \to \infty, \tau_n/\sqrt{n} \to 0$ as $n \to \infty$ [25].

3.3. Quantiles of \mathbb{E}_m as e-values. When true signals are weak, the above method of variable selection leads to very conservative estimates of non-zero coefficient indices, i.e. a large number of false positives in a sample setting. This happens because even though at the population level $\mu(\mathbb{E}_*)$ separates the population means of inadequate model sampling distributions and those of adequate models, for weak signals bootstrap estimates of adequate model distributions almost overlap with those of the full model.

Figure 3.1 demonstrates this phenomenon in our simulation setup. Here we analyze data on 250 families with monozygotic twins, each individual being genotyped for 50 SNPs. Four of these 50 SNPs are causal: each having a heritability of h/6% with respect to the total error variation present. The four panels show density plots of $\hat{\mathbb{E}}_{-j}$ for $j=1,\ldots,p$, as well as $\hat{\mathbb{E}}_*$: based on resampling schemes with four different values of the standard deviation parameter $s \equiv s_n = \tau_n/\sqrt{n}$. While smaller values of s are able to separate out the bootstrap estimates of \mathbb{E}_{-j} for inadequate and adequate models, all the density plots are to the left of the curve corresponding to the full model.

However, notice that the inadequate and adequate model distributions have different tail behaviors for smaller values of s, and setting an appropriate upper threshold to tail probabilities for a suitable fixed quantile of these distributions with respect to the full model distribution can possibly provide a better separation of the two types of distributions. For this reason we use tail quantiles as e-values.

We denote the q^{th} population quantile of \mathbb{E}_m by $c_q(\mathbb{E}_m)$. For this we have equivalent results to Theorem 3.1 as $n \to \infty$:

THEOREM 3.2. Given that the estimator $\hat{\theta}$ satisfies conditions (P1) and (P2), and the evaluation map satisfies conditions (E1)-(E4), we have

$$(3.3) c_q(\mathbb{E}_*) \to c_{q,\infty} < \infty$$

(3.4)
$$|c_q(\mathbb{E}_m) - c_q(\mathbb{E}_*)| \to 0 \text{ when } \mathcal{M} \text{ is adequate}$$

(3.5)
$$c_q(\mathbb{E}_m) \to 0 \text{ when } \mathcal{M} \text{ is inadequate}$$

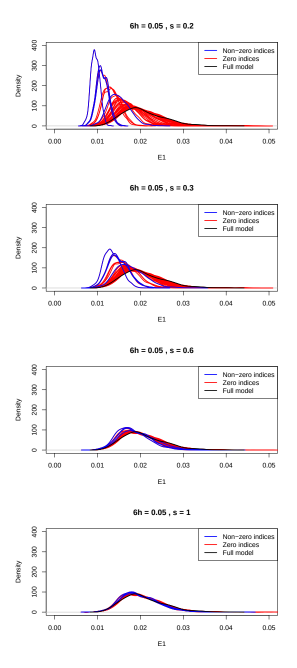


Fig 3.1: Density plots of bootstrap approximations for \mathbb{E}_* and \mathbb{E}_{-j} for all j in simulation setup, with s=0.2,0.3,0.6,1

When the q^{th} quantile is taken as the e-value instead of the mean, we set a lower detection threshold than the same functional on the full model, i.e. choose all j such that $c_q(\mathbb{E}_{-i}) < c_{qt}(\mathbb{E}_*), 0 < t < 1$ to be included in the model. The choice of t potentially depends on several factors such as the value of quantile evaluated, the statistical model used, sample size and degree of sparsity of parameters in the data generating process. We illustrate this point on simulated data in Section 4.

3.4. Bootstrap procedure. We use generalized bootstrap [4] to obtain approximations of the sampling distributions \mathbb{E}_{-i} and \mathbb{E}_* . It calculates bootstrap equivalents of the parameter estimate $\hat{\theta}$ by minimizing a version of the estimating equation in (3.1) with random weights:

(3.6)
$$\hat{\theta}_w = \arg\min_{\theta} \sum_{i=1}^n \mathbb{W}_i \Psi_i(\theta, B_i)$$

The resampling weights $(\mathbb{W}_1, \dots, \mathbb{W}_n)$ are non-negative exchangeable random variables chosen independent of the data, and satisfy the following conditions:

(3.7)
$$\mathbb{E}\mathbb{W}_1 = 1; \quad \mathbb{V}\mathbb{W}_1 = \tau_n^2 \uparrow \infty; \quad \tau_n^2 = o(a_n^2)$$

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$$\mathbb{E}W_1 = 1; \quad \mathbb{V}W_1 = \tau_n^2 \uparrow \infty; \quad \tau_n^2 = o(a_n^2)$$

(3.8) $\mathbb{E}W_1W_2 = O(n^{-1}); \quad \mathbb{E}W_1^2W_2^2 \to 1; \quad \mathbb{E}W_1^4 < \infty$

with $W_i := (\mathbb{W}_i - 1)/\tau_n$; $i = 1, \ldots, n$ being the centered and scaled resampling weights. Under standard regularity conditions on the estimating functional $\Psi(.)$ [4, 25] and conditional on the data, $(a_n/\tau_n)(\hat{\theta}_w - \hat{\theta})$ converges to the same asymptotic distribution as $a_n(\hat{\theta} - \theta_0)$, i.e. \mathbb{T}_0 .

We use empirical quantiles of the full model bootstrap samples as the quantile e-value estimates. Specifically, we go through the following steps:

- 1. Fix $q, t \in (0, 1)$;
- 2. Generate two independent set of bootstrap weights, of size R and R_1 , and obtain the corresponding approximations to the full model sampling distribution, say $[\hat{\theta}_r]$ and $[\hat{\theta}_{r_1}]$;
- 3. For $j = 1, 2, \dots p$ and estimate the e-value of the j^{th} predictor as the empirical q^{th} quantile of $\hat{\mathbb{E}}_{-i} := [E(\hat{\theta}_{r,-i}, [\hat{\theta}_{r,-i}])], \text{ with } \hat{\theta}_{r,-i} \text{ obtained}$ from $\hat{\theta}_r$ by replacing the j^{th} coordinate with 0;
- 4. Estimate the set of non-zero covariates as

$$\hat{\mathcal{S}}_0 = \{ j : c_q(\hat{\mathbb{E}}_{-j}) < c_{qt}(\hat{\mathbb{E}}_*) \};$$

Conditions (3.7) and (3.8) on the resampling weights ensure bootstrapconsistent approximation of the evaluation map quantiles:

THEOREM 3.3. Given the estimator $\hat{\theta}$ and evaluation map E in Theorem 3.2, and a generalized bootstrap scheme satisfying (3.7) and (3.8), we get

(3.9)
$$|c_q(\hat{\mathbb{E}}_m) - c_q(\hat{\mathbb{E}}_*)| \stackrel{P_n}{\to} o_P(1) \text{ when } \mathcal{M} \text{ is adequate}$$

(3.10)
$$c_q(\hat{\mathbb{E}}_m) \stackrel{P_n}{\to} o_P(1)$$
 when \mathcal{M} is inadequate

where P_n is probability conditional on the data.

Generalized bootstrap covers a large array of resampling procedures, for example the m-out-of-n bootstrap and a scale-enhanced version of the bayesian bootstrap. Furthermore, given that $\Psi_i(.)$ are twice differentiable in a neighborhood of θ_0 and some other conditions in [4], there is an approximate representation of $\hat{\theta}_w$:

(3.11)
$$\hat{\theta}_w = \hat{\theta} - \frac{\tau_n}{a_n} \left[\sum_{i=1}^n W_i \Psi_i''(\hat{\theta}, B_i) \right]^{-1} \sum_{i=1}^n W_i \Psi_i'(\hat{\theta}, B_i) + \mathbf{R}_{wn}$$

with $\mathbb{E}_w \|\mathbf{R}_{wn}\|^2 = o_P(1)$.

Given the full model estimate $\hat{\theta}$, and the score vectors $\Psi'_i(\hat{\theta}, B_i)$ and hessian matrices $\Psi''_i(\hat{\theta}, B_i)$, (3.11) allows us to obtain multiple copies of $\hat{\theta}_w$ through Monte-Carlo simulation of several arrays of bootstrap weights. This bypasses the need to fit the full model for each bootstrap sample, resulting in extremely fast computation of e-values.

We adapt the approximation of (3.11) to the LMM in (2.1). We first obtain the maximum likelihood estimates $\hat{\beta}_g$, $\hat{\sigma}_a^2$, $\hat{\sigma}_c^2$, $\hat{\sigma}_e^2$ through fitting the LMM. Then we replace the variance components in (2.2) with corresponding estimates to get $\hat{\mathbf{V}}_i$ for i^{th} pedigree, and aggregate them to get the covariance matrix estimate for all samples:

$$\hat{\mathbf{V}} = \operatorname{diag}(\hat{\mathbf{V}}_1, \dots, \hat{\mathbf{V}}_m)$$

We take m random draws from Gamma(1,1) - 1, say $\{w_{r1}, \ldots, w_{rm}\}$, as resampling weights in (3.11), using the same weight for all members of a pedigree. Consequently, the bootstrapped coefficient estimate $\hat{\beta}_{rg}$ has the following representation:

(3.12)
$$\hat{\beta}_{rg} \simeq \hat{\beta}_g + \frac{\tau_n}{\sqrt{n}} (\mathbf{G}^T \hat{\mathbf{V}}^{-1} \mathbf{G})^{-1} \mathbf{W}_r \mathbf{G}^T \hat{\mathbf{V}}^{-1} (y - \mathbf{G} \hat{\beta}_g)$$

with $\mathbf{G} = (\mathbf{G}_1^T, \dots, \mathbf{G}_m^T)^T$ and $\mathbf{W}_r = \operatorname{diag}(w_{r1}\mathbf{I}_4, \dots, w_{rm}\mathbf{I}_4)$. Finally we repeat the procedure for two independent sets of resampling weights, say of sizes R and R_1 , to obtain two collections of bootstrapped estimates $\{\hat{\beta}_{1g}, \dots, \hat{\beta}_{Rg}\}$.

4. Simulation. We now evaluate the performance of the above formulation of quantile e-values in a simulation setup. For this, consider the model in (2.1) with no environmental covariates. We consider familes with MZ twins and first generate the SNP matrices G_i . We take a total of $p_g = 50$ SNPs, and generate them in correlated blocks of 6, 4, 6, 4 and 30 to simulate correlation among SNPs in the genome. We set the correlation between two SNPs inside a block at 0.7, and consider the blocks to be uncorrelated. For each parent we generate two independent vectors of length 50 with the above correlation structure, and entries within each block being 0 or 1 following Bernoulli distributions with probabilities 0.2, 0.4, 0.4, 0.25 and 0.25 (Minor Allele Frequency or MAF) for SNPs in the 5 blocks, respectively. The genotype of a person is then determined by taking the sum of these two vectors: thus entries in G_i can take the values 0, 1 or 2. Finally we set the common genotype of the twins by randomly choosing one allele vector from each of the parents and taking their sum.

We repeat the above process for m=250 families. In GWAS there are generally a small number of causal SNPs, each explaining small proportions of the overall variability in response variable. To reflect this in our simulation setup, we assume that the first entries in each of the first four blocks above are causal, and each of them explains $h/(\sigma_a^2 + \sigma_c^2 + \sigma_e^2)\%$ of the overall variability. The term h is known as the heritability of the corresponding SNP (and can of course vary across SNPs). The value of the non-zero coefficient in k-th group: k=1,...,4, say β_k is calculated using the formula:

(4.1)
$$\beta_k = \sqrt{\frac{h}{(\sigma_a^2 + \sigma_c^2 + \sigma_e^2).2\text{MAF}_k(1 - \text{MAF}_k)}}$$

We fix the following values for the error variance components: $\sigma_a^2 = 4$, $\sigma_c^2 = 1$, $\sigma_e^2 = 1$, and generate pedigree-wise response vectors y_1, \ldots, y_{250} using the above setup. To consider different SNP effect sizes, we repeat the above setup for $h \in \{10, 7, 5, 3, 2, 1, 0\}$, generating 1000 datasets for each value of h.

- 4.1. Methods and metrics. For this simulated data, we compare our evalue based approach using the evaluation maps E_1 and E_2 in (3.2) with two other methods:
- (1) Model selection on linear model: Here we ignore the dependency structure within families by training linear models on the simulated data and selecting SNPs with non-zero effects by backward deletion using a modification of the BIC called mBIC2. This has been showed to give better results than

single-marker analysis in GWAS for unrelated individuals [11] and provides approximate False Discovery Rate (FDR) control at level 0.05 [2].

(2) Single-marker mixed model: We train single-SNP versions of (2.1) using a fast approximation of the Generalized Least Squares procedure (named Rapid Feasible Generalized Least Squares or RFGLS: [23]), obtain marginal p-values from corresponding t-tests and use the Benjamini-Hochberg (BH) procedure to select significant SNPs at FDR = 0.05.

With the e-value being the q^{th} quantile of the evaluation map distribution, we set the detection threshold value at the t^{th} multiple of q for some 0 < t < 1. This means all indices j such that q^{th} quantile of the bootstrap approximation of \mathbb{E}_{-j} is less than the tq^{th} quantile of the bootstrap approximation of \mathbb{E}_* get selected as the set of active predictors. To enforce stricter control on the selected set of SNPs we repeat this for $q \in \{0.5, 0.6, 0.7, 0.8, 0.9\}$, and take the SNPs that get selected for all values of q as the final set of selected SNPs.

Since the above procedure depends on the bootstrap standard deviation parameter s, we repeat the process for $s \in \{0.3, 0.15, \dots, 0.95, 2\}$, and take as the final estimated set of SNPs the SNP set $\hat{\mathcal{S}}_t(s)$ that minimizes fixed effect prediction error (PE) on an independently generated test dataset $\{(y_{test,i}, \mathbf{G}_{test,i}), i = 1, \dots, 250\}$ from the same setup above:

$$PE_{t}(s) = \sum_{i=1}^{250} \sum_{j=1}^{4} \left(y_{test,ij} - g_{test,ij}^{T} \hat{\beta}_{\hat{\mathcal{S}}_{t}(s)} \right)^{2};$$
$$\hat{\mathcal{S}}_{t} = \arg\min_{s} PE_{t}(s)$$

We use the following metrics to evaluate each method we implement: (1) True Positive (TP), which is the proportion of causal SNPs detected; (2) True Negative (TN), which is the proportion of non-causal SNPs undetected; (3) Relaxed True Positive (RTP), which is the: proportion of detecting any SNP in each of the 4 blocks with causal SNPs, i.e. for the selected index set by some method m, say \hat{S}_m ,

$$RTP(\hat{\mathcal{S}}_m) = \frac{1}{4} \sum_{i=1}^{4} \mathbb{I}(Block \ i \cap \hat{\mathcal{S}}_m \neq \emptyset)$$

and finally (4) Relaxed True Negative (RTN), which is the proportion of SNPs in block 5 undetected. We consider the third and fourth metrics to cover situations in which the causal SNP is not detected itself, but highly correlated SNPs with the causal SNP are. This is common in GWAS [11]. Finally, we average all the above proportions over 1000 replications, and repeat the process for two different ranges of t for E_1 and E_2 .

4.2. Results. We present the simulation results in table 4.1. For all heritability values, applying mBIC2 on linear models performs poorly compared to applying RFGLS and then correcting for multiple testing. This is expected because the linear model ignores within-family error components.

Our method works better than the two competing methods for detecting true signals across different values of h: the average TP rate going down slowly than other methods across the majority of choices for t. Both mBIC2 and RFGLS+BH have very high true negative detection rates, which is matched by our method for higher values of q. Since all reduced model distributions reside on the left of the full model distribution, we expect the variable selection process to turn more conservative at lower values of t. This effect is more noticeable for lower q, indicating that the right tails of evaluation map distributions are more useful for this purpose. Finally for h=0, we report only TN and RTN values since no signals should ideally be detected: in terms of this a value of q=0.9 or q=0.5 leads to the same TN and RTN performance as RFGLS+BH for all choices of t.

RTP performances for all methods are better than the corresponding TP/TN performances. However, for mBIC2 this seems to be due to detecting SNPs in the first four blocks by chance since for h=0 its RTN is less than TN. Also E_2 seems to perform slightly better than E_1 , in the sense that it yields a higher TP (or RTP) while having the same TN (or RTN) rates.

5. Analysis of the MCTFR datas. We now apply the above methods on SNPs from the MCTFR dataset. We assume a nuclear pedigree structure, and for simplicity only analyze pedigrees with MZ and DZ twins. After setting aside samples with missing response variables, we end up with 1019 such 4-member families. We look at the effect of genetic factors behind the response variable pertaining to the amount of alcohol consumption, which is highly heritable in this dataset according to previous studies [28]. We analyze SNPs inside some of the most-studied genes with respect to alcohol abuse: GABRA2, ADH1A, ADH1B, ADH1C, ADH4-ADH7, SLC6A3, SLC6A4, OPRM1, CYP2E1, DRD2, ALDH2, and COMT [8] through separate gene-level models. The ADH genes did not contain many SNPs individually, so we club all seven of them together. We include sex, birth year, age and generation (parent or offspring) of individuals as covariates to control for their potential effect.

For model selection we use E_2 as the evaluation function because of its slighty better performance in the simulations. For each gene-level model, We train the LMM in (2.1) on 75% of randomly selected families, perform our

E_2	E_1	 ¤	E_2	E_1	 ש
t = 0.8 t = 0.74 t = 0.68 t = 0.62 t = 0.56 t = 0.5	$t = \exp(-1)$ $t = \exp(-2)$ $t = \exp(-3)$ $t = \exp(-4)$ $t = \exp(-5)$	Method mBIC2 RFGLS+BH	t = 0.8 t = 0.74 t = 0.68 t = 0.62 t = 0.56 t = 0.5	$t = \exp(-1)$ $t = \exp(-2)$ $t = \exp(-3)$ $t = \exp(-4)$ $t = \exp(-5)$	Method mBIC2 RFGLS+BH
0.97/0.98 0.96/0.98 0.95/0.99 0.95/0.99 0.94/0.99 0.92/0.99	0.95/0.98 0.94/0.99 0.94/0.99 0.92/0.99 0.9/0.99	$h = 10 \\ 0.84/0.99 \\ 0.96/0.99$	0.97/0.98 0.96/0.98 0.95/0.99 0.95/0.99 0.94/0.99 0.92/0.99	0.95/0.98 0.94/0.98 0.94/0.99 0.92/0.99 0.9/0.99	h = 10 0.79/0.99 0.95/0.92
0.91/0.97 0.89/0.98 0.87/0.98 0.85/0.99 0.83/0.99 0.79/0.99	0.87/0.97 0.85/0.98 0.83/0.99 0.8/0.99 0.75/0.99	$h = 7 \\ 0.66/0.99 \\ 0.83/0.99$	0.9/0.97 0.88/0.97 0.87/0.98 0.84/0.98 0.82/0.99 0.79/0.99	0.87/0.97 0.85/0.98 0.82/0.98 0.79/0.99 0.75/0.99	h = 7 0.59/0.99 0.82/0.95
0.8/0.96 0.76/0.97 0.73/0.98 0.69/0.98 0.66/0.99 0.61/0.99	0.75/0.97 0.71/0.98 0.67/0.99 0.62/0.99 0.56/0.99	h = 5 $0.48/0.99$ $0.64/0.99$	0.79/0.96 0.75/0.97 0.72/0.98 0.68/0.98 0.65/0.99 0.6/0.99	0.74/0.97 $0.69/0.98$ $0.65/0.98$ $0.61/0.99$ $0.55/0.99$	h = 5 $0.41/0.99$ $0.62/0.97$
0.57/0.96 0.51/0.97 0.48/0.98 0.42/0.99 0.38/0.99 0.32/0.99	0.5/0.97 0.45/0.98 0.39/0.99 0.33/0.99 0.27/1	h = 3 $0.26/0.99$ $0.32/0.99$	0.54/0.96 0.48/0.97 0.45/0.98 0.4/0.99 0.36/0.99 0.31/0.99	0.47/0.97 $0.43/0.98$ $0.37/0.99$ $0.32/0.99$ $0.26/1$	h = 3 0.2/0.99 0.29/0.98
0.38/0.97 0.33/0.98 0.29/0.98 0.24/0.99 0.2/0.99 0.17/1	0.32/0.98 0.28/0.98 0.22/0.99 0.18/0.99 0.14/1	h = 2 $0.16/0.99$ $0.16/1$	0.34/0.97 0.29/0.98 0.26/0.98 0.26/0.99 0.19/0.99 0.16/1	0.28/0.97 $0.25/0.98$ $0.2/0.99$ $0.17/0.99$ $0.13/1$	$h = 2 \\ 0.11/0.99 \\ 0.14/0.99$
0.2/0.98 0.15/0.98 0.12/0.99 0.11/0.99 0.08/0.99 0.06/1	0.15/0.98 0.12/0.99 0.09/0.99 0.07/1 0.05/1	h = 1 $0.08/0.99$ $0.05/1$	0.15/0.98 0.12/0.98 0.1/0.99 0.09/0.99 0.07/1 0.05/1	0.12/0.98 0.09/0.99 0.07/0.99 0.06/1 0.04/1	h = 1 $0.05/0.99$ $0.04/1$
-/0.97 -/0.98 -/0.98 -/0.99 -/0.99	-/0.98 -/0.98 -/0.99 -/1 -/1	h = 0 -/0.98	-/0.99 -/0.99 -/0.99 -/0.99 -/1 -/1	-/0.99 -/0.99 -/1 -/1 -/1	h = 0 -/0.99 -/1

Table 4.1: (Top) Average True Positive (TP)/ True Negative (TN) rates for mBIC2, RFGLS+BH and the e-values method with E_1 and E_2 as evaluation maps and different values of t over 1000 replications, and (Bottom) Average Relaxed True Positive (RTP) and Relaxed True Negative (RTN) rates

Gene	Total no.	No. of SNPs detected by				
	of SNPs	e-value	RFGLS+BH	mBIC2		
GABRA2	11	5	0	0		
ADH	44	3	1	0		
OPRM1	47	25	1	0		
CYP2E1	9	5	0	0		
ALDH2	6	5	0	1		
COMT	15	14	0	0		
SLC6A3	18	4	0	0		
SLC6A4	5	0	0	0		
DRD2	17	0	0	1		
Table 5.1						

Table of analyzed genes and number of detected SNPs in them by the three methods

e-values procedure for $s = 0.2, 0.4, \dots, 2.8, 3, t = 0.1, 0.15, \dots, 0.75, 0.8$; and select the set of SNPs that minimizes fixed effect prediction error on the data from the other 25% of families over this grid of (s, t).

As seen in Table 5.1, our e-value based technique detects a much higher number of SNPs than the two competing methods. Our method selects all but one SNP in the genes ALDH2 and COMT. These are small genes of size 50kb and 30kb, respectively, thus SNPs within them have more chance of being in high Linkage Disequilibrium (LD). On the other-hand, it does not select any SNPs in SLC6A4 and DRD2. Variants of these genes are known to interact with each other and are jointly associated with multiple behavioral disorders [17, 37].

A number of SNPs we detect (or SNPs situated close to them) have known associations with alcohol-related behavioral disorders. We summarize this in Table 5.2. Prominent among them are rs1808851 and rs279856 in the GABRA2 gene, which are at perfect LD with rs279858 in the larger, 7188-individual version of our twin studies dataset [16]. This SNP is the marker in GABRA2 that is most frequently associated in the literature with alcohol abuse [9], but was not genotyped in our sample. A single SNP RFGLS analysis of the same twin studies data that used Bonferroni correction on marginal p-values missed the SNPs we detect [16]: highlighting the advantage of our approach. We give a gene-wise discussion of associated SNPs, as well as information on all SNPs, in the supplementary material.

We plot the 90th quantile e-value estimates in Figures 5.1, 5.2 and 5.3. We obtained gene locations, as well as the locations of coding regions of genes, i.e. exons, inside 6 of these 9 genes from annotation data extracted from the UCSC Genome Browser database [31]. Exon locations were not available for OPRM1, CYP2E1 and DRD2. In general, SNPs tend to get selected in groups with neighboring SNPs, which suggests high LD. Also

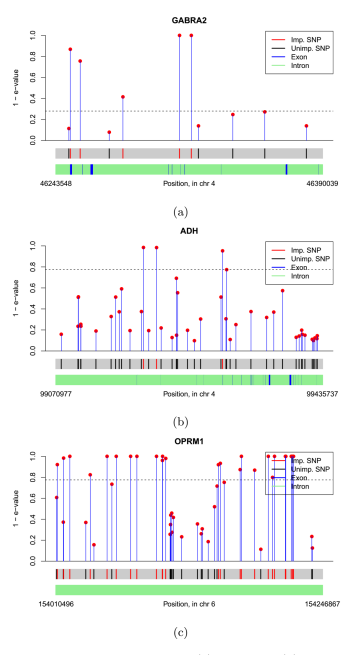


Fig 5.1: Plot of e-values for genes analyzed: (a) GABRA2, (b) ADH1 to ADH7, (c) OPRM1. For ease of visualization, 1-e-values are plotted in the y-axis.

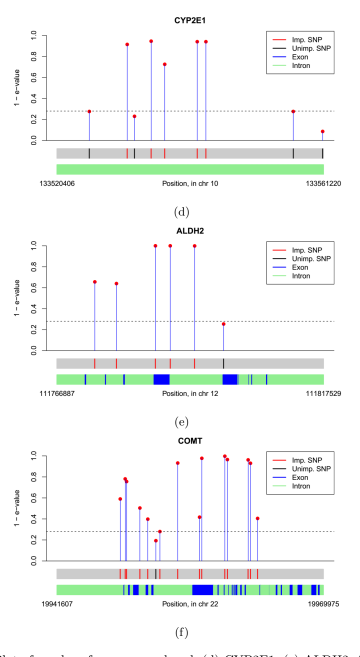


Fig 5.2: Plot of e-values for genes analyzed: (d) CYP2E1, (e) ALDH2, (f) COMT

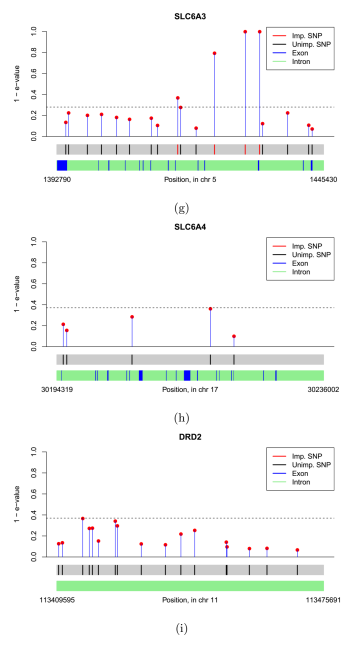


Fig 5.3: Plot of e-values for genes analyzed: (g) SLC6A3, (h) SLC6A4, (i) DRD2

Gene	Detected SNPs with	Reference for
	known associations	associated SNP
GABRA2	rs1808851, rs279856: close to	[9]
	rs279858	
ADH genes	rs17027523: 20kb upstream of	Multiple studies (https:
	rs1229984	//www.snpedia.com/
		index.php/Rs1229984)
OPRM1	rs12662873: 1 kb upstream of	Multiple studies (https:
	rs1799971	//www.snpedia.com/
		index.php/Rs1799971)
CYP2E1	rs9419624: 600b downstream of	[24]
	rs4646976; rs9419702: 10kb up-	
	stream of rs4838767	
ALDH2	rs16941437: 10kb upstream of rs671	Multiple studies
		(https://www.snpedia.
		com/index.php/Rs671)
COMT	rs4680, rs165774	[36]
SLC6A3	rs464049	[13]

Table 5.2
Table of detected SNPs with known references

most of the selected SNPs either overlap or in close proximity to the exons, which underline their functional relevance,

6. Discussion and conclusion. To expand the above approach to a genome-wide scale, we need to incorporate strategies for dealing with the hierarchical structure of causal SNPs: there are a few causal genes behind a quantitative phenotype, which can be further attributed to a proportion of SNPs inside each gene. To apply the e-values method here, it is plausible to start with an initial screening step to eliminate evidently non-relevant genes. Methods like the grouped Sure Independent Screening [22] and min-P test [38] can be useful here. Following this, in a multi-gene predictor set, there are several possible strategies to select important genes and important SNPs in them. Firstly, one can use a two-stage e-value based procedure. The first stage is same as the method described in this paper, i.e. selecting important SNPs from each gene using multi-SNP models trained on SNPs in that gene. In the second stage, a model will be trained using the aggregated set of SNPs obtained in the first step, and a group selection procedure will be run on this model using e-values. This means dropping groups of predictors (instead of single predictors) from the full model, checking the reduced model e-values, and selecting a SNP group only if dropping it causes the e-value to go below a certain cutoff. Secondly, one can start by selecting important genes using an aggregation method of SNP-trait associations (e.g. [21]) and then run the

e-value based SNP selection on the set of SNPs within these genes. Thirdly, one can also take the aggregated set of SNPs obtained from running the e-values procedure on gene-level models, then use a fast screening method (e.g. RFGLS) to select a subset of those SNPs.

We plan to study merits and demerits of these strategies and the computational issues associated with them in detail through synthetic studies as well as in the GWAS data from MCTFR. Finally, the current evaluation map based formulation requires the existence of an asymptotic distribution for the full model estimate. We plan to explore alternative formulation of evaluation maps under weaker conditions to bypass this, thus being able to tackle high-dimensional (n < p) situations.

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