FHSA-SED: Two-Locus Model Detection for Genome-Wide Association Study with Harmony Search Algorithm

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Abstract

Motivation

Two-locus model is a typical significant disease model to be identified in genome-wide association study (GWAS). Due to intensive computational burden and diversity of disease models, existing methods have drawbacks on low detection power, high computation cost, and preference for some types of disease models.

Method

In this study, two scoring functions (Bayesian network based K2-score and Gini-score) are used for characterizing two SNP locus as a candidate model, the two criteria are adopted simultaneously for improving identification power and tackling the preference problem to disease models. Harmony search algorithm (HSA) is improved for quickly finding the most likely candidate models among all two-locus models, in which a local search algorithm with two-dimensional tabu table is presented to avoid repeatedly evaluating some disease models that have strong marginal effect. Finally G-test statistic is used to further test the candidate models.

Results

We investigate our method named FHSA-SED on 82 simulated datasets and a real AMD dataset, and compare it with two typical methods (MACOED and CSE) which have been developed recently based on swarm intelligent search algorithm. The results of simulation experiments indicate that our method outperforms the two compared algorithms in terms of detection power, computation time, evaluation times, sensitivity (TPR), specificity (SPC), positive predictive value (PPV) and accuracy (ACC). Our method has identified two SNPs (rs3775652 and rs10511467) that may be also associated with disease in AMD dataset.

Introduction

With the advent of high-throughput sequencing technology, it is possible to measure all of single-nucleotide polymorphisms (SNPs) from thousands of individuals [1]. The genome wide
association studies (GWAS), that aim to detect the casual relationship between SNPs and disease status and explore multiple SNPs synergistic effect on diseases status in a population, play a very important role in identifying the causes of disease [2] [3] [4], which have successfully identified many SNP genetic markers associated with a wide range of diseases and quantitative traits [5] [6]. Around 30 schizophrenia associated loci have been identified through GWAS techniques [7–10]. However, it is also an enormous challenge in calculation capability to detect the casual relationship between multi-SNPs and disease status at a whole-genome scale due to the enormous computational burden imposed by a very high-dimensional search space: a brute force search method is infeasible to evaluate the entire multi-locus model in genome wide scale. For identifying multi-locus disease models, there have been a number of algorithms proposed to search the multi-locus models in recent years. These algorithms can be categorized as exhaustive combinatorial search, stochastic search, heuristic search and machine learning based technique [11–12].

The exhaustive search approach, in which all possible multi-locus SNP combinations are evaluated on the strength of their associations with disease states, is very simple and can be realized through a parallel mechanism for detecting SNPs combinations for non-genome-wide association study [13–16]. However, current calculation technique is usually limited, and it is infeasible to detect the multi-locus epistasis models using the exhaustive search algorithm for GWAS.

Heuristic search algorithm [17–20] is an approximate search algorithm, which can expedite the search process by reducing the search space. Stochastic search algorithm [21–22] works by using probabilistic methods to search the optimal solution. However, both heuristic search and stochastic search cannot ensure discovering the global optimal solution. Machine learning based technique [23–25] is also adopted widely in computational biology, which can be categorized as classification for difference analysis and regression analysis, which is usually combined with feature selection technique for selecting a group of features (such as SNPs, genes) that affect significantly the phenotypes or traits, but it can not determine the true causal relationship between genotype and phenotype.

In recent years, swarm intelligent optimization algorithms, inspired by natural phenomenon or biological system, have attracted considerable attention for genetic interactions [1, 4, 26–29]. For example, AntEpiSeeker [29] introduced a two-stage ant colony optimization (ACO) algorithm for the detection of epistatic model. M Aflakparast et al. (2014) [30] proposed a cuckoo search epistasis (CSE) algorithm which combined Bayesian scoring with cuckoo search (CS) algorithm [31] for detecting the multi-locus disease-causing models. Jing and Shen (2014) [4] proposed a Multi-objective Ant Colony Optimization algorithm for SNP Epistasis Detection (MACOED), in which both Bayesian network scoring and logistical regression scoring are combined as evaluation criterions for SNP interactivities. However, these methods have drawbacks on low detection power and high computation cost.

It is very important to develop or choose appropriate methods for identifying the multi-locus disease-causing models for genome-wide study. There has been remarkable activity in the development of methodology (e.g. Bayesian methods, regression-based methods, linkage disequilibrium (LD) and haplotype-based methods) [32] for the detection of epistasis in the past ten years. However, they perform inconsistently usually with different disease models [4] because they were conceived merely based on part of detective models of epistasis. Some multi-objective detection methods were proposed to improve the performance for detecting the multi-locus epistasis models, such as multi-filter enhanced genetic ensemble (MF-GE) system [23] and multi-objective ant colony optimization algorithm (MACOED). the MF-GE algorithm requires diverse and accurate classifiers to achieve better accuracy and requires configuring parameters properly for each classifier, which is a very large challenge for MF-GE method; MACOED simultaneously employs the Bayesian-based K2-score and regression-based AIC-
score as evaluation indexes in the filter stage, in which, however, the two-fold scoring method would increase the computation burden and make some models failed to pass the screening stage due to overly strict evaluation methods. Although the two-fold scoring method could decrease the false positive rate (Type I error) in MACOED, it is apparent that the false negative rate (Type II error) increases; in addition, the regression-based AIC-score methods require an iteration process to optimize the regression coefficients, which is often computationally unaffordable for SNP datasets with very large number of markers. To tackle these drawbacks (preference to some types of disease models, high computation cost), we propose a two stages (screening and testing) intelligent search algorithm named FHSA-SED (Harmony Search Algorithm with two scoring functions for SNP Epistasis Detection)” to detect two-locus disease models. To quickly identify various disease models, in the FHSA-SED algorithm, two evaluation criteria (Bayesian network based K2-score and Gini-score) are employed to enhance the ability for identifying various disease models, Harmony Search Algorithm (HSA) is improved to speed up the process of detecting disease models and a local search algorithm with two-dimensional tabu table is presented to avoid repeatedly evaluating (overcoming the premature convergence) some disease models which have strong main effects.

In this study, our central goal is to detect as various disease models as possible, and to enhance the power of identifying disease models by employing two complementary methods (K2-score and Gini-score). Our method is divided into two stages: in the 1st stage, we want to quickly obtain some most likely two-locus disease models (candidate solutions) using harmony search algorithm; in the 2nd stage, we adopt the G-test method to test the candidate solutions.

Some terms (Joint effect, Evaluation times, Computation time and two-locus disease model) are explained in Box 1.

Outline

A flow chart of our method is illustrated in Fig 1, in which the detection process of two-locus disease models in FHSA-SED algorithm is divided into two stages: “screening” and “testing”. In the screening stage, an improved harmony search algorithm (HSA) (Z.W. Geem, 2001) [35] is employed to search two-locus models that might be associated with phenotype, and two criteria (Bayesian network based K2-score and Gini-score) are respectively used to evaluate the causality between the two-locus models and phenotype. Some two-locus models with highest K2-score are stored in harmony memory HM1, and some models with highest Gini-score are stored in HM2. Next, HM1 and

Box 1

Terms:

1. **Joint effect (Synergy effect)** denotes k SNP locus act jointly to have a particular phenotypic effect, which includes additive effect, statistical interaction effect and so on.
2. **Evaluation times** represent the number that k-locus models are evaluated using Bayesian scoring criterion and Gini scoring criterion.
3. **Computation time** denotes the time spent executing algorithm in the program.
4. **two-locus disease model** is defined as by penetrance table, in which a two-way SNP genetic combination is referred to as collective association with the dichotomous phenotype (disease status) if the genotype distribution at the two SNPs is different significantly between cases and controls, and it may be responsible for significantly increasing the risks of complex diseases [33–34].
HM2 are merged into a union set HM (= HM1∪HM2) as shortlisted candidates. In the Testing stage, these shortlisted candidates are further checked using a G-test statistical method.

**Methods**

Let a set of SNP variables $X = \{x_1, x_2, \cdots, x_N\}$ indicate $N$ SNP markers for $L$ individuals (samples), $Y$ be the phenotype variable with values of $\{y_1, y_2, \cdots, y_J\}$; we represent the homozygous major
allele, heterozygous allele and homozygous minor allele as 0, 1 and 2, respectively. For a $k$-loci combination model, $I$ denotes the number of genotype combinations (there are $3^k$ SNP genotype combinations), $J$ is the number of phenotype states $Y$ (which is equal to 2 for a case-control dataset). $n_i$ is the number of cases in the dataset with SNP nodes taking the $i$-th genotype combination, $n_{ij}$ represents the number of cases that belong to the phenotype state $j$ where the $k$-way SNPs variables have $i$-th genotype combination.

Bayesian network scoring criterion and Gini index criterion

It is a vital factor to design new effective method for identifying the disease models successfully. Some existing methods [4] [32] usually prefer one type of disease models to others. To tackle the preference problem to various disease models, we employ two evaluation criteria (Bayesian network based K2-score and Gini-score) to improve the power for identifying various disease models.

**Bayesian network scoring criterion.** A Bayesian network (BN) is a kind of statistical model which represents a set of random variables and their conditional dependencies by using a directed acyclic graph (DAG). In the DAG, nodes denote random variables, and edges represent conditional dependences between two linked nodes.

There are more than 20 kinds of BN models [22] [36–37] that have been developed to find causal relationships, perform explanatory analysis, describe the causal influence and make predictions. In GWAS studies, BN model is also used to detect the interaction effect among SNP markers, which can represent the causal relationship between genetic variants and disease status.

In a DAG of Bayesian network for representing the relationships of SNP markers and disease states, there are only directed edges linking from the SNP markers to diseases status, and there is no edge connected from disease state to SNP markers and also no linkage among SNP markers. In the DAG, if and only if SNP $x_i$ is a direct cause of phenotype state $y_j$, there is a direct edge linking from node $x_i$ to phenotype $y_j$.

According the theorem 1 that is given in [38] (more detail interpretation about Bayesian network scoring method are introduced in S1 File), the K2-Score based on Bayesian network scoring criterion [37] can be described as Eq (1),

$$K2 - Score = \prod_{i=1}^{I} \left( \frac{(J-1)!}{(n_i + J-1)^{J}} \prod_{j=1}^{J} n_{ij}^{j} \right)$$

**Gini index criterion.** Gini index (Gini coefficient) is a measure of statistical dispersion (http://en.wikipedia.org/wiki/Gini_coefficient#cite_note-1) [39–42], which can be used to measure the impurity of a data partition or the inequality among values of a frequency distribution.

For a binary classification case-control problem, the Gini index is a diversity index [43] which is defined as Eq (2),

$$Gini - score = \sum_{i=1}^{I} P_i \cdot \left(1 - \sum_{j=1}^{J} p_{ij}^{2} \right)$$

where, $p_{ij}$ ($p_{ij} = \frac{n_{ij}}{n_i}$) is the estimated probability that the $i$-th genotype combination actually associated with phenotype $y_j$, $\left(1 - \sum_{j=1}^{J} p_{ij}^{2} \right)$ means the estimated probability that genotype combination is misclassified as phenotype $y_j$, $P_i$ ($P_i = \frac{n_i}{L}$) is the percentage of $i$-th genotype combination in sample set.
Proposed FHSA algorithm for two-locus disease model detection

Detecting multi-locus models at a whole-genome scale is a non-trivial task since it takes too much time to detect all models from hundreds of millions of SNPs. In this approach, we propose a fast harmony search algorithm (HSA) to accelerate the detection process of disease models, without an exhaustive search.

Standard HSA (see S2 File) is a meta-heuristic algorithm, which mimics the process of improvising a musical harmony. Compared with traditional mathematical optimization algorithms, HSA does not require substantial gradient information and is not dependent to initialization, making it widely applied in the fields of combinatorial optimization.

In the standard HSA, harmony memory is a set of harmonies. By evaluating their fitness with some criterion, some harmonies in the set are substituted with some other harmonies which are supposed to be with more fitness. Such a process continues until some finishing criterion is satisfied.

In the proposed FHSA-SED algorithm, each harmony denotes a \( k \)-way (\( k \)-locus) model that is a combination of \( k \) different SNP markers (\( k = 2 \) in this study) and we employ two harmony memories: HM1 and HM2. The harmony in HM1 is evaluated with Bayesian network scoring criterion, and the Gini scoring criterion is used to evaluate the harmony in HM2. [Fig B1 in S2 File presents the flow chart of fast harmony search algorithm (FHSA)].

The pseudo code of FHSA is as Algorithm-1.

Algorithm-1: harmony search algorithm for SNP Epistasis Detection with two scoring criterion (K2-Score and Gini-Score)

Input: maximum model evaluation times (MMEs) of SNP-pairs model, HS parameters: HMCR, PAR and HMS

Output: HM1, HM2, and fitness values of each harmony in HM1 and HM2

1. Initialize harmony memory HM1 and HM2 randomly.
   For I = 1:HMS
   HM1(I, 1:k) = (r1, r2, ..., rk); // \( r_j \in \{1, 2, ..., M\} \) (\( r_1 < r_2 < ... < r_k \)),
   HM2(I, 1:k) = (s1, s2, ..., sk); // \( s_j \in \{1, 2, ..., M\} \) (\( s_1 < s_2 < ... < s_k \))
   End

2. Calculate the fitness value of each harmony in HM1 using Bayesian network scoring function (\( f_1 \)), and the fitness value of each harmony in HM2 using Gini scoring function (\( f_2 \)), respectively.
   For I = 1:HMS
   Score1(I) = \( f_1 \) (HM1(I, 1:k));
   Score2(I) = \( f_2 \) (HM2(I, 1:k));
   End

3. Generate a new harmony \( H_{new} \) as follows:
   for i = 1:k
   if rand(0,1)<HMCR
   \( a = [\text{rand}(0,1) \times \text{HMS} \times 2]; \)
   if a<HMS
   \( H_{new}(i) = \text{HM1}(a, i); \)
   if rand(0,1)<PAR
   \( H_{new}(i) = H_{new}(i) + (\text{rand}(0,1)-0.5) \times |\text{HM1(idbest1,i)}-\text{HM1}(r, i)|; \)
   end
   else
   \( H_{new}(i) = H_{new}(i); \)
   if rand(0,1)<PAR
   \( H_{new}(i) = H_{new}(i) + (\text{rand}(0,1)-0.5) \times |\text{HM2(idbest2,i)}-\text{HM2}(r, i)|; \)
   end
   end
   else
   
(Please see the computational process of an example in Table A1 in S1 File.)
\[ H_{\text{new}}(i) = \lfloor \text{rand}(0,1) \times N \rfloor; \]
end
end
If \( H_{\text{new}} \) has been visited before
execute the local search algorithm in the neighborhood of \( H_{\text{new}} \);
end

4. **Calculate the fitness of \( H_{\text{new}} \) using scoring functions \( f_1 \) and \( f_2 \) respectively:**
\[
\text{score}_1 = f_1(H_{\text{new}}), \quad \text{score}_2 = f_2(H_{\text{new}});
\]

5. **Determine whether \( H_{\text{new}} \) can replace the worst harmony in \( \text{HM1} \) or \( \text{HM2} \):**
   if score1 is better than Score1(idworst1)
   \( \text{HM1}(\text{idworst1,:}) = H_{\text{new}}; \)
   end
   if score2 is better than Score2(idworst2)
   \( \text{HM2}(\text{idworst2,:}) = H_{\text{new}}; \)
   end

6. **If termination conditions meet, output \( \text{HM1} \) and \( \text{HM2} \), otherwise, turn to step 4.**

In algorithm1, \( r_1 \) and \( r_2 \) are random integer between 1 and HMS; idbest1/ idworst1 denotes the index of best/worst harmony in the \( \text{HM1} \); idbest2 and idworst2 denote the indexes of best and worst harmonies in the \( \text{HM2} \), respectively.

**Local search algorithm for FHSA.** As a heuristic search algorithm, HSA is also easy to trap into a local search and repeatedly evaluate some solution (sampling with repetition) in solving the combinational optimization problems, which causes time-consuming due to these repeated calculation (repeated sampling). To tackle this problem, we establish a tabu table (TT) to store the evaluation state of each SNP-pair (If a SNP-pair has not been evaluated, its evaluation state equals ‘0’, otherwise, it is ‘1.’) and a local search algorithm is proposed to discover new disease models that have not been evaluated. The TT is different from frequently-used linear tabu list, which is a two-dimensional table for marking the state of each two-locus model, if a two-locus model has been evaluated, its corresponding value on TT is set to “1”; otherwise it is equal to “0”. The advantage of the two-dimensional tabu table compared to linear tabu list is that it can get the evaluation state of each two-locus model using one times search (time complexity is \( O(1) \)); however, linear tabu list requires a sequential search whose time complexity is \( O(n) \).

Local search algorithm is used to obtain a closest solution (that has not been visited) in the neighborhood of current solution, for example Fig 2, if a new generated solution \( H_{\text{new}} = (X_7,X_3) \) has been visited, then one of the nearest solutions that have not been evaluated will replace it as new solution \( H_{\text{new}} = (X_8,X_2) \) to be evaluated. The local search algorithm has two advantages: First, it can avoid evaluating the same one two-locus model twice; second, it can achieve the same performance as exhaustive search if we set the maximum model evaluation times (MMEs) equal to \( \binom{N}{k} \). Therefore, the proposed FHSA algorithm is a global search algorithm for detecting two-locus disease model.

However, when most of the elements of TT have been marked, the efficiency of local search algorithm will decrease because most of solutions nearby current solution \( H_{\text{new}} \) have been evaluated, which will increase search times for near solutions. Thus we transform the two-dimensional TT into a link Table. For each element \( TT_{ij} \) in two-dimensional Table can be denoted with a link table element \( Y(k) \). The transformational formula is expressed as Eq (3). The Fig 2 can be transformed as Fig 3.

\[
Y\left( N \times (i-1) - \frac{i(i-1)}{2} + j - i \right) \leftarrow TT_{ij}\]

(3)

where, \( N \) is the number of SNP.
Fig 2. Local search algorithm based on two-dimensional tabu table (TT). In Fig 2, $X_i$ is the $i^{th}$ SNP locus, ‘1’ denote the two-locus model has been evaluated. ‘0’ is otherwise.

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Fig 3. Doubly linked Table as tabu table (TT). All adjacent elements are linked each other. (b) When an element is just evaluated, one of near solutions of the element is selected with a random step for evaluating in the next time.

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In Fig 3, we establish a doubly linked list for storing Y, in which adjacent elements in row major order are linked with doubly links at first (see Fig 3(A)). When one element (solution) has been evaluated, it will be removed from the doubly linked list. Meantime, in the doubly linked list, one of near elements of the solution will be chosen with a random step BW, which is evaluated in the next time (see Fig 3(B)). The BW is changed dynamically with the increasing of iterations, which is expressed as Eq (4). It can be seen from Eq (4) that the BW is between 1 and 10. In the beginning stage, there are most of the elements that have not been evaluated, thus at this time it has a large probability that BW possesses a large value. Conversely, in the later stage, it has a large rate that BW possesses a small value.

\[
BW = \min(10, \max(rand(0, 1) \times \#E, 1))
\]

where, #E denotes the number of elements having not been evaluated in doubly linked list.

**G-test**

G-test is a likelihood-ratio test that is being progressively applied in different significance tests (http://en.wikipedia.org/wiki/G-test#cite_note-2) [44]. The G-test and chi-squared (χ²) test will lead to the same conclusions for a reasonable sample size with the Person chi-squared tests. However, the Pearson χ² test is inferior to the approximation to the theoretical chi-squared distribution for the G-test [42]. And for testing goodness-of-fit, G-test statistical method is more efficient than Pearson χ² test method [45–47].

The general formula for the value of G is as follows

\[
G = 2 \sum_{i=1}^{O} O_i \cdot \ln \frac{O_i}{E_i}
\]

where, \(O_i\) is the observed frequency, \(E_i\) is the expected frequency under the null hypothesis, \(\ln\) denotes the natural logarithmic function.

For \(k\)-loci model detection, an \(I \times J\) contingency table requires be adopted for calculating the G value with the follow formula

\[
G = 2 \sum_{i=1}^{I} \sum_{j=1}^{J} O_{ij} \cdot \ln \frac{O_{ij}}{E_{ij}}
\]

where, \(O_{ij}\) and \(E_{ij}\) are respectively the observed numbers and expected number of genotypes when phenotype takes the state \(y_j\) and genotypes take \(i\)-th \(k\)-combination. We can get the observed number \(O_{ij}\) from dataset by using simple counting statistics method. The expected number \(E_{ij}\) of genotype frequency could be obtained according to Hardy-Weinberg principle [48].

The null hypothesis is that the \(k\)-combination of SNP set has no association with the phenotype. If the \(P\)-value of the G-test statistic is smaller than a significance level \(\alpha_0\), the alternative hypothesis is accepted, which means the \(k\)-combination of SNPs has a certain association with phenotype. In order to control false positive rate (Type I error rate), we adopt Bonferroni-corrected significance level \(\alpha = \frac{\alpha_0}{N/k}\) to deal with multiple testing. Because sometimes the number of some genotypes equals zero or very small (less than \(e\), \(e\) is a small integer) we do a
minor modification for calculating G-test value as follows,

\[ G = \sum_{i=1}^{I} \sum_{j=1}^{J} O_{ij} \cdot P_{ij} \]

\[ P_{ij} = \begin{cases} 
\ln \frac{O_{ij}}{E_{ij}}, & O_{ij} > 0 \\
0, & \text{otherwise} 
\end{cases} \]

The degree of freedom \(d\) (\(d = (I-1)(J-1)\)) is also modified as follow:

For each \(i\) (\(i = 1, 2, \cdots, I\))

If \(\sum_{j=1}^{J} O_{ij} < \epsilon\), then the degree of freedom \(d = d-1\).

End

Experiments and Results

Parameters and Environments Setting

To investigate of FHSA-SED algorithm, we evaluated its performance using 82 simulation datasets with different type of disease models and compared its performance with two excellent intelligent optimization algorithms (MACOED, CSE). The MACOED and CSE algorithm have advantages over AntEpiSeeker, BEAM and BOOST on the detection of multi-locus disease models in terms of power, sensitivity (true positive rate: TPR) or specificity (SPC) (true negative rate: TNR). The Matlab source codes of MACOED[4] and CSE[30] algorithms can be downloaded from http://www.csbio.sjtu.edu.cn/bioinf/MACOED/ and http://lbb.ut.ac.ir/Download/LBBsoft/CSE separately, in which we made some minor revisions on the source codes of two methods in order to perform a fair comparison; the main body of the source codes was unchanged.

In the experiments, parameters setting for the compared algorithms are shown in Table 1. To make a fair comparison, we set the same termination condition and the same runtime environment for three compared algorithms, where maximum model evaluation times (MMEs) is less than the number by using exhaustive search algorithm. All the experiments were performed on Windows XP 64 system with Intel(R) Xeon(R) CPU E5504 @2.0GHz, 8 GB memory, and all the program codes were written in MATLAB R2014b (the source code of FHSA-SED is in S5 File).

Performance evaluation criteria

In order to investigate the performance of the FHSA-SED algorithm comprehensively on detecting two-locus disease models which is associated with disease states, we adopt seven

| Algorithms | Parameters |
|------------|------------|
| **FHSA-SED** | HMCR=0.9; PAR=0.35; ||HM1||=100; ||HM2||=100; P-value=0.01/Ck; || || denotes the size of set; MMEs = 4500 for 100SNP markers; MMEs = 300000 for 1000SNP markers |
| **MACOED** | \(\tau_0 = 1; \ T_0 = 0.8; \ \beta = 0.9; \ \lambda = 2; \ \text{Ant number} = 100; \ \text{P-value} = 0.01/C_k; \ MMEs = 4500 \ \text{for 100SNP markers}; \ MMEs = 300000 \ \text{for 1000SNP markers} |
| **CSE** | \(\text{MaxLe}vy\text{StepSize} = 1; \ \text{Number of SNPs in each Group is 5}; \ \text{Fraction of eggs discarded each generation is equal to 0.25}; \ \text{The number of nest equals 30}; \ MMEs = 4500 \ \text{for 100SNP markers}; \ MMEs = 300000 \ \text{for 1000SNP markers} |

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metrics: power, evaluation times, computation time, sensitivity (true positive rate: TPR), specificity (SPC) (true negative rate: TNR), Positive predictive value (PPV) and Accuracy (ACC).

1) Maximum Model evaluation times (MMEs): in the experiment, we set Maximum Model evaluation times (MMEs) of SNP combinations as the terminal condition of algorithm, in other words, the harmony search algorithm will be terminated if the current evaluation times of two-locus models have been larger than MMEs. If the known disease-causing models have been found, the searching algorithm would be terminated early, the number that two-locus models have been evaluated currently is defined as Model evaluation times (MEs) and the elapsed time from start to end is denoted as computation time.

2) The TPR, SPC, PPV and ACC are defined as follows

\[
\begin{align*}
\text{TPR} & = \frac{TP}{TP + FN} \\
\text{SPC} & = \frac{TN}{FP + TN} \\
\text{PPV} & = \frac{TP}{TP + FP} \\
\text{ACC} & = \frac{(TP + TN)}{(TP + TN + FN + FP)}
\end{align*}
\]

where TP, FP, TN and FN denote the number of true positives, number of false positives, number of true negatives and number of false negatives, respectively.

The TPR, SPC, PPV and ACC in this study are employed to measure the statistical precision of hypothesis testing method for having found disease-models in the screening stage. The TP is equal to the number of disease-models that have passed threshold of testing method, FN is the number of disease-models failed to pass the threshold of testing method. FP is the number of non-disease-models passed the threshold, TN equals the number of non-disease-models failed to pass the threshold.

3) The power is defined as follow

\[
\text{power} = \frac{\#(S)}{\#T}
\]

where \#T denotes the number of datasets that are generated by the same model parameters (\#T = 100 in our experiment), \#(S) is the number of datasets in which the true disease-causing models are found and passed the corresponding evaluation criteria among all \#T datasets. The power of screening stage (1st power) denotes the rate that the true disease models have been put into the candidate set in 1st stage. The power of testing stage (2nd power) is the rate that the true disease models have been passed the significant threshold of G-test, which is equal to TP/\#T.

Simulation datasets

Model-based data. We perform experiments on 82 simulated data sets to investigate the performance of FHSA-SED algorithm. These data sets are divided into two categories: disease loci with main effects (DME 1- DME 12) and disease loci without main effects (DNME 1 – DNME 70).

1) Simulation 1 (disease loci with main effects: DME).

The DME model has both main effects and interaction effects. Twelve disease models (Model 1-Model 12) [4], which are composed of multiplicative model, threshold model and concrete model, are adopted in Simulation 1.

DME 1- DME 4 (H^2 = 0.005, MAF = 0.05, 0.1, 0.2 and 0.5) are multiplicative models with two disease locus, in which the disease prevalence given the frequency of genotype combination...
increases multiplicatively with the incremental presence of the disease. The genetic heritability ($H^2$) of DME 1-4 are all equal to 0.005, minor allele frequencies (MAF) of them equal 0.05, 0.1, 0.2 and 0.5, respectively.

It is very difficult to identify the disease locus from the four DME models due to having very low genetic heritability. The fitness landscape of DME 1 is shown in [Fig E1 in S3 File], in which the fitness value of disease-causing SNP-pair is more or less similar to those of some non-pathogenic SNP-pairs. As seen from [Fig E1 in S3 File] that the disease-causing SNP-pair (10, 80) has not very significant difference with some other two-locus models, which makes the search algorithm easy to be deviated from correct direction and leads to the miss of the disease-causing two-locus model.

DME 5-8 ($H^2 = 0.02$, MAF = 0.05, 0.1, 0.2 and 0.5) are the threshold models in which the prevalence of genotype frequency does not increase until the number of disease alleles pass the threshold. [Fig E2 in S3 File] is the fitness landscape of DME 8, which has strong marginal effect and interaction effect. From [Fig E2 in S3 File], a SNP marker with strong marginal effect (e.g. SNP marker 10 and SNP marker 80) would form many false disease models with other SNP markers that are not truly associated with the phenotype state.

DME 9-12 ($H^2 = 0.02$, MAF = 0.05, 0.1, 0.2 and 0.5) are the concrete model [49]. [Fig E3 in S3 File] is the fitness landscape of DME 12, which shows the model with low marginal effect and strong interaction effect. It can be seen from [Fig E3 in S3 File] that a SNP-pair with very weak marginal effect is just like an isolated point.

In Simulation 1, the parameters and the values of penetrance of 12 models are given in [Table E-1 in S3 File]. The corresponding data sets are generated using the software GAMETES_2.0 [50]. The disease loci of all generated datasets with GAMETES_2.0 are on the last two SNP markers. In order to avoid position preference for an optimization algorithm, we exchange the places of disease locus to other positions randomly. In each data set, 100 SNPs and 1000 SNPs are respectively simulated.

(2) Simulation 2 (disease loci with no main effects: DNME)

The DNME model only has the interaction effects without the marginal effects. We adopt 70 epistatic models which have different genetic heritability $H^2$ (0.01, 0.025, 0.05, 0.1, 0.2, 0.3 and 0.4), MAF (0.2 and 0.4) and different penetrance values. The data corresponding to the 70 models was downloaded from http://discovery.dartmouth.edu/epistatic_data [51]. These data sets have 1000 attributes, the first two being functional, the remainder randomly generated. [Fig E4 in S3 File] is the fitness landscape of a DNME model (MAF = 0.4, $H^2 = 0.025$). It can be seen from [Fig E4 in S3 File], the disease-causing SNP-pair is almost an isolated point without any associated neighborhood that would make the heuristic search algorithm difficultly to find the veritable disease model. The penetrance Tables of 70 DNME models are provided in [Table E-2 in S3 File].

Results comparison and analysis on model-based data. In Simulation 1, we compare FHSA-SED algorithm with MACOED and CSE.

Figs 4–7 present the power, evaluation times and computation time of three algorithms on 12 DME models for the datasets which have 100 SNP markers and quantitative comparisons are also presented in Table 2. In order to further evaluate the performance of FHSA-SED algorithm, we compared four performance metrics (TPR, SPC, PPV and ACC) of FHSA-SED and MACOED algorithms on the DME models. Our results are presented in Fig 8 and Table 3.

In Fig 4 and Table 2, the powers of HS+ (K2-Score), HS+ (Gini-Score) and 1st FHSA-SED are respectively equal to $\frac{S^1_{1st}}{#T}$, $\frac{S^2_{1st}}{#T}$ and $\frac{S^2_{1st}}{#T}$, where $#T$ denotes the number of datasets that are generated by the same parameters ($#T = 100$ in our experiment), $S^1_{1st}$, $S^2_{1st}$ and $S^3_{1st}$ denote the
numbers of the true two-locus disease models (in #T simulate datasets) having been put into HM1, HM2 and HM, respectively. Likewise, 1st MACOED is the union power of ACO + (K2-Score and ACI-Score) in the screening stage. The powers of FHSA-SED and MACOED are the rate that the true disease models have been passed the significant threshold $P$-value of G-test and Chi-square test respectively.

It is indicated from Fig 4 and Table 2 that the FHSA-SED algorithm outperforms MACOED and CSE methods on all 12 DME models, in which the HS algorithm with K2 Scoring criterion (HS+(K2-Score) has a higher power than HS+(Gini-Score) on DME 1–2, however, HS+(Gini-Score) is more powerful on DME 3 and DME 4 than HS+(K2-Score) algorithm. This illustrates that the two scoring criterions in 1st FHSA-SED can complement each other. We can found from column 4 (1st FHSA-SED) and column 5 (FHSA-SED) in Table 2 that the power of FHSA-SED, for DME 1 ~ DME 4, is lower than that of 1st FHSA-SED because part of short-listed candidates of 1st FHSA-SED failed to pass the significant threshold of G-test, resulting in type II errors. Which because, for DME 1 ~ DME 4, there are very small significant difference

Fig 4. The power comparison on three DME models: (a) The left figure is the multiplicative model ($H^2 = 0.005$); (b) The middle figure is threshold model ($H^2 = 0.02$); (c) The right figure is the concrete model ($H^2 = 0.02$).

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Fig 5. The evaluation times on DME1 ~ DME 12 for three algorithms: FHSA-SED, MACOED and CSE. (1) The left figure illustrate the evaluation time of three algorithm on DME 1~DME 4 ($H^2 = 0.005$, $P(D) = 0.1$).(2) The middle figure presents the evaluation time of three algorithm on DME 5~DME 8 ($H^2 = 0.02$, $P(D) = 0.1$).(3) The right figure presents the evaluation time of three algorithm on DME 9~DME 12 ($H^2 = 0.02$, $P(D) = 0.1$).

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between case data and control data. If the significant threshold of G-test is relaxed, some false disease models might pass the significant threshold for DME 7 ~ DME 12 (type I errors), which error is generally even less acceptable. Nevertheless, the shortlisted candidates in 1st FHSA-SED that have failed to pass the significant threshold of G-test are worth studying further by employing or developing effective approaches.

As is illustrated in Fig 5, Fig 6, Fig 7 and Table 2, the evaluation times and the computation time of our method are significantly less than other two methods. For three type of DME models (multiplicative model: DME 1–4, threshold model: DME 5–8 and concrete model: DME 9–12), the mean evaluation times of our method are less than 1500, 300 and 600, respectively, and the mean computation time is less than 2s, 0.6s and 1s.

Fig 6. The computation time on DME1-DME 12 for three algorithms: FHSA-SED, MACOED and CSE. (1) The left figure illustrate the computation time (s) of three algorithm on DME 1–DME 4 (H² = 0.005, P(D) = 0.1). (2) The middle figure presents the computation time (s) of three algorithm on DME 5–DME 8 (H² = 0.02, P(D) = 0.1). (3) The right figure presents the computation time (s) of three algorithm on DME 9–DME 12 (H² = 0.02, P(D) = 0.1).

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Fig 7. The statistical box plots of FHSA-SED algorithm. Illustrating the distribution of evaluation times and computation time (s). (1) The left figure illustrates the statistical distribution of evaluation times for 12 DME models for 100*5 datasets (100 datasets for each model, and FHSA-SED runs 5 times repeatedly for each data set). (2) The right figure illustrates the corresponding statistical distribution of computation times.

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A detailed view of Table 3 shows that the TPR of FHSA-SED on most of DME models is larger than that of MACOED. Yet the TPR on DME-4 is relatively smaller than that of MACOED, which is because some SNP-pairs that have been obtained in the 1st FHSA-SED are rejected in testing stage (see the Table 2, the powers of 1st FHSA-SED on DME 1~DME 4 are equal to 93%, 89%, 93% and 100%, which are much larger than powers of 1st MACOED), which makes the false negative rate a little high because the significantly difference (in multiplicative models: DME 1~ 4) between case data and control data is not very obvious. However, on DME 5 ~ DME 12, the TPR, SPC, PPV and ACC of FHSA-SED algorithm are all better than or not significant differences with those of MACOED.

As can be noticed in Table 3, for some models, the values of TPR are larger than or equal to the values of power, which because some disease models have not been found in the 1st screening stage. For example, if in the 1st FHSA-SED, 55 true disease models have been found from 100 datasets (1st power = 55%), and 2 models among these 55 disease models failed to pass the

![Figure 8](https://example.com/figure8.png)

**Fig 8. The performance (TPR, SPC, PPV and ACC) on DME1 -DME 12 for FHSA-SED and MACOED algorithms.** TPR, SPC, PPV and ACC, which are all multiplied by the corresponding power of each algorithm for 12 DME models, are shown in four sub figures in Fig 8.

Table 2. Powers, evaluation times and computation time for FHSA-SED, MACOED and CSE algorithms (100 SNP markers).

| Model | HS + (K2-Score) | HS + (Gini-Score) | 1st FHSA-SED | 1st MACOED | FHSA-SED | MACOED | CSE | FHSA-SED | MACOED | CSE |
|-------|----------------|-------------------|--------------|------------|----------|--------|-----|----------|--------|-----|
| DME-1 | 93%            | 68%               | 93%          | 57%        | 53%      | 0%     | 16% | 1475.00  | 2618.33 | 4389.9 | 1.82 | 38.64 | 26.65 |
| DME-2 | 89%            | 85%               | 89%          | 32%        | 61%      | 2%     | 18% | 1408.72  | 2560.04 | 4290.9 | 1.74 | 36.31 | 24.90 |
| DME-3 | 85%            | 93%               | 93%          | 50%        | 61%      | 13%    | 18% | 1149.08  | 2448.03 | 4367.4 | 1.41 | 35.73 | 25.18 |
| DME-4 | 83%            | 100%              | 100%         | 44%        | 62%      | 23%    | 22% | 460.65   | 2495.70 | 4200.0 | 0.58 | 35.60 | 24.04 |
| DME-5 | 100%           | 100%              | 100%         | 99%        | 92%      | 53%    | 20% | 495.77   | 2446.50 | 4232.0 | 0.61 | 36.80 | 24.30 |
| DME-6 | 100%           | 100%              | 100%         | 98%        | 93%      | 91%    | 21% | 303.11   | 2610.45 | 4245.9 | 0.37 | 39.48 | 23.14 |
| DME-7 | 100%           | 100%              | 100%         | 100%       | 90%      | 90%    | 23% | 248.43   | 2377.67 | 4233.3 | 0.30 | 35.82 | 24.11 |
| DME-8 | 100%           | 100%              | 100%         | 100%       | 85%      | 85%    | 30% | 232.16   | 2731.15 | 4114.5 | 0.29 | 41.78 | 24.80 |
| DME-9 | 100%           | 100%              | 100%         | 100%       | 93%      | 93%    | 16% | 241.46   | 2548.91 | 4362.0 | 0.30 | 40.20 | 25.91 |
| DME-10| 100%           | 100%              | 100%         | 100%       | 91%      | 91%    | 23% | 247.45   | 2519.15 | 4321.5 | 0.30 | 35.77 | 26.22 |
| DME-11| 100%           | 100%              | 100%         | 100%       | 91%      | 91%    | 28% | 693.27   | 2391.45 | 4101.9 | 0.85 | 34.40 | 25.32 |
| DME-12| 100%           | 100%              | 100%         | 100%       | 91%      | 96%    | 19% | 295.48   | 2689.01 | 4287.3 | 0.42 | 40.61 | 25.78 |

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threshold of G-test in 2nd FHSA-SED (TP = 53, FN = 2), then TPR = TP / (TP + FN) = 96%, power = 53%, and TPR > power.

As also can be found in Table 3, the CSE algorithm has not been contained, which because the goal of CSE is to find the disease models using Cuckoo search algorithm and Bayesian evaluation criterion. For each simulation dataset, the output of CSE is Yes (if the only disease-causing has been found) or No (if the disease-model has not been found), and CSE does not contain any statistical analysis for the output results. Therefore, to be fair, the TPR, SPC, PPV and ACC are not included in CSE algorithm.

In order to make a fair comparison, we multiply TPR, SPC, PPV and ACC by corresponding power of each algorithm for each model. Results are presented in Fig 8, indicating that our method is more effective than MACOED on all 12 DME models.

We also perform our algorithm on 12 DME models for datasets which have 1000 SNP markers; Fig 9 and Fig 10 present the power, evaluation times and computation time of three algorithms on 12 DME models and quantitative comparisons are presented in Table 4 and Table 5.

As shown in Fig 9, Fig 10 and Table 4, the FHSA-SED has much higher power than MACOED and CSE for most of DME models, and the evaluation times and runtime of FHSA-SED are also far less than those of MACOED and CSE. We can found from Table 5 that,

Table 3. The performance (TPR, SPC, PPV, FDR, ACC) comparisons for FHSA-SED and MACOED (100 SNP markers).

| Model   | FHSA-SED | MACOED |
|---------|----------|--------|
|         | power    | TPR    | SPC    | PPV    | ACC    | power    | TPR    | SPC    | PPV    | ACC    |
| DME-1   | 57%      | 51.16%  | 99.28% | 98.61% | 75.22% | 0%       | 0.00%  | 100.00%| 0.00%  | 81.56% |
| DME-2   | 32%      | 20.27%  | 98.76% | 94.23% | 59.51% | 2%       | 4.55%  | 100.00%| 100.00%| 85.00% |
| DME-3   | 50%      | 41.34%  | 98.63% | 96.80% | 69.99% | 13%      | 33.33% | 99.20% | 85.71% | 90.91% |
| DME-4   | 44%      | 29.00%  | 98.12% | 93.91% | 63.56% | 23%      | 68.75% | 94.98% | 62.86% | 92.10% |
| DME-5   | 99%      | 98.00%  | 98.87% | 98.86% | 98.43% | 53%      | 57.78% | 100.00%| 100.00%| 81.82% |
| DME-6   | 98%      | 98.00%  | 95.75% | 95.84% | 96.87% | 91%      | 96.81% | 95.50% | 94.79% | 96.10% |
| DME-7   | 100%     | 100.00% | 88.73% | 89.87% | 94.36% | 90%      | 100.00%| 90.99% | 89.81% | 94.71% |
| DME-8   | 100%     | 100.00% | 78.72% | 82.46% | 89.36% | 85%      | 98.94% | 83.78% | 83.78% | 90.73% |
| DME-9   | 100%     | 100.00% | 84.42% | 86.52% | 92.21% | 93%      | 97.83% | 81.74% | 81.08% | 88.89% |
| DME-10  | 100%     | 100.00% | 87.63% | 88.99% | 93.81% | 91%      | 97.85% | 79.83% | 79.13% | 87.74% |
| DME-11  | 100%     | 100.00% | 98.63% | 98.65% | 99.32% | 91%      | 97.83% | 100.00%| 100.00%| 99.06% |
| DME-12  | 100%     | 100.00% | 95.36% | 95.57% | 97.68% | 96%      | 98.94% | 88.29% | 87.74% | 93.17% |

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Fig 9. The power comparison on 12 DME models with 1000 SNP markers.

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for some models (e.g. DME 5 and DME 6), the TPR of MACOED is higher than that of FHSA-SED, which because only part of disease-models with significant difference between case and control have been discovered in 1st stage of MACOED, but many other unobvious disease-models that have low significant difference between case data and control data have not been found. However, if we think about the value of power, the power×TPR in FHSA-SED is much higher than that in MACOED, which illustrates that the FHSA-SED is more powerful in detecting various disease-models than MACOED.

In Simulation 2, performance comparisons on 70 DNME models are performed. [Fig E5-E6 in S3 File] display the powers of three algorithms, Fig 11 and [Table E-3 in S3 File] present the evaluation times and computation time for three algorithms when the number of SNP markers equals 100. Other four performance metrics (TPR, SPC, PPV and ACC) are also shown in [Table E-4 in S3 File].

From Fig E5 and [Fig E6 in S3 File], we can find that the power of 1st FHSA-SED is higher than CSE for all 70 DNME models, and its power is better than that of 1st MACOED for most of DNME models. FHSA-SED method has distinct advantages over MACOED and CSE.

### Table 4. Powers, evaluation times and computation time for FHSA-SED, MACOED and CSE algorithms (1000 SNP markers).

| Model | power | Mean evaluation Times | computation time (s) |
|-------|-------|------------------------|----------------------|
|       | HS+(K2-Score) | HS+(Gini-Score) | 1st FHSA-SED | 1st MACOED | CSE | FHSA-SED | MACOED | CSE | FHSA-SED | MACOED | CSE |
| DME-1 | 37% | 25% | 37% | 13% | 20% | 0% | 0% | 67088.1 | 159627 | 300000 | 137.3 | 1598 | 995 |
| DME-2 | 42% | 39% | 43% | 0% | 0% | 0% | 11% | 66366.9 | 294749 | 288150 | 135.8 | 2744 | 1360 |
| DME-3 | 56% | 66% | 66% | 21% | 20% | 0% | 13% | 43360.8 | 146310 | 282450 | 93.7 | 1460 | 1427 |
| DME-4 | 56% | 82% | 82% | 4% | 20% | 0% | 9% | 23487.6 | 190546 | 298100 | 50.1 | 1917 | 1477 |
| DME-5 | 98% | 98% | 98% | 77% | 50% | 10% | 15% | 24765.8 | 205019 | 290100 | 37.2 | 2055 | 1459 |
| DME-6 | 99% | 99% | 99% | 98% | 71% | 55% | 47% | 9811.6 | 182828 | 231100 | 15.3 | 1848 | 761 |
| DME-7 | 100% | 100% | 100% | 100% | 72% | 72% | 43% | 2122.4 | 191243 | 231900 | 3.1 | 2184 | 764 |
| DME-8 | 100% | 100% | 100% | 100% | 81% | 81% | 27% | 2171.6 | 185136 | 267450 | 3.1 | 1850 | 902 |
| DME-9 | 100% | 100% | 100% | 100% | 63% | 63% | 18% | 2366.8 | 208807 | 277950 | 3.7 | 1850 | 956 |
| DME-10 | 100% | 100% | 100% | 100% | 89% | 89% | 22% | 2559.5 | 208357 | 279150 | 3.7 | 1850 | 1135 |
| DME-11 | 100% | 100% | 100% | 100% | 51% | 51% | 18% | 3602.1 | 231101 | 277350 | 5.2 | 2243 | 1324 |
| DME-12 | 100% | 100% | 100% | 100% | 63% | 63% | 21% | 57318.2 | 238678 | 238350 | 89.9 | 2383 | 1242 |

For some models (e.g. DME 5 and DME 6), the TPR of MACOED is higher than that of FHSA-SED, which because only part of disease-models with significant difference between case and control have been discovered in 1st stage of MACOED, but many other unobvious disease-models that have low significant difference between case data and control data have not been found. However, if we think about the value of power, the power×TPR in FHSA-SED is much higher than that in MACOED, which illustrates that the FHSA-SED is more powerful in detecting various disease-models than MACOED.
algorithm on power for DNME-1–DNME-5 and DNME-36–DNME-40 (the genetic heritability $H^2 = 0.01$). For other DNME models, the FHSA-SED and MACOED have nearly equal powers.

In addition, Fig 11 indicates that the FHSA-SED algorithm takes very less computation time than MACOED and CSE. MACOED spends most time among three algorithms, which is almost 10 times what FHSA-SED spends. It is indicated from Fig 11 that FHSA-SED requires slightly less evaluation times than MACOED for DNME models.
As shown in Table E-4 in S3 File, the FHSA-SED algorithm has a close performance to the MACOED algorithm for most of DNME models. However, for DNME-2, DNME-36–DNME-40, the TPR of FHSA-SED is lower than that of MACOED, which is because 1st FHSA-SED has much higher power than 1st MACOED (See Table E-3 in S3 File), and small significant threshold value (P-value = 0.01/4950) make some candidate solutions prone to obstructed pass the threshold of G-test in the testing stage, which means some true candidate solutions fail to pass the G-test due to small significant threshold (P-value) although they have successfully passed the screening in 1st FHSA-SED (these candidate solutions maybe filtered out in 1st MACOED). This illustrates that candidate solutions in 1st FHSA-SED are very worth studying further, and which are called for a good testing method that embrace the complex disease models in future work.

In order to investigate the performance of FHSA-SED on solving DNME models, we test it using 70 DNME models which have 1000 SNP markers. The results are illustrated in S3 File (Fig E7 ~ Fig E8 and Table E-5).

Experiments on AMD real data

According to the previous analysis for simulation experiments, the proposed algorithm has a good performance on 70 simulation models. In this section, we conduct experiments on a real data set (AMD: Age-related macular degeneration) [52] using our proposed algorithm. The AMD dataset contains 103611 SNPs genotyped for 50 controls and 96 cases. Our goal of the experiment is to find quickly the disease-causing two SNP loci in AMD dataset using FHSA-SED algorithm.

Firstly, SNP loci with p-values from G-test less than 0.3 are removed from AMD dataset. Subsequently, 31341 SNP loci remain in the AMD dataset.

The setting of parameters for FHSA-SED algorithm is as follows:

- $\|H_M1\| = 500$; $\|H_M2\| = 500$;
- maximum evaluation times for SNP-pairs is equal to $3E+6$;
- The p-value threshold for SNP-pairs equals $\frac{0.05}{\binom{31341}{2}}$;
- Other setting of parameters is the same as those of Table 1.

The experiment took 4 hours approximately. There are 638 SNP-pairs (See S4 File) survived in the final output set.

All these 638 SNP-pairs are displayed in Fig 12, and the corresponding gene-pairs (mapped from SNP) are presented in Fig 13. It can be seen evidently from Fig 12 that three SNPs 'rs380390', 'rs1329428' and 'rs10272438' are associated with more other SNPs. In Fig 13, CFH, NA and BBS9 are linked with more other genes, where NA is not a gene, which means many SNPs are not in a gene region.

Similar to literatures [53–54], we select 26 top SNPs whose frequency is larger than 5 in the 638 SNP-pairs. In Table 6, the top two highest frequency SNPs ('rs380390' and 'rs1329428') which are all in an intron gene CFH, have been widely believed to be significantly associated with AMD [54–55]. Eight high frequency SNPs ranked from third to tenth may be also genetic factor contributing to the underlying mechanism of AMD. To our knowledge, 'rs10272438', 'rs1740752', 'rs1394608', 'rs1363688', 'rs7006908' and 'rs10492272' have been reported before; however, 'rs3775652' and 'rs10511467' have not been reported before, which need further to be studied and confirmed whether these SNPs are truly associated with AMD by developing a more efficient test method or using large scale samples.
In Table 7, top-20 SNP-pairs are presented in terms of P-value of G-test. It is noted that there are 15 SNP-pairs associated with three SNPs: 'rs380390', 'rs1329428' and 'rs10272438', other five SNP-pairs are associated with two unreported SNPs 'rs3775652' and 'rs10511467'.

**Discussion**

**Relationship between FHSA-SED and MACOED**

In this study, we proposed a HS algorithm using Screening and Testing to identify the SNP-pair disease models among all SNP-pairs, which has nearly the same algorithmic framework as MACOED. The key differences lie between FHSA-SED and MACOED:

1. MACOED employs two scoring functions (Bayesian network-based K2-score and logical regression-based AIC-score) to screen the disease models in the first stage, in which logic "and" operation is carried out between the two scoring functions. FHSA-SED also adopts two scoring criteria (K2-score and Gini-score) to evaluate the association of two-locus
models with disease status, and the logic "or" operation is performed between two scoring criteria. Therefore, in the screening stage, the MACOED algorithm adopts stricter criteria to screen the disease models than FHSA-SED algorithm; however, MACOED will make some true disease models be filtered out.

2. MACOED is intended to search disease models via ACO algorithm (employing large population size). In FHSA-SED, HS algorithm is employed to detect disease-causing SNP-pairs and a local search algorithm is presented to discover no-visited solutions in constant time. In MACOED, logical regression-based AIC-score requires some iteration to calculate regression coefficients, which take much more time than the Gini-scoring in FHSA-SED.

3. In MACOED, Pearson’s $\chi^2$ test is performed on the no-dominant solutions obtained in the screening stage. FHSA-SED employs the G-test to test the candidate solutions in the testing stage.

We investigate the performance of FHSA-SED algorithm via three simulation experiments:

1. 12 DME models: the disease loci have both main effects and interaction effects.
2. 70 DNME models: the disease loci have only the interaction effects without the main effects.
3. AMD dataset that contains 103611 SNPs genotyped for 50 controls and 96 cases.
Results of DME models indicate that FHSA-SED is more effective in seven performance metrics than MACOED and CSE, especially, it takes very fewer evaluation times of SNP-pairs and much less computation time than MACOED.

The simulation experiment on DNME models demonstrates that the performances of our method on power, evaluation times, TPR, SPC, PPV, and ACC are better than or equivalent to those of MACOED, and computation time of our method is much less than that of MACOED.

The real data AMD experiment also indicates that our method has found out the known disease loci successfully and also discovered some new suspected disease loci.

Advantages and Limitations of FHSA-SED

Advantages. FHSA-SED is a fast swarm intelligent optimization algorithm and is a model-free method that assumes neither any prior distribution nor any particular disease models. Two scoring functions in FHSA-SED can complement with each other and enhance the detection power of the two-locus disease models. Our algorithm detects the two-locus disease models without evaluating all genotype combinations by using a tabu table, and it can achieve the search performance of exhaustive search algorithm when maximum model evaluation times (MMs) is equal to the number of genotype combinations. So it is a global optimization algorithm for the detection of two-locus disease models. FHSA-SED can be easily implemented using parallel computing via splitting the tabu table (TT) into some small tabu table and each computing can be performed independently in a small tabu table.

Table 6. Top 26 high-frequency SNPs in 638 SNP-pairs on AMD dataset.

| Order | SNP     | P-value | Chromosome | Gene    | Frequency | Reported in ref |
|-------|---------|---------|------------|---------|-----------|-----------------|
| 1     | rs380390| 6.2E-07 | 1          | CFH     | 121       | [23, 28,54–58]  |
| 2     | rs1329428| 5.99E-06| 1          | CFH     | 98        | [23, 28,54–58]  |
| 3     | rs10272438| 9.67E-06| 7          | BBS9    | 57        | [56]            |
| 4     | rs1740752| 4E-05   | 10         | NA      | 40        | [54]            |
| 5     | rs3775652| 3.73E-07| 4          | INPP4B  | 38        | no reported     |
| 6     | rs1394608| 4.21E-05| 5          | SGCD    | 38        | [54,57–58]      |
| 7     | rs1363688| 3.84E-05| 5          | NA      | 36        | [28,54]         |
| 8     | rs10511467| 2.91E-05| 9          | NA      | 24        | no reported     |
| 9     | rs7006908| 0.000138| 8          | NA      | 19        | no reported     |
| 10    | rs10492272| 0.000259| 12         | ANKS1B  | 19        | [57]            |
| 11    | rs6053291| 0.000196| 20         | PROKR2  | 14        | [62]            |
| 12    | rs10512413| 0.000211| 9          | ABL1    | 13        | no reported     |
| 13    | rs10512174| 0.000194| 9          | ISCA1   | 13        | [28, 54, 58–61] |
| 14    | rs7104698| 0.000159| 11         | NA      | 12        | [58]            |
| 15    | rs6104678| 0.000212| 20         | NA      | 11        | [58]            |
| 16    | rs200642| 0.000368| 20         | TSHZ2   | 11        | no reported     |
| 17    | rs10254116| 0.00014 | 7          | BBS9    | 11        | [23,56,59]      |
| 18    | rs713392| 0.001531| 7          | IMMPP2L | 10        | no reported     |
| 19    | rs3915771| 0.000772| 5          | NA      | 9         | no reported     |
| 20    | rs3914244| 1.44E-05| 12         | NA      | 9         | no reported     |
| 21    | rs1233255| 0.000472| 2          | PMS1    | 8         | [60]            |
| 22    | rs10485193| 0.004187| 10         | NA      | 8         | no reported     |
| 23    | rs1930022| 2.37E-06| 9          | NA      | 7         | no reported     |
| 24    | rs10507949| 0.000574| 13         | NA      | 7         | [28]            |
| 25    | rs9294603| 9.7E-05 | 6          | NA      | 7         | no reported     |
| 26    | rs206695 | 0.001028| 6          | LOC728275| 5        | no reported     |

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Limitations. FHSA-SED consumes much large memory due to the considerable size of tabu table (TT). The current version of FHSA-SED cannot deal with the detection of multi-SNPs (>2) disease models. Facing various type of disease models, the balance between type I errors and type II errors has not yet to be satisfactorily solved. For the DME models with small genetic heritability H2 or minor allele frequency (MAF), the type II errors might occur, and for the models with strong marginal effects, the type I errors might be generated.

Future work. To our knowledge, there does not exist a very powerful approach in detecting high-order disease models at GWAS, therefore, at this moment, multi-loci interaction detection have many room to explore. In addition, powerful identification algorithms and statistical methods are very needed for high-order disease models. We are also developing a fast niche harmony search algorithm with small size of tabu table for detecting high-order disease models.

Supporting Information

S1 File. The method for Bayesian network scoring criteria and Gini scoring criteria. Including the detail description of Bayesian network scoring and Gini index criteria.

S2 File. Standard Harmony algorithm. Including the introduction of standard Harmony algorithm and the flow chart of FHSA-SED algorithm.

S3 File. The experiments results. All the supplementary experiment data, experiment results and figures.
S4 File. 638 SNP-pairs having strongest association with AMD. All 638 SNP-pairs that passed the Screening in 1st stage and the Testing in 2nd stage.

(XLS)

S5 File. Matlab source code of FHAS-SED algorithm.

(ZIP)

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Author Contributions

Conceived and designed the experiments: ST. Performed the experiments: ST. Analyzed the data: ST. Contributed reagents/materials/analysis tools: ST. Wrote the paper: ST JZ. Proposed the FHSA-SED algorithm firstly: ST. Put forward many constructive ideas: JZ. Gave some good ideas for this work: XY YZ ZL.

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