

Potpourri: An Epistasis Test Prioritization Algorithm via Diverse SNP Selection

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ABSTRACT

Genome-wide association studies explain a fraction of the underlying heritability of genetic diseases. Investigating epistatic interactions between two or more loci help closing this gap. Unfortunately, sheer number of loci combinations to process and hypotheses to test prohibit the process both computationally and statistically. Epistasis test prioritization algorithms rank likely-epistatic SNP pairs to limit the number of tests. Yet, they still suffer from very low precision. It was shown in the literature that selecting SNPs that are individually correlated with the phenotype and also diverse with respect to genomic location, leads to better phenotype prediction due to genetic complementation. Here, we hypothesize that an algorithm that pairs SNPs from such diverse regions and carefully ranks the pairs can detect statistically more meaningful pairs and can improve prediction power. We propose an epistasis test prioritization algorithm which optimizes a submodular set function to select a diverse and complementary set of genomic regions that span the underlying genome. SNP pairs from these regions are then further ranked w.r.t. their co-coverage of the case cohort. We compare our algorithm with the state-of-the-art on three GWAS and show that (i) we substantially improve precision (from 0.003 to 0.652) while maintaining the significance of selected pairs, (ii) decrease the number of tests by 25 folds, and (iii) decrease the runtime by 4 folds. We also show that promoting SNPs from regulatory/coding regions improves the precision (up to 0.8). Potpourri is available at <http://ciceklab.cs.bilkent.edu.tr/potpourri>.

INTRODUCTION

Genome-wide association studies (GWAS) have been an important tool for susceptibility gene discovery in genetic disorders (1, 2, 3). Analyzing single loci associations have provided many valuable insights but they alone do not account for the full heritability (4). Investigating the interplay among multiple loci with has helped to bridge the missing heritability gap. Such statistically significant interactions between two or more loci is called epistasis and it has a major role in complex genetic traits such as cancer (5) and neurodevelopmental disorders (6).

Exhaustive identification of interacting loci, even just pairs, is intractable computationally. Moreover, such an approach

lacks statistical power due to multiple hypothesis testing. Several methods have been developed to circumvent this problem. TEAM and BOOST are exhaustive models which exploit data structures and efficient data representation to improve the brute force performance (7, 8). However, these methods still perform many tests and do not scale for large tasks. For instance, BOOST reports a runtime of 60 hours for 360k SNPs. Another approach is to reduce the search space by filtering pairs based on a type of statistical threshold. Examples include SNPHarvester (9), SNPRuler (10) and IBBFS (11). Despite their advantages, these methods mostly do not follow a biological reasoning and tend to detect interactions that are in linkage disequilibrium (LD) as noted in (12). On another track, incorporating biological background and testing the SNP pairs that are annotated has also proven useful (13, 14, 15, 16, 17). Yet, this approach requires most SNPs to be discarded as many are quite far away from any gene to be associated. Moreover, this introduces a literature bias in the selections of the algorithms.

A rather more popular approach is to prioritize the tests to be performed rather than discarding pairs from the search space and controlling for Type-I error. In this approach, the user can keep performing tests in the order specified by the algorithm until a desired number of significant pairs are found. The idea is to provide the user with a manageable number of true positives (statistically significant epistatic pairs) while minimizing the number of tests to ensure statistical power. The first algorithm of this kind is iLOCi (12), which ranks SNP pairs by performing a dependence test and avoiding pairs that are unrelated to disease but might seem related due to LD. This work was followed by (18) who proposed testing pairs of SNPs in *population covering locus sets* - POCOs. First, the algorithm greedily selects multiple exclusive groups of SNPs that *cover* all affected individuals. That is, each case sample has to have at least one SNP in each POCO. Epistasis tests then are performed across POCOs with the hope that independent coverage of the cases will lead different POCOs to include complementary and epistatic SNPs (18, 19). Finally, Cowman and Koyuturk, introduced the LINDEN algorithm (20). First, in a bottom-up fashion, the method generates SNP trees on greedily selected genomic regions (LD forest). Each node represents the genotypes of cases and controls for the SNPs in that node. Then, by comparing the roots of these trees, it decides if this pair of regions is a promising candidate for epistasis test. Nodes in lower levels are continued to be checked and leaf pairs (individual SNPs) are tested only if the estimation at higher levels is promising. LINDEN was shown to achieve the state of the art results. Despite using various heuristics, all methods still have high false discovery rates. For instance, the FDR of LINDEN ranges from 0.96 to 0.998 on

three GWAS from Wellcome Trust Case Control Consortium (WTCCC) - the ratio of significant pairs to the number of detected (reciprocally) significant epistatic pairs (20).

Linkage disequilibrium is an important source of information for epistasis prioritization algorithms. Two SNPs that appear to be interacting statistically, might not be biologically meaningful if they are on the same haplotype block. For this reason, all three methods mentioned above focus on such regions and aim at avoiding testing pairs that are located in close vicinity of each other. In an orthogonal study, (21) propose a feature (SNP) selection algorithm which avoids LD for better phenotype prediction. Authors show that while looking for a small set of loci (i.e., 100) that is the most predictive of a continuous phenotype, selecting SNPs that are far away from each other, results in better predictive power. This method, SPADIS, is designed for feature selection for multiple regression. As the SNP set it generates contains diverse and complementary SNPs, it results in better R^2 values by covering more biological functions.

Inspired by this idea, we conjecture that selecting pairs of SNPs from genomic regions that (i) harbor individually informative SNPs, and (ii) are diverse in terms of genomic location would avoid LD better and yield more functionally complementing and more epistatic SNP pairs compared to the current state of the art since no other algorithm exploits this information. We propose a new method that for the first time incorporates the genome location diversity with the population coverage density. Specifically, our proposed method, *Potpourri*, maximizes a submodular set function to select a set of genomic regions (i) that include SNPs which are individually predictive of the cases, and (ii) that are topologically distant from each other on the underlying genome. Epistasis tests are performed for pairs across these regions, such that pairs that densely co-cover the case cohort are given priority.

We validate our hypothesis and show that *Potpourri* is able to detect statistically significant and biologically meaningful epistatic SNP pairs. We perform extensive tests on three Wellcome Trust Case Control Consortium (WTCCC) GWA studies (Type 2 Diabetes (T2D), Bipolar Disorder (BD), and Hypertension (HT)), and compare our method with the state-of-the-art LINDEN algorithm. First, we *guide* LINDEN by pruning its search space using *Potpourri*-selected-SNPs to show that (i) it is possible to significantly improve the precision (from 0.003 up to 0.302) and (ii) that our diversification approach is sound. Then, we show that the ranking of the diverse SNPs by the co-coverage of the case cohort further improves the prediction power and the precision (up to 0.652 in the selected setting). *Potpourri* drastically reduces the number of hypothesis tests to perform (from $\sim 380k$ to $\sim 15k$), and yet is still able to detect more epistatic pairs with similar significance levels in all three GWA studies considered. The running time is also cut by 4 folds in the selected settings. Another problem with the current techniques is the biological interpretation of the obtained epistatic pairs. Once the most significant SNP pairs returned are in the non-coding regions and are too isolated to be associated with any gene, the user can hardly make sense of such a result despite statistical significance. We investigate the advantage of promotion of SNPs falling into regulatory and non-coding regions for testing and propose three techniques. We show

that these techniques further improve the precision (up to 0.8) with similar number of epistatic pairs detected. Finally, we investigate the biological meaning of the detected SNP pairs. We find (i) a SNP pair which supports the hypothesis of a shared genetic architecture between T2D and chronic kidney disease; and (ii) a pair which suggests a new candidate risk gene (*NPW*) for HT which has loose indications only in rat studies.

MATERIALS AND METHODS

Notation

A GWAS dataset consists of genotypes of a set of samples S who are associated with a binary phenotype: *Case* or *Control*. Let $f(s)$ be an indicator function that corresponds to phenotype of a sample $s \in S$, i.e.: $f(s)=1$, if s is a case sample, and $f(s)=0$, otherwise. Function h which represents the genotype of sample $s \in S$ at locus $v \in V$ is encoded as:

$$h(s,v) = \begin{cases} 0, & \text{if genotype of sample } s \text{ at } v \text{ is Homozygous reference} \\ 1, & \text{if genotype of sample } s \text{ at } v \text{ is Heterozygous} \\ 2, & \text{if genotype of sample } s \text{ at } v \text{ is Homozygous alternative} \end{cases}$$

One can generate an undirected SNP-SNP network $G(V', E)$, where $V' \subseteq V$ and $v_i \in V'$ if $\exists s \in S$ where $h(s, v_i) \neq 0$. $e_{v_i, v_j} \in E$ where $v_i, v_j \in V$ and they are *related*. The definition of *relatedness* might change with respect to the application and the biological definition. In our setting, two SNPs are related if they are neighbors on the underlying genome.

Selection of Diverse and Informative SNP Regions

Regions harboring informative SNPs (ones that are correlated with the phenotype) and also far away on the underlying genome with respect to a SNP-SNP network is likely to yield a diverse and explanatory SNP set without introducing a literature bias. Our hypothesis is that those selected SNPs are likely to be epistatic because SNPs selected using this technique leads to better phenotype prediction due to better genomic complementation (21). Hence, carefully picking pairs from such a set should yield highly epistatic SNP pairs and is well situated for epistasis test prioritization. In our application the goal is not to predict the phenotype. Our SNP set selection approach differs from (21) in the calculation of the SKAT scores (22) as in epistasis test prioritization we are analyzing a GWA study in which the phenotype is dichotomous. Moreover, we introduce an hyper-parameter which can assign extra artificial price to SNPs that fall in regulatory or coding regions to give priority to SNPs which are more likely to have functional effect. Also, node prizes are normalized by the set size to be able to compare performances of selected sets of different sizes.

First, for every SNP v_i , a score c_{v_i} is calculated using SKAT method which works by regressing phenotype on the covariates using a flexible semiparametric linear model (22). Instead of directly associating genotypes of the variants with the phenotype, SKAT uses a nonparametric function of the genotypes that is possibly contained in a vector space generated by a positive semi-definite kernel function. Given

a user specified number SNPs to select (k), the second step selects a subset of loci $EP \subset V'$ such that it maximizes the sum of selected SNP scores while penalizing SNPs that are in close vicinity of each other. In particular the function F as shown in Equation 1 is maximized:

$$F(EP) = \sum_{v_i \in EP} \frac{c_{v_i} \omega_{v_i}}{k} + \beta \left(1 - \sum_{v_i, v_j \in EP; v_i \neq v_j} \frac{D - d(v_i, v_j)}{2kD} \right) \quad (1)$$

In this equation, $D \in \mathbb{R}_{>0}$ is a parameter that sets the upper limit on the distance that two SNPs are considered *close*. $d(v_i, v_j)$ is the distance between vertices $v_i, v_j \in EP$ on graph G . In this application function $d(v_i, v_j)$ denotes the shortest path distance between v_i, v_j only if $d(v_i, v_j) \leq D$, otherwise $d(v_i, v_j) = D$ to cancel out the penalty term. The parameter $\beta \in \mathbb{R}_{\geq 0}$ adjusts the relative magnitudes of the prices and the penalties. Finally, the parameter $\omega_{v_i} \in \mathbb{R}_{\geq 1}$ rewards $v_i \in V'$ if it falls into a regulatory or coding region of interest. If this information is not taken into consideration $\omega_{v_i} = 1, \forall v_i \in V'$ - see Subsection *Promoting SNPs in Regulatory and Coding Regions*. The subset EP^* that maximizes F is selected as the seed set for epistasis prioritization as shown in Equation 2.

$$EP^* = \underset{EP \subseteq V' \subseteq V, |EP|=k}{\operatorname{argmax}} F(EP) \quad (2)$$

Given the set of SNPs V' , the above-mentioned function $F: 2^{V'} \rightarrow \mathbb{R}$ is a submodular function (see Supplementary Text 1.1 for the proof). Submodular optimization is NP-hard. However, the greedy algorithm given in Supplementary Text 1.2, which basically iteratively adds the next *best* SNP to the set that maximizes F at each step, ensures a $(1 - \frac{1}{e})$ -factor approximation to the optimum solution (23). This algorithm requires the submodular set function to be also monotone non-decreasing and non-negative, which are proven in Supplementary Text 1.3.

Note that using only the SNPs in EP^* would restrict the epistasis testing to SNPs which are individually correlated with the trait. We rather use those as anchors to detect genomic regions of interest. Let R be a set of consecutive SNPs that fall into a region in the genome. After EP^* is set, for every SNP $v_i \in EP^*$, a region R_i is formed such that it includes, v_i and m upstream and m downstream SNPs of v_i . Thus, $|R_i| = 2m + 1$. \mathbf{R} is the set of sets (regions) $R_i, \forall v_i \in EP^*$. Note that $\bigcup_{R_i \in \mathbf{R}} R_i = 2mk + k$.

Prioritization of the Tests

The region set \mathbf{R} generates a pruned search space for finding likely epistatic SNP pairs. Within region tests are avoided in order to avoid LD. That is, SNPs v_i and v_j are not tested if both $v_i \in R_x$ and $v_j \in R_x$ such that $R_x \in \mathbf{R}$. Still, the number of possible tests is $\frac{k * (k-1)}{2} (2m+1)^2$ and a prioritization scheme is needed to rank candidate pairs for testing. We employ two strategies as explained next.

Guiding LINDEN for Ranking SNP Pairs In this ranking strategy, we use LINDEN to rank the pairs selected in \mathbf{R} . That is, we use Potpourri selected SNPs to guide LINDEN. Thus, in this strategy, Potpourri acts as a preprocessing step for LINDEN to prune its large search space. Doing so, we test if our diverse SNP selection scheme is sound and whether it improves the performance of LINDEN.

LINDEN is input with only SNPs that fall into the regions in \mathbf{R} . Then, the algorithm is run as described in (20). That is, the algorithm forms LD trees over greedily selected regions. The leaves of these trees represent actual SNPs each of which are represented by a sample genotype vector (i.e., the genotype of this SNP for every sample). The tree is constructed in bottom-up fashion and nodes are merged to generate higher levels. During this process, sample genotype vectors are merged and ambiguous indices (i.e., samples with different genotypes) are assigned a *NIL* value. This merging step continues until a threshold d is met that denotes the fraction of *NIL* values allowed. This is a dynamically adjusted threshold goes up with the number of iterations. Then, the tree pairs are tested with respect to the sample genotype vectors starting from the root, ignoring the *NIL* values. Lower levels are tested only if the significance of the chi-squared test meets a certain threshold. In parallel with our hypothesis on diversification of the regions, we prohibit LINDEN to merge SNPs from different regions. Thus, the tests are performed across regions in \mathbf{R} .

Population Co-covering For Ranking SNP Pairs We propose a new strategy which aims at maximizing the population coverage of co-occurring SNPs. Population cover for epistasis test prioritization was first proposed by (18). This approach selects multiple exclusive groups of SNPs (*POCOs*) that *covers* the case cohort. That is, the union of the samples with the SNPs in a POCO should be equal to the case cohort. Epistasis tests are performed across POCOs and the idea is that the independent coverage of the cases across POCOs will result in detecting complementary SNPs. We take a different approach in terms of covering the population. We would like the SNP pairs to be tested to co-cover the population. This is intuitive; the diverse selection step finds complementing regions and avoids LD, but for a SNP pair to be epistatic they also need to be observed together in cases. More formally, let $p(v_x, v_y)$ be a function that scores the SNP pair $v_x, v_y \in V'$ for testing. Given three SNPs v_x, v_y and $v_z \in V'$, $p(v_x, v_y) > p(v_x, v_z)$ if and only if v_x is observed more frequently with v_y in cases as compared to v_z . p is formally defined as follows:

$$p(v_x, v_y) = \sum_{s \in S; f(s)=1} \mathbb{1}_{(h(s, v_x) \neq 0 \wedge h(s, v_y) \neq 0)} \quad (3)$$

Potpourri computes the pairwise population co-covering of all SNP pairs (SNP_i, SNP_j) such that $SNP_i \in R_x, SNP_j \in R_y, \forall R_x, R_y \in \mathbf{R}, x \neq y$ and $i \neq j$. Testing is then performed in the descending population co-cover order. The algorithm is restricted to test top v pairs among all region pairs in \mathbf{R} .

Promoting SNPs in Regulatory and Coding Regions SNPs can alter gene expression and the downstream function depending on their genomic location. Those that fall on to regulatory

regions can affect mRNA levels and those that fall onto coding regions can alter the structure and function of the protein. Since such SNPs are likely to alter the function and more likely to be related to the phenotype, we conjecture that we can find more statistically significant and biologically meaningful SNP pairs via promoting mutations in regulatory and coding regions. While this might introduce a literature bias, for a life scientist who would like to narrow down the search space using functional regions this might be plausible and investigate the usefulness of this approach. However, one should not totally disregard the unannotated parts of the genome as most of the variation exists in such regions. Thus, we seek a balance.

We employ 3 techniques to *promote* regulatory/coding variants. In the first approach (Potpourri RC1), we assign an artificial prize to SNPs in the diverse SNP selection phase of the algorithm as described in the *Selection of Diverse and Informative SNP Regions* Section. This approach favors the regions in R to be regulatory/coding regions. Then, the pairs are tested with respect to the population co-cover ranking. The second approach (Potpourri RC2), uses $\omega_i = 1, \forall v_i \in V'$. That is, the SNP selection step does not promote selection of coding/regulatory SNPs. Rather, it first gathers SNP pairs such that at least one SNP falls into regulatory/coding regions and then rank these pairs with respect to the population co-cover. After all such pairs are tested, the algorithm test remaining pairs with respect to their population co-cover ranking. The final strategy (Potpourri RC3) employs both RC1 and RC2 strategies at the same time.

We use three functional region types which are (i) distant-acting transcriptional enhancer regions, (ii) promoter regions (1kb downstream and upstream of the transcription start sites - TSS) and (iii) coding regions.

RESULTS

Datasets

In order to benchmark Potpourri, we used three GWAS datasets obtained from Wellcome Trust Case Control Consortium (WTCCC): (i) Type 2 diabetes (T2D), (ii) Hypertension (HT), and (iii) Bipolar disorder (BD). We use the 1958 Birth (58C) cohort as control for all datasets (24). As exercised in several articles (25, 26), we took the following quality control and preprocessing steps using PLINK tool (27):

- **Gender assignment check:** The subjects, in which the chromosome X data conflicts with the gender reported, were removed from the datasets.
- **Removing individuals with high missing genotype data:** We removed the individuals with more than 10 percent missing genotype rate from the datasets.
- **Removing rare SNPs:** Once we excluded the individuals with poor quality, we removed SNPs with less than 5 percent minor allele frequency.
- **Removing SNPs with high missing genotype rate:** We removed SNPs with more than 10 percent missing genotype rate from the datasets.

- **Removing SNPs that fail Hardy-Weinberg Equilibrium (HWE) test:** We removed SNPs that fail to pass HWE with a nominal p-value threshold $1e-6$.

After preprocessing the datasets, remaining number of SNPs, as well as number of cases and number of controls used are given in Supplementary Table 1.

We used the following resources to obtain the regulatory and coding region information. We obtained the distant-acting transcriptional enhancer dataset from VISTA Enhancer Browser (28). VISTA enhancer dataset contains 1912 human noncoding fragments with gene enhancer activity. Transcription start sites (TSSs) and coding region coordinates were obtained from UCSC Genome Browser (29). We chose Ensembl Genes as gene annotation track (30) and the March 2006 NCBI36/hg18 assembly of the human genome which matches the WTCCC datasets. The number and types of genes obtained from the Ensembl dataset are given in Supplementary Table 2. We defined 1 Kb downstream and upstream of the TSSs as a regulatory region. The coding region for a gene begins from the first base of the first exon and continues to the last base of the last exon.

Potpourri operates on a SNP-SNP interaction network. In this study, we used the genomic sequence (GS) network as defined in (31). In this network, SNPs are connected if they are adjacent on the genome. This was the network of choice as it shown to provide the best regression performance in (21).

Experimental Setup

For Potpourri, the parameters were selected using a nested 10-fold cross validation. First, the distance parameter D was selected via a line search in among 7 values (log-scale) between 1 and D_{max} (a value for which the distance penalty for all SNPs in the selected set is 0). D value that maximizes the L2-regularized logistic regression performance was picked. Then, 16 β values between 10^{-4} and a maximum $\beta = 2kD_{max}$ were tried. Again, the β value that maximizes the classification performance was picked. We experimented with the following k values: 500, 750, 1000, 1500, and 2000. Overall, best results were obtained when $k=750$ and it was set as the default parameter setting. Then, we added $m=9$ upstream and $m=9$ downstream neighbors SNPs of the for further coverage analysis. Unless otherwise stated, we set the $w_i = 1, \forall v_i \in V'$. Once the regulatory and coding regions were considered by the diverse SNP selection part, $w_i > 1, \forall i \in RC$ where RC contains SNPs that fall into regulatory and coding regions. We experimented with the three ω values: $1+10^{-0.5}$, $1+10^{-0.25}$, and 2.

Unless otherwise stated, we used the suggested settings for LINDEN as described by (20). That is, we set the parameter d to 0.45, which determines the fraction of ambiguous samples; and parameter b to 10, which determines the extent of LD. We run LINDEN by setting the maximum number of threads to 20 in parallel setting. The input data for LINDEN was also preprocessed as described in the *Datasets* Section.

When guiding LINDEN with our approach, we used default parameters for LINDEN. The only exception was that, we limited the extent of LD in the first two iterations of the merging procedure of LINDEN as explained in the *Guiding LINDEN for Ranking SNP Pairs* Section. That is, LINDEN's

merging step could merge SNPs from trees only within selected regions that contain of $2m+1$ SNPs ($m=9$ in our experiments). We used the above-mentioned parameter selection techniques for Potpourri. For the population co-cover, we performed epistasis test for the top $v=10$ SNP pairs among regions in **R**.

To quantify the performance of the proposed algorithms, we used precision as the evaluation metric in which true positives (TP) refer to the reciprocally significant epistatic pairs that pass the Bonferroni-adjusted statistical significance threshold. False positives (FP) refer to failed tests: the reciprocally significant epistatic pairs that fail to pass the aforementioned threshold. We use the definition of *reciprocally significant epistatic pair* from (20). As the authors argue, most epistatic pairs are dominated by some hub SNPs and this leads to detection of redundant pairs. On the other hand, SNPs in a reciprocal pair are the most epistatic partner for each other. We also use this definition to measure the performance of our algorithm. We set the significance level as 10% throughout experiments and adjust the significance level using the Bonferroni correction based on the number of test performed by each approach. Epistasis testing is performed via a chi-squared test on a 9×2 contingency table of all genotype combinations between cases and controls for a selected SNP pair ($df = 8$) as also done by (20). All tests are performed on Intel(R) Xeon(R) CPU E5-2650 v3 with two 2.30GHz processors. 251 GB RAM is used in the parallel setting.

Guiding LINDEN with Diverse and Informative Regions Improves Precision

We quantified the improvement in precision when LINDEN is guided by Potpourri-selected regions. First, we ran LINDEN on all datasets and it detected 1786, 885, and 1022 reciprocally significant epistatic pairs for T2D, BD, and HT datasets, respectively. Only 5, 35, and 2 pairs were statistically significant at the 10% level after Bonferroni correction, respectively. These numbers correspond to precision values of 0.003, 0.04, and 0.002, respectively. These results set our baseline. Then, we ran Potpourri on these datasets with top k SNPs, where $k=500, 750, 1000, 1500$ and 2000. We obtained 5 **R** sets. We then *guide* LINDEN with these regions and ran it as explained in the *Guiding LINDEN for Ranking SNP Pairs* Section.

Complete results are shown in Table 1, Supplementary Table 3, and Supplementary Table 4 for T2D, BD and HT datasets, respectively. The guidance of Potpourri improves the precision substantially, from 0.003 to 0.302 when $k=750$ in the T2D dataset (up to .421 when $k=500$). This is achieved by drastically reducing the number of false positives, and also increasing the number of true positives. Our pipeline outperforms LINDEN for all k values on all datasets, but we observe that the ideal k values are 500, 750 and 1000. Similar precision increases are also observed for BD and HT datasets.

The significance levels of each performed epistasis test by LINDEN and Potpourri-guided LINDEN are shown in Figure 1, Supplementary Figure 1, and Supplementary Figure 2 for the T2D, BD and HT datasets, respectively. The green lines denote the significance level (0.1) to be passed for each approach ($k=750$) after Bonferroni correction. Each point represents a SNP pair and the ones below threshold

are false positives and the ones above are true positives (reciprocally significant epistatic SNP pairs). It is clear that the pipeline drastically reduces the number of false positives while increasing the number true positives. Also, we can observe the importance of the number of tests performed by looking at the difference between Bonferroni corrected significance thresholds. Since Potpourri provides a pruned search space for LINDEN by eliminating SNPs that are most likely irrelevant to the trait, it also reduces number of tests that will be performed during epistasis test. Indirectly, it eliminates the negative effect of multiple hypothesis testing which reduces the statistical power. As seen in the figures, due to low Bonferroni threshold, the pipeline is able to discover more true positives compared to LINDEN. We also show that our pipeline not only minimizes the number of false positives but is also able to maintain the significance level of the returned pairs. While LINDEN's top SNP pair is more significant, *guided* LINDEN's top SNP pairs still stand out in terms of their p-values.

Table 1. Results for T2D dataset that compares LINDEN, Potpourri-guided LINDEN and Potpourri (with population co-cover). Number of pairs reported is the total number of reciprocally significant pairs returned by each method for varying number of selected SNPs. The number in parentheses denotes the significant pairs passing significance threshold (0.1) after Bonferroni correction based on the number of tests performed by each method. Bold denotes the best result for a given k value. The table shows that the guidance of Potpourri substantially improves the precision of LINDEN. For all considered k values, Potpourri provides the best precision values.

Method		# Tested Loci	# Reported (Rec. Sig.)	Precision
LINDEN		378016	1786 (5)	0.003
Potpourri guided LINDEN	$k = 500$	9250	19 (8)	0.421
	$k = 750$	14146	43 (13)	0.302
	$k = 1000$	18943	55 (21)	0.382
	$k = 1500$	28014	79 (18)	0.228
	$k = 2000$	37927	95 (11)	0.116
Potpourri	$k = 500$	9250	12 (8)	0.667
	$k = 750$	14145	23 (15)	0.652
	$k = 1000$	18943	31 (20)	0.645
	$k = 1500$	28014	50 (21)	0.420
	$k = 2000$	37927	74 (18)	0.243

In short, by just providing a guidance to LINDEN using Potpourri's diverse and informative SNP region selection, we were able to improve the performance of the state-of-the-art with substantially smaller number of tests performed. Next, we provide the results of the complete Potpourri pipeline which uses population co-cover strategy for prioritization, instead of LINDEN's tree based strategy.

Comparison of Potpourri with the State-of-the-art

In this section, we evaluate the performance of the Potpourri pipeline with population co-covering technique for prioritizing the tests. Again, we compare the performance with the LINDEN algorithm. Table 1 shows the results of the algorithm for different k values on the T2D dataset. We observe that population co-cover ranking strategy results in even further performance improvements. The precision is moved up to 0.652, as 15 out of 23 reciprocally significant pairs passes the Bonferroni-corrected threshold ($k=750$). See Supplementary Tables 3 and 4 for the results on BD and HT datasets, respectively, which follow the same pattern.

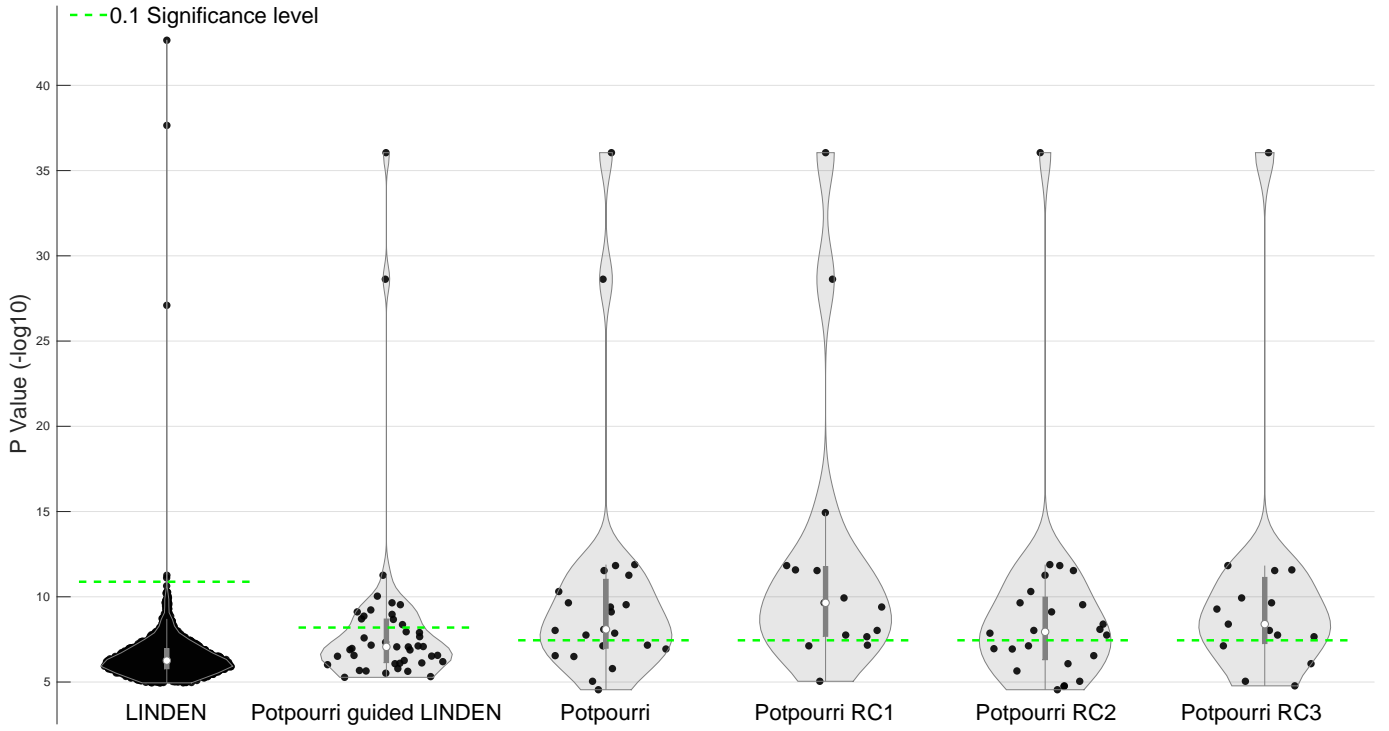


Figure 1. On the T2D dataset, this figure compares the p-values of the selected SNP pairs by the following methods: (i) LINDEN, (ii) Potpourri *guided* LINDEN, (iii) Potpourri; and then also the variants of Potpourri which further promotes SNP pairs in regulatory and coding regions: (iv) Potpourri RC1, (v) Potpourri RC2, and finally (vi) Potpourri RC3. We show the significance levels (y-axis) of each reported pair (dots) given the Bonferroni-corrected significance threshold (0.1, green dashed lines). X – axis is just randomly assigned values to pairs for better visualization. Potpourri is run with $k = 750$ for all 5 related subplots. For RC1 and RC3 strategies ω is set to 1.31623, for Potpourri and RC2, it is set to 1. LINDEN is run with default parameters. For Potpourri *guided* LINDEN, tree formation is restricted to be done on distinct regions in Potpourri-selected regions **R** as described in the *Guiding LINDEN for Ranking SNP Pairs* Section.

LINDEN performs 7,629,272,394 tests (378,016 tests for leaves) as opposed to 14,146 tests performed by Potpourri. This sets a much conservative significance threshold. One could argue that the precision gain is only due to the reduced significance threshold. However, Figure 1 shows that it is not the case. Third panel shows the significance levels of Potpourri-selected SNP pairs on the T2D dataset. We see that 6 out of 15 Potpourri-selected reciprocally significant pairs are significant even when the Bonferroni-corrected threshold of LINDEN is considered (dashed green line on the left-most panel which is far more conservative). Note that LINDEN only detects 5 reciprocally significant pairs. We also observe that the number of false positives are even further decreased compared to Potpourri-guided LINDEN. See Supplementary Figures 1 and 2 for the results on BD and HT datasets, respectively, which again show a similar pattern.

We also checked if adjusting LINDEN's parameters to make it more conservative would improve the precision. Increasing d to .9 and to .99 enforced it to perform a smaller number of tests but still too many to get close to Potpourri's efficiency, as shown in Supplementary Table 5.

Promoting Regulatory and Coding Regions Improves Precision

In this section, we investigate the advantage of promoting coding and regulatory regions in the Potpourri pipeline. For the T2D dataset, Table 2 compares the performance of original Potpourri pipeline with the three variants (RC1, RC2

and RC3) that promote selection of SNPs from coding and regulatory regions. Results show that in the suggested setting ($k = 750$) RC1 technique can increase the precision up to 0.8 and the original Potpourri pipeline cannot achieve a better result in any of the k values. Last 3 panels in Figure 1 show the significance levels of tested pairs. For the RC1 technique, we see that while we keep the most significant pairs detected by the original pipeline, we also discover new significant pairs (i.e., $p < 10^{-15}$). Moreover, the 2 out of 3 pairs that failed to pass the significance threshold are borderline. 7 out of 15 reciprocally significant pairs are also significant with respect to LINDEN's stringent significance threshold while LINDEN was able to detect only 5 such pairs and Potpourri was able to detect 6. Supplementary Tables 10 and 13 provide similar results for the BD and HT datasets, respectively. The margin of improvement is relatively low in the HT dataset (up to ~ 0.2). Supplementary Figures 1 and 2 compare the significance levels of the detected pairs for the BD and HT datasets, respectively.

We experimented with three ω values and picked the 1.31623 as the best performing and suggested parameter. It is used to produce above-mentioned figures and tables. See Supplementary Tables 6 - 9, 11, 12 for all results obtained using other parameter choices. Finally, selected SNP pairs by all Potpourri variant techniques with suggested settings ($k = 750$ and $\omega = 1$ or 1.31623) are listed in Supplementary Tables 14 - 25.

Table 2. Comparison of the Potpourri pipeline with three strategies (RC1, RC2, and RC3) to promote regulatory and coding regions on the T2D dataset. For RC1 and RC3 ω is set to 1.31623. Number of pairs reported is the total number of reciprocally significant pairs. The number in parentheses denotes the significant pairs passing significance threshold (0.1) after Bonferroni correction based on the number of tests performed by each method. Results show that promoting regulatory and coding regions improves Potpourri results substantially in all strategies, RC1 performing the best. Bold denotes the best result for a given k value.

Method		# Tested Loci	# Reported (Rec. Sig.)	Precision
Potpourri	$k = 500$	9250	12 (8)	0.667
	$k = 750$	14145	23 (15)	0.652
	$k = 1000$	18943	31 (20)	0.645
Potpourri RC1	$k = 500$	9489	17 (13)	0.765
	$k = 750$	13727	15 (12)	0.800
	$k = 1000$	15500	21 (12)	0.571
Potpourri RC2	$k = 500$	9250	13 (7)	0.538
	$k = 750$	14145	24 (14)	0.583
	$k = 1000$	18943	33 (23)	0.697
Potpourri RC3	$k = 500$	9489	18 (11)	0.611
	$k = 750$	13727	15 (11)	0.733
	$k = 1000$	15500	20 (13)	0.650

Novel Epistatic Pairs

One interesting pair discovered by Potpourri is the *rs12548378* - *rs10787472* pair that is detected for T2D. The former falls into *TCF7L2* gene and the latter falls into long intergenic non-coding RNA LINC01111. On one hand, *TCF7L2* is a well known type 2 diabetes susceptibility gene, which affects the blood glucose balance by regulating of proglucagon gene expression over the WNT signaling pathway (32). Its other epistatic interactions were also reported in the literature (33). On the other hand, LINC0111 was shown to be related to urine creatinine level in UK Biobank samples (genome-wide significant, $p = 5.28E - 09$) (34). Elevated levels of urine albumin-to-creatinine ratio (UACR) is a biomarker for diabetic nephropathy (DN.) DN occurs as a result of the damage to renal nerves due to high glucose levels in blood (35). While, T2D causes nephropathy over nerve damage, it has also been long debated that two might have overlapping genetic architectures. At least four T2D loci are associated with kidney function in American Indians, and two are related to kidney disease partially independently of T2D (36). (37) reports that a variant in *TCF7L2* causes renal dysfunction but this affect is not independent of T2D in south Indian population. (38) reports that a variant in *TCF7L2* is significantly more frequent in patients with diabetic nephropathy compared to non-diabetic nephropathy. Finally, an interaction between *EGFR* and *RXRG* genes was reported which increases the risk of DN in a T2D cohort (39). Thus, this new epistatic interaction further supports the hypothesis of a shared genetic architecture between these two diseases and that T2D related loci can also genetically increase risk for DN.

In the HT data, we detect an epistatic interaction between *rs34585560* in *MECOM* gene and *rs8051877* in *NPW* gene. *MECOM* is a transcriptional regulator and is a well-known risk gene for high blood pressure as detected in a large scale GWAS (200k samples) (40). However, not much is known about the mechanism over which *MECOM* affects

Table 3. Time performances of LINDEN, Potpourri-guided LINDEN and Potpourri on the T2D dataset.

		T2D	
Method		CPU Time (log10)	Run Time (hh:mm:ss)
LINDEN		4.54	01:04:53
Potpourri guided LINDEN	$k = 500$	3.73	00:17:39
	$k = 750$	3.67	00:16:50
	$k = 1000$	3.79	00:24:40
	$k = 1500$	3.85	00:28:18
	$k = 2000$	3.88	00:30:20
Potpourri	$k = 500$	3.69	00:14:54
	$k = 750$	3.77	00:19:16
	$k = 1000$	3.75	00:20:47
	$k = 1500$	3.84	00:22:45
	$k = 2000$	3.89	00:22:30

blood pressure (41). *NPW* is a G-coupled protein activator which regulates neuroendocrine signals. While its ties to hypertension in humans is loose, (42) and (43) demonstrate in rats that exogenously applied *NPW* increases mean arterial pressure. Thus, this new epistatic pair finding can suggest *NPW* as a new risk gene candidate for HT coupled with the effect of *MECOM*.

Runtime Comparison

In this section, we compare the runtime of each approach (LINDEN, Potpourri-guided LINDEN and Potpourri) on the three datasets in terms of CPU time and clock-time. The results are provided in Table 3, Supplementary Tables 26 and 27, for T2D, BD and HT datasets, respectively. While epistasis test prioritization is an offline task and is not time critical, our results show that Potpourri decreases the time requirement 4 folds (in the default setting, $k=750$) while providing substantial improvements in precision for all datasets. Additional time required by Potpourri to consider regulatory and coding regions is negligible for all schemes.

DISCUSSION

We compared Potpourri with the state-of-the-art, and have shown that in almost all settings epistasis detection power has been substantially improved with our technique. We also show that promoting the testing of SNPs in regulatory and coding regions increases the precision and interpretability of the results. This is actually one of the biggest challenges in epistasis test prioritization. It is just a numbers game if one only considers the p-values and ignores the fact that SNPs in unannotated regions might not lead to any biological insight. Interpreting the biological meaning is an important issue to be reckoned with. One extreme is restricting the analysis to coding and regulatory regions for interpretability (also to limit the number of tests) and performing gene/pathway/network level analyses (13). This approach totally discards variants in the non-coding regions. However, the top pairs detected by Potpourri in all three datasets were in non-coding regions which could have been missed this way. Thus, a balance is needed. In this study, we propose a trade off between analyzing the exome and the genome. We show that our techniques to promote functional regions helps increase the precision. We were also able to detect 2 novel interactions

using the annotations provided which suggest new risk candidates.

One drawback of our scheme is the large number of parameters to be optimized. Thus, we elected to optimize β and D parameters with respect to the phenotype prediction performance and let k and ω to be supplied by the user. We think this is a fair choice since the ultimate goal of epistasis test prioritization is to provide the user with a manageable number of candidates. The optimum k value might yield a very large set of SNPs and stringent significance threshold. Similarly, promotion of regulatory/coding regions introduces a literature bias as discussed earlier and it makes sense for a user to be able to tune it themselves. Finally, our framework does not consider SNP effect sizes and only rely on univariate correlations of SNPs with the phenotype. Incorporating this information can further improve the performance, which is left as a future work.

CONCLUSION

Detecting epistatic interactions is a promising direction for understanding the genetic underpinnings of complex traits, but the sheer number of possible hypotheses to test prohibits brute force techniques. We proposed a new test prioritization technique and showed that selecting individually informative and topologically diverse SNPs in terms of genomic location leads to detecting statistically significant epistatic interactions. Our approach performs favorably compared to the state of the art.

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