A Bayesian Method for Evaluating and Discovering Disease Loci Associations

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Abstract

Background: A genome-wide association study (GWAS) typically involves examining representative SNPs in individuals from some population. A GWAS data set can concern a million SNPs and may soon concern billions. Researchers investigate the association of each SNP individually with a disease, and it is becoming increasingly commonplace to also analyze multi-SNP associations. Techniques for handling so many hypotheses include the Bonferroni correction and recently developed Bayesian methods. These methods can encounter problems. Most importantly, they are not applicable to a complex multi-locus hypothesis which has several competing hypotheses rather than only a null hypothesis. A method that computes the posterior probability of complex hypotheses is a pressing need.

Methodology/Findings: We introduce the Bayesian network posterior probability (BNPP) method which addresses the difficulties. The method represents the relationship between a disease and SNPs using a directed acyclic graph (DAG) model, and computes the likelihood of such models using a Bayesian network scoring criterion. The posterior probability of a hypothesis is computed based on the likelihoods of all competing hypotheses. The BNPP can not only be used to evaluate a hypothesis that has previously been discovered or suspected, but also to discover new disease loci associations. The results of experiments using simulated and real data sets are presented. Our results concerning simulated data sets indicate that the BNPP exhibits both better evaluation and discovery performance than does a p-value based method. For the real data sets, previous findings in the literature are confirmed and additional findings are found.

Conclusions/Significance: We conclude that the BNPP resolves a pressing problem by providing a way to compute the posterior probability of complex multi-locus hypotheses. A researcher can use the BNPP to determine the expected utility of investigating a hypothesis further. Furthermore, we conclude that the BNPP is a promising method for discovering disease loci associations.

Introduction

The advent of high-throughput technologies has enabled genome-wide association studies (GWAS). A GWAS can involve examining a million representative single-nucleotide polymorphisms (SNPs) in individuals from some population. Often GWAS are conducted on cases and controls, where cases are individuals with a disease and controls are individuals without the disease. We then investigate the statistical association of each SNP with the disease. In doing so, a million hypotheses (disease-SNP relationships) or more may be investigated.

GWAS studies provide researchers unprecedented opportunities to investigate the complex genetic basis of diseases such as cancer. For example, GWAS have indicated that alleles in the FGFR2 gene are associated with sporadic postmenopausal breast cancer [1]; that five loci are associated with breast cancer including the plausible causative genes FGFR2, TNRC9, MAP3K1, and LSP1 [2]; and that GAB2 alleles may modify Alzheimer’s risk in APOE ε4 carriers [3]. Studies investigating SNPs in tumorous and nontumorous tissue have revealed somatic mutations possibly associated with cancer. For example, recent studies showed eight genes somatically mutated in glioblastoma tumors [4], and 26 genes somatically mutated in lung adenocarcinoma [5]. The 1000 Genomes Project plans to produce sequence coverage that will extend the list of human genetic variation [6], and gene-environment-wide association studies are emerging [7], both of which will increase the number of hypotheses investigated. Epistasis is the interaction between two or more genes to affect a phenotype such as disease susceptibility. 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. Researchers now believe that epistasis may account for a significant portion of the dark matter of genetic risk for disease [8], and it is becoming increasingly commonplace for researchers to investigate epistasis using GWAS data sets [9,10], which dramatically increases the number of hypotheses investigated. For example, if we only
considered all 2-SNP interactions when there are 500,000 SNPs, we would have $1.25 \times 10^{11}$ additional hypotheses.

These exciting possibilities for learning potential disease risk from high-dimensional data sets presents us with a challenge - namely how do we analyze and interpret our results when there are possibly billions of hypotheses? The hypothesis testing involved here is substantially different than that involved in a typical analysis where we might analyze the effect of a new drug. In this latter case, we are analyzing only one hypothesis, and the drug has a fairly high prior probability of being effective, otherwise the study would not have been considered. In discovery studies involving many hypotheses, each hypothesis has a very low prior probability.

Historically, the most common strategy for handling this multiple hypotheses testing problem has been to control type I error (false discovery) by using the Bonferroni correction to constrain the family-wise error rate. For example, the results in [3] were reported as being significant with Bonferroni correction. However, these corrected results often fail to duplicate across studies [8]. More recently developed techniques include the false discovery rate [11], false positive report probability [12], and Bayesian false discovery probability [13].

These methods all have the same purpose, namely to provide us with a way to decide which SNPs to “flag as noteworthy for further investigation” [13]. A difficulty with these methods is that they are not applicable to a complex multi-locus hypothesis, which has several competing hypotheses rather than only a null hypothesis. However, as mentioned above, it is becoming increasingly commonplace to investigate gene-gene interactions. So, a method that computes the posterior probability of a complex multi-locus hypothesis (and thereby flags the SNPs in the hypothesis as noteworthy) is a pressing need. In the Methods section we present a fully Bayesian method called the Bayesian network posterior probability (BNPP) method that is able to handle multi-locus hypotheses by computing the posterior probability of a hypothesis; it does so by assigning prior probabilities over all the hypotheses and computing the likelihoods of specialized Bayesian network structures [14], as explained below.

The Results section shows results of experiments illustrating the effectiveness of the BNPP at both evaluation and discovery, using both simulated and real data sets. In the remainder of this section we briefly review current methods and point out difficulties that they encounter.

When testing multiple hypotheses as in a GWAS, one of the hypotheses is likely to have a significant $p$-value by chance. As a result, researchers often use the Bonferroni correction to control the family-wise error rate by multiplying the $p$-value by the number of hypotheses $n$. For example, if $p = 2.1 \times 10^{-7}$ for a given SNP-outcome association and $n = 500,000$, then the Bonferroni-corrected $p$-value is $p \times n = 500,000 \times 2.1 \times 10^{-7} = 0.105$. This result would not be deemed significant by most standards, and the null hypothesis would not be rejected. A related correction is the Šidák correction, which is $1-(1-p)^n$.

Wakefield [13] notes that in the case of a GWAS the Bonferroni correction will often be an overly conservative procedure since at least in the current early stages of such studies we are more concerned with avoiding missed associations, and making some false discoveries is not too high a cost to pay to find real associations. Neapolitan [15] has a more fundamental problem with the Bonferroni correction. He argues that it is a misguided practice, and that the significance we attach to a result concerning a particular hypothesis cannot depend on the number of hypotheses we happen to test along with that hypothesis.

Regardless of one’s stance on this matter, there are clear difficulties in applying the Bonferroni correction in GWA studies. Suppose that one study investigates 100,000 SNPs while another investigates 500,000 SNPs. Suppose further that the data concerning a particular SNP and the disease is identical in the two studies. Due to the different corrections, that SNP could be reported as significant in one study but not the other. Yet the data concerning the SNP is identical in the two studies. As noted earlier, in GWAS results are often not duplicated across studies.

One reason may be the practice of using different corrections across different studies. Initially GWAS data were analyzed by investigating only 1-SNP models (hypotheses). We use the terms “model” and “hypothesis” interchangeably. Strictly speaking, the hypothesis is the statement that the model is correct.

So if there were 500,000 SNPs, there would be 500,000 hypotheses. Based on these results, quite a few results have been reported as significant with correction [3,16–18]. It is becoming increasingly popular to also investigate 2-SNP models in the effort to identify epistatic relationships [8–10]. As mentioned above, if there are 500,000 SNPs, there are about $1.25 \times 10^{11}$ 2-SNP models. If the researchers who previously reported significant results had also investigated the 2-SNP models, the corrections would have been based on many more hypotheses and the results likely would not have been reported as significant.

Realizing these problems, some researchers [17,19] have suggested that we uniformly use a Bonferroni correction assuming $10^6$ independent tests in GWA studies. This value was arrived at based on assuming only 1-SNP models are tested. If this were done, the problem concerning using different corrections when analyzing 1-SNP models in different studies would be addressed, but the problem of analyzing 1-SNP models along with 2-SNP models would not. We could correct the significance of a 1-SNP model assuming $10^6$ tests, and then perhaps we could correct the significance of a 2-SNP model assuming a much larger number of tests. However, even if we did all this we would still have the problem identified by Wakefield [13] concerning the correction being overly conservative, and the problem that we have ignored the probability of the data given the alternate hypothesis (power).

Benjamini and Hochberg [11] concluded that a desirable error rate to control is the expected proportion of errors among the rejected null hypotheses, which they termed the false discovery rate (FDR). That is, the FDR is $E(V/R)$, where $E$ denotes expected value, $R$ is the number of null hypotheses rejected and $V$ is the number of true null hypotheses rejected (Recall that we make a discovery when we reject a null hypothesis). They prove the following theorem, which enables us to control the FDR in practice: Suppose we have $n$ hypotheses with corresponding $p$-values $p_1 \leq p_2 \leq \ldots \leq p_n$. Denote by $H_i$ the null hypothesis corresponding to $p_i$. Let

\[ k \text{ be the largest } i \text{ such that } p_i \leq \frac{i}{n} q. \]

Then if we reject $H_1, H_2, \ldots, H_k$, the FDR is $\leq q$.

Storey [20] gave the FDR a Bayesian interpretation, showed that the $E(V/R|R>0)$ (called the positive FDR) does not depend on $n$, and defined the $q$-value. Storey and Tibshirani [21] develop an empirical method for estimating the $q$-value from the observed distribution of $p$-values.

Storey et al. [22] developed a method that computes the posterior probability that a locus is in the true model given the data, without ever estimating the entire true model. However, this method is applicable to the situation in which we are investigating many phenotype traits simultaneously rather than a single trait. The FDR was used to correct for multiple comparisons in this method.
The next two methods discussed concern the following analysis. We test \( H_0 : \theta = 0 \) vs. \( H_1 : \theta > 0 \) where \( \theta \) is the log odds ratio. For example, if \( H_1 \) is the hypothesis that a particular SNP \( S \) is associated with disease \( D_i \):

\[
\theta = \ln \left( \frac{\text{Odds}(D = 1 | S = 1)}{\text{Odds}(D = 1 | S \neq 1)} \right),
\]

where by \( D = 1 \) we mean the disease is present, and by \( S = 11 \) we mean that an individual has two copies of the mutant allele 1. We have assumed in this example that the wild type is dominant. The model assumes a test statistic \( T \) with \( E(T) = \theta \). For example, we may fit a logistic regression model so that \( T \) is the maximum likelihood estimate of the log odds ratio.

The false positive report probability (FPRP) [12] is defined as follows:

\[
\text{FPRP} = P(H_0 | \text{Data}) = \frac{p \times P(H_0)}{p \times P(H_0) + \text{power} \times P(H_1)},
\]

where \( p \) is the \( p \)-value, the \( \text{Data} \) are assumed to be the result that \( |T| > t_{\text{obs}} \) where \( t_{\text{obs}} \) is the observed value of \( T \), and \( \text{power} = P(\text{Data} | \theta_1) \) is evaluated at a pre-specified \( \theta_1 \).

There are a number of difficulties with this method:

1. Information is being lost by considering the data as being the result that \( |T| > t_{\text{obs}} \) rather than the point value \( (t_{\text{obs}}) \) we observed.
2. How do we decide on a particular value of \( \theta_1 \)? Perhaps we should consider a range of values.
3. The odds ratio only considers two possibilities, either a condition is present or it is not. However, we may want to model that there could be a different effect on disease for each of the three values a SNP can obtain.
4. We can only consider a null hypothesis \( H_0 \) and an alternative hypothesis \( H_1 \). However, if we model 1-SNP, 2-SNP, 3-SNP models, etc., there are several competing models (hypotheses) besides the one whose probability we are computing and each has a different likelihood. This issue is discussed in more detail in the Methods section.

As an alternative to the FPRP, Wakefield [13] developed the Bayesian false discovery probability (BFDP) which addresses several of the difficulties just presented. We do not go into its details here, but only mention that it does not attend to Difficulties 3 and 4.

A Bayesian method was used to compute the strength of association of a finding obtained using GWAS data in the Methods section. We test \( H_0 \) vs. \( H_1 \) where \( \theta \) denotes the parameters in the model. For all three models a logistic regression model was used for the likelihood \( P(\text{Data} | \theta, H_i) \). The log of the Bayes factor was reported for both \( H_1 \) and \( H_2 \). This method addresses Difficulty 3 to some extent by considering three values of the genotype. However, it does not concern multi-locus hypotheses and address Difficulty 4.

Zhang and Liu [24] developed Bayesian epistasis association mapping (BEAM) for the purpose of discovery. However, the method does assign prior probabilities to loci being associated with the disease and reports posterior probabilities. It does not consider multiple competing hypotheses.

Sebastiani et al. [25] computed the posterior probabilities of individual SNPs using a likelihood like the one presented here. These researchers performed a GWAS concerning 298,734 SNPs with the purpose of developing a system for predicting extended longevity (EL). In the first stage of their investigation they computed the posterior probability of each SNP being associated with EL, where the prior probability was assumed to be 0.5. They used these posterior probabilities to rank the SNPs and thereby flag SNPs to include in the second phase of the investigation, which was to decide which SNPs to include in the predictive model.

Bayesian networks have previously been used to discover disease loci interactions using likelihoods [10,26–28]. However, we know of no previous research that used them to determine the posterior probability of a complex multi-SNP model being associated with the disease.

### Methods

We developed the BNPP method specifically to enable us to compute the posterior probability of multi-locus models, which addresses Difficulty 4 above; however, it also attends to the other three difficulties.

A 1-locus model is the model that a single locus by itself is associated with a phenotype such as a disease, a 2-locus model is the model that two loci together are associated with a phenotype, and so on. The BNPP method represents such models using particular types of Bayesian network structures and computes the posterior probability of a model based on the likelihoods of these structures and their prior probabilities.

The BNPP was designed for the purpose of flagging SNPs for further investigation; that is, it is intended to compute the posterior probability of a model that was already discovered or conjectured. We previously used Bayesian networks for discovery of disease loci associations [10,26,27]. However, we only computed the likelihoods of the models; we did not consider their prior probabilities. A bigger model (more loci) will sometimes have a higher likelihood, but be less probable because of its smaller prior probability. The BNPP accounts for this situation, whereas a method that only looks at likelihoods does not. So, the BNPP is also a new, promising technique for discovery.

Before describing the BNPP algorithm, we first review Bayesian networks on which the algorithm is based.
Bayesian Networks

Bayesian networks [14] have been used for modeling and knowledge discovery in many domains, including applications to bioinformatics [29]. A Bayesian network (BN) consists of a directed acyclic graph (DAG) \( G \) whose set of nodes \( V \) contains random variables and a joint probability distribution \( P(V) \) that satisfies the Markov condition with \( G \). We say that \((G, P)\) satisfies the Markov condition if for each variable \( X \in V \), it holds that \( X \) is conditionally independent in \( P \) of the set of all its nondescendents in \( G \) given the set of all its parents in \( G \). It is a theorem [14] 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 for 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 this theorem, BNs are often developed by first defining 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.

Figure 1 shows a BN representing the causal relationships among gene expression levels. The expression levels have been discretized into two values, low and high. 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 \( C = \text{low} \) and \( S = \text{high} \), we can determine the probability of \( A \) being low and of \( D \) being low. That is, we can compute

\[
P(A = \text{low}|C = \text{low}, S = \text{high}) = \] 

and \( P(D = \text{low}|C = \text{low}, S = \text{high}) \).

Methods have been developed both for learning the parameters in a BN and the structure (called a DAG model) from data. The research discussed here concerns structure learning, which we discuss next. 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 and their expression levels of the genes shown in Figure 1, we might be able to learn the DAG in Figure 1 from data. When the edges represent causal influences, this means we can learn causal influences from data under assumptions. In the score-based structure learning approach, we assign a score to a DAG based on how well the DAG fits the data.

Cooper and Herskovits [30] developed the Bayesian score for discrete variables, which is the probability of the Data given the DAG. This score uses a Dirichlet distribution to represent our prior belief for each conditional probability distribution in \( G \) and contains hyperparameters that represent these prior beliefs. The score is as follows:

\[
score_{Bayes}(G : \text{Data}) = P(\text{Data}|G) = \prod_{i=1}^{n} \prod_{k=1}^{r_i} \frac{\Gamma(\sum_{j=1}^{q_i} a_{ijk})}{\Gamma(a_{ijk} + s_{ijk})} = \prod_{i=1}^{n} \prod_{k=1}^{r_i} \frac{\Gamma(a_{ijk} + s_{ijk})}{\Gamma(a_{ijk})},
\]

where

1. \( n \) is the number of variables in the DAG model \( G \);
2. \( r_i \) is the number of states of \( X_i \);
3. \( q_i \) is the number of different values that the parents of \( X_i \) in \( G \) can jointly assume;
4. \( a_{ijk} \) is our assessed prior belief from previous experience (before obtaining the current data) of the number of times \( X_i \) took its \( k \)th value when the parents of \( X_i \) took their \( j \)th value;
5. \( s_{ijk} \) is the number of times in the data that \( X_i \) took its \( k \)th value when the parents of \( X_i \) took their \( j \)th value.

The Bayesian score does not necessarily 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 \( X \rightarrow Y \) and \( X \leftarrow Y \) are Markov equivalent. Heckerman et al. [31] show that if we determine the values of the hyperparameters from a single parameter \( z \) called the prior equivalent sample size then Markov equivalent DAGs obtain the same score. If we use a prior equivalent sample size \( z \) and want to represent a prior uniform distribution for each variable in the network, then for all \( i, j, k \) and \( s_{ijk} \) are defined as above. When we use a prior equivalent sample size \( z \) in the Bayesian score, the score is called the Bayesian Dirichlet equivalent (BDe) score. When we also represent a prior uniform distribution for each variable, the score is called the Bayesian Dirichlet equivalent uniform (BDeu) score and is given by the following formula, which is a special case of Equation 1:

\[
P(\text{Data}|G) = score_{BDe}(G : \text{Data}) = \prod_{i=1}^{n} \prod_{j=1}^{r_i} \frac{\Gamma(z/q_i)}{\Gamma(z/q_i + s_{ijk})} = \prod_{i=1}^{n} \prod_{j=1}^{r_i} \frac{\Gamma(z/q_i + s_{ijk})}{\Gamma(z/r_i q_j)}.
\]

Figure 1. A Bayesian network showing possible relationships among gene expression levels. The levels have been discretized to the values low and high. The network is for illustration purposes only; it is not meant to accurately portray real relationships.

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Posterior Probabilities of Disease-SNP Models

In what follows, for simplicity we refer to variables that might be associated with a disease as SNPs. However, in general they could be any genetic information or environmental factors. We can represent the relationship between a disease and SNPs using simple DAG models like those shown in Figure 2. The first model represents that SNP $S_i$ is associated with disease $D$. The second model represents that $S_i$ and $D$ together are associated with $D$ (this could happen because each individually is associated with $D$ or because together they are associated with $D$ due to an epistatic interaction), and the fourth model represents that SNPs $S_i$, $S_j$, and $S_k$ together are associated with $D$.

Our goal is to compute the posterior probability of a model $M$ given Data. We can do that using Bayes' Theorem as follows:

$$ P(M|\text{Data}) = \frac{P(\text{Data}|M)P(M)}{P(\text{Data})}. \tag{3} $$

The $P(\text{Data}|M)$ term can be computed using the BDeu score (Equation 2) with a particular choice of $\alpha$. The BDeu score has been used successfully to learn epistatic interactions from real GWAS data sets, and in one analysis [26] it has been shown to more often identify the model generating the data than other methods discussed previously.

Computing the BNPP

Consider first a 1-SNP model. Let $M_i$ be the model that $S_i$ is associated with $D$ and $M_0$ be the model that it is not (see Figure 3). Then the posterior probability of $M_i$ is given by

$$ P(M_i|\text{Data}) = \frac{P(\text{Data}|M_i)P(M_i)}{P(\text{Data}|M_0)P(M_0) + P(\text{Data}|M_0)P(M_0)}. \tag{4} $$

Note that the model in Figure 3 is not just that $S_i$ is associated with the disease, but rather that it is associated all by itself. That is, if $S_i$ was involved in an epistatic interaction with no marginal effects, the model would be false. Note further that $S_i$ can have any number of discrete values in the model. We are not restricted to only two values as in some of the methods discussed previously. So we can represent all three values of a SNP, or if we are representing an environmental feature with multiple values we can represent all of them. If the environmental feature is continuous, we can discretize it. So we overcome Difficulty 3 mentioned in the introduction (recall that Difficulty 3 is that the odds ratio only considers two possibilities, either a condition is present or it is not).

Figure 4 shows the model $M_j$ that $S_i$ and $S_j$ together are associated with $D$ (without needing other interacting SNPs). Note that this model includes the possibility that there is epistasis with no marginal effects, as well as the possibility that each SNP by itself has an association with $D$. The three competing models are on the right. Note further that the model denoted as $M_i$ is not the same as the model $M_j$ in Figure 3. Model $M_i$ in Figure 4 represents that $S_i$ is not associated with $D$ either by itself (other than possibly through $S_j$) or together with $S_i$, whereas model $M_j$ in Figure 3 says nothing about $S_j$.

No other method discussed in the introduction considers these multiple competing hypotheses. They would only consider the null hypothesis $M_0$ in which no association with $D$ holds versus $M_j$. However, if either model $M_i$ or $M_j$ were the correct model, we would observe an association of the two SNPs together with $D$ (and therefore reject $M_0$) even though $M_j$ is incorrect. An example of this situation is the relationship between $APOE$ and rs41377151, which will be discussed when we analyze the data in Figure 3. Then the posterior probability of $M_j$ is given by

$$ P(M_j|\text{Data}) = \frac{P(\text{Data}|M_j)P(M_j)}{P(\text{Data}|M_j)P(M_j) + P(\text{Data}|M_0)P(M_0) + \sum_k P(\text{Data}|M_k)P(M_k)} \tag{5} $$

where $k$ sums over the two 1-SNP models.

Figure 5 shows a 3-SNP model and the competing models. The number and complexity of the competing models increases with the size of the model. However, we need not identify all the competing models because we have developed the following recursive algorithm for computing $P(\text{Data})$ for an arbitrary number of SNPs, which is the denominator in the formula for the posterior probability of a model:

**Algorithm:** Compute $P(\text{Data})$.

The SNPs in the model being evaluated are $S_1$, $S_2$, ..., $S_n$. $\text{prior}[m]$ is the prior probability of an $m$-SNP model.
We assume all $m$-SNP models have the same prior probability, but this assumption is not necessary.

$$P(Data) = 0;$$

for $m = 0$ to $n$

$$\text{likelihood} = 0;$$

$M = \emptyset;$

ComputeLikely($1$, $m$);

$$P(Data) = P(Data) + \text{likelihood} \times \text{prior}[m];$$

endfor

procedure ComputeLikely($k$, $m$) // $m$ is the size of the model being considered.

if $m = 0$

$$\text{likelihood} = \text{likelihood} + P(Data|M);$$ // $\text{likelihood}$ and $M$ are global to this procedure.

else

for $i = k$ to $n - m + 1$

add $S_i$ to $M$;

ComputeLikely($i+1$, $m-1$);

remove $S_i$ from $M$;

endif

There are $n$ SNPs in the model being analyzed. The algorithm proceeds by calling procedure ComputeLikely for every $m \leq n$. For each value of $m$ this routine then computes the contribution of all $m$ SNP models to the likelihood by recursively visiting all such models. Since every subset of the $n$ SNPs determines a competing model, the likelihoods for $2^n$ models are computed. However, since ordinarily there are at most 5 SNPs in a model, this computation is feasible.

There are various possibilities for the data structure we could use in representing a model. We currently choose to represent a model simply as an $n$-element array $M$, where $M[i]$ contains the index of the $i$th SNP in the model. For example, if $n = 3$ and $S_2$, $S_4$, and $S_{10}$ are the SNPs in the model, then $M[1] = 2$, $M[2] = 4$, and $M[3] = 10$.

**Results**

Next we present results of evaluating the BNPP using both simulated and real data sets. All experiments were done using a
Macbook Pro notebook with a 2.66 GHz processor and 8 GB of RAM. For the sake of focus, in what follows we will always refer to the phenotype as a disease.

Simulated Data

Velez et al. [33] created 70 epistasis models that are described in Supplementary Table one to that paper. Each model represents a probabilistic relationship in which two SNPs together are statistically associated with the disease, but neither SNP is individually predictive of disease. The relationships represent various degrees of penetrance, heritability, and minor allele frequency. Data sets were generated with case-control ratio of 1:1. To create one data set they fixed the model. Based on the model, they then generated data concerning the two SNPs that were related to the disease in the model, 10 other unrelated SNPs, and the disease. For each of the 70 models, 100 data sets were generated for a total of 7000 data sets. This procedure was followed for data set sizes equal to 200, 400, 800, and 1600. The data sets were generated separately. See http://discovery.dartmouth.edu/epistatic_data to obtain these data sets.

For each of these data sets, we computed the posterior probability of each 1-SNP, 2-SNP, and 3-SNP model using the BNPP, making a total of 1350 models investigated per data set. As discussed in Supporting Information S1, researchers estimate that in an agnostic study the prior probability of an individual SNP being associated with a disease is between 0.0001 and 0.00001. An agnostic study is an explorative study in which we have no special prior belief concerning any particular locus. Lower and upper posterior probabilities were obtained using each of these priors for an individual SNP being associated with a disease, and using the strategy for determining model priors based on individual SNP priors, which is also presented in Supporting Information S1. To compute the likelihoods the BDeu score (Equation 2) was used. The hyperparameter \( \alpha \) was set equal to 54 because this value yielded good epistasis discovery in a previous study [27] using the Velez and other data.

Table 1 shows the results. The average probability of the true models is much higher than that of the false models. Furthermore, this average probability is moving toward 1 as the sample size increases, whereas that of the false models remains quite small. Finally, the results for the two different priors are not substantially different. This robustness result is encouraging because the assessment of priors is arguably the most contentious part of a Bayesian analysis.

We repeated the analysis using \( p \)-values obtained from Pearson's chi-square test. Table 2 shows the results. These \( p \)-values are uncorrected since a Bonferroni or Šidák correction would be the same for all of them, and therefore not change the relative order. Notice that the average \( p \)-value of the best false models is smaller than that of the true models (recall that smaller \( p \)-values are more significant). Table 1 shows that the average posterior probability of the best false models is smaller than that of the true models (larger posterior probabilities are more significant).

The performance of an evaluation method can be judged by how high it ranks true models and how low it ranks false models. The previous results support that the BNPP algorithm exhibits better evaluation performance than the method based on \( p \)-values.

Next we address discovery. Figure 6 shows ROC curves concerning the posterior probabilities when the individual SNP prior is 0.00001 and the \( p \)-values for the simulated data sets. The results for the posterior probabilities were almost identical when the prior was 0.0001; so we do not show them. A receiver operating characteristic (ROC) curve plots the true positive rate (sensitivity) on the y-axis and the false positive rate (1 - specificity) on the x-axis. It is obtained by considering various threshold probabilities as being binary indicators of discovery. For example, the point (0.014, 0.449) appears on the curve for the posterior probability in Figure 6 (a) because 0.014 fraction of the false models have posterior probabilities exceeding a threshold (in this case \( 2.95 \times 10^{-5} \)), while 0.449 fraction of the true models have posterior probabilities exceeding this threshold. The point (0.16, 1.0) appears on the curves in Figures 6 (a), (b), and (c) and the point (0.13, 1.0) appears on the curve in Figure 6 (d). This means that if we were using the posterior probability as a binary indicator of discovery in the case of samples sizes of 200, 400, or 800, we could discover all the true models with a false discovery rate of about 16% (based on this analysis). On the other hand, the true positive rate for the \( p \)-value does not reach 1 until the false positive rate reaches 1. This is true even when the data set has size 1600 (this is not noticeable in the display of its ROC curve). This result supports the effective discovery performance of the BNPP.

Velez et al. [33] showed that models 55-59 in the Velez Data are the most difficult models to learn. They have the weakest broad-sense heritability (0.01) and a minor allele frequency of 0.2. These models are arguably most like relationships we might find in nature. ROC curves concerning only these models appear in Figure 7. Although the curves for the posterior probability are not that much worse than when we consider all models, the ones for the \( p \)-value are substantially worse except when the sample size is 1600. The worst possible ROC curve is a straight line from (0,0) to (1,1). The \( p \)-value ROC curve when the sample size is 200 is not much better than that line.

We can perhaps apply different corrections to different sized models and stay in the framework in which the correction is applied by arguing that 1-SNP models, 2-SNP models, and 3-SNP models are different families of models and we should apply different corrections for each of these families. Since there were 20 SNPs total in the simulations, we applied the Šidák correction using 20 for 1-SNP models, \( \binom{20}{2} = 190 \) for 2-SNP models, and \( \binom{20}{3} = 1140 \) for 3-SNP models. The resultant curves appear with a dotted line in Figure 7. Although we have improved the results, they are still not as good as those for the posterior probability. Also, if we did a study with a different number of SNPs, we would need to apply a different correction, and we would expect to obtain different results. On the other hand, the BNPP model suggests using the same prior probabilities across all agnostic studies.

The average times to compute the posterior probabilities and the \( p \)-values for all one to three SNP models were 1.8 seconds and 0.7 seconds respectively.

Real Data

Alzheimer's Data set. Reiman et al. [3] analyzed a GWAS late onset Alzheimer's disease (LOAD) data set on 312,317 SNPs from an Affymetrix 500K chip, plus the measurement of a locus in the \( APOE \) gene, which is known to be predictive of LOAD. The data set consists of three cohorts containing a total of 1411 participants. Of the 1411 participants, 861 had LOAD and 550 did not. In addition, 644 participants were \( APOE \) e4-carriers, who carry at least one copy of the \( APOE \) e4 genotype and 767 were \( APOE \) e4 non-carriers. See http://www.tgen.org/neurogenomics/data concerning this data set. Reiman et al. found the \( APOE \) gene is significantly associated with LOAD. The \( GAB2 \) gene is not significantly associated with LOAD, the \( GAB2 \) gene is significantly associated with LOAD in \( APOE \) e4 carriers, and the \( GAB2 \) gene is not significantly associated with LOAD in the \( APOE \) e4 non-
A Bayesian Method for Disease Loci Evaluation

carriers. These results indicate that APOE and GAB2 may interact epistatically to affect LOAD. Using these same data, we computed the posterior probability of each locus being associated with LOAD (1-locus models), and the posterior probability of each locus together with APOE being associated with LOAD (2-locus models).

The average posterior probability of all 1-locus models was $2.85 \times 10^{-5}$ for the individual SNP prior equal to 0.0001, and $9.18 \times 10^{-6}$ for that prior equal to 0.00001. Furthermore, the numbers of models (loci) with posterior probabilities less than 0.01 were respectively $312,301$ and $312,273$ for the two priors. Figure 8 shows bar charts depicting the results concerning the remaining loci. Table 3 shows the loci in the 10 most probable models. APOE has a posterior probability of $\sim 1$, regardless of the prior, as does SNP rs41377151. SNP rs41377151 is on the APOC1 gene, which is in strong linkage disequilibrium with APOE and for which previous studies have indicated that they predict LOAD equally well [34]. The 3rd most probable locus is rs1082930, which is on the PRKG1 gene. There are a number of previous studies associating this gene with LOAD [35,36]. Of the seven remaining probable loci, there is some previous evidence linking four of them to LOAD [37].

As mentioned in Supporting Information S1, as more genome-wide association studies are carried out we will better be able to assess appropriate priors. These results indicate that 0.00001 may be more appropriate than 0.0001 since the latter prior resulted in fairly high posterior probabilities for three SNPs that have no known previous association with LOAD; nonetheless, these might be valid predictors of LOAD that have not been appreciated previously.

The average posterior probability of all 2-locus models, in which one of the loci was APOE, was $1.05 \times 10^{-4}$ for the individual SNP prior equal to 0.0001 and $1.41 \times 10^{-5}$ for that prior equal to 0.00001. Furthermore, the numbers of models with posterior probabilities less than 0.01 were respectively 312,267 and 312,028 for the two priors. Figure 9 shows bar charts depicting the results concerning the remaining models. Table 4 shows the loci in the ten most probable models. Eight of those loci are SNPs located on the GAB2 gene. The prior probability of a 2-SNP model is $6 \times 10^{-10}$ when the individual SNP prior is 0.0001 and $6 \times 10^{-12}$ when that prior is 0.00001 (See Supporting Information S1). We see from Table 4 that the posterior probabilities of 2-locus models containing APOE and a GAB2 SNP are much greater than these prior probabilities. On the other hand, the 1-locus models containing GAB2 SNPs had posterior probabilities about equal to their prior probabilities. These results together indicate GAB2 by itself does not affect LOAD, but that GAB2 interacts with APOE to affect LOAD.

The two loci in the top ten 2-locus models that are not on GAB2, namely SNPs rs6784615 and rs12162084, are among the 10 most probable 1-locus models (see Table 3). These results together indicate that each of these SNPs may affect LOAD independently of APOE. As indicated in Table 3, previous studies have associated these SNPs with LOAD.

Another interesting result is that APOE and rs41377151 (the two loci with posterior probabilities about equal to 1 in Table 3), when considered together, had posterior probabilities of $1.25 \times 10^{-4}$ and $1.25 \times 10^{-5}$ for the individual SNP priors of 0.0001 and 0.00001 respectively. This result indicates that the model containing both loci is incorrect. As mentioned above, SNP

### Table 1. The posterior probability results for the simulated data sets.

<table>
<thead>
<tr>
<th>prior probability</th>
<th>sample size</th>
<th>times true model was highest</th>
<th>avg. posterior probability of true models</th>
<th>avg. posterior probability of best false models</th>
<th>avg. posterior probability of all false models</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00001</td>
<td>200</td>
<td>2783</td>
<td>0.2009</td>
<td>$4.5 \times 10^{-4}$</td>
<td>$4.5 \times 10^{-7}$</td>
</tr>
<tr>
<td>0.00001</td>
<td>400</td>
<td>4265</td>
<td>0.4555</td>
<td>0.0019</td>
<td>$1.72 \times 10^{-6}$</td>
</tr>
<tr>
<td>0.00001</td>
<td>800</td>
<td>5433</td>
<td>0.6502</td>
<td>0.0053</td>
<td>$4.62 \times 10^{-6}$</td>
</tr>
<tr>
<td>0.00001</td>
<td>1600</td>
<td>6191</td>
<td>0.8128</td>
<td>0.0069</td>
<td>$5.72 \times 10^{-6}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>200</td>
<td>3135</td>
<td>0.2651</td>
<td>0.0036</td>
<td>$3.85 \times 10^{-6}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>400</td>
<td>4549</td>
<td>0.5024</td>
<td>0.0095</td>
<td>$9.27 \times 10^{-6}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>800</td>
<td>5580</td>
<td>0.6910</td>
<td>0.0136</td>
<td>$1.32 \times 10^{-5}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>1600</td>
<td>6298</td>
<td>0.8351</td>
<td>0.0134</td>
<td>$1.25 \times 10^{-5}$</td>
</tr>
</tbody>
</table>

The 1st column shows whether the smaller or larger priors were used; the 3rd column shows the number of times (out of 7000 data sets) the true (i.e., the data-generating) model had the highest posterior probability; the 4th column shows the average posterior probability of the true models; the 5th column shows the average posterior probability of the most probable false models (in each of the 7000 data sets); and the last column shows the average posterior probabilities of all false models.

doi:10.1371/journal.pone.0022075.t001

<table>
<thead>
<tr>
<th>prior probability</th>
<th>sample size</th>
<th>times true model was highest</th>
<th>avg. posterior probability of true models</th>
<th>avg. posterior probability of best false models</th>
<th>avg. posterior probability of all false models</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0001</td>
<td>200</td>
<td>2783</td>
<td>0.2009</td>
<td>$4.5 \times 10^{-4}$</td>
<td>$4.5 \times 10^{-7}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>400</td>
<td>4265</td>
<td>0.4555</td>
<td>0.0019</td>
<td>$1.72 \times 10^{-6}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>800</td>
<td>5433</td>
<td>0.6502</td>
<td>0.0053</td>
<td>$4.62 \times 10^{-6}$</td>
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<td>0.0001</td>
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<td>0.0069</td>
<td>$5.72 \times 10^{-6}$</td>
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<td>0.0001</td>
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<td>$3.85 \times 10^{-6}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>400</td>
<td>4549</td>
<td>0.5024</td>
<td>0.0095</td>
<td>$9.27 \times 10^{-6}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>800</td>
<td>5580</td>
<td>0.6910</td>
<td>0.0136</td>
<td>$1.32 \times 10^{-5}$</td>
</tr>
<tr>
<td>0.0001</td>
<td>1600</td>
<td>6298</td>
<td>0.8351</td>
<td>0.0134</td>
<td>$1.25 \times 10^{-5}$</td>
</tr>
</tbody>
</table>

The values are like those in Table 1 except they concern the p-values obtained using Pearson’s chi-square test.

doi:10.1371/journal.pone.0022075.t002

### Table 2. The p-value results for the simulated data sets.

<table>
<thead>
<tr>
<th>sample size</th>
<th>avg. p-value of true models</th>
<th>avg. p-value of best false models</th>
<th>avg. p-value of all false models</th>
</tr>
</thead>
<tbody>
<tr>
<td>200</td>
<td>0.0605</td>
<td>0.0026</td>
<td>0.478</td>
</tr>
<tr>
<td>400</td>
<td>0.0271</td>
<td>0.0011</td>
<td>0.4829</td>
</tr>
<tr>
<td>800</td>
<td>0.0083</td>
<td>$3.92 \times 10^{-4}$</td>
<td>0.486</td>
</tr>
<tr>
<td>1600</td>
<td>0.0012</td>
<td>$7.87 \times 10^{-5}$</td>
<td>0.487</td>
</tr>
</tbody>
</table>
rs41377151 is on the APOC1 gene, and previous investigations have shown that APOE and APOC1 are in linkage disequilibrium and each of them predicts LOAD as well as the other [34]. However, we know of no previous study substantiating that the two loci identify the same single causal mechanism of LOAD. This result could not have been obtained with a method that only considered the null hypothesis that the two loci together are not associated with LOAD, and the alternative hypothesis that they are. For example, using Pearson’s chi-square test, we obtained $p$-values all equal to $\sim 0$ for APOE alone, rs41377151 alone, and APOE and rs41377151 together (the 2-locus model). The BNPP determined that the 2-locus model is improbable because it also evaluated the competing hypotheses that only one locus is directly causative of LOAD. To learn that the 2-locus model is not significantly better than the 1-locus model using commonly applied frequentist statistics, we would need to perform an analysis such as stepwise regression or regression on the two loci followed by an investigation of the coefficients.

The three interesting results just discussed (the first concerning GAB2, the second rs6784615 and rs12162084, and the third rs41377151) follow from our computing the posterior probabilities of all 1-locus models and all 2-locus models containing APOE. It was not necessary to suspect any of them ahead of time or perform a focused analysis.

The running times were 196 seconds and 193 seconds to investigate all 312,318 1-locus models using individual SNP priors $0.0001$ and $0.00001$, respectively. The corresponding running times to investigate all 2-locus models containing APOE were 593 seconds and 584 seconds.

**Breast Cancer Data set.** Hunter et al. [1] conducted a GWAS concerning 546,646 SNPs and breast cancer as part of the National Cancer Institute *Cancer Genetic Markers of Susceptibility* (CGEMS) Project. (see http://cgems.cancer.gov/.) They determined the significance of each SNP using logistic regression with two degrees of freedom. Two of the six most significant SNPs were on the FGFR2 gene. Furthermore, two other FGFR2 SNPs were among the 16 most significant SNPs. Previously, it was known that FGFR2 is amplified and overexpressed in breast cancer [38,39]. Furthermore, a large, three-stage GWAS of breast cancer had identified SNPs in FGFR2 as the strongest of its associations [2].

---

Figure 6. ROC curves concerning the posterior probabilities when the prior is 0.00001 and the $p$-values for the simulated data sets. The curve for the posterior probability is a solid line, while the one for the $p$-values is a dashed line. 1-specificity is on the x-axis and the sensitivity is on the y-axis.

doi:10.1371/journal.pone.0022075.g006
Figure 7. ROC curve concerning the posterior probabilities when the prior is 0.00001 and the \( p \)-values for models 55–59. The curve for the posterior probability is a solid line, the one for the \( p \)-value is a dashed line, and the one for the \( p \)-value with the Šidák correction is a dotted line. 1-specificity is on the x-axis and the sensitivity is on the y-axis.

doi:10.1371/journal.pone.0022075.g007

Figure 8. Bar charts showing the number of 1-locus models in each posterior probability range. The posterior probability is that of the model in which a single locus is associated with LOAD.

doi:10.1371/journal.pone.0022075.g008
Table 3. Results concerning the 10 most probable 1-locus models in the LOAD study in [3].

<table>
<thead>
<tr>
<th>locus</th>
<th>posterior probability</th>
<th>previous LOAD association</th>
</tr>
</thead>
<tbody>
<tr>
<td>APOE (1, 1)</td>
<td>(1, 1)</td>
<td>Yes</td>
</tr>
<tr>
<td>rs41377151 (1, 1)</td>
<td>(1, 1)</td>
<td>Yes</td>
</tr>
<tr>
<td>rs10824310 (0.124, 0.586)</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>rs4356530 (0.086, 0.485)</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>rs17330779 (0.066, 0.416)</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>rs6784615 (0.048, 0.335)</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>rs10115381 (0.027, 0.222)</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>rs12162084 (0.027, 0.217)</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>rs4862146 (0.024, 0.192)</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>rs249153 (0.017, 0.152)</td>
<td>Yes</td>
<td></td>
</tr>
</tbody>
</table>

Based on their results and these previous findings, Hunter et al. [1] investigated FGFR2 in three additional studies and found further support for an association of FGFR2 with breast cancer.

Using this same GWAS data set, we computed the posterior probability of all 1-locus models using the agnostic individual SNP priors of 0.00001 and 0.00001 and the informative priors of 0.01 and 0.1. The average posterior probability of all 1-locus models was 3.81 × 10⁻⁷ for the prior equal to 0.00001 and 3.78 × 10⁻⁶ for the prior equal to 0.0001. Furthermore, the numbers of models (loci) with posterior probabilities less than 0.01 were respectively 546,645 and 546,637 for the two priors. Table 5 shows results concerning the ten most probable models. Columns 2–5 show posterior probabilities in the second range

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Kuschel et al. [42] investigated 16 SNPs in seven genes involved in the repair of double-stranded DNA breaks and breast cancer in a case-control study involving 2200 cases and 1900 control subjects. Using standard significance testing, they found two polymorphisms in XRCC3 and one polymorphism in each of XRCC2 and LIG4 to be the most significant. They also performed a haplotype analysis investigating the effect of the genetic variants in the XRCC3 gene on breast cancer. Wacholder et al. [12] analyzed these same data using the FPRP method. Statistical power in their analysis was the power to detect an odds ratio of 1.5 for the homozygote with the rare genetic variant and an odds ratio of 1.0 for the homozygote with the common variant. Based on previous findings [40,41], Wacholder et al. [12] assigned a prior range of 0.01 to 0.1.

We analyzed these same data using the BNPP algorithm to obtain the posterior probabilities of the models. Table 6 shows the results. The last two columns show posterior probabilities of association with breast cancer; to make comparisons easier, we show 1-FPRP in columns 3 and 4. The p-values in the second column were computed using the chi-square test with two degrees of freedom.

The FPRP and BNPP exhibit similar results concerning the four SNPs and the haplotype, however, the results for BNPP are more

Figure 9. Bar charts showing the number of models in each posterior probability range. The posterior probability is that of the 2-locus model in which each locus together with APOE is associated with LOAD.

doi:10.1371/journal.pone.0022075.g009
concerning the rare homozygote in the case of the latter SNP has a smaller XRCC2 are somewhat larger than those for A Decision Analytic Approach to Using the BNPP probabilities (using both the FPRP and BNPP) for the only conditions on the models being true. Note that the posterior value would result in smaller posterior probabilities. The BNPP was used for statistical power in the case of the FPRP. A larger associated with breast cancer is either result in Table 5 that the posterior probability of rs10510126 being probable models are not very high. For example, consider the GWAS data sets, often the posterior probabilities of even our most very low. So, given the limited number of samples in current agnostic GWAS investigation the prior probabilities are ordinarily 

doi:10.1371/journal.pone.0022075.t005

<table>
<thead>
<tr>
<th>locus</th>
<th>posterior probability range</th>
<th>GAB2</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1007837</td>
<td>(0.266, 0.784)</td>
<td>No</td>
</tr>
<tr>
<td>rs7101429</td>
<td>(0.214, 0.731)</td>
<td>No</td>
</tr>
<tr>
<td>rs901104</td>
<td>(0.201, 0.715)</td>
<td>Yes</td>
</tr>
<tr>
<td>rs4291702</td>
<td>(0.139, 0.617)</td>
<td>Yes</td>
</tr>
<tr>
<td>rs4945261</td>
<td>(0.144, 0.564)</td>
<td>Yes</td>
</tr>
<tr>
<td>rs12162084</td>
<td>(0.144, 0.563)</td>
<td>No</td>
</tr>
<tr>
<td>rs7115850</td>
<td>(0.108, 0.547)</td>
<td>Yes</td>
</tr>
<tr>
<td>rs10793294</td>
<td>(0.099, 0.523)</td>
<td>Yes</td>
</tr>
<tr>
<td>rs2450130</td>
<td>(0.088, 0.491)</td>
<td>Yes</td>
</tr>
<tr>
<td>rs6784615</td>
<td>(0.081, 0.462)</td>
<td>No</td>
</tr>
</tbody>
</table>

doi:10.1371/journal.pone.0022075.t004

A Decision Analytic Approach to Using the BNPP

The question remains as to what to with BNPP results. In an agnostic GWAS investigation the prior probabilities are ordinarily very low. So, given the limited number of samples in current GWAS data sets, often the posterior probabilities of even our most probable models are not very high. For example, consider the result in Table 5 that the posterior probability of rs10510126 being associated with breast cancer is either 0.0118 or 0.1185 depending on whether the prior probability is 0.00001 or 0.0001. The average of these values, namely 0.0652, can be used to represent our posterior belief in the validity of this association. This value is not very high, and so one may ask whether it is significant. In general, statistics cannot tell us whether a result is significant; it can only change our belief. It has become a controversial practice by some to consider a p-value of 0.05 or smaller to be significant largely because of R.A. Fisher’s [43] statement in 1926 that “it is convenient to draw the line at about the level at which we can say: Either there is something in the treatment, or a coincidence has occurred such as does not occur more than once in twenty trials.” However, as has been often discussed, there is nothing special about the value 0.05 for a p-value that enables a dichotomous announcement, just as there is nothing special about a particular posterior probability.

The value 0.0652 represents our belief concerning the truth of the model based on our knowledge concerning the model, namely our prior belief and the data. Although we cannot dichotomously announce whether the value is significant, we can use it to make a decision about what to do. We should report the finding concerning model M if the expected utility of not reporting M is less than the expected utility of reporting M. Let UTD be the utility of a true discovery, which is the utility of reporting a true model, UFD be the utility of a false discovery, which is the utility of reporting a false model (and which is therefore negative), UTND be the utility of a true non-discovery, which is the utility of not reporting a false model, and UFNĐ be the utility of a false non-discovery, which is the utility of not reporting a true model (and which is therefore negative). We should report model M if

\[ U_{\text{FND}}(P(M|Data)) + U_{\text{TN}}(1-(P(M|Data)) \]

or

\[ U_{\text{TD}}(P(M|Data)) + U_{\text{FD}}(1-(P(M|Data))) \]

or

\[ U_{\text{TN}} - U_{\text{FD}} - U_{\text{TD}} - U_{\text{FND}} \leq P(M|Data) \leq Odds(M|Data). \]

In the current analysis, Odds(M|Data) = 0.0652/(1 - 0.0652) = 0.0697. So we should report the finding (and therefore investigate the model further) if (UTND - UFD)/(UTD - UFNĐ) < 0.0697.

If we take this decision-analytic approach to using the BNPP, we conclude that it provides researchers with a useful tool for guiding how they should proceed based on their findings.

Table 5. Results concerning the ten most probable models in the breast cancer study in [1].

<table>
<thead>
<tr>
<th>SNP</th>
<th>prior = 0.0001</th>
<th>prior = 0.0001</th>
<th>prior = 0.01</th>
<th>prior = 0.1</th>
<th>p-value</th>
<th>Šidák</th>
<th>previous BC association</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs10510126</td>
<td>0.0118</td>
<td>0.1185</td>
<td>0.9967</td>
<td>0.9992</td>
<td>2.4 x 10^{-6}</td>
<td>0.7307</td>
<td>No</td>
</tr>
<tr>
<td>rs17157903</td>
<td>0.0031</td>
<td>0.0306</td>
<td>0.9693</td>
<td>0.9968</td>
<td>8.8 x 10^{-6}</td>
<td>0.9919</td>
<td>Yes</td>
</tr>
<tr>
<td>rs2420946 (FGFR2)</td>
<td>0.0022</td>
<td>0.0218</td>
<td>0.9570</td>
<td>0.9955</td>
<td>1.5 x 10^{-3}</td>
<td>0.9997</td>
<td>Yes</td>
</tr>
<tr>
<td>rs1219648 (FGFR2)</td>
<td>0.0021</td>
<td>0.0209</td>
<td>0.9552</td>
<td>0.9953</td>
<td>1.2 x 10^{-5}</td>
<td>0.9986</td>
<td>Yes</td>
</tr>
<tr>
<td>rs7696175</td>
<td>0.0013</td>
<td>0.0131</td>
<td>0.9298</td>
<td>0.9925</td>
<td>1.5 x 10^{-5}</td>
<td>0.9997</td>
<td>Yes</td>
</tr>
<tr>
<td>rs197275</td>
<td>0.0012</td>
<td>0.0123</td>
<td>0.9256</td>
<td>0.9920</td>
<td>Not Avl.</td>
<td>Not Avl.</td>
<td>No</td>
</tr>
<tr>
<td>rs12505800</td>
<td>0.0012</td>
<td>0.0123</td>
<td>0.9255</td>
<td>0.9920</td>
<td>8.1 x 10^{-6}</td>
<td>0.9881</td>
<td>No</td>
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<tr>
<td>rs210739</td>
<td>0.0011</td>
<td>0.0114</td>
<td>0.9204</td>
<td>0.9914</td>
<td>Not Avl.</td>
<td>Not Avl.</td>
<td>Yes</td>
</tr>
<tr>
<td>rs10779967</td>
<td>0.0011</td>
<td>0.0113</td>
<td>0.9194</td>
<td>0.9913</td>
<td>Not Avl.</td>
<td>Not Avl.</td>
<td>No</td>
</tr>
<tr>
<td>rs2981579 (FGFR2)</td>
<td>0.0008</td>
<td>0.0083</td>
<td>0.8933</td>
<td>0.9882</td>
<td>Not Avl.</td>
<td>Not Avl.</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Columns 2–5 show posterior probabilities for various priors, while Columns 6 and 7 shows p-values (obtained from [1]) and Šidák-corrected p-values.

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Wakefield [13] proposed a formula similar to Equation (5), but only considered the $U_{ID}$ and the $U_{IFD}$. That is, the utilities of a true discovery and of a true non-discovery were not factored into the decision.

**Discussion**

We identified four difficulties with many current methods for computing the posterior probability of a model analyzed using a GWAS data set. Most importantly, they only consider a null hypothesis $H_0$ and an alternative hypothesis $H_1$. So, they cannot handle a complex multi-locus hypothesis which has several competing hypotheses. Yet it is becoming increasingly commonplace to investigate multi-locus hypotheses. We developed the BNPP method which enables us to compute the posterior probability of such hypotheses, and which also attends to the other difficulties. We illustrated its effectiveness by applying it to both simulated and real data sets. We showed how the BNPP can be used to obtain a decision analytic solution as to when to report a finding.

The greatest difficulty in most Bayesian analyses is arguably the assessment of prior probabilities. The early rejection and now the slow acceptance of the Bayesian approach has been due in large part to the perceived arbitrary nature of these assessments. For example, in 1921 R.A. Fisher [44] stated that “The Bayesian approach depends upon an arbitrary assumption, so the whole method has been widely discredited.” However, the Bayesian approach does provide an elegant and general solution to the multiple hypothesis testing problem. Let $\pi_1$ denote the prior probability that the model is correct. Wakefield [13] points out that “as more genome-wide association studies are carried out lower bounds on $\pi_1 = 1 - \pi_0$ will be obtained from the confirmed ‘hits’ - it is a lower bound since clearly many non-null SNPs for which we have low power of detection will be missed.” We agree that in time results will help us to determine priors. However, one avenue of research that builds on the results presented here, would be to hasten this process by performing a comprehensive literature search to investigate current beliefs concerning agnostic priors. Supporting Information S1 proposes initial prior probabilities based on beliefs reported in two articles.

The BNPP was designed for the purpose of flagging SNPs for further investigation; that is, it is intended to compute the posterior probability of a model that was already discovered or conjectured. However, as mentioned at the beginning of the Methods section, the BNPP is also a promising technique for loci-disease association discovery. Indeed, in order to illustrate the effectiveness of the BNPP we showed results in which it was used for discovery. Future research can further investigate its discovery capability and compare its performance to other related discovery methods such as those appearing in [10,24,26–28].

An immediate plan we have for using the BNPP is the following. We will expand our previous work on discovery to develop a system that outputs likely one to five SNP models in the first stage. The BNPP will then be used in the second stage to compute the posterior probability of each model consisting of a subset of the SNPs from each of these models. The most probable models will be reported. This method will be compared to using the BNPP directly for discovery.

### Supporting Information

**Supporting Information S1 Assessing Prior Probabilities**

**Author Contributions**

Conceived and designed the experiments: XJ. Performed the experiments: XJ. Analyzed the data: XJ GFC MMB MJB. Wrote the paper: XJ. All authors read and approved the final manuscript.

**References**


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