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Ensemble gene selection by grouping for microarray data classification

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ABSTRACT

Selecting relevant and discriminative genes for sample classification is a common and critical task in gene expression analysis (e.g. disease diagnostic). It is desirable that gene selection can improve classification performance of learning algorithm effectively. In general, for most gene selection methods widely used in reality, an individual gene subset will be chosen according to its discriminative power. One of deficiencies of individual gene subset is that its contribution to classification purpose is limited. This issue can be alleviated by ensemble gene selection based on random selection to some extend. However, the random one requires an unnecessary large number of candidate gene subsets and its reliability is a problem. In this study, we propose a new ensemble method, called ensemble gene selection by grouping (EGSG), to select multiple gene subsets for the classification purpose. Rather than selecting randomly, our method chooses salient gene subsets from microarray data by virtue of information theory and approximate Markov blanket. The effectiveness and accuracy of our method is validated by experiments on five publicly available microarray data sets. The experimental results show that our ensemble gene selection method has comparable classification performance to other gene selection methods, and is more stable than the random one.

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1. Introduction

The rapid advances in high-throughput technologies such as gene expression microarray not only provide new insights into biological phenomena, but also allow simultaneous measurement of activities of thousands of genes (features), thereby improving the prospects for effective and reliable disease diagnosis and prognosis [1]. Among various scientific tasks about microarray data (e.g. disease discovery), the identification of arrays with evidently different expression (or gene maker) is extensively studied by researchers. This is also known as classification problem in machine learning community. To effectively process microarray data, many machine learning methods, such as decision tree and nearest neighbor, have been introduced into bioinformatics [2–4], and several comparison experiments (see, e.g. [5]) have demonstrated their effectiveness.

Microarray data set (or gene expression profile), which is obtained by experimenting on several samples [1], is usually organized as a two-dimensional matrix M with n rows and m columns. These m columns represent different genes G and each row is a sample of expressions of different genes. The purpose of classification on microarray data is to separate or distinguish one

type of samples (e.g. healthy patients) from other types (e.g. cancer patients), so as to further predict response to therapy. This kind of data analysis is especially important in early tumor and cancer discovery because its result can effectively help cancer diagnosis and clinical treatment [3,4]. For microarray data, one of its characters is that the number *n* of samples collected is very small (typically less than 100), while the number m of genes is relatively huge (usually thousands or tens of thousands). This, however, poses a great challenge to traditional classification algorithms. With such an overwhelming number of genes, the efficiencies of traditional learning algorithms will be very low and at the same time, classifiers built upon them will be prone to over-fitting. Moreover, the presence of inherent noise raised from complex scientific procedures makes it even worse, when the sample size is small. To alleviate this so-called high-dimensional small-sample problem [6], an effective solution, called gene selection, is introduced.

Gene selection refers to the process of removing irrelevant or noise genes from microarray data and preserving those informative genes to predict classes or diseases. Since it can bring several advantages to classifier, such as reducing computational expense, improving prediction performance and yielding more compact and conveniently interpreted results for diagnostic task, gene selection has now attracted increasing interests in bioinformatics and many outstanding gene selection methods have been developed [7–9]. Generally, gene selection methods can be grouped into three categories, i.e., filter, wrapper and hybrid methods, depending on

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whether their evaluation criterion involves classification models [7]. Filter method is independent of classification or learning algorithm. It chooses salient genes on the ground of discriminant criterion that only relies on the characteristics of data. A representative example of this kind is mRMR [10], which maximizes relevancy with the response classes and minimizes redundancy with the selected genes by virtue of mutual information. Shen et al. [11] measured the significant degree of gene by the suitability score. Contrastively, wrapper model embeds classifier within search and evaluation procedure. As a typical illustration, SVM-RFE and its extensions adopt support vector machine (SVM) to evaluate and eliminate redundant genes recursively [12]. Besides, genetic algorithm (GA) has also been used to optimize the search procedure of gene subset. For example, Yeh [13] applied GA to identify a group of relevant genes from cancer microarray data, and then fed these genes into classifiers. While Zhu et al. [14] integrated Markov blanket with mimetic operators in GA to efficiently eliminate redundant genes. Due to the limitation of space, here we do not list them one by one. Interesting readers can refer to up-to-date surveys (see, e.g. [7–9]) to get more information about gene selection. One may observe that both filters and wrappers have their respective limitations. For instance, filters have relatively poor prediction performance, while wrappers require much more computational cost, making them inappropriate to high-dimensional data [7].

Since traditional gene selection methods provide limited contributions to classification, many researchers resort to hybrid or sophisticated techniques to choose interesting gene. For example, Au et al. [15] employed k-mean-like method to cluster genes into several groups and then selected an informative one from each cluster. Similarly, Yu et al. [16] grouped genes with similar property by kernel dense estimation. In addition, random forest has also been utilized to evaluate and select important genes, because it can take interactions between genes into account explicitly and free distributional assumptions [17,18]. To further improve prediction performance, recently many efforts have been made on ensemble technique. A typical example is ECRP [19], where different gene subsets are firstly chosen by random partitions and then used to construct classifiers. Instead of random selection. Cho and Won [20,21] obtained gene subsets by seven different correlation coefficients. However, Saeys et al. [22] utilized different filter methods to identify multiple gene subsets, while Wang et al. [23] chose a relevant gene subset in terms of class-dependent criterion for each class. Additionally, Okun and Priisalu [24] took data set complexity into consideration, and the L least complex gene subsets would be further processed. Yan and Zhang [25] obtained multi-gene subsets by backward elimination strategy on small random sets of genes one at a time with information measures. After that, the genes were ranked based on their aggregated return frequencies.

Unlike other methods, in this paper, we propose a group-based ensemble gene selection method for microarray data classification. The rationale behind it is that given a microarray profile, there is a moderate quantity of different gene subsets having the same or similarly good prediction performance, notwithstanding many of them have only a few genes in common [26]. Specifically, our method consists of three steps. At the beginning, it divides genes into several groups by approximate Markov blanket. After this grouping procedure, similar genes are bound within the same group, while dissimilar genes belong to different groups with respect to information correlation coefficient. In the second stage, for each group, one representative gene is randomly picked out to compose a gene subset. Since this subset summarizes the patterns seen across the entire data set, it is an informative one. Furthermore, multiple subsets are constructed in the same way to improve robustness. Thus they can provide complementary information about classification. Finally, classifiers are trained with these obtained subsets and an ensemble one is formed by the

majority voting strategy. The structure of the rest is organized as follows. Section 2 provides the framework of our ensemble gene selection algorithm by grouping genes. Experimental results conducted to evaluate the usefulness and effectiveness of our approach is presented in Section 3. Finally, conclusions and future works are given in the end.

2. Methods

As mentioned above, classification is the process of labeling samples with pre-defined classes in terms of available information, which is often embodied in genes. This implies that to achieve better classification performance, the common way for classifiers is to select those genes with the most discriminative capability. To measure this capability of gene, many metrics, such as discriminative contribution [27], Pearson's correlation coefficient, Spearman's correlation coefficient, Euclidean distance, cosine coefficient, signal to noise ratio and *t*-statistic [20,28], have been proposed. However, these metrics are linear ones and some of them are parametric. As a result, they do not robustly capture the non-parameterized structure shared among genes.

Unlike other criteria, information metric based on entropy are nonlinear and non-parametric [29]. Since entropy is capable of exactly quantifying the uncertainty of variable and no assumption about the distribution of data is made, information metric has attracted much attention and seems to be widely studied in practice [30]. Additionally, several empirical studies (see, e.g. [31,32]), have demonstrated that information metric is superior to others for classification in many cases. In this paper, we also place our focus on information metric. Before we delve into the details of our method, let us turn our attention to information entropy.

2.1. Information correlation coefficient

In information theory, information entropy is a fundamental concept [29]. Let X be a discrete random variable. p(x) and dom(X) denote the marginal probability distribution and domain of X, respectively. Its information amount (or uncertainty) is represented as *entropy* H(X), where

$$H(X) = -\sum_{x \in dom(X)} p(x) \log p(x). \tag{1}$$

Note that information entropy does not depend on the actual values of variable X, but only its probability distribution. For continuous variable, its alternative is often taken as an integral form. Since the estimation of probability density of continuous variable is difficult and cost, here we only deal with discrete variables with finite values for the sake of simplification, and if there is no ambiguity, dom(X) will be dropped. In a similar vein, the *joint entropy* between two variables X and Y is

$$H(X,Y) = -\sum \sum p(x,y) \log p(x,y). \tag{2}$$

Mutual information is another important concept to represent relevance between two variables. It mainly scales the average reduction of entropy (uncertainty) of one variable under the context of another one. Given two variables *X* and *Y*, their *mutual information* is

$$I(X;Y) = \sum \sum p(x,y) \log \frac{p(x,y)}{p(x)p(y)}.$$
 (3)

From this equation, one may notice that the larger their mutual information is, the higher the relevant degree between two variables is. Just owing to this, it has been extensively studied in gene selection and taken as evaluation criterion to measure the significance of genes [30]. However, an unfavorable situation for gene

selection based on mutual information is that genes with more values will be chosen in higher priority. The reason is that the value of I(X; Y) increases with the number of possible values of X and Y. Here mutual information will be normalized by the joint entropy.

Given two variables X and Y, if $H(X, Y) \neq 0$, their information correlation coefficient (*ICC*) is defined as

$$ICC(X;Y) = \frac{I(X;Y)}{H(X,Y)}. (4)$$

Otherwise, ICC(X;Y) = 1. Clearly, $0 \le ICC(X;Y) \le 1$, and if ICC(X;Y) = 1, X and Y are strictly dependent and relevant. Contrastively, ICC(X;Y) = 0 indicates that they are statistically independent or irrelevant to each other. With this notion we can express a correlation degree by stating that X is relevant to Y with degree ICC(X;Y). Moreover, ICC(X;Y) meets reflexive, symmetric and monotonic properties. Unlike mutual information, the monotonic property guarantees that this metric is not prone to choosing those genes with more values in gene selection.

2.2. Grouping genes by approximate Markov blanket

According to previous discussion, information correlation coefficient can be used to represent the gene-class relevant degree, and the higher ICC(g;C) is, the more discriminative information embodied in gene g about the target classes C. Thus, a naive solution for the problem of gene selection is to evaluate each candidate gene in terms of this metric, and select those genes with the highest correlation coefficient to build a classifier. Such method of selecting top genes has been proven effective in producing good classification performance for microarray data [33]. Unfortunately, this method only involves contribution of individual gene to classes and does not take into consideration the relevancies between selected genes. In fact, when two selected genes highly correlate to each other, their respective class-discriminative power will not change much after one of them has been removed [10]. In this scenario, it is a wise choice to consider both the gene-class and gene-gene correlation degrees in evaluating candidate genes, and a gene is good enough if it is highly correlated to classes and uncorrelated to already selected genes.

Similarly, the information correlation coefficient can also be adopted to represent the gene-gene relevant degree. If $ICC(g_1; g_2)$ is larger enough, gene g₁ is considered to be highly correlated with gene g_2 , and in some way, they have similar class-discriminative power in classification predication. That is to say, once g_1 has been selected in advance, g_2 should not be in consideration later. This gives us a good indication for selecting informative genes, i.e., before selecting genes, we can cluster genes which are highly correlated with each other into the same group, and others into different groups. Under this context, genes in the same group are highly relevant to each other and share certain biological pathway, while genes in different groups have their respective characteristics. During the procedure of selection, if gene g_i is chosen, all genes in its corresponding group are redundant and should not be considered again. By picking one representative gene out from each group, a gene subset is produced. This gene subset has good discriminate capability to the target classes, because its members come from different groups.

For gene clustering, various techniques, such as k-means and hierarchical clustering, are available [34]. However, they are often restricted by parameters and require much more training time. Inspiring from FCBF [35], we also adopt approximate Markov blanket technique to group similarity genes. Given two genes g_i and g_j , g_i is an approximate Markov blanket of g_j , if and only if $ICC(g_i, C) \ge ICC(g_j, C)$ and $ICC(g_i, g_j) \ge ICC(g_j, C)$. This definition implies that for any gene g_i , it has more relevant to its approximate

Markov blanket genes than others, because the approximate Markov blanket genes subsume not only the information that g has about C, but also about others. That is to say, the gene shares more similarity with its approximate Markov blanket genes and it can be packed into the same group where its approximate Markov blanket genes are contained. On the contrary, if a gene has no approximate Markov blanket, it forms a new group. Based on this principle, our gene grouping algorithm is shown as follows:

Algorithm1. Gene grouping algorithm using approximate Markov blanket.							
Input:	A microarray data set D with gene set G and the target						
	classes C;						
Output:	A set of gene groups GS;						
Step 1:	Initialize the set of gene groups GS as an empty set;						
Step 2:	For each gene $g \in G$, calculate its information						
	correlation coefficient $ICC(g; C)$ with the classes C and						
	sort them in a descending order;						
Step 3:	Assign the first gene to a new group G_1 and label it as						
	the center of this group;						
Step 4:	For each gene g in the sorted list, if there is a group						
	$G_i \in GS$ such that the center of G_i is its approximate						
	Markov blanket, then group g into G_i ; Otherwise,						
	insert g into a new group G_j and label g as its center;						
Step 5:	Repeat Step 4, until all genes in the list have been						
	grouped into one of gene groups;						
Step 6:	Return the set of gene groups $GS = \{G_1, \dots, G_k\}$;						

This grouping algorithm works in a straightforward way. Firstly, the gene groups GS is initialized as an empty set, and then the gene-class correlation coefficient for each gene is calculated and sorted in a descending order. The purpose of sorting is to determine whether a gene has an approximate Markov blanket in advance and facilitate to mark it as the center of a new gene group if it has no. After that, the grouping procedure for each gene continues. At this stage, each gene will be assigned into either one of existing groups or a new group, depending on whether it has approximate Markov blanket or not. The time complexity consists of two components: sorting and grouping. Assume that the number of genes is m, the time complexity of the sorting procedure is $O(m \log m)$, while the grouping one is O(mk), where k is the number of groups in GS. In the worst case where each gene forms a group containing only itself, k = m. Besides its efficiency, another advantage of this grouping algorithm is that the number k of groups is adaptively determined.

2.3. Ensemble gene selection

Once gene groups have been generated, a gene subset can be formed by picking one representative gene out from each group. Meanwhile, a corresponding classifier can be constructed by binding pre-specified learning algorithm with this gene subset. Intuitively, the center gene is a representative one for each group. However, it may loss its center position if training samples are changed. Hence, the classifier constructed in such way is not enough robust to new samples. To alleviate this issue, Ein-Dor et al. [26] suggested that many different gene subsets can lead to better classification performance, although less genes contained in these subsets are common. This is a kind of ensemble learning, which initially is a method of combining several decisions induced by individual classifiers into one in some way to improve the performance of the overall system [36]. Since ensemble learning can effectively improve classification performance, reliability and stability, it is now becoming a popular technique in machine learning community [36].

Based on this principle, our solution is to construct multiple gene subsets in the same way and then integrate them into an overall one for classification purpose. The general framework of our method is illustrated as Fig. 1, where m, k and p are the quantities of genes, gene groups and subsets, respectively. Usually, $k\neq p$ and k is adaptively achieved by Markov blanket. In the first stage, all available genes G in a microarray data set D are grouped into k disjointed gene groups GS by Algorithm 1, where |GS| = $k, GS_i \cap GS_i = \emptyset$ and $\cup GS_i = G$. After that, p gene subsets Gss can be formed from these gene groups by choosing one representative gene from each group. That is to say, for each gene subset Gss_i, it consists of k genes, i.e., $|Gss_i| = k$, and the jth gene in Gss_i comes from the *i*th gene group *GS_i*. It is observed that the center of gene group GS_i is the approximate Markov blanket of genes in this group. Although each gene in GS_i has similar discriminative power, its center has the most discriminative power, and the more near (i.e., relevant) to the center, the more information gene has. Therefore, in picking the jth gene out from GS_i , we randomly select a representative one around the center of GSi. Finally, for each gene subset Gss_i, a classifier is built and then these p base classifiers are combined by ensemble technique with majority voting manner.

A successful ensemble classifier highly relies on two aspects: diversity and combining strategy [37]. The diversity is mainly represented as different samples, genes and base classifiers. In our ensemble method, we generate p different gene subsets from gene groups to increase its diversity. Additionally, for each gene group, we randomly select representative gene only from the top t genes near around its center. It is noticeable that t is a counterpoise and its value should be determined by specific problems at hand. If t is large, the diversity increases while the stability decreases. Contrarily, if t is too small, the diversity is a problem and ensemble technique may loss its advantage. In our experiments, we found that our method worked well when t = 15.

3. Results and discussion

3.1. Data sets

To evaluate the performance of our proposed method, five publicly available gene microarray data sets were selected from literatures. These data sets are often used to validate the performance of classifiers and gene selectors. Their brief specifications are presented in the following.

Breast cancer [38] the quantities of genes and samples in this data set are 24,481 and 97, respectively. Among these samples, 46 of which are from patients who had labeled as *relapse*, the rest 51 samples are from patients who remained healthy and regarded as

non-relapse.

CNS (central nervous system) [39] records embryonal tumor patients in the central nervous system. It contains 60 patient samples with 7129 genes, where 21 are survivors (who are alive after treatment) and 39 are failures (who succumbed to their disease).

Colon cancer [40] consists of 62 samples collected from coloncancer patients. Among them, 40 patients suffer colon cancer and the remaining are normal. Although originally expression levels have 6000 genes, 2000 genes were selected in this study for the confidence in the measured expression levels.

Leukemia [1] contains the expression levels of 7129 genes for 72 bone marrow samples labeled with two classes: 47 acute lymphoblastic leukemia (ALL) and 25 acute myeloid leukemia (AML).

Prostate [41] comprises the expression levels of 12,600 genes. It contains in total 102 samples with two classes: tumor and non-tumor. Among them, 52 and 50 patients belong to tumor and non-tumor samples, respectively.

In these data sets, all genes are expressed as numerical values at different measurement levels. For convenience, we normalized each gene so that its mean and standard deviation are zero and one, respectively. Followed by Ding and Peng's suggestion in [10], each gene is further discretized into three disjointed partitions, i.e., $(-\infty, -0.5]$, (-0.5, 0.5) and $[0.5, +\infty)$, and each segment corresponds to a discrete value (e.g. 0, 1 and 2).

3.2. Results and discussion

We compared our methods (EGSG) with three model-free gene selection methods: FCBF [35], mRMR [10] and ECRP [19]. The reason of choosing them is that they are typical and popular selectors. FCBF [35] measures the relevance between genes by symmetric uncertainty and eliminate irrelevant genes by virtue of approximate Markov blanket. In mRMR [10], only those genes that may bring more relevance to the classes and less redundancy to the selected genes at the same time will be selected. ECRP [19] is an ensemble gene selector. It picks multiple gene subsets out from the original space by the manner of random partitions. Based on these gene subsets, classifiers are constructed and then integrated in majority voting way.

In our experiments, the same quantity of genes was selected for each selector to make an impartial comparison. Additionally, two classical learning algorithms, Naive Bayes (NBC) [42] and k-near-est-neighbor (kNN) [43], were adopted to build classifiers on the selected gene subsets. For kNN, k=3 and its distance was calculated by Euclidean formula in our experiments. In assessing the performance of selectors, bootstrap and re-substitution have better

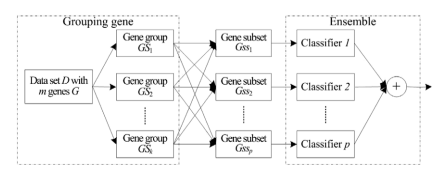


Fig. 1. A framework of ensemble gene selection by grouping technique, where k and p are the quantities of gene groups and subsets.

performance than k-fold cross-validation [44]. Nevertheless k-fold cross-validation seems more popular in literatures. There is no exception to our experiments, where leave-one-out cross-validation (LOOCV) is adopted because it is relatively steady and accurate in obtaining an unbiased performance estimation. All source codes were implemented with VC++ 6.0 and experiments were conducted on a Pentium IV PC with 2.4 GHz processor and 512 MB main memory.

3.2.1. Classification error

Table 1 summarizes the classification errors of NBC and 3NN by using four gene selectors, where the number of base classifiers (*nc*) in EGSG and ECRP is 30. From this table, one can observe that the classification errors induced by our method are not higher than those by other selectors in both classifiers, and ECRP is relatively poor. It is true because EGSG is an ensemble method, but FCBF is not, notwithstanding they share similar characteristics. For ECRP, its performance is dependent on two aspects: the number of selected genes and the intrinsic dimensionality of data set. If the number of selected genes is larger and the intrinsic dimensionality of data set is low, the classification performance of ECRP will be high. For example, in the NBC classifier, the error of ECRP on *Leuke-mia* is 1.39%. This is slightly higher than other selectors.

For the *Leukemia* data set, all samples were correctly recognized by classifiers with EGGS, FCBF and mRMR. Perhaps the reason is that gene selectors chose too many genes. To demonstrate this, we conducted extra experiments by selecting fewer genes. The result was that under the context of zero error, the quantities of genes selected by EGSG, FCBF and mRMR in NBC (3NN) were 14(15), 19(27) and 8(12), respectively. At this point, mRMR outperforms our method on this data set.

In [45], the classification performance of NBC with the RBF selector on the *Colon, Leukemia* and *Breast* were 88.71%, 98.61% and 93.81%, respectively, and the quantities of selected genes were 4, 16 and 34. Meanwhile, the corresponding accuracies of ReliefF were recorded as 85.48%, 97.22% and 79.38%. However, if we selected the same number of genes on these data sets, our accurate rates were 91.94%, 100% and 100%, respectively. Au et al. [15] proposed a gene selection algorithm based on attribute clustering, called ACA. In their paper, the classification errors of NBC with ACA on *Colon* and *Leukemia* were 35.5% and 38.2% with 7 and 50 selected genes, and these are all higher than ours: one was 8.06% and another was 0.0%.

In the 3NN classifier, Yang et al. [46] gave the classification errors achieved by their method on *Leukemia* and *Prostate* with 30 selected genes were 5.6% and 6.9%, respectively. Under the same condition, our errors were 1.39% and 4.9%. Additionally, the performance of EGSG is also superior to that of ACA on *Colon* and *Leukemia*. The similar situation can be found in comparing to DRAGS [16], which also adopts grouping technique to select informative genes. The difference with our method is that DRAGS groups genes into several clusters by virtue of the similarity of kernel dense estimation.

Table 1The LOOCV classification errors of NBC and 3NN, using four gene selectors, EGSG, FCBF, mRMR and ECRP, on five microarray data sets. For EGSG and ECRP, the number of gene subsets was 30.

Data sets	NBC				3NN				
	EGSG	FCBF	mRMR	ECRP	EGSG	FCBF	mRMR	ECRP	
Breast	0.00	2.06	0.00	17.53	3.09	7.22	5.15	25.77	
CNS	0.00	0.00	1.67	13.33	8.33	15.00	11.67	40.00	
Colon	6.45	8.06	11.21	8.06	9.68	11.29	22.58	29.03	
Leukemia	0.00	0.00	0.00	1.39	0.00	0.00	0.00	13.89	
Prostate	0.98	2.94	1.96	34.31	1.96	2.94	2.94	32.35	

Besides ECRP, our method is also competitive in comparing with other ensemble methods. For example, the recognition rates of 3NN with 25 genes on *Leukemia* and *Colon* in [20] were not more than 97.1% and 83.9%, respectively. Due to its robustness, random forest (RF) has also been applied to select or rank genes [18]. To demonstrate EGSG outperforms the RF in [18], we selected the same quantities of genes and built the same quantities of classifiers (i.e., 2000) on the *Colon*, *Prostate* and *Leukemia* data sets. The results show that our method achieved lower errors.

To illustrate the impact of validation manner, we carried out several added experiments with gene selectors by 5-fold and 10-fold cross-validations. The classification errors of 3NN are given in Table 2. The results tell us that our method can still work well under different validate manners, and there is no significant difference. Similar cases can be found in the NBC classifier, which will not be presented here due to space limitation.

3.2.2. The ensemble quantity and stability

The quantity of base classifiers in ensemble is an important factor that should be taken into consideration in its implementation. Generally, the classification performance of ensemble method increases along with the number of base classifiers (nc). Hence, in order to achieve better performance, a simple solution is to combine more base classifiers. However, when the number of base classifiers within an ensemble reaches a certain point, its classification performance increases faintly. On the other hand, it requires much more training time. To validate this assume, we performed EGSG on these five data sets 10 times. The mean accuracies are presented as Fig. 2. From this illustration, one can observe that the classification performance were changed slightly as the number of base classifiers reached a point, and for different classifiers, the threshold was different. Summarily, it is reasonable for EGSG that the threshold was assigned to 30 on these five data sets in our experiments.

Table 2The classification errors of 3NN with four gene selectors by 5-fold and 10-fold cross-validations on five microarray data sets.

Data sets	5-fold				10-fold				
	EGSG	FCBF	mRMR	ECRP	EGSG	FCBF	mRMR	ECRP	
Breast	3.09	11.34	5.15	27.84	3.09	7.22	5.15	25.77	
CNS	6.67	15.00	11.67	35.00	8.32	13.33	11.67	35.00	
Colon	8.06	11.29	22.58	29.03	8.06	11.29	24.19	30.65	
Leukemia	1.39	0.00	0.00	16.67	0.00	0.00	0.00	15.28	
Prostate	1.96	2.94	2.94	31.37	2.94	2.94	2.94	30.39	

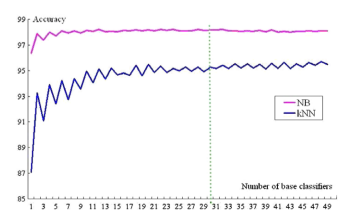


Fig. 2. The relationship between the mean performance and the number of classifiers in EGSG conducted on five data sets 10 times.

Table 3The standard deviations of 3NN with EGSG and ECRP conducted on five data sets 10 times.

	2	4	6	8	10	12	14	16	18	20	22	24
Breast												
EGSG	1.75	2.57	1.67	1.95	1.77	1.74	1.42	1.01	1.20	1.13	1.40	1.15
ECRP	3.64	4.40	3.97	4.20	3.02	3.30	3.10	1.90	3.45	4.14	3.78	3.36
CNS												
EGSG	3.96	4.33	3.00	2.71	2.60	3.06	3.08	3.16	3.86	3.64	1.70	1.71
ECRP	7.39	6.40	6.28	8.37	6.67	5.92	5.07	4.31	3.67	3.58	4.31	4.56
Colon												
EGSG	2.50	1.52	1.64	2.10	1.29	1.34	1.34	1.45	1.26	1.41	1.03	1.97
ECRP	4.09	4.19	3.37	3.90	4.88	5.86	5.73	5.50	4.85	4.23	4.67	5.97
Leukemia												
EGSG	0.69	0.56	0.42	0.00	0.00	0.00	0.00	0.00	0.00	0.42	0.42	0.00
ECRP	7.08	6.53	3.99	2.50	2.22	2.55	2.06	2.33	1.76	1.86	1.96	1.45
Prostate												
EGSG	1.98	1.32	1.55	1.48	0.99	0.96	0.98	0.90	0.90	0.63	0.63	0.65
ECRP	4.21	4.49	3.43	3.56	2.80	3.28	3.19	3.80	2.68	3.03	3.01	2.42
	26	28	30	32	34	36	38	40	42	44	46	48
Breast												
EGSG	1.05	1.24	1.31	0.92	0.80	0.92	0.81	0.81	0.65	0.56	0.72	0.66
ECRP	4.07	3.57	4.17	4.38	3.89	3.33	3.43	3.48	3.43	3.39	3.89	3.57
CNS												
EGSG	1.89	2.17	2.11	1.89	1.38	2.03	1.33	1.17	1.83	1.89	1.80	2.01
ECRP	3.59	4.73	4.01	3.83	4.29	4.64	3.93	3.14	4.33	3.87	2.29	2.81
Colon												
EGSG	1.62	1.97	1.91	1.07	1.21	1.07	1.26	1.26	1.45	1.08	1.07	1.07
ECRP	5.46	5.20	5.45	4.09	4.14	3.94	3.71	3.49	2.55	3.23	2.81	3.34
Leukemia												
EGSG	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
ECRP	1.50	1.89	1.65	1.94	1.81	1.25	1.50	1.50	1.36	1.25	1.45	1.52
Prostate												
EGSG	0.63	0.65	0.65	0.48	0.49	0.45	0.63	0.48	0.65	0.48	0.48	0.48
ECRP	2.25	2.90	2.60	2.63	1.76	1.82	2.28	1.98	2.28	2.30	1.85	2.40

Besides accuracy, stability is another aspect associated with gene selection algorithms. It mainly derives from two facts: selecting a minimum subset of genes and relatively small number of samples in data set [16,47]. To measure the stability of gene selector, various metrics have been proposed. Most of them calculate the frequencies of genes occurring in the selected gene subsets (see, e.g. [16,18,46]). However, they are not suitable for ensemble gene selection, and it is unfair to evaluation the stability of ECRP with this way because its selection operation is fully random and the total number of genes is large. Like the notion of bias-variance decomposition, which is often used to measure the stability of ensemble method in machine learning community [48], we made a comparison on the stability between EGSG and ECRP by using the variance of classification accuracy. In experiments, EGSG and ECRP were conducted on data sets with different number of gene subsets 10 times and the standard deviations of accuracies are listed in Table 3.

According to the standard deviations in this table, we can conclude that EGSG is more stable than the random one. Moreover, when the number of gene subsets (i.e., the number of base classifiers) near around 30, the standard deviations of EGSG on these five data sets (except *Leukemia*) drops sharply. That is to say, 30 is the turning point for the number of gene subsets in EGSG over four data sets. Additionally, another interesting fact is that the more gene subsets, the more stability of ensembles. This, however, can be interpreted by the relationship between the classification performance and the number of base classifiers discussed above.

4. Conclusions

In this paper, we proposed a three-stage ensemble gene selection method by grouping technique for microarray data classification, i.e., grouping genes, selecting subsets and ensemble. The

simulation results on five real data sets indicate that EGSG is competitive and effective. Under the same context, it not only leads to better classification accuracies, but also has higher stability. However, one of limitations of EGSG is that comparing with several gene selection methods, it tends to choose more genes than mRMR (e.g. the *Leukemia* data set in our experiments). Additionally, the optimal value of parameter t (i.e., the selection bound of genes) is hard to be determined in advance. Thus, our future work will be dedicated to cope with these issues. Besides, we will further validate the performance of EGSG on more data sets and under the conditions of bootstrap and re-substitution sampling.

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