Mining Pure, Strict Epistatic Interactions from High-Dimensional Datasets: Ameliorating the Curse of Dimensionality

Xia Jiang¹*, Richard E. Neapolitan²

1 Department of Biomedical Informatics, University of Pittsburgh, Pittsburgh, Pennsylvania, United States of America, 2 Division of Health and Biomedical Informatics, Department of Preventive Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois, United States of America

Abstract

Background: The interaction between loci to affect phenotype is called *epistasis*. It is strict *epistasis* if no proper subset of the interacting loci exhibits a marginal effect. For many diseases, it is likely that unknown epistatic interactions affect disease susceptibility. A difficulty when mining epistatic interactions from high-dimensional datasets concerns the *curse of dimensionality*. There are too many combinations of SNPs to perform an exhaustive search. A method that could locate strict epistasis without an exhaustive search can be considered the *brass ring* of methods for analyzing high-dimensional datasets.

Methodology/Findings: A *SNP pattern* is a Bayesian network representing SNP-disease relationships. The *Bayesian score* for a SNP pattern is the probability of the data given the pattern, and has been used to learn SNP patterns. We identified a bound for the score of a SNP pattern. The bound provides an upper limit on the Bayesian score of any pattern that could be obtained by expanding a given pattern. We felt that the bound might enable the data to say something about the promise of expanding a 1-SNP pattern even when there are no marginal effects. We tested the bound using simulated datasets and semi-synthetic high-dimensional datasets obtained from GWAS datasets. We found that the bound was able to dramatically reduce the search time for strict epistasis. Using an Alzheimer's dataset, we showed that it is possible to discover an interaction involving the APOE gene based on its score because of its large marginal effect, but that the bound is most effective at discovering interactions without marginal effects.

Conclusions/Significance: We conclude that the bound appears to ameliorate the curse of dimensionality in high-dimensional datasets. This is a very consequential result and could be pivotal in our efforts to reveal the dark matter of genetic disease risk from high-dimensional datasets.

Citation: Jiang X, Neapolitan RE (2012) Mining Pure, Strict Epistatic Interactions from High-Dimensional Datasets: Ameliorating the Curse of Dimensionality. PLoS ONE 7(10): e46771. doi:10.1371/journal.pone.0046771

Editor: Xiaofeng Wang, Cleveland Clinic Lerner Research Institute, United States of America

Received July 1, 2012; Accepted September 7, 2012; Published October 12, 2012

Copyright: © 2012 Jiang, Neapolitan. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was supported by [grant number 1K99LM010822-01] from the National Library of Medicine at the United States National Institutes of Health. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: xij6@pitt.edu

Introduction

In Mendelian diseases, a genetic variant at a single locus may give rise to a disease [1]. However, in many common diseases, it is likely that manifestation of the disease is due to genetic variants at multiple loci, with each locus conferring modest risk of developing the disease. For example, there is evidence that gene-gene interactions may play an important role in the genetic basis of hypertension [2], sporadic breast cancer [3], and other common diseases [4]. The interaction between two or more genes to affect a phenotype such as disease susceptibility is called *epistasis*. Biologically, epistasis likely arises from physical interactions occurring at the molecular level. Statistically, epistasis refers to an interaction between multiple loci such that the net effect on phenotype cannot be predicted by simply combining the effects of the individual loci. Often, the individual loci exhibit weak marginal effects; sometimes they may exhibit none. We say it is a *pure epistatic* interaction if no single locus exhibits a marginal effect, and we say it is a strict

epistatic interaction if no subset of the interacting loci exhibits a marginal effect.

The ability to identify epistasis is important in understanding the inheritance of many common diseases. For example, studying genetic interactions in cancer is essential to further our understanding of cancer mechanisms at the genetic level. Many cancerassociated mutations and interactions among the mutated loci remain unknown. For example, highly penetrant cancer susceptibility genes, such as BRCA1 and BRCA2, are linked to breast cancer [5]. However, only about 5 to 10 percent of breast cancers can be explained by germ-line mutations in these single genes. Most women with a family history of breast cancer do not carry germ-line mutations in the single highly penetrant cancer susceptibility genes, yet familial clusters continue to appear with each new generation [6]. This kind of phenomenon is not yet well understood, and undiscovered mutations or undiscovered interactions among mutations are likely responsible. The most common genetic variation is the *single nucleotide polymorphism (SNP)* that results when a nucleotide that is typically present at a specific location on the genomic sequence is replaced by another nucleotide. In most cases a SNP is biallelic, that is it has only two possible values among A, G, C, and T (the four DNA nucleotide bases). The less frequent (rare) allele must be present in 1% or more of the population for a site to qualify as a SNP [7]. The human genome is estimated to contain 15–20 million SNPs. In what follows we will refer to SNPs as the loci investigated when searching for a correlation of some loci with a phenotype such as disease susceptibility.

The advent of high-throughput technologies has enabled genomewide association studies (GWAS). A GWAS involves genotyping representative SNPs in individuals sampled from a population. A GWAS dataset can concern millions of SNPs and may soon concern billions. Data in which each record has such a large number of attributes is called *high-dimensional*. In case-control GWAS we identify the disease status along with the values of the SNPs. Such studies provide researchers unprecedented opportunities to investigate the complex genetic basis of diseases. By looking at single-locus associations, researchers have identified over 150 risk loci associated with 60 common diseases and traits [8], [9], [10], [11], [12], [13].

However, single-SNP investigations could not detect complex epistatic interactions in which each locus by itself exhibits little or no marginal effect. To fully exploit genomic data and possibly reveal a great deal of the dark matter of genetic risk, it is critical that we analyze such data using multi-locus methods, which investigate k-SNP patterns. Briefly, a k-SNP pattern is a Bayesian network (BN) model that describes the relationship between k SNPs and disease status, and is described in detail in the Methods Section.

When investigating SNP patterns, in some way we must score the patterns to determine which patterns are most noteworthy. Standard techniques such as linear regression may not work well because both the predictors and the target are discrete. One wellknown technique is *Multifactor Dimensionality Reduction (MDR)* [14]. MDR combines two or more variables into a single variable (hence leading to dimensionality reduction); this changes the representation space of the data and facilitates the detection of nonlinear interactions among the variables. MDR has been successfully applied to detect epistatic interactions in diseases such as sporadic breast cancer [3] and type II diabetes [15]. Using 28,000 simulated datasets and a real Alzheimer's GWAS dataset, Jiang et al. [16] evaluated the performance of 22 BN scoring criteria and MDR when scoring SNP patterns. They found that several of the BN scoring criteria performed substantially better than other scores and MDR. The BN scores that performed best were ones that computed the Bayesian score, which is the probability of the data given the pattern.

A difficulty when learning SNP patterns from high-dimensional GWAS datasets concerns the *curse of dimensionality*. For example, if we only investigated all 0, 1, 2, 3 and 4-SNP patterns when there are 500,000 SNPs, we would need to investigate 2.604×10^{21} patterns. Therefore, researchers have worked on developing heuristic search methods that investigate multiple loci using a GWAS dataset. Lasso, which is a shrinkage and selection method for linear regression [17,18], was applied to this task [19,20]. However, linear regression obviously has difficulty handling nonlinear epistatic interactions. Other methods include permutation testing [21,22], the use of ReliefF [23,24], random forests [25], predictive rule inference [26], a variational Bayes algorithm [27], a Bayesian marker partition method [28], an MCMC approximate model averaging technique [29], a Markov blanket

method [30], the use of maximum entropy [31], the use of Bayesian networks and greedy search [32], and an ensemble-based method that uses boosting [33].

Each of these methods has at least one of the following shortcomings: 1) It only investigates two-locus interactions and still requires quadratic time; or 2) It has only been shown to detect interactions in which only one interacting locus has no significant marginal effect. Many of the methods proceed in stages, using the first stage to identify promising SNPs, which in some way are investigated further in the second stage. Strict epistasis constitutes the worst-case in terms of detecting disease associations because such associations are only observable if all interacting SNPs are included in the disease model. None of these two-stage methods made any progress towards detecting strict epistasis. So, Evans et al. [34] conclude that "it is preferable to perform an exhaustive two-locus search across the genome rather than either of the twostage procedures that we examined. Otherwise, investigators risk discarding significant loci that only exhibit small effects at the margins.'

An exhaustive search is not possible when there are millions of SNPs. So some researchers turned their efforts to reducing the search space based on ancillary knowledge. You et al. [35] performed a two-stage application of MDR. The first stage is a within-gene search in which all combinations of SNPs allocated to the same gene are investigated. Briggs et al. [36] identified promising regions harboring epistatic candidates by looking for concordance in affected sibling pairs. Jiang et al. [37] investigated all 2-loci combinations where one of the loci was previously known to be associated with the disease. Perhaps the most promising technique for reducing the search space is to restrict the search space for candidate gene sets by using knowledge about molecular pathways [38].

However, once the search space is reduced, we can still be left with a large number of SNPs, prohibiting an exhaustive search of even the pruned dataset. Furthermore, in an agnostic study we are searching for possible interactions for which we have no previous knowledge. Therefore, a multi-stage technique that can effectively locate strict epistatic interactions could still be considered the *brass ring* of methods for analyzing high-dimensional datasets.

Initially it might seem that it is not possible to successfully prune our search for strict epistatic k+1-SNP interactions by investigating k-SNP patterns. This is likely the case when we are scoring the patterns. However, we have identified a bound (not a score) for SNP patterns. The bound provides an upper limit on the Bayesian score of any pattern that could be obtained by expanding a given pattern. For example, it gives an upper limit on the Bayesian score of any 2-SNP pattern that could be obtained from a particular 1-SNP pattern. We speculated that this bound might enable the data to tell us something about the promise of expanding a 1-SNP pattern even when there are no marginal effects. We tested the bound using 6000 simulated datasets developed from models of strict epistasis. We also injected epistatic interactions in two real GWAS datasets to create 2400 high-dimensional semi-synthetic datasets. In the case of both the simulated datasets and the semisynthetics datasets, the bound was able to significantly reduce the search time for the epistatic interaction. This was the case regardless of the heritability used to generate the interaction and the dimension of the dataset. The average fraction of the search space investigated before finding the true pattern (the one representing the interjected interaction) was as little as 0.0004.

Methods

We developed both simulated and semi-synthetic datasets based on models of strict epistasis. We compared the performance of the bound and Bayesian score in their ability to efficiently locate the true pattern. After providing background on Bayesian networks, we discuss specialized Bayesian networks called SNP patterns that represent relationships among SNPs and a disease. Then we show the algorithm used for the comparisons, and we describe the datasets we developed.

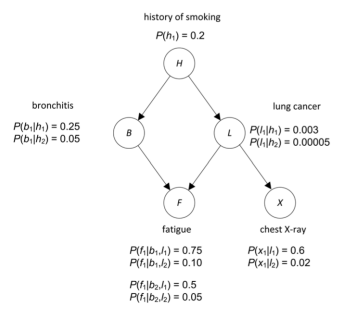
Bayesian Networks

Let V be a set of random variables, P be a joint probability distribution of these random variables, and G = (V, E) be a *directed acyclic graph* (*DAG*) where V is the set of nodes in G and E is the set of edges among these nodes. We say that (G, P) satisfies the *Markov condition* if for each variable $X \in V$, X is conditionally independent of the set of all its nondescendents in G given the set of all its parents. If (G, P) satisfies the Markov condition, we call (G, P) a *Bayesian network* (*BN*).

It is a theorem [39] that (G, P) satisfies the Markov condition (and therefore is a BN) if and only if P is equal to the product of its conditional distributions of all nodes given their parents in G, whenever these conditional distributions exist. That is, if our variables are X_1, X_2, \ldots, X_n , and PA_i is the set of parents of X_i , then

$$P(X_1, X_2, \ldots, X_n) = \prod_{i=1}^n P(X_i | PA_i).$$

Due to the theorem just mentioned, a BN is often developed by first identifying a DAG that satisfies the Markov condition relative to our belief about the probability distribution of the nodes in the DAG, and then determining the conditional probability distributions for this DAG. Often the DAG is a causal DAG, which is a DAG in which there is an edge from X to Y if and only if X is a direct cause of Y relative to the other nodes in the DAG. See [39] for a discussion as to why a causal DAG should often satisfy the Markov condition with the probability distribution of the variables





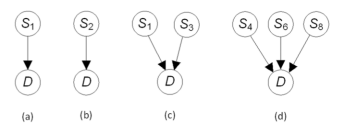


Figure 2. Four SNP patterns. doi:10.1371/journal.pone.0046771.g002

in the DAG. Figure 1 shows a BN representing the causal relationships among variables related to lung disorders. In this network, h_1 denotes an individual has a smoking history and h_2 denotes that the individual does not. The other variables have similar denotations. Using this BN, we can determine conditional probabilities of interest using the BN and a BN inference algorithm. For example, if a given individual has a smoking history, a positive chest X-ray, and is fatigued, we can determine the conditional probability of the individual having lung cancer. That is, we can compute $P(l_1|h_1, x_1, f_1)$. These inference algorithms exploit Bayes' Theorem and are efficient for a large class of BNs [39].

Methods have been develop both for learning the parameters in a BN and the structure (DAG) from data. Pierrier et al. [40] provide a recent review of the literature concerning BN structure learning. We review structure learning using the Bayesian score because that method is relevant to the research described here.

A DAG model consists of a DAG G = (V, E) where V is a set of random variables, and a parameter set θ whose members determine conditional probability distributions for G, but without specific numerical assignments to the parameters. The task of learning a unique DAG model from data is called *model selection*. As an example, if we had data on a large number of individuals concerning lung disorders we might be able to learn the DAG in

Table 1. Properties of the strict epistatic models used in the simulated datasets.

# SNPs	MAF	Low Heritability	High Heritability
2	0.05	0.05	0.1
2	0.1	0.05	0.2
2	0.2	0.05	0.2
2	0.3	0.05	0.2
2	0.4	0.05	0.2
3	0.05	0.005	0.01
3	0.1	0.05	0.1
3	0.2	0.05	0.2
3	0.3	0.05	0.2
3	0.4	0.05	0.2
4	0.05	0.001	0.002
4	0.1	0.005	0.01
4	0.2	0.05	0.1
4	0.3	0.05	0.2
4	0.4	0.05	0.2

doi:10.1371/journal.pone.0046771.t001

Table 2. Properties of the strict epistatic models used in the semi-synthetic datasets.

// CND-		l la sita hilita a	
# SNPs	MAF	Heritability	
2	0.05	0.1	
2	0.1	0.2	
2	0.15	0.3	
2	0.2	0.4	
3	0.05	0.01	
3	0.1	0.04	
3	0.15	0.1	
3	0.2	0.2	
4	0.05	0.001	
4	0.1	0.01	
4	0.15	0.04	
4	0.2	0.1	

doi:10.1371/journal.pone.0046771.t002

Figure 1 from data. When the edges represent causal influences, this means that we can learn causal influences from data.

In the constraint-based structure learning approach [41], we try to learn a DAG model from the conditional independencies that the data suggest are present in the generative probability distribution P. By the generative probability distribution P we mean the underlying joint probability distribution of the random variables, making the assumption that such a distribution exists. In the score-based structure learning approach [39], we assign a score to a DAG based on how well the DAG fits the data.

A straightforward score, called the *Bayesian score*, is the probability of the DAG model given the *Data*, which is as follows:

$$P(G|Data) = K \times P(Data|G)P(G)$$

where K is a normalizing constant. If we assign a uniform prior probability distribution to the DAG models, we can score them using the likelihood P(Data|G), and that is the approach taken here. So we will simply refer to the likelihood as the score. For a DAG G containing a set of discrete random variables $V = \{X_1, X_2, \ldots, X_n\}$ and Data such that each data item is a vector of values of all random variables in V, Cooper and Herskovits [42] develop the following formula for this likelihood:

$$P(Data|G) = \prod_{i=1}^{n} \prod_{j=1}^{q_i} \frac{\Gamma(\sum_{k=1}^{r_i} a_{ijk})}{\Gamma(\sum_{k=1}^{r_i} a_{ijk} + \sum_{k=1}^{r_i} s_{ijk})} \prod_{k=1}^{r_i} \frac{\Gamma(a_{ijk} + s_{ijk})}{\Gamma(a_{ijk})}, \quad (1)$$

where r_i is the number of states of X_i , q_i is the number of different instantiations of the parents of X_i , a_{ijk} is the ascertained prior belief concerning the number of times X_i took its kth value when the parents of X_i had their *j*th instantiation, and s_{ijk} is the number of times in the data that X_i took its kth value when the parents of X_i had their *j*th instantiation.

The likelihood in Equation 1 assumes that our prior belief concerning each unknown parameter in each DAG model is represented by a Dirichlet distribution, where the hyperparameters a_{ijk} are the parameters for this distribution. Cooper and Herskovits [42] suggested setting the value of every hyperparameters a_{ijk} equal to 1, which amounts to assigning a prior uniform distribution to the value of each parameter (prior ignorance as to its value). They called this the K2 score.

However, Heckerman et al. [43] showed that the K2 score does not assign the same score to Markov equivalent DAG models (two DAGs are *Markov equivalent* if they entail the same conditional independencies). For example, the DAGs $X_1 \rightarrow X_2$ and $X_2 \rightarrow X_1$ do not obtain the same score. They suggested determining the values of the hyperparameters from a single parameter α called the *prior equivalent sample size*. If we want to use a prior equivalent sample size and represent a prior uniform distribution for each variable (not parameter) in the network, for all *i*, *j*, and *k* we set $a_{ijk} = \alpha/r_i q_i$ where r_i is the number of states of the *i*th variable and

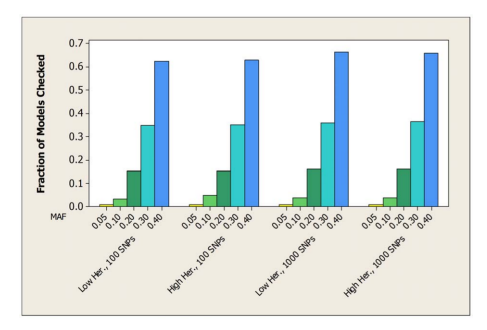


Figure 3. For the simulated datasets developed from 2-SNP models, the average fraction of the total number of patterns checked before the bound finds the true pattern. doi:10.1371/journal.pone.0046771.g003

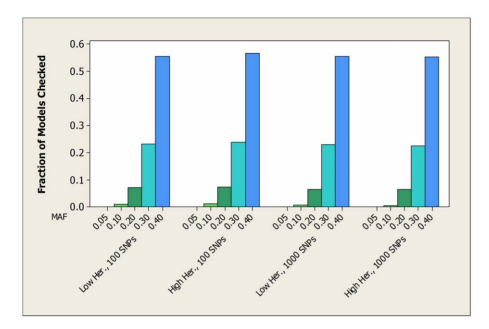


Figure 4. For the simulated datasets developed from 3-SNP models, the average fraction of the total number of patterns checked before the bound finds the true pattern. doi:10.1371/journal.pone.0046771.g004

 q_i is the number of different instantiations of its parents. When we do this, the Bayesian score is called the *Bayesian Dirichlet equivalent uniform (BDeu)* score. That score is as follows:

$$P(Data|G) = \prod_{i=1}^{n} \prod_{j=1}^{q_i} \frac{\Gamma(\frac{\alpha}{q_i})}{\Gamma(\frac{\alpha}{q_i} + \frac{r_i}{k=1} s_{ijk})} \prod_{k=1}^{r_i} \frac{\Gamma(\frac{\alpha}{r_i q_i} + s_{ijk})}{\Gamma(\frac{\alpha}{r_i q_i})}.$$

Heckerman et al. [43] showed that if we use a prior equivalent sample size, then Markov equivalent DAGs have the same score.

The Bayesian score does not explicitly include a DAG penalty. However, the penalty is implicitly determined by the hyperparameters a_{ijk} . Silander et al. [44] show that if we use the BDeu score, then the DAG penalty decreases as α increases.

The Bayesian score decomposes into the product of local scores, one for each node X_i in the DAG. In the case of the BDeu score, the local score for X_i with parent set PA_i is given by

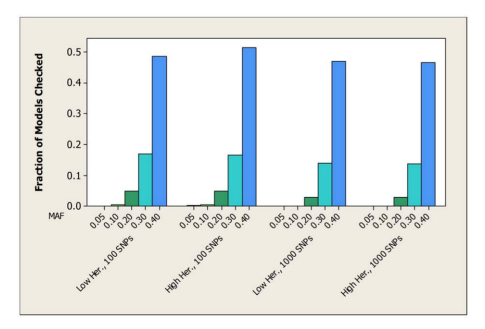


Figure 5. For the simulated datasets developed from 4-SNP models, the average fraction of the total number of patterns checked before the bound finds the true pattern. doi:10.1371/journal.pone.0046771.g005

Table 3. For the simulated datasets developed from 2-SNP models, average fraction over all 100 datasets of the total number of patterns checked before finding the true pattern.

	100 Data Iter	ns			1000 Data	a Items		
	Low Heritabi	lity	High Heritabi	ility	Low Herit	ability	High Heri	tability
MAF	Bound	Score	Bound	Score	Bound	Score	Bound	Score
0.05	0.007	0.042	0.007	0.042	0.008	0.044	0.008	0.048
0.1	0.033	0.133	0.047	0.134	0.037	0.148	0.037	0.139
0.2	0.151	0.356	0.153	0.385	0.159	0.386	0.159	0.404
0.3	0.348	0.594	0.349	0.554	0.359	0.571	0.362	0.555
0.4	0.623	0.641	0.629	0.696	0.662	0.673	0.656	0.714

doi:10.1371/journal.pone.0046771.t003

$$score(X_i, PA_i) = \prod_{j=1}^{q_i} \frac{\Gamma(\frac{\alpha}{q_i})}{\Gamma(\frac{\alpha}{q_i} + \frac{r_i}{k=1} s_{ijk})} \prod_{k=1}^{r_i} \frac{\Gamma(\frac{\alpha}{r_i q_i} + s_{ijk})}{\Gamma(\frac{\alpha}{r_i q_i})}, \quad (2)$$

where r_i is the number of states of X_i , and q_i is the number of different instantiations of the nodes in PA_i . When learning a DAG model from data, we could try to maximize our local scores by looking for an optimal parent set PA_i for each node X_i . We could prune our search space if we had an upper bound on the local score for any parent set that could be obtained by adding parents to a current parent set PA_i . Singh and Moore [45] proved the following Theorem relevant to such a bound.

Theorem 1. If in Equation 2 we fix the value of $r_i = s_{ijk}$ for each j, then $score(X_i, P_i)$ is maximized if for each j, $s_{ijkj} = r_i = s_{ijk}$ for some k_j . That is, for every combination of values of the parents of X_i , all data items that have those values have the same value of X_i .

This theorem could be used to obtain a bound on the local score that could be obtained by adding parents to a given parent set. However, the bound is a very loose bound (i.e. the bounds are much greater than the scores), and therefore the bound has not proven to be useful in pruning the search space.

SNP Patterns and a Bound for SNP Patterns

We could develop a DAG model that represents many factors that affect phenotype including inheritable allele variation, somatic mutations in alleles, environmental factors, and epigenetic phenomena such as DNA methylation. A subnetwork of that model contains only variables that represent inheritable allele variation (the variables whose values are obtained in a GWAS). If, for example, we have a 5-way interaction, two 3-way interactions, and two 2-way interactions, there are 15 SNPs in this subnetwork, all which have edges to the disease node D. We have neither the data nor the computational time to score such networks. So we must settle for trying to learn pieces of the network, such as particular interactions, separately. These small subnetworks are the focus of this paper, and we call them SNP patterns. Examples of such patterns appear in Figure 2. The first two represent that a single SNP by itself is associated with the disease, the third one represents that two SNPs together are associated with the disease, and the fourth one represents that three SNPs together are associated with the disease. It is important to recognize that, for example, the pattern in Figure 2 (c) does not entail that S_1 and S_3 are interacting to affect D. Each could be affecting it separately. Without making specialized mathematical assumptions, it is difficult to distinguish these two situations from data alone. Jiang et al. [37] provide one way to do this.

In the case of a SNP pattern G and *Data* concerning the SNPs and disease in the pattern, the local BDeu score (Equation 2) is as follows:

$$score(D, P_D) = \prod_{j=1}^{q} \frac{\Gamma\left(\frac{\alpha}{q}\right)}{\Gamma\left(\frac{\alpha}{q} + \frac{2}{k-1}s_{jk}\right)} \prod_{k=1}^{2} \frac{\Gamma\left(\frac{\alpha}{2q} + s_{jk}\right)}{\Gamma\left(\frac{\alpha}{2q}\right)}, \quad (3)$$

where P_D is the parent set of D in the pattern, q is the number of

Table 4. For the simulated datasets developed from 3-SNP models, average fraction over all 100 datasets of the total number of patterns checked before finding the true pattern.

	100 Data Iten	ns			1000 Data	a Items		
	Low Heritabil	ity	High Heritabi	ility	Low Herit	ability	High Heri	tability
MAF	Bound	Score	Bound	Score	Bound	Score	Bound	Score
0.05	0.001	0.015	0.001	0.017	0.0005	0.006	0.0009	0.014
0.1	0.010	0.075	0.012	0.079	0.006	0.032	0.005	0.031
0.2	0.072	0.312	0.073	0.309	0.064	0.316	0.063	0.307
0.3	0.232	0.557	0.238	0.566	0.228	0.557	0.225	0.579
0.4	0.554	0.699	0.566	0.770	0.554	0.705	0.552	0.738

doi:10.1371/journal.pone.0046771.t004

Table 5. For the simulated datasets developed from 4-SNP models, average fraction over all 100 datasets of the total number of patterns checked before finding the true pattern.

	100 Data Ite	ms		1000 Data Items					
	Low Heritabi	Low Heritability		High Heritability		Low Heritability		High Heritability	
MAF	Bound	Score	Bound	Score	Bound	Score	Bound	Score	
0.05	0.001	0.007	0.002	0.012	0.0004	0.004	0.0005	0.015	
0.1	0.005	0.052	0.005	0.053	0.001	0.036	0.001	0.037	
0.2	0.048	0.271	0.048	0.275	0.028	0.262	0.028	0.283	
0.3	0.170	0.554	0.165	0.544	0.139	0.534	0.137	0.552	
0.4	0.486	0.674	0.514	0.751	0.471	0.751	0.466	0.713	

doi:10.1371/journal.pone.0046771.t005

different instantiations of the parents of D, and s_{jk} is the number of times in the data that D took its kth value when the parents of D had their *j*th instantiation.

Using Theorem 1, we can obtain a bound on the local score that could be obtained by adding parents to a given parent set. Given a SNP pattern and the score in Equation 3, an upper limit on the BDeu score of any SNP pattern obtained by adding more SNPs is as follows (*t* is the number of combined states of the SNPs added):

$$bound(D, P_D) = \prod_{j=1}^{q} \prod_{k=1}^{2} \frac{\Gamma(\frac{\alpha}{qt})}{\Gamma(\frac{\alpha}{qt} + s_{jk})} \times \frac{\Gamma(\frac{\alpha}{2qt} + s_{jk})}{\Gamma(\frac{\alpha}{2qt})}$$

If all variables are binary, and we are looking only for a bound on adding one SNP, t=2, if it is a bound on adding two SNPs, t=4, and if it is a bound on adding three SNPs, t=8.

As noted earlier, this is a very loose bound and would not be useful for provably pruning the search space. However, in the case of searching for strict epistatic interactions, perhaps it is asking too much to hope for provable results. We would achieve valuable progress if we could just often heuristically locate such interactions without an exhaustive search. We conjectured that the bound might enable us to do that, and that is what is investigated here.

Algorithm

Suppose we have a dataset concerning SNPs and a disease, and our goal is to find a 2-SNP pattern with a particular BDeu score (Equation 3). This pattern could be the highest scoring pattern or it could be the true pattern representing an epistatic interaction used to generate the data. Often these two patterns are the same. We search for that pattern as follows. First, all the 1-SNP patterns are sorted by either their 2-SNP scores or their 2-SNP bounds. Let $S_1, S_2, S_3, S_4,...$ be the resultant sorted list of SNPs. We then investigate 2-SNP patterns according to the following order:

S_1S_2 ,

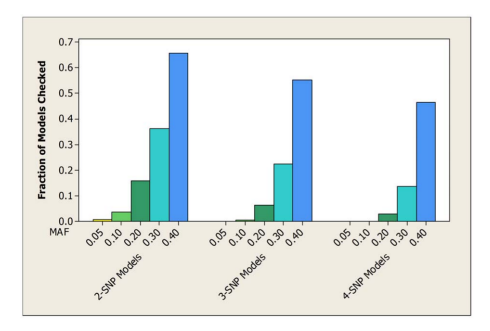


Figure 6. For the simulated datasets with 1000 SNPs and high heritability, the average fraction of the total number of patterns checked before the bound finds the true pattern. doi:10.1371/journal.pone.0046771.g006

Table 6. Approximation of the average running time before finding the true pattern.

	1000 SNPs	1000 SNPs		10,000 SNPs		
	Bound	Exhaustive	Bound	Exhaustive	Bound	Exhaustive
2-SNP	10 sec	12 min	16 min	20 hours	28 hours	84 days
3-SNP	7 min	3 days	5 days	8 years	14 years	7,662 years
4-SNP	1 day	2 years	14 years	19,146 years	$4\!\times\!10^5$ years	$2\!\times\!10^8$ years

doi:10.1371/journal.pone.0046771.t006

$$S_1S_3, S_2S_3,$$

$S_1S_4, S_2S_4, S_3S_4,$

$$S_1 S_5, \ldots$$

We stop when we locate the pattern for which we are searching, and we keep track of the number of patterns checked before we find that pattern.

This strategy extends readily to search for k-SNP patterns. The following is an algorithm for the general case:

Algorithm 1 Find_Pattern

n = total number of SNPs;

k = number of SNPs in the patterns we are investigating;

S = array containing the SNPs in sorted sequence;

scoresearch = score of pattern we are trying to locate; *numpatterns* = 0;

scorepatterns(k, n);

procedure *scorepatterns*(int *start*, int *end*);

if start = 0

numpatterns = numpatterns + 1;

score = Bayesianscore(V);if score = scoresearchoutput (numpatterns); halt; endif else for i = start to end V[t - start + 1] = S[i]; scorepatterns(start - 1, i - 1);endfor endelse

Algorithm 1 constructs each candidate pattern in the array V. When V contains k SNPs, the pattern is scored. If its score is equal to the score of the pattern for which we are searching, the algorithm outouts the number of patterns checked and halts. Note that if we are searching for a pattern representing a particular epistatic interaction, it will halt when we find any pattern with the same score as that pattern. However, in our experiments this unlikely event never occurred.

Simulated Data Sets

Fisher et al. [46] created GAMETES, which is a software package for generating strict epistatic models with random architectures' We used GAMETES to develop 2-SNP, 3-SNP, and 4-SNP models of strict epistatic interaction. The software

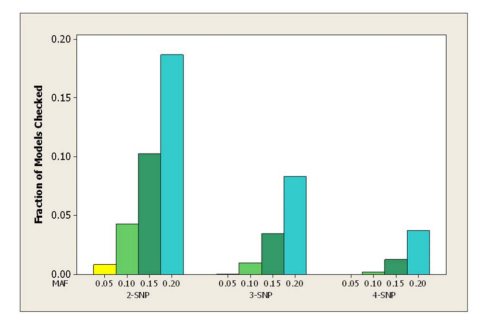


Figure 7. For the LOAD datasets, the average ratio of the number of patterns checked by the bound to the number of patterns checked by the score before each finds the true pattern. doi:10.1371/journal.pone.0046771.g007

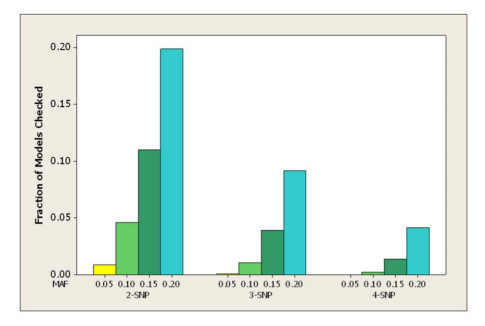


Figure 8. For the breast cancer datasets, the average fraction of the total number of patterns checked before the bound finds the true pattern. doi:10.1371/journal.pone.0046771.g008

allows the user to specify the *heritability* and the *minor allele frequency* (MAF). We wanted to test a full range of MAFs (0.05 to 0.4) and relatively low and high heritabilities (0.05 and 0.2). However, we found that to obtain models with a low MAF, it was sometimes necessary to decrease the heritability. The actual MAFs and heritabilities used in the simulations are shown in Table 1.

For each of the 30 combinations of MAF and heritability, we developed datasets in which there were 100 SNPs and 1000 SNPs. Each dataset had 1000 cases and 1000 controls. For each of the 60 variations, 100 datasets were generated, making a total of 6000 datasets. We used the BDeu score with $\alpha = 9$ to score the SNP patterns and to compute their bounds using a given dataset. For each of the datasets, we first sorted the SNPs by their bounds. For the 2-SNP interactions, it was the bound on adding one SNP; for the 3-SNP interactions, it was the bound on adding two SNPs; and for the 4-SNP interactions, it was the bound on adding three SNPs. We then used Algorithm 1 to determine how many patterns were checked before the true pattern representing the interaction was discovered. In the case of the 1000 SNP datasets it was not computationally feasible to actually run Algorithm 1. Rather we simply determined the locations where the interacting SNPs appeared in the sorted list, and then using these locations we

Table 7. For the LOAD datasets, average fraction over all 100 datasets of the total number of patterns checked before finding the true pattern.

MAF 2-SNP			3-SNP	3-SNP		4-SNP	
	Bound	Score	Bound	Score	Bound	Score	
0.05	0.008	0.054	0.0003	0.017	0.0001	0.006	
0.1	0.043	0.170	0.009	0.085	0.002	0.044	
0.15	0.102	0.274	0.035	0.206	0.012	0.124	
0.2	0.187	0.393	0.083	0.353	0.037	0.281	

doi:10.1371/journal.pone.0046771.t007

computed where Algorithm 1 would find the true pattern representing the interacting SNPs. We repeated this procedure with the SNPs sorted by their scores.

We did not vary the number of cases and controls because that number does not have a significant effect on the running time. The data needs to preprocessed to obtain the counts needed in the computation of the bound and the score. The only effect that the number of cases and controls has on the running time is that the pre-processing time increases linearly with the total number of cases and controls.

Semi-Synthetic Datasets

Reiman et al. [13] developed a GWAS late onset Alzheimer's disease (LOAD) dataset on 312,317 SNPs from an Affymetrix 500 K chip, plus the measurement of a locus in the APOE gene, which is known to be predictive of LOAD. The dataset consists of 859 cases and 552 controls. See http://www.tgen.org/ neurogenomics/data concerning this dataset. Hunter et al. [47] conducted a GWAS concerning 546,646 SNPs and breast cancer as part of the National Cancer Institute Cancer Genetic Markers of Susceptibility (CGEMS) Project. The dataset consists of 1145

1	
	Table 8. For the breast cancer datasets, average fraction over
	all 100 datasets of the total number of patterns checked
	before finding the true pattern.

MAF	2-SNP	2-SNP		3-SNP		4-SNP	
	Bound	Score	Bound	Score	Bound	Score	
0.05	0.008	0.045	0.0008	0.014	0.00009	0.004	
0.1	0.045	0.137	0.010	0.079	0.002	0.045	
0.15	0.109	0.272	0.038	0.173	0.013	0.120	
0.2	0.199	0.370	0.092	0.283	0.041	0.258	

doi:10.1371/journal.pone.0046771.t008

 Table 9.
 The 1-SNP score, 2-SNP bound and 2-SNP score for real loci and injected loci.

Locus	1-SNP Score	2-SNP Bound	2-SNP score
APOE	-836.08	-13.74	-
rs1007837 (GAB2)	-947.88	-17.50	-831.33
rs7101429 (GAB2)	-947.16	-17.53	-830.58
rs901104 (GAB2)	-947.69	-17.57	-830.69
rs4291702 (GAB2)	-947.36	-17.66	-830.51
rs4945261 (GAB2)	-947.83	-17.59	-831.68
rs7115850 (GAB2)	-945.19	-17.89	-827.261
rs10793294 (GAB2)	-947.24	-18.39	-830.84
rs2450130 (GAB2)	-948.28	-17.60	-830.49
S1 (0.05)	-949.25	-14.60	-836.31
S2 (0.05)	-949.12	-14.42	-836.31
S1 (0.1)	-964.67	-16.42	-761.12
S2 (0.1)	-964.50	-16.24	-761.12
S1 (0.15)	-950.29	-17.56	-668.48
S2 (0.15)	-950.16	-17.54	-668.48
S1 (0.20)	-950.56	-18.42	-612.31
S2 (0.20)	-950.36	-18.60	-612.31

The 2-SNP scores for the real loci are the scores of the patterns in which the other locus is APOE. The 2-SNP scores for the injected loci are the scores of the injected patterns. For the injected SNPs, the MAFs are shown in parentheses. doi:10.1371/journal.pone.0046771.t009

cases and 1142 controls. See http://cgems.cancer.gov/ concerning this dataset.

We developed 2-SNP, 3-SNP, and 4-SNP models of strict epistatic interactions using GAMETES, and used the models to inject interacting SNPs into each of the real GWAS datasets resulting in semi-synthetic datasets. The models generated have the properties shown in Table 2. For each of the 12 models, 100 datasets were developed.

We used the BDeu score with $\alpha = 9$ to score the SNP patterns and to compute their bounds using a given dataset. We used Algorithm 1 to search for the true pattern representing the interacting SNPs using both the bound and the score to sort the SNPs.

Results

Simulated Datasets

For the simulated datasets developed from the 2-SNP models, Table 3 shows the average over all 100 datasets of the fraction of patterns checked by both the bound and the score before the true pattern is found. Tables 4 and 5 show the same information for the datasets developed from the 3-SNP and 4-SNP models. Figures 3, 4, and 5 show the average fraction of patterns checked by the bound in graphical format.

These tables and figures show that the bound usually performs much better than we would expect by chance, and also performs substantially better than the score. When the MAF is small (0.05) the average fraction of patterns checked by the bound is at most around 0.008. By chance alone we would expect that average to be around 0.5. In general, the average fraction increases as the MAF increases. On the other hand, the heritability and the dimension of the dataset have little effect on the average fraction. It is somewhat surprising that the bound performs about as well when the heritability is low as when it is high. Since unknown genetic risk might confer low heritability, this is an encouraging result.

The other variable that affects the performance of the bound is the number of SNPs involved in the epistatic interaction. That is, the performance improves as the number of SNPs increases. Figure 6 illustrates this using the experiments in which there were 1000 SNPs and the heritability was high.

In order to make these results more transparent, we did an analysis using actual times. As noted above, the performance does not seem to degrade as the dimension of the datasets increases (i.e., the results are about the same for the 100 SNP datasets and for the 1000 SNP datasets). If we assume that this result holds true for larger dimensions, then Table 6 shows the approximate average running needed to locate the true pattern using both the bound and a blind exhaustive search. The times were based on using an Intel Xeon CPU 2.66 GHz, which takes about 0.0029 seconds to compute the bound for one pattern. In the exhaustive search it is assumed that the true model will be found after checking half of the models on the average.

Notice that the bound always performs about an order of magnitude better than the exhaustive search as far as the dimension that it can handle in an acceptable amount of time. That is, in the case of the 2-SNP models the bound can handle 100,000 SNPs whereas the exhaustive search can only handle 10,000 SNPs; in the case of the 3-SNP models the bound can handle 10,000 SNPs whereas the exhaustive search can only handle 10000 SNPs; and in the case of the 4-SNP models the bound can handle 1000 SNPs; whereas the exhaustive search can only handle 1000 SNPs; and in the case of the 4-SNP models the bound can handle 1000 SNPs whereas the exhaustive search cannot.

These results indicate that in the case of pure epistasis the bound can often locate the true pattern much sooner than would be expected by chance, and offers a substantial improvement over using the score to locate that pattern.

Semi-Synthetic Datasets

For the LOAD datasets, Table 7 shows the average over all 100 datasets of the fraction of patterns checked by both the bound and the score before the true pattern was found. Figure 7 show the average fraction of patterns checked by the bound in graphical format. Table 8 and Figure 8 show the same information for the breast cancer datasets.

The results are similar to those for the simulated datasets. In the case of both the LOAD and the breast cancer datasets, when the MAF is small (0.05 or 0.1) the bound never checks more than 5% of the patterns before finding the pattern representing the interacting SNPs, and it always checks substantially fewer patterns than the score.

Consider the following example. In the case of the LOAD dataset, there are $4.877 \, 1 \times 10^{10} \, 2$ -SNP patterns. When the MAF is 0.05, the bound checks only 3.9016×10^8 of these patterns. Using an Intel Xeon CPU 2.66 GHz, it takes about 0.0029 seconds to compute the score or the bound for one pattern. So an exhaustive could be expected to find the true pattern in about 2.24 years. However, the bound can be expected to find the true pattern in about 13.09 days.

It is well known that the APOE gene is associated with LOAD [12,13]. The APOE gene has three common variants $\epsilon 2$, $\epsilon 3$ and $\epsilon 4$. The least risk is associated with the $\epsilon 2$ allele, while each copy of the $\epsilon 4$ allele increases risk. As mentioned above, Reiman et al. [13] developed a GWAS late onset Alzheimer's disease (LOAD) dataset on 312,317 SNPs and APOE status. These researchers investigated pairs of loci, where one locus is APOE status, in order to possibly learn epistatic interactions of other loci with APOE. They performed separate analyses in APOE $\epsilon 4$ carrier and APOE $\epsilon 4$ non-carriers. They found that 10 of the 25 SNPs with the most

significant LOAD association in APOE $\varepsilon 4$ were located in the GAB2 gene. They did not find that the GAB2 gene is significantly associated with LOAD in the APOE $\varepsilon 4$ non-carriers. Reiman et al. [13] provide biological evidence that GAB2 modifies LOAD risk in APOE $\varepsilon 4$ carriers.

Using this same dataset, Jiang et al. [37] scored all 2-SNP patterns where one of the loci is the APOE gene and the other loci is one of the 312, 317 SNPs being investigated. Eight of the ten highest scoring patterns contained a GAB2 SNP, substantiating the results in [13].

This discovery of a possible interaction between GAB2 and APOE was achieved because of APOE's high marginal effect. Table 9 compares this sort of result to our result concerning the discovery of the strict epistatic interactions we injected into the LOAD dataset. The average score and bound over all SNPs were -950.46 and -17.898 respectively. We see from Table 9 that APOE has an extremely high score (by far higher than any other 1-SNP pattern), and that the GAB2 SNPs have both unremarkable scores and bounds. However, due to APOE's high score, it was reasonable and computationally efficient to investigate the interaction of every SNP with APOE. By so doing, it was discovered that patterns including APOE and GAB2 were quite likely. Our injected SNPs have unremarkable scores. Little discovery would be expected by looking at their scores alone. However, the bounds for the SNPs are quite high (almost as high as APOE's bound) when the MAF is 0.05, and they decrease as the MAF increases. This result is consistent with the results shown in Table 7. So the bounds enabled us to discover the strict epistatic interactions.

It might seem odd that the average bounds for the interacting SNPs are around -18 when the MAF is 0.2 (see Table 9), which is worse than the average bound, but yet we find the pattern containing those SNPs after investigating only about 0.187 fraction of the 2-SNP patterns (see Table 7). The explanation for this is that both SNPs' bounds ordinarily ranked around the mid-point of the sorted list of all bounds. So Algorithm 1 would find the pattern containing them fairly early. The following example illustrates why this is the case. If there were 7 SNPs, there would be 21 patterns, and if the SNPs ranked as the 3rd and 4th SNPs, Algorithm 1 would discover the pattern containing them after investigating only 6 patterns.

References

- Bateson W (1909) Mendel's principles of heredity. Cambridge University Press.
 Moore JH, Williams SM (2002) New strategies for identifying gene gene
- interactions in hypertension. Annals of Medicine 34: 88–95.
 Ritchie MD, Hahn LW, Roodi N, Bailey LR, Dupont WD, et al. (2001) Multifactor-dimensionality reduction reveals high-order interactions among estrogen-metabolism genes in sporadic breast cancer. American Journal of Human Genetics 69: 138–147.
- Nagel RI (2005) Epistasis and the genetics of human diseases. C R Biologies 328: 606–615.
- Armes JE, Egan AJ, Southey MC, Dite GS, McCredie MR, et al. (2000) The histologic phenotypes of breast carcinoma occurring before age 40 years in women with and without BRCA1 or BRCA2 germline mutations. Cancer 83: 2335–2345.
- National Cancer Institute. Available: http://www.cancer.gov/cancertopics/ understandingcancer/cancergenomics.
- Brookes AJ (1999) The essence of SNPs. Gene 234: 177–186.
- Manolio TA, Collins FS (2009) The HapMap and genome-wide association studies in diagnosis and therapy. Annual Review of Medicine 60: 443–456.
- Herbert A, Gerry NP, McQueen MB (2006) A common genetic variant is associated with adult and childhood obesity. Journal of Computational Biology 312: 279–384.
- Spinola M, Meyer P, Kammerer S, Falvella FS, Boettger MB, Hoyal CR, et al. (2001) Association of the PDCD5 locus with long cancer risk and prognosis in smokers. American Journal of Human Genetics 55: 27–46.

Finally, note that the bound is a very loose bound. Therefore, it would not be useful in a best-search first algorithm that prunes SNPs based on their bounds and which is able to guarantee that we discovered the highest scoring pattern. However, as we have seen, the bound can be quite effective for guiding a heuristic search for high scoring patterns. In general, when we are searching for possible epistatic interactions, our concern is with finding likely patterns that we can then further investigate for biological plausibility. It is not necessary that we know that a discovered interaction has the highest score of all patterns.

Discussion

We identified a bound on the Bayesian score of any SNP pattern that could be obtained by expanding a given SNP pattern. Using simulated datasets based on models of strict epistasis, we showed that the bound can locate the true pattern, when searching moderate-dimensional datasets, much faster than can be expected by chance. Using semi-synthetic datasets based on models of strict epistasis, we showed that the bound can locate the true pattern, when searching high-dimensional GWAS datasets, much faster than can be expected by chance. The average fraction of patterns checked before finding the true pattern was as little as 0.0004. These results indicate that the bound can be an extremely useful tool in algorithms that search high-dimensional datasets for strict epistatic interactions.

We used an algorithm that sorts the SNPs by their bounds and by their scores to test the effectiveness of the bound. Although this was an effective way to compare the bound to the score, in practice it may not be the most effective way to use the bound in a heuristic search. First, the algorithm assumes we know the number of interacting SNPs up front. Second, if, for example, we were looking for a 4-SNP interaction, we would be able to visit very few different SNPs. We plan to incorporate the bound into other algorithms we have developed such as MBS [32]. Our final algorithm will use both the bound and the score to guide our search.

Author Contributions

Conceived and designed the experiments: RN XJ. Performed the experiments: XJ. Analyzed the data: RN. Contributed reagents/materials/analysis tools: RN XJ. Wrote the paper: RN XJ.

- Lambert JC, Heath S, Even G, Campion S, Sleegers K, et al. (2009) Genomewide association study identifies variants at CLU and CR1 associated with Alzheimer's disease. Nature Genetics 41: 1094–1099.
- Coon KD, Myers AJ, Craig DW, Webster JA, Pearson JV, et al. (2007) A highdensity whole-genome association study reveals that APOE is the major susceptibility gene for sporadic late-onset Alzheimer's disease. Journal of Clinical Psychiatry 68: 613–618.
- Reiman EM, Webster JA, Myers AJ, Hardy J, Dunckley T, et al. (2007) GAB2 alleles modify Alzheimer's risk in APOE carriers. Neuron 54: 713–720.
- Hahn LW, Ritchie MD, and Moore JH (2003) Multifactor dimensionality reduction software for detecting gene-gene and gene-environment interactions. Bioinformatics 19: 376–382.
- Cho YM, Ritchie MD, Moore JH, Park JY, Lee KU, et al. (2004) Multifactor dimensionality reduction reveals a two-locus interaction associated with type 2 diabetes mellitus. Diabetologia 47: 549–554.
- Jiang X, Neapolitan RE, Barmada MM, Visweswaran S (2011) Learning genetic epistasis using Bayesian network scoring criteria. BMC Bioinformatics 12(89): 1471–2105.
- Tibshirani R. Regression shrinkage and selection via the lasso. (1996) J Royal Statist Soc B 58(1): 267–288.
- Chen S, Donoho DL, Saunders MA (1998) Atomic decomposition by basis pursuit. SIAM Journal on Scientific Computing 20:33–61.
- Wu TT, Chen YF, Hastie T, Sobel E, Lange K (2009) Genome-wide association analysis by lasso penalized logistic regression. Genome Analysis 25: 714–721.

- Wu J, Devlin B, Ringguist S, Trucco M, Roeder K (2010) Screen and clean: A tool for identifying interactions in genome-wide association studies. Genetic Epidemiology 34: 275–285.
- Wongseree W, Assawamakin A, Piroonratana T, Sinsomros S, Limwongse C, et al. (2009) Detecting purely epistatic multi-locus interactions by an omnibus permutation test on ensembles of two-locus analyses. BMC Bioinformatics 10:294.
- 22. Zhang X, Pan F, Xie Y, Zou F, Wang W (2009) COE: a general approach for efficient genome-wide two-locus epistasis test in disease association study. In: Proceedings of the 13th Annual International Conference on Research in Computational Molecular Biology (RECOMB.
- Moore JH, White BC (2007) Turing Relief for genome-wide genetic analysis. In: Marchiori E, Moore JH, Rajapakee JC, editors. Proceedings of EvoBIO 2007, Berlin: Springer-Verlag.
- Epstein MJ, Haake P (2008) Very large scale ReliefF for genome-wide association analysis. In: Proceedings of IEEE Symposium on Computational Intelligence in Bioinformatics and Computational Biology.
- Meng Y, Yang Q, Cuenco KT, Cupples LA, Destefano AL, et al. (2007) Twostage approach for identifying single-nucleotide polymorphisms associated with rheumatoid arthritis using random forests and Bayesian networks. BMC Proc. 1 (Suppl 1): S56.
- Wan X, Yang C, Yang Q, Xue H, Tang N, et al. (2010) Predictive rule inference for epistatic interaction detection in genome-wide association studies. Bioinformatics 26(1): 30–37.
- Logsdon BA, Hoffman GE, Mezey JG (2010) A variational Bayes algorithm for fast and accurate multiple locus genome-wide association analysis. BMC Bioinformatics 11: 58.
- Zhang Y, Liu JS (2007) Bayesian inference of epistatic interactions in case control studies. Nature genetics 39: 1167–1173.
- Verzilli CJ, Stallard N, Whittaker JC (2006) Bayesian graphical models for genomewide association studies. The American Journal of Human Genetics 79: 100–112.
- Han B, Park M, Chen X (2009) A Markov blanket-based method for detecting causal SNPs in GWAS. In: Proceeding of IEEE International Conference on Bioinformatics and Biomedicine.
- Miller DJ, Zhang Y, Yu G, Liu Y, Chen L, et al. (2009) An algorithm for learning maximum entropy probability models of disease risk that efficiently searches and sparingly encodes multilocus genomic interactions. Bioinformatics 25(19): 2478–2485.
- Jiang X, Neapolitan RE, Barmada MM, Visweswaran S, Cooper GF (2010) A fast algorithm for learning epistatic genomics relationships. AMIA Annu Symp Proc 2010: 341–345.

- Li J, Horstman B, Chen Y (2011) Detecting epistatic effects in association studies at a genomic level based on an ensemble approach. Bioinformatics 27 (13): 222– 229.
- Evans DM, Marchini J, Morris A, Cardon LR (2006) Two-stage two-locus models in genome-wide association. PLOS Genetics 2(9): e157.
- Oh S, Lee J, Kwon MS, Weir B, Ha K, et al. (2012) A novel method to identify high order gene-gene interactions in genome-wide association studies: Genebased MDR. BMC Bioinformatics (Suppl 9): S5.
- Briggs FB, Ramsay PP, Madden E, Norris JM, Holers VM, et al. (2010) Supervised machine learning and logistic regression identifies novel epistatic risk factors with PTPN22 for rheumatoid arthritis. Genes and Immunity 11: 199– 208.
- Jiang X, Barmada MM, Cooper GF, Becich MJ (2011) A Bayesian method for evaluating and discovering disease loci associations. PLoS ONE 6(8): e22075.
- Iossifov I, Zheng T, Baron M, Gilliam TC, Rzhetsky A (2008) Genetic-linkage mapping of complex hereditary disorders to a whole-genome molecularinteraction network. Genome Res. 18(7): 1150–1162.
- Neapolitan RE (2004) Learning Bayesian Networks. Upper Saddle River, NJ: Prentice Hall.
- Perrier E, Imoto S, Miyano S (2008) Finding optimal Bayesian network given a super-structure. Journal of Machine Learning Research 9: 2251–2286.
- Spirtes P, Glymour C, Scheines R (2000) Causation, Prediction, and Search. New York; Springer-Verlag; 1993; second edition: Boston, MA; MIT Press.
- Cooper GF, Herskovits E (1992) A Bayesian method for the induction of probabilistic networks from data. Machine Learning 9: 309–347.
- Heckerman D, Geiger D, Chickering D (1995) Learning Bayesian networks: the combination of knowledge and statistical data. Technical Report MSR-TR-94-09. Microsoft Research.
- 44. Silander T, Kontkanen P, Myllymäki P (2007) On sensitivity of the MAP Bayesian network structure to the equivalent sample size parameter. In Proceedings of the Proceedings of the 23rd Conference Annual Conference on Uncertainty in Artificial Intelligence (UAI-07). Corvallis, Oregon: AUAI Press.
- Singh AP, Moore AW (2005) Finding optimal Bayesian networks by dynamic programming. CMU-CALCD-05-0106.
- 46. Fisher J, Urbanowicz R, Kiralis J, Sinnott-Armstrong N, Heberling T, et al. (2012) GAMETES: a fast, direct algorithm for generating pure, strict, epistatic models with random architectures. under review. Software: GAMETES (Version 1.0 Beta). Available: http://sourceforge.net/projects/gametes/.
- Hunter DJ, Kraft P, Jacobs KB, Cox DG, Yeager M, et al. (2007) A genomewide association study identifies alleles in *FGFR2* associated with risk of sporadic postmenopausal breast cancer. Nature Genetics 39: 870–874.