

Dear Editor,

Enclosed please find our substantially revised manuscript “MGRFE: multilayer recursive feature elimination based on embedded genetic algorithm for cancer classification”. In this revised manuscript, we have carefully addressed all the concerns by the reviewers. We greatly appreciate the Referee’s comments on our manuscript. The following is our point-by-point response to each comment of the reviewers. Furthermore, I would like to take this opportunity to thank you for handling the review of our manuscript and provide us the chance to modify our manuscript again.

Our responses to the review comments are in blue.

Yours sincerely,

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### **Response to Editor Comments**

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Editor Comments

Associate Editor

Comments to the Author:

The manuscript was reviewed by the original reviewers.

Although Reviewer 1 is satisfied with the revised version, Reviewer 2 gives very critical comments.

Therefore, I recommend the authors to revise the manuscript with taking all comments into account.

Since I understand that giving theoretical justification is difficult,  
it is enough to give some discussions.

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**Response:** Thanks very much for providing us this valuable opportunity to revise our manuscript again. We have carefully revised the previous manuscript considering all the review comments. The point-to-point response to each comment of the reviewer 2 in the detail are provided as follows.

## Response to Comments of Reviewer 2

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Reviewer: 2

Recommendation: Reject

Comments:

The selection of highly informative genes in cancer patients is a standard problem with many techniques in existence. The paper presents yet another approach based on an embedded genetic algorithm. In my previous review I had raised a number of queries, which have essentially been dismissed by the authors in their revised version. My queries have NOT been addressed satisfactorily.

My original comment 3 is that there is no validation on an independent data set. The authors state in their rebuttal that "Thus, the currently published gene selection algorithms on microarrays are commonly validated within each microarray benchmark dataset."

I am sorry to say that this is incorrect. I have published several papers in computational cancer biology, and ALL of them had validations on independent data sets. I am not persuaded by the authors' argument.

"For microarray benchmark datasets about same disease, the features and sample classes are usually different. Different microarray datasets usually have different gene features for the gene probes vary among different microarray analysis platform. For example, on the leukemia related datasets of Leuk and MLL used in this study, the gene probes are very different for generating from different microarray platforms."

This is PRECISELY the reason why validation on an independent data set is so crucial. It is true that two different databases of the same form of cancer may have different genes under study. The way to handle this is to study only those genes that are common to both databases. One can also convert microarray values to Z-scores by subtracting the sample mean and dividing by the sample variance. The authors don't even try to do this.

"Thus, the currently published gene selection algorithms on microarrays are commonly validated within each microarray benchmark dataset."

This is not correct. The authors are simply trying to justify why they did not do any validation on an independent dataset.

If they have managed to do cross-validation on another dataset for leukemia, then that should be in the main paper, not in the supplementary material.

My comment 4 was that their method lacked theoretical justification and compared it to SVM-RFE. Here again the authors simply explain away my objection. They say that their GA (genetic algorithm) works faster than that of Kar et al. That was not my point at all.

In short, I believe that the authors have not adequately addressed my previous comments. Without either theoretical justification or validation on independent datasets, there is very little merit in the paper.

#### Response:

Thank you very much for your constructive and valuable comments. We also do appreciate your patient and detailed explanation on the issue that we did not understand well, which not only provide us the great help in this process of our revision, but also in our future research. As you said, in the previous revision, we indeed did not addressed your queries well due to our incorrect understanding. In this revision, we have supplemented more experiments and revised our manuscript again. We sincerely hope you can provide us another chance to review our revision.

#### # Response for previous comment 3:

According to your suggestion, we have totally added 10-group cross-validation experiments on independent datasets (Please see **Table 1**), the later seven validation datasets are collected from GEO data repository. For each experiment, firstly, the selected gene probe features by MGRFE from the first dataset were transformed into the official gene symbols; secondly, the obtained gene symbols were transformed into corresponding gene probe Ids in the second dataset; thirdly, a kind of classifier were used to perform 10 times  $k$ -fold cross validation using the samples and selected gene probe features on the second dataset; and fourthly, the performance of three different classifiers, Naive Bayes (NB), Support Vector Machine (SVM) and Random Forest (RF), on each validation dataset were recorded. Particularly, no feature mapping between Gas1 and Gas2 for they are generated simultaneously and have identical feature set [1].

Below **Table 1** becomes the TABLE 7 in the revised manuscript.

**Table 1.** Independent validation of selected gene features by MGRFE with 10-time  $k$ -fold cross validation.

Feature From / #Features	Feature Tested / #Features	#Samples	Classifier	$Sn$	$Sp$	$Acc$	$Avc$	$MCC$	$AUC$
Leuk / 2	MLL / 4	52	NB	<b>0.963</b>	<b>0.955</b>	<b>0.961</b>	<b>0.959</b>	<b>0.929</b>	<b>0.993</b>
			SVM	0.935	0.887	0.913	0.911	0.844	0.975
			RF	0.960	<b>0.955</b>	0.959	0.958	0.925	0.977
Gas1 / 2	Gas2 / 3	124	NB	<b>0.968</b>	<b>0.966</b>	<b>0.967</b>	<b>0.967</b>	<b>0.937</b>	<b>0.993</b>
			SVM	0.952	<b>0.982</b>	<b>0.967</b>	<b>0.967</b>	<b>0.937</b>	0.992
			RF	0.957	0.931	0.944	0.944	0.895	0.987
Gas2 / 3	Gas1 / 2	144	NB	<b>0.949</b>	0.968	<b>0.958</b>	<b>0.958</b>	<b>0.920</b>	<b>0.975</b>
			SVM	0.941	<b>0.972</b>	0.956	0.956	0.916	0.970
			RF	0.936	0.958	0.947	0.947	0.900	0.974

DLBCL / 3	GSE56315 / 7	88	NB	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
			SVM	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
			RF	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
Prostate / 4	GSE8511 / 5	41	NB	<b>0.884</b>	<b>0.852</b>	<b>0.870</b>	<b>0.868</b>	<b>0.753</b>	<b>0.935</b>
			SVM	<b>0.900</b>	0.665	0.806	0.783	0.582	0.900
			RF	0.868	0.752	0.822	0.810	0.646	0.917
Gastric / 3	GSE2685 / 1	30	NB	0.919	<b>0.650</b>	<b>0.846</b>	<b>0.785</b>	<b>0.584</b>	0.861
			SVM	<b>0.990</b>	0.440	0.843	0.715	0.464	<b>0.865</b>
			RF	0.862	0.500	0.765	0.681	0.365	0.686
Gastric / 3	GSE66229 / 7	400	NB	0.903	<b>0.896</b>	0.902	0.900	0.764	0.961
			SVM	<b>0.955</b>	0.864	0.932	0.909	0.823	<b>0.971</b>
			RF	0.950	0.894	<b>0.936</b>	<b>0.922</b>	<b>0.835</b>	<b>0.971</b>
Adenoma / 1	GSE8514 / 3	15	NB	0.900	0.800	0.867	0.850	0.700	<b>0.960</b>
			SVM	0.900	0.500	0.767	0.700	0.400	0.920
			RF	<b>0.910</b>	<b>0.820</b>	<b>0.880</b>	<b>0.865</b>	<b>0.730</b>	0.950
Colon / 6	GSE44076 / 23	148	NB	<b>0.988</b>	0.950	<b>0.976</b>	<b>0.969</b>	<b>0.948</b>	0.996
			SVM	0.969	0.952	0.963	0.961	0.924	0.995
			RF	0.977	<b>0.960</b>	0.972	<b>0.969</b>	0.940	<b>0.998</b>
ALL1 / 1	GSE2604 / 4	14	NB	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
			SVM	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
			RF	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>

On the tested datasets with sample number greater than 50, 10-time 10-fold cross validation were performed with different random seeds. Meanwhile, 10-time 5-fold cross validation were performed on datasets with samples less than 50.

The latter seven validation datasets are retrieved from GEO and named as their GEO accessions.

NB, SVM and RF represent Naive Bayes, Support Vector Machine and Random Forest classifiers, respectively.

The bold face values denote the highest performance achieved by the three classifiers.

The extra pre-processing on below four GEO datasets should be explained.

1. Dataset GSE8511 has three kinds of samples: "Benign Prostate", "Local Prostate Cancer" and "Metastatic Prostate Cancer". The latter two kinds of samples are combined together as "Prostate Cancer" samples in validation.
2. Dataset GSE44076 has three kinds of samples: "Mucosa sample from healthy Normal donor", "Normal paired sample from patient" and "Tumor sample from patient". The first two kinds of samples are combined together as "Normal" samples in validation.
3. On dataset GSE8511, there are total seven gene probe features mapped from Prostate dataset, but two of them contain Null values and have been abandoned. Thus, only five gene features are used on GSE8511.
4. On dataset GSE2604, there are 36 samples in total, but the 22 samples with Null values have been removed. Thus, only 14 samples are used on GSE2604.

On tested datasets GSE56315 and GSE2604 with gene features from DLBCL and ALL1 respectively, NB, SVM and RF classifiers all achieved 1.0 cross validation accuracy in each test. In particular, there are only 14 samples totally on GSE2604, which means the classifiers were trained on merely about 10 samples in each 5-fold cross validation. Thus, the selected unique gene *CD3D* is one ideal discrimination for B-cell acute lymphoblastic leukemia (ALL) and T-cell ALL. On GSE2685, there

is only one gene probe Id mapped from the selected gene *LIFR* on Gastric dataset, and no mapping items for selected genes of *GATA6-AS1* and *HHIP*. Meanwhile, the sample number of GSE2685 is merely 30. But NB and SVM still achieved acceptable cross validation accuracies over 0.8. Except three tested datasets of GSE8511, GSE2685 and GSE8514 with samples less than 50, the prediction accuracies of three classifiers are above 0.9 on all other seven datasets in cross validation. The independent validation results proved that the selected genes features by MGRFE in each dataset have strong association with the disease phenotype and can be selected as the candidates for biomarkers.

#### # Response for previous comment 4:

We are sorry not to address the comment 4 very well during the previous response and modification. In this revision, we try to discuss and compare in depth the SVM-RFE and GA-RFE from experiment to theory.

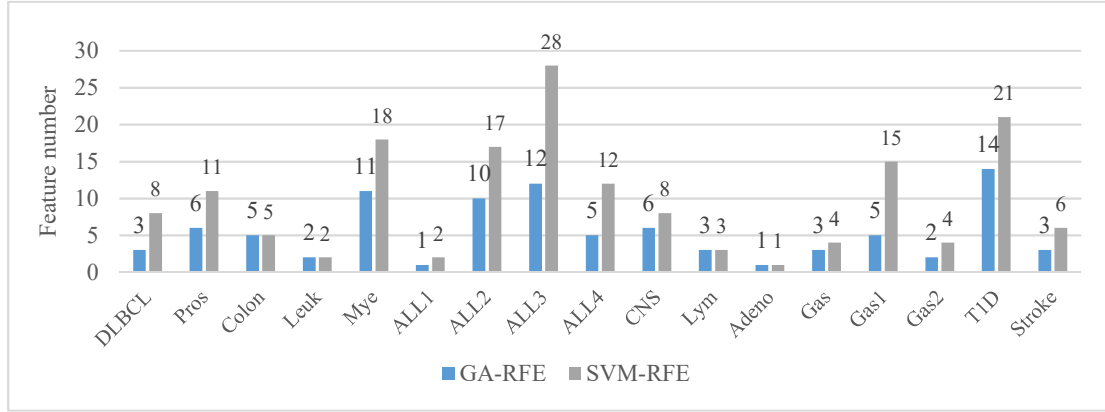
To provide a fair comparison between GA-RFE and SVM-RFE, comprehensive experiments were performed on all the 17 binary classification datasets and below details have been considered.

- Isabelle Guyon [2] pointed out that the feature normalization in pre-processing is of great importance to SVM-RFE. For each feature, we have subtracted its mean and then divided the result by its standard deviation as suggested. As a result, the feature scales are comparable within a dataset.
- Both GA-RFE and SVM-RFE use SVM model with linear kernel as the embedded classifier in the whole process. The penalty parameter  $C$  is set as 100 as in the original paper.
- Feature filter process are performed on each datasets to provide GA-RFE and SVM-RFE with same initial high quality features.
- Only one GA-RFE process is used to do comparison with SVM-RFE on each dataset. The multi-layer iteration manner is abandoned here for fairness.

**Table 2.** Both GA-RFE and SVM-RFE can achieve 100% 5-fold cross validation accuracies on 17 binary classification datasets. But GA-RFE used more compact gene subsets with smaller sizes.

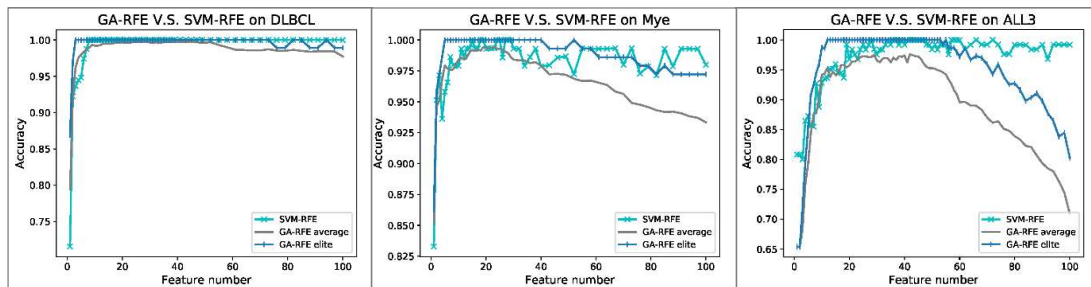
Dataset	GA-RFE		SVM-RFE	
	#Genes	Accuracy	#Genes	Accuracy
<b>DLBCL</b>	3	1.0	8	1.0
<b>Pros</b>	6	1.0	11	1.0
<b>Colon</b>	5	1.0	5	1.0
<b>Leuk</b>	2	1.0	2	1.0
<b>Mye</b>	11	1.0	18	1.0
<b>ALL1</b>	1	1.0	2	1.0
<b>ALL2</b>	10	1.0	17	1.0
<b>ALL3</b>	12	1.0	28	1.0
<b>ALL4</b>	5	1.0	12	1.0
<b>CNS</b>	6	1.0	8	1.0
<b>Lym</b>	3	1.0	3	1.0
<b>Adeno</b>	1	1.0	1	1.0

<b>Gas</b>	3	1.0	4	1.0
<b>Gas1</b>	5	1.0	15	1.0
<b>Gas2</b>	2	1.0	4	1.0
<b>T1D</b>	14	1.0	21	1.0
<b>Stroke</b>	3	1.0	6	1.0



**Figure 1.** Sizes of selected gene subsets by GA-RFE and SVM-RFE for achieving 100% cross validation accuracy on 17 datasets.

The performance of GA-RFE and SVM-RFE on 17 datasets are recorded in **Table 2**. Both GA-RFE and SVM-RFE could achieve 100% 5-fold cross validation accuracies on all these datasets. **Figure 1** provides the histogram graph of the used feature number by these two methods for better visual illustration. On 14 datasets, GA-RFE could find more compact feature subsets to achieve the same performance as SVM-RFE did.



**Figure 2.** Performance comparison of GA-RFE and SVM-RFE with varying feature number. The accuracy is calculated in 5-fold cross validation. These two methods have been provided with an initial feature set containing 500 genes generated by feature filter process on each dataset. Then, GA-RFE begin its evolution from some randomly sampled feature subsets 100 in length to save time. SVM-RFE starts with the exactly provided feature set.

To better analyze the reason why GA-RFE is more effective than SVM-RFE in finding the minimal discriminatory feature subset, we plot their dynamic performance on three datasets where GA-RFE could achieve obvious smaller feature subsets than SVM-RFE. **Figure 2** shows the performance of GA-RFE and SVM-RFE in their iterations with feature size ranging from 1 to 100 on datasets DLBCL, Mye and ALL3. SVM-RFE begins its iteration from the initial 500 features provided by

the filter process. GA-RFE starts the iteration with randomly sampled feature subsets 100 in length. From the three sub-plots in **Figure 2**, it could be noticed that GA-RFE need a convergence process to find some elites in population with 100% accuracies. For example, on ALL3 and Mye, the average performance in population (GA-RFE average) and the performance of population elites (GA-RFE elite) are constantly increased in the process of reducing the feature range from 100 to ~50. In fact, in this process, the GA evolution number is set very small (1 or 2) and RFE step is set relatively large (3 or 2) for each feature length to save time. But when some GA elites find 100% accuracy feature subsets, this best performance is constantly kept for a long feature range. The performance of SVM-RFE is relatively stable when the feature size is larger than its finally selected feature subset. But when the feature range is smaller than its finally selected feature subset, the performance usually decline obviously. GA-RFE is more robust than SVM-RFE for the elites in GA population could maintain 100% accuracy to smaller gene subset.

SVM-RFE by Isabelle Guyon [2] followed the Structural Risk Minimization (SRM) principal and has been widely recognized as the classical feature selection method on microarray data and other related problems. Its key procedure can be briefly described as follows.

- a. Train a SVM classifier with linear kernel using current feature set.  
A weight vector  $\mathbf{w}$  and a bias value  $\mathbf{b}$  are learnt in optimizing the loss function. The obtained decision function is  $D(\mathbf{x}) = \mathbf{w} \cdot \mathbf{x} + \mathbf{b}$ , where  $\mathbf{x}$  is an input train sample.
- b. Compute the ranking criterion for all features. The weight of  $i$ -th feature is  $(\mathbf{w}_i)^2$ .
- c. Remove the feature with smallest ranking weight. The removed feature is added to the head of a feature ranking list, an empty list at the start.
- d. Repeat steps a~c until the feature set is empty.
- e. Output the feature ranking list.

SVM-RFE has below remarkable advantages compared to its previous methods.

- 1) The mutual information between features are considered in SVM training. This avoids the implicit orthogonality assumption of features in simple feature filter methods.
- 2) The selecting of useful samples (support vectors of SVM model) and useful features could be connected together.

Meanwhile, it should be noted that there are also some potential shortcomings of SVM-RFE.

- 1) Its greedy feature elimination manner makes it not very robust in achieving the minimal discriminatory feature subset. The  $(\mathbf{w}_i)^2$  is only the estimation for the influence of a feature on the loss function. When a feature with currently lowest  $(\mathbf{w}_i)^2$  is removed, it don't have chance to be considered again.
- 2) No validation of feature subset. From beginning to end, the SVM model is trained on the same training dataset and the weight vector  $\mathbf{w}$  is used for computing feature importance, without any independent test or validation.
- 3) The classifier models that can be used are limited. Only the classifiers that can infer feature importance could be considered for replacing SVM in the SVM-RFE framework.
- 4) Though SVM-RFE is fast, the time cost is still high when the feature range is very large. SVM-RFE is a sequentially executed algorithm in essence. When there are tens of thousands of features in dataset, it is too time consuming to train same number of SVM models sequentially

to generate the feature ranking.

The original idea of our algorithm comes from combining the RFE framework with swarm intelligence method of GA to build a more robust and flexible feature selection method. The feature filter process is used to quickly reduce the feature range. The multi-layer iteration manner is designed to help improve stability. As a result, the former 4 potential shortcomings of SVM-RFE could be eased or improved.

- 1) The population of feature subsets in GA could bring about more robustness. When a feature is removed from one feature subset, it could still exist in other feature subsets. The population evolution of GA is more robust and fault-tolerant than the feature refining process on a single feature subset in original SVM-RFE.
- 2) For each feature subset, cross validation could be performed to evaluate its performance. The generation and evaluation processes of feature subsets are separated in GA-RFE.
- 3) All kinds of classifiers could be embedded in our method to get their most suitable feature subsets.
- 4) Two filter methods, *t*-test and MIC, are employed to quickly generate an obviously reduced feature subset with high quality feature candidates. The later RFE process are performed based on the limited candidate features. Thus, time cost is notably reduced compared with direct RFE process on initial feature set.

In our algorithm, GA is employed to generate different feature subsets. As a kind of bio-inspired algorithm, the applications of GA on various problems are far ahead of its theoretical researches. There are two basic theoretical analysis ways for GA: Schema Theorem and Markov chain. The Schema Theorem formalized by Holland is a mile stone in theory analysis for GA [3]. Simply stated, he try to prove that in the evolution process of a canonical GA, the good schemas with performance higher than the average performance in population are exponentially increased in expectation. However, there are several shortcomings in his proof which lead to more modern approaches [4-6]. For the information of GA population in the next generation usually only rely on the population in the current generation, Markov chain could be used naturally to model the behavior of GA evolution. Goldberg and Segrest present a finite Markov chain analysis to a single-locus, binary-coded finite population GA [7]. Eiben et al. employed Markov chain to prove that GA with elitism preservation mechanism has global convergence ability [8]. The elitism preservation mechanism has been adopted in the design of GA-RFE.

As you have pointed out, the swarm intelligence based optimization methods like GA have some well-known shortcomings, including the lack of precise theoretical analysis, sensitive to parameters and time cost problem. But on the other hand, it should also be noticed that the evolution calculation based method is one of the main branches for gene selection on microarray. Many related leading methods have been constantly published in recent years and still cause high concerns [9-13]. This is because these swarm intelligence based methods have their own advantages in this feature selection problem. The feature combinations in microarray is exponential correlated with the feature number, thus make this problem NP-hard. But swarm intelligence based methods are widely known for their effectiveness in solving many NP-hard and complex optimization problems.



Re-think the motivation for identifying compact discriminatory gene features in the microarray, the below two points should be noticed.

1. Structural risk minimization. Structural risk minimization is an inductive principle for model selection used for learning from finite training data sets. It describes a tradeoff between the empirical error in training data and hypothesis space complexity of a learning model. On microarray data, there are usually several thousand to tens of thousands of gene features but only dozens or hundreds of samples. Thus, the features used by the prediction model must be limited to control the model complexity. By selecting relatively small number of gene features, the learnt model could avoid the overfitting problem and have better generalization ability on unseen data. In Recursive Feature Elimination process, the number of gene features is reduced step by step, thus the corresponding learnt models are arranged in order of decreasing complexity. In minimizing both the empirical error and capacity of a model, the idea of SRM is clearly embodied.
2. Finding disease related genes and potential biomarkers. The selected minimal discriminatory gene subset has high correlation with the disease phenotype on microarray data. Thus, they are biomarker candidates for the specific disease and may provide researchers with insights into the genetic nature of the disease and mechanism behind it. Therefore, the discriminatory genes are well worth of further biological analysis.

By introducing GA into the RFE framework, our designed algorithm is fault-tolerant in feature elimination and flexible in classifier selection and feature subset evaluation. GA-RFE is robust and effective in finding the minimal discriminatory feature subset. The independent validation experiments proved that the selected gene features have good generalization performance, thus could be regarded as biomarker candidates for corresponding disease. The shortcomings of our algorithm including lack of precise theoretical analysis and kind of time consuming.

In doing the comparison experiments between GA-RFE and SVM-RFE, we find that the combination of feature normalization process and SVM model with linear kernel is more effective than our previously used Gauss Naive Bayes. The performance records in **Table 2** are higher than our previous performance records on 17 binary datasets in manuscript. Updating these records means that all the selected gene features on these datasets need to be updated simultaneously, thus many cascaded analysis and experiments, like independent validation, need to be checked and performed again. We plan to update these records and all the affected table and figure results in our next revision.

We do appreciate your rigorous academic style and responsible attitude for review work. Your comments have obviously enhanced our algorithm and manuscript. We sincerely hope that you can provide us another chance to improve our manuscript.

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# MGRFE: multilayer recursive feature elimination based on an embedded genetic algorithm for cancer classification

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**Abstract**—Microarray gene expression data have become a topic of great interest for cancer classification and for further research in the field of bioinformatics. Nonetheless, due to the “large  $p$ , small  $n$ ” paradigm of limited biosamples and high-dimensional data, gene selection is becoming a demanding task, which is aimed at selecting a minimal number of discriminatory genes associated closely with a phenotype. Feature or gene selection is still a challenging problem owing to its nondeterministic polynomial time complexity and thus most of the existing feature selection algorithms utilize heuristic rules. A multilayer recursive feature elimination method based on an embedded integer-coded genetic algorithm, MGRFE, is proposed here, which is aimed at selecting the gene combination with minimal size and maximal information. On the basis of 19 benchmark microarray datasets including multiclass and imbalanced datasets, MGRFE outperforms state-of-the-art feature selection algorithms with better cancer classification accuracy and a smaller selected gene number. MGRFE could be regarded as a promising feature selection method for high-dimensional datasets especially gene expression data. Moreover, the genes selected by MGRFE have close biological relevance to cancer phenotypes. The source code of our proposed algorithm and all the 19 datasets used in this paper are available at <https://github.com/Pengeace/MGRFE-GaRFE>.

**Index Terms**—Gene selection, Genetic algorithm, Recursive feature elimination, Microarray data, Cancer classification.

## 1 INTRODUCTION

ONE chief challenge in bioinformatics is the “large  $p$  small  $n$ ” paradigm [1], on account of ever-increasing high-dimensional data and limited available experimental samples. In particular, for gene expression data, the sample number is distinctively small compared with several thousand to tens of thousands of genes. For the analysis of high-dimensional data, feature selection is essential, which is designed to remove irrelevant and redundant features, thus cutting down the dimensionality and improving the predictive performance and model interpretability. On the other hand, due to its nondeterministic polynomial (NP) time complexity, feature selection is still a challenging and extensively studied problem in the machine learning and data mining fields. As for the field of bioinformatics, there are numerous high-dimensional biological data in sequence analy-

sis, microarray analysis, and spectral analysis. This situation makes feature selection more important and challenging. On the basis of the process of choosing features for classification, feature selection methods can be roughly subdivided into three categories: filter, wrapper, and hybrid techniques [2].

Filter algorithms generally evaluate features according to the inherent characteristic of a dataset, then rank all the features and preserve only an optimal subset of the original features. Up to now, lots of filter algorithms have been designed, such as the methods based on the  $t$ -test [3],  $\chi^2$  test [4], mutual information [5], maximal information coefficient (MIC) [6], and signal-to-noise ratio [7]. The  $t$ -test is the frequently used and efficient statistical approach to detecting differentially expressed genes in microarray analysis [8], [9], [10], [11], [12], [13], [14]. MIC is an information theory-based measurement for capturing a wide range of associations, which has shown excellent performance on detecting novel associations in large datasets [15]. The recent study of McTwo [16], which is based on MIC for selection of a gene subset in a microarray data, has outperformed most of existing algorithms. In addition, MIC may offer

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more convenience in dealing with multiclass datasets. Hence, the  $t$ -test and MIC are practicable and qualified approaches for selecting statistically significant discriminative genes and thus are mostly used in the feature preprocessing stage to generate a candidate gene set in the analysis of microarray. Because there is no classification algorithm involved in the filter algorithm, its computational speed is high and suitable for large datasets. On the other hand, the filter techniques for gene feature selection also have some limitations. First, filter methods are likely to add redundant features into the chosen subsets, which will lead to inaccessibility of optimal results. Second, the mutual information between features is ignored for the implicit orthogonality assumption of features [17]. Third, the features top-ranked by a filter algorithm are not always the best features for classification [16].

Wrapper algorithms usually employ classification models and contain heuristic rules to select feature subsets guided by the classification performance on the feature subsets being used, which are usually superior to filter algorithms but more time-consuming. A variety of wrapper algorithms have emerged involving simulated annealing, randomized hill climbing [18], regularized random forest (RRF) [19], particle swarm optimization (PSO) [20], [21], and genetic algorithm (GA) [22]. With the rapid development of heuristic rules and evolutionary strategies commonly present in wrapper techniques, various swarm intelligence algorithms have been applied to the optimization of feature selection. Kar *et al.* have proposed a particle swarm optimization method based on adaptive K-nearest neighborhood (KNN) to identify a minimum meaningful gene subset [23]. Moosa *et al.* have presented a modified artificial bee colony algorithm (ABC) to select a minimum number of genes with high predictive accuracy for cancer classification [24]. Oreski *et al.* have designed a hybrid GA with neural networks to identify an optimal feature subset with high classification accuracy and scalability for credit risk assessment [25]. Jung and Zscheischler have described a guided hybrid GA to minimize the number of cost function evaluations [26]. Nevertheless, all these feature selection methods based on swarm intelligence algorithms use the binary encoding method and lack an explicit reduction in the feature number. The feature number only changes in the randomized evolution operation like mutation and crossover. Thus, these methods lack the precise control over the gene features in the individuals and can not explicitly remove genes to decrease the feature number. Meanwhile, it has been verified that only a minimal number of informative genes is enough for effective diagnosis of different phenotypes in microarray gene

datasets [16], [17], [24], [27]. The feature selection using binary encoding has three main shortcomings in finding an optimal gene combination: (1) The fixed chromosome length for the encoding length must be equal to the gene range to represent all the genes. This arrangement can result in impossibility of the explicit reduction in the gene number and unnecessary space occupation when there are only several 1s among lots of 0s. (2) There are different numbers of actual existing genes in different individuals. Because there are different numbers of 1s, the actual number of genes varies among individuals and cannot be controlled precisely. (3) The convergence speed is usually low and the time cost is high to generate the minimal informative gene combination. The sizes of the optimal gene combinations in most of datasets are usually below 10. The evolution-based feature selection algorithms using binary encoding lack of an explicit feature reduction mechanism, which results in low probability and high time cost to generate the optimal minimal gene combination among the several thousand to tens of thousands of genes in each dataset. Recursive feature elimination (RFE) is a popular strategy that yields an explicit recursive feature reduction by recursively removing features with the least weights [17], [28], [29], [30].

Hybrid algorithms are the combination of filter and wrapper strategies [2]. First, the filter algorithms are applied to remove irrelevant features and narrow the search space. Second, the wrapper algorithms are performed on the pre-selected subsets to accomplish optimal feature selection. Hybrid algorithms can take advantage of both filter and wrapper techniques.

A multilayer recursive feature elimination method with an embedded integer-coded genetic algorithm, MGRFE, is proposed here, which can be categorized into a hybrid algorithm. On the one hand, MGRFE uses the  $t$ -test and MIC to obviously reduce the feature range and generate a candidate feature set. On the other hand, MGRFE combines the advantages of both evolution calculation of GA and the explicit feature elimination of RFE to achieve the minimum discriminative gene subset with optimal classification ability. To validate the performance of the proposed method, we performed comprehensive experiments on 19 benchmark gene expression datasets including multiclass and imbalanced datasets and compared the performance with other various feature selection methods. The comparison results show that our method outperforms most of state-of-the-art feature selection algorithms for selecting a smaller gene subset but yielding the same or higher classification accuracy. By validation experiments on independent datasets, the generalization ability of selected feature subset by

MGRFE is verified. Furthermore, the specific biomedical relevance of the selected genes to the related cancer phenotypes has also been confirmed by text mining in Pubmed. The whole work flow of this study is presented in Fig. 1.

## 2 MATERIALS AND METHODS

### 2.1 Materials

This study involves 19 benchmark microarrays including binary, multiclass, balanced, and imbalanced datasets, which are subdivided into two large datasets. Dataset One consists of the 17 binary classification datasets used in ref. [16], which includes diffuse large B-cell lymphoma (DLBCL) [31], Prostate (Pros) [32], acute lymphoblastic leukemia (ALL; subdivided into four subtypes based on different phenotypes) [33], central nervous system embryonal tumor (CNS) [34], Lymphoma (Lym) [35], Adenoma (Adeno) [36], Colon [37], Leukaemia (Leuk) [38], Myeloma (Mye) [39], Gastric (Gas) [40], and Gastric1/Gastric2 (Gas1/Gas2) cancer [41] as well as type 1 diabetes (T1D) [42], and Stroke [43]. Among them, DLBCL, Colon, Leukaemia, Myeloma, ALL1-4, and CNS datasets are imbalanced. Dataset Two is composed of the three typical benchmark microarray datasets used in ref. [23], including two multiclass datasets of small-round blue-cell tumor (SRBCT) [44] and mixed lineage leukemia (MLL) [45] and one binary dataset of acute lymphoblastic leukemia and acute myeloid (ALL\_AML). Many previous experiments are conducted on these three datasets [44], [46], [47], [48], [49], [50], [51], [52], [53], [54], [55], [56], [57], [58], [59], [60]. The performance comparison between these methods and MGRFE will be provided in the Results section. It should be mentioned that the widely used benchmark Leukaemia was tested in both refs. [16] and [23] named as Leuk and ALL\_AML respectively, but they are same actually. The information on the 19 datasets in the two large datasets is given in Tables 1 and 2 in Supplementary Material. All these datasets can be downloaded directly from <https://github.com/Pengeace/MGRFE-GaRFE>.

### 2.2 Classification performance measurements

On the 17 binary classification datasets, we performed six widely used measurements to compare the performance: Accuracy ( $Acc$ ), Sensitivity ( $Sn$ ), Specificity ( $Sp$ ), Average accuracy ( $Avc$ ), Matthews Correlation Coefficient ( $MCC$ ), and  $AUC$ .  $AUC$  is the area under the receiver operating characteristic (ROC) curve, and the formulas of the other five measurements are presented in Equation (1). In Equation (1),  $P$  and  $N$  represent the numbers of positive and negative

samples;  $TP$  and  $TN$  mean the number of correctly predicted positive and negative samples; and  $FP$  and  $FN$  denote the wrongly predicted positive and negative samples, respectively. For the two multiclass datasets, for consistency and convenience, only  $Acc$  is used.

$$\begin{aligned} Sn &= \frac{TP}{TP + FN}, & Sp &= \frac{TN}{TN + FP}, \\ Acc &= \frac{TP + TN}{P + N}, & Avc &= \frac{Sn + Sp}{2}, \\ MCC &= \frac{TP \cdot TN - FP \cdot FN}{\sqrt{(TP + FP)(TP + FN)(TN + FP)(TN + FN)}} \end{aligned} \quad (1)$$

### 2.3 Method

As shown in Fig. 2, the proposed MGRFE is divided into three stages, which are (1) Search space reduction; (2) Precise wrapper search and (3) Multiple k-fold cross-validation (CV). MGRFE is a multilayer iterative feature selection method with GA-RFE acting as the feature selection unit in every layer. GA-RFE is a recursive feature elimination process including embedded GA. The Pseudocodes 1, 2 and 3 in Supplementary Material provide the pseudocodes of the processes MGRFE, GA-RFE and embedded GA.

#### 2.3.1 Stage 1: Search space reduction

At Stage 1, two filter methods, the  $t$ -test and MIC, are used to decrease the gene range and offer a candidate gene set for later precise wrapper search stage. First, we perform the  $t$ -test on all genes and subject them to ascending sorting according to their  $p$  values, then the top-ranked statistically significant features with  $p$  values less than 0.05 are preserved. Next, the upper limit of the features kept after the  $t$ -test is set to 1000, that is, when there are more than 1000 features having  $p$  values less than 0.05, only the top 1000 with lower  $p$  values would be kept. If the preserved features after the  $t$ -test screening are fewer than 500, they are all kept directly and definitively to form the candidate gene set without MIC screening; otherwise, the MIC-based selection will be followed. Second, we carry out MIC calculation on the preserved genes and resort them according to their MIC values, then the candidate gene set is generated from the top 500 genes with higher MIC values. For the two multiclass datasets, a candidate gene set is generated based only on the descending order of MIC values of all genes for which the multivariate  $t$ -test cannot be performed directly. In the Table 5 in Supplementary Material, the number of statistically significant genes with  $t$ -test-based  $p$ -values less than 0.05 on each binary-class dataset are listed. In the "S8" section of Supplementary Material, we also give a simple comparison of  $t$ -test+MIC with other filter combinations.

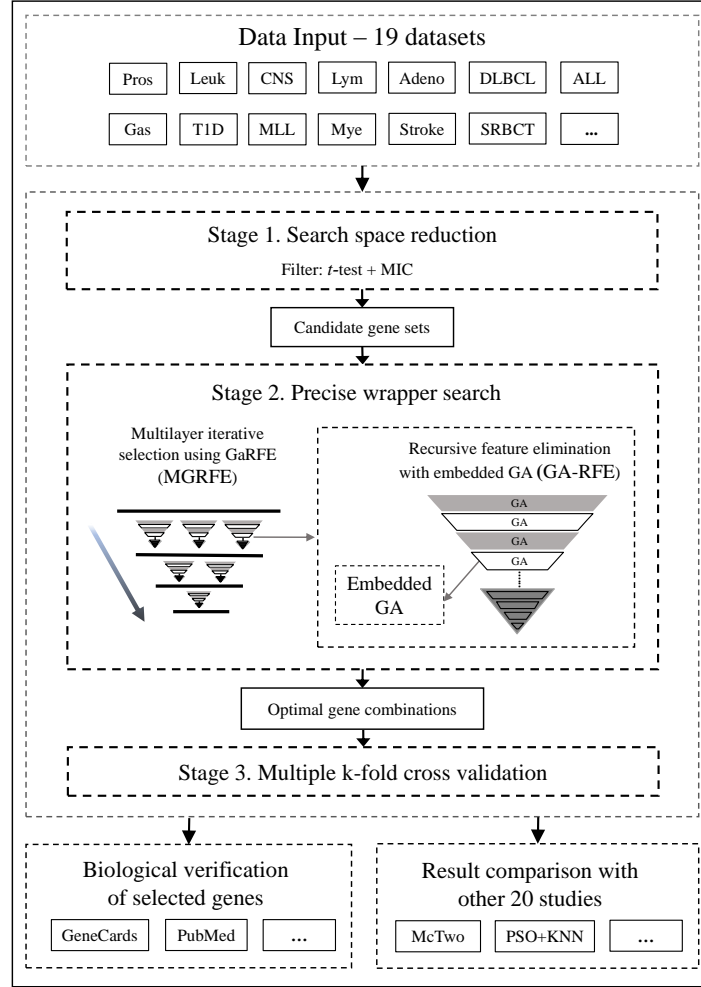


Fig. 1: A flowchart of the whole MGRFE procedure in this study.

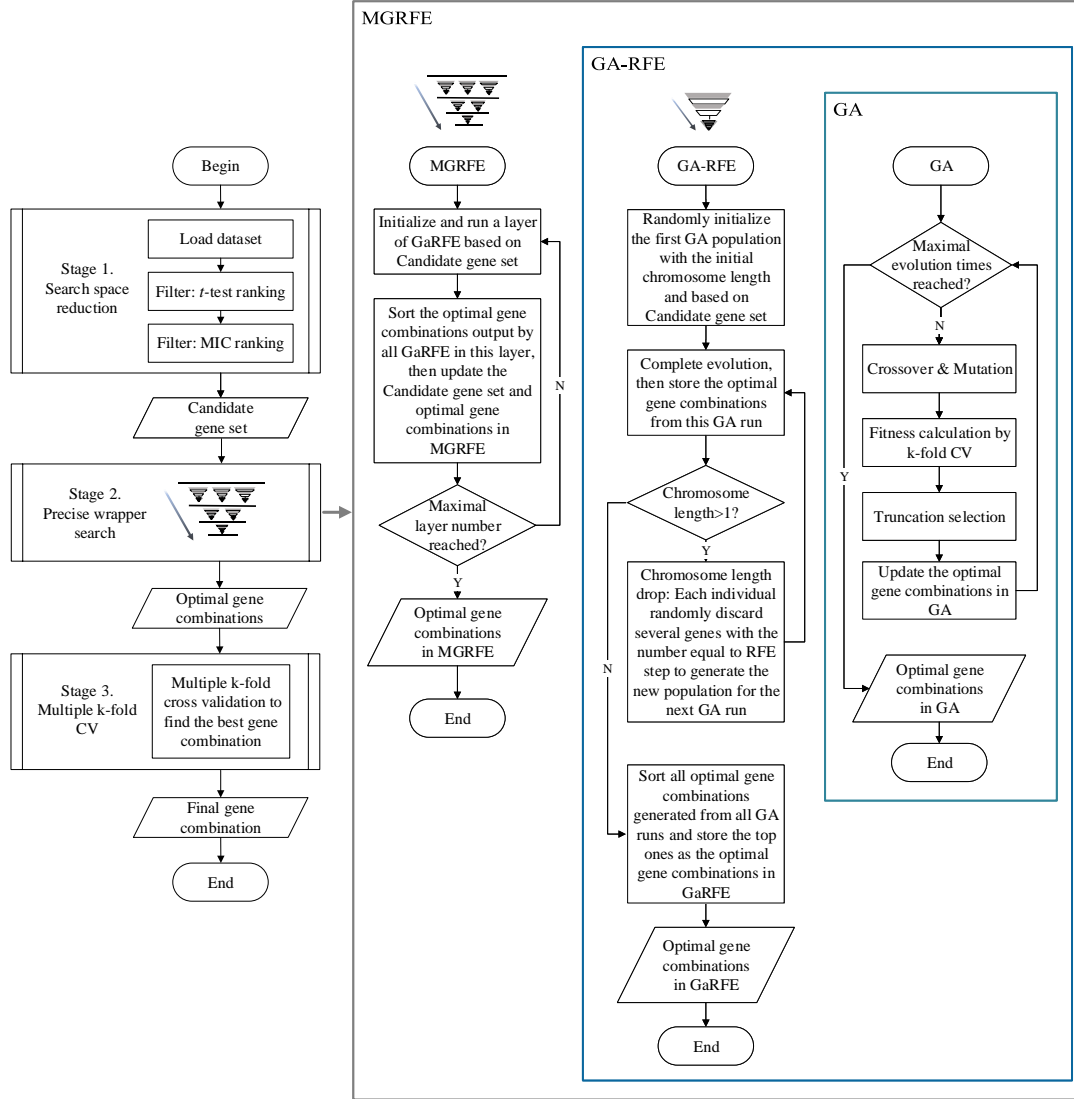
### 2.3.2 Stage 2: Precise wrapper search

At Stage 2, we search the candidate gene set obtained from Stage 1 and compute the optimal gene combinations for further selection at Stage 3. MGRFE is a multilayer iterative feature selection method and its selection unit in each layer is a GA-RFE process. GA-RFE, the inverted triangle in MGRFE as shown in Fig. 1, is the recursive feature elimination process with every stair being embedded GA. Embedded GA is the integer-coded genetic algorithm with a dynamic-length chromosome. The key feature of MGRFE is GA-RFE in each layer, in which embedded GA is responsible for generating optimal gene combinations, and the RFE process is responsible for cutting down the gene number. Therefore, our method can find gene combinations with both significantly reduced sizes and excellent classification performance.

#### Embedded GA:

In our method, the modified GA using variable-length integer-coded chromosome is embedded in the RFE process as each stair in the inverted triangle of GA-RFE. The embedded GA includes the following steps. First, we initialize the GA population by a certain amount of individuals representing gene combinations with the same sizes. Then, we perform fitness calculation and genetic operators including mutation, crossover, and selection until the stopping criterion is satisfied. In the end, we return the best individuals that represent the best gene combinations to GA-RFE. The stopping criterion of embedded GA is iteration time, which is set to 1 to 3.

To embed GA in the RFE process and achieve the minimal informative genes, some modifications are made in the original GA. The embedded GA uses variable-length integer-coding technique for the chromosome in a GA individual, and each individual has



**Fig. 2:** The flowchart of MGRFE, which is divided into 3 stages: search space reduction, precise wrapper search, and multiple k-fold CV. Stage 2 is the core of MGRFE, which includes two key processes: GA-RFE and embedded modified GA.

a set of integers representing different genes to make up a gene combination. In every run of GA, the gene combinations represented by different individuals all have a fixed size. Between two adjacent GA runs in the RFE process, every individual sheds the same number of genes from its chromosome.

A truncation selection method is used as the selection operator in embedded GA [61], which simply ranks all individuals and selects the top ones to form the next generation. The elitism preservation mechanism is used to save the currently generated best individuals. The mutation and crossover operators for generating new individuals are adjusted to adapt to the variable-length integer-encoding technique. One main challenge that should be addressed in these two

processes is avoidance of duplicated genes in every individual, which leads to the decline of the number of actual existing genes. Based on our encoding technique, the mutation operation for an individual is randomly changing some genes to others. It should be ensured that new genes do not exist in this individual previously to avoid repetitions. Crossover in embedded GA also involves single-point crossover which is the most widely used crossover method in binary encoding. Specifically, a random position is selected in the chromosome, and two parent individuals split themselves at this crossover point and then exchange chromosome tails to generate children individuals. After the crossover, the potential duplicate genes in the children individuals are replaced with other genes

from their parents to avoid decreasing the gene number. Fitness ( $F$ ) of an individual is defined in Equation (2):

$$F = \begin{cases} Acc & , \text{balanced dataset} \\ \alpha Acc + (1 - \alpha) Avc & , \text{imbalanced dataset} \end{cases} \quad (2)$$

, where  $\alpha$  is an adjustment coefficient to deal with the imbalanced datasets. For imbalanced datasets, fitness defined as  $\alpha Acc + (1 - \alpha) Avc$  can adjust the trend of predicting samples as abundant classes for  $Avc = (Sn + Sp)/2$  takes the correct prediction proportion of both sample classes into consideration. In our experiments, we take  $\alpha$  0.6 for imbalanced datasets. For balanced datasets,  $F$  is simply defined as  $Acc$ .  $F$  is calculated by 5-fold CV, and the employed classifier is Gaussian Naive Bayes classifier (NB) [62]. We sort different GA individuals based on two metrics,  $F$  and gene number. The individual with higher  $F$  is superior. For two individuals with the same  $F$  values, the one with a smaller gene number is superior. MGRFE and GA-RFE also use the above-mentioned sorting rule to rank different gene combinations.

#### Recursive feature elimination with embedded GA:

GA-RFE as shown in Fig. 2 is designed as an explicit recursive feature elimination process with embedded GA to find minimal discriminatory gene combinations. First, we randomly generate the initial GA population based on a certain candidate gene set and chromosome length. Then, we implement a chromosome length drop and a GA run in turns until the chromosome length in GA drops to 1. Finally, we sort the optimal gene combinations from all GA runs and then return the overall top-ranked gene combinations to MGRFE. The chromosome length drop means that every individual in the current GA population randomly discards the same number of genes to generate the new GA population for the next run. The number of discarded genes between two GA runs, the RFE step, is set from 1 to 3 according to the current chromosome length. A larger decline step is set for larger chromosome length to avoid time cost and a smaller decline step set for smaller chromosome length to do precise searching.

#### Multilayer iterative selection:

MGRFE is designed as a multilayer iterative feature selection method with the selection unit in each layer being GA-RFE. In every iteration layer, the GA-RFE processes analyze the current candidate gene set and return their obtained optimal gene combinations. Then the candidate gene set is reduced and subjected to the next layer of iterative selection. The candidate gene set used by the first layer of MGRFE is from the search space reduction stage. After each iteration layer, all optimal gene combinations in MGRFE will be sorted and the top-ranked ones will form the up-

dated reduced candidate gene set. After the specified layers of iteration, MGRFE sorts all the optimal gene combinations and provides the top-ranked gene combinations for Stage 3 to execute further validation.

#### 2.3.3 Stage 3: Multiple $k$ -fold CV to select the final gene combination

Stage 3 is aimed at finding the optimal gene combination with the best classification performance and minimal variance among different CV processes.  $K$ -fold CV is used for calculating the fitness of a GA individual. Multiple  $k$ -fold CV based on different random seeds is performed to further validate and select the final optimal gene combination.

## 3 RESULTS

In this section, comprehensive experiments on total 19 datasets are performed to validate the performance of MGRFE. Furthermore, the independent validation and biological verification of the selected genes are provided.

### 3.1 Results on Dataset One

The results of MGRFE on Dataset One including 17 binary datasets are given in Table 1, where six measurements calculated by 5-fold CV and the  $t$ -test and MIC-based gene rankings are listed. For 17 datasets,  $Acc$  values are all above 0.9 within 10 genes. Moreover, for 8 of 17 datasets (DLBCL, Leuk, ALL1, Lym, Adeno, Gas, Gas2, and Stroke),  $Acc$  reached 1.0 with gene number less than 5. MGRFE also show the strong robustness in dealing with imbalanced datasets like DLBCL, Colon, Leuk, ALL1, ALL4, and CNS, for which  $Sn$ ,  $Sp$ ,  $Avc$ ,  $MCC$ , and  $AUC$  are all above 0.95 without being influenced by the data imbalance. According to the  $t$ -test and MIC-based gene ranking, the best gene feature subset is not always the highest-ranked features in the filter method, thus the filter algorithm alone cannot generate the optimal feature combination. It could be noted that the relative positions of selected genes in the two ranking methods are consistent on most datasets. The top-ranked genes in the  $t$ -test are also top-ranked in the MIC sorting (e.g. the selected gene on ALL1 is the top one in both  $t$ -test and MIC ranking). For 5 of 17 datasets, the top one gene according to the  $t$ -test appeared in the final selected gene subsets. Generally, the selected informative genes are top-ranked by the  $t$ -test and MIC methods. Therefore, the filter techniques are qualified for the search space reduction task. Moreover, MGRFE achieves relatively stable classification performance in 10 repetitions of 10-fold CV as depicted in Fig. 3.



TABLE 1: Results of MGRFE on 17 datasets in Dataset One

Datasets	Pos/Neg	Genes/Total	$S_n$	$S_p$	$Acc$	$Avc$	$MCC$	$AUC$	t-test/MIC-based gene rankings
DLBCL	58/19	3/7129	1.0	1.0	<b>1.0</b>	1.0	1.0	1.0	[13/8, 39/24, 54/52]
Pros	52/50	4/12625	0.980	0.982	<b>0.981</b>	0.981	0.963	0.98	[1/1, 15/47, 74/49, 694/618]
Colon	40/22	6/2000	1.0	0.960	<b>0.985</b>	0.980	0.969	0.97	[15/6, 58/21, 176/297, 225/80, 240/555, 495/482]
Leuk	47/25	2/7129	1.0	1.0	<b>1.0</b>	1.0	1.0	1.0	[4/3, 7/5]
Mye	137/36	7/12625	0.963	0.839	<b>0.937</b>	0.901	0.816	0.95	[3/3, 15/103, 83/142, 143/13, 378/217, 404/644, 569/707]
ALL1	95/33	1/12625	1.0	1.0	<b>1.0</b>	1.0	1.0	1.0	[1/1]
ALL2	65/35	8/12625	0.914	0.908	<b>0.910</b>	0.911	0.829	0.94	[1/80, 52/395, 78/3040, 80/1297, 522/2448, 687/2038, 737/920, 760/1449]
ALL3	24/101	8/12625	0.830	0.950	<b>0.927</b>	0.890	0.785	0.93	[4/500, 52/3437, 75/3010, 142/393, 488/443, 510/795, 715/1551, 770/1321]
ALL4	26/67	6/12625	1.0	0.986	<b>0.990</b>	0.993	0.978	0.99	[1/2, 6/45, 39/356, 282/226, 535/497, 754/1377]
CNS	39/21	7/7129	1.0	1.0	<b>1.0</b>	1.0	1.0	0.98	[9/907, 53/542, 130/620, 131/519, 272/57, 273/454, 520/49]
Lym	22/23	3/4026	1.0	1.0	<b>1.0</b>	1.0	1.0	1.0	[4/7, 5/4, 669/135]
Adeno	18/18	1/7457	1.0	1.0	<b>1.0</b>	1.0	1.0	1.0	[468/27]
Gas	29/36	3/22645	1.0	1.0	<b>1.0</b>	1.0	1.0	1.0	[22/1, 77/32, 306/36]
Gas1	72/72	3/22283	0.986	0.973	<b>0.980</b>	0.980	0.961	0.99	[132/74, 248/167, 717/500]
Gas2	62/62	2/22283	1.0	1.0	<b>1.0</b>	1.0	1.0	1.0	[38/6, 89/62]
T1D	57/44	7/54675	0.911	0.912	<b>0.911</b>	0.912	0.826	0.94	[14/2229, 25/1579, 113/1287, 559/1282, 578/353, 680/426, 978/1728]
Stroke	20/20	4/54675	1.0	1.0	<b>1.0</b>	1.0	1.0	1.0	[1/3, 23/115, 129/543, 276/539]

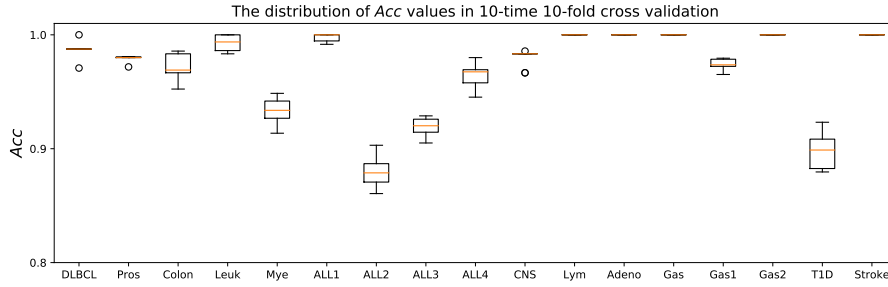


Fig. 3: The distribution of  $Acc$  values in 10-time 10-fold CV for the selected gene combinations of 17 datasets in Dataset One.

### 3.2 Comparison with other methods on Dataset One

McTwo [16] thoroughly tested all the datasets in Dataset One and demonstrated satisfactory performance. Here, we present the performance comparison between McTwo and MGRFE. Table 2 lists the overall maximal  $Acc$  and numbers of selected genes on total 17 datasets for MGRFE and McTwo. On all 17 datasets, MGRFE can obtain equal or better prediction performance compared to McTwo. On five datasets ALL2, ALL3, ALL4, Stroke and CNS, MGRFE achieves distinctly better classification performance than McTwo with relatively more genes. For a fairer and more specific comparison, the  $Acc$  values of the two algorithms are listed when the gene number of MGRFE is equal to McTwo as shown in Table 3. The results indicate that MGRFE still outperforms McTwo. Nonetheless, the  $Acc$  values associated with the usage of the gene numbers fall behind our optimal  $Acc$  values on these datasets. Thus, MGRFE selected somewhat more genes to achieve the optimal results.

### 3.3 Results on Dataset Two

Here we present the results of MGRFE on Dataset Two including three benchmark datasets, where two datasets are multiclass datasets. MGRFE selects five,

two, and three genes in SRBCT, ALL\_AML, and MLL respectively and the overall maximal  $Acc$ s are all 1.0 in 5-fold CV. In our experiments, we notice that the  $Acc$  values of the best GA individuals are kept at 1.0 in the majority of gene number ranges and only begin to drop when the gene number is significantly reduced. We also carried out 10 repetitions of 10-fold CV to further validate the final selected gene combinations in Dataset Two as shown in Fig. 4. The mean of  $Acc$ s for SRBCT, ALL\_AML, and MLL are 1.0, 0.982, and 0.997, respectively, with standard deviations being 0.0, 0.008, and 0.006. The results confirm that MGRFE has high classification stability.

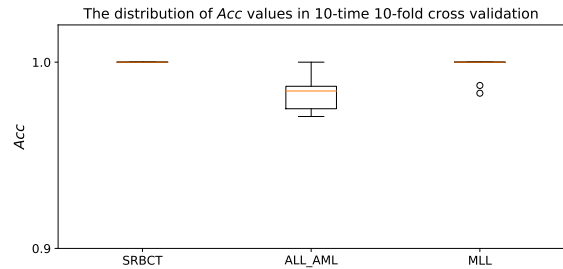


Fig. 4: The distribution of  $Acc$  values in 10-time 10-fold CV for the selected gene combinations in Dataset Two.

**TABLE 2:** Performance comparison between McTwo and MGRFE on 17 datasets in Dataset One

	DLBCL	Pros	Colon	Leuk	Mye	ALL1	ALL2	ALL3	ALL4	CNS	Lym	Adeno	Gas	Gas1	Gas2	T1D	Stroke
MGRFE <i>Acc</i>	1.0	0.981	0.985	1.0	0.937	1.0	0.910	0.927	0.990	1.0	1.0	1.0	1.0	0.980	1.0	0.911	1.0
McTwo <i>Acc</i>	1.0	0.95	0.9	1.0	0.85	1.0	0.75	0.8	0.88	0.85	1.0	1.0	0.97	0.95	1.0	0.81	0.85
MGRFE Genes	3	4	6	2	7	1	8	8	6	7	3	1	3	3	2	7	4
McTwo Genes	4	3	6	2	7	1	2	5	2	4	4	2	3	4	2	6	1

**TABLE 3:** Performance comparison on five datasets between MGRFE and McTwo when MGRFE uses the same gene numbers as McTwo does

Datasets	Methods	Genes	<i>Acc</i>
ALL2	MGRFE	2	0.760
	McTwo	2	0.75
ALL3	MGRFE	5	0.874
	McTwo	5	0.8
ALL4	MGRFE	2	0.896
	McTwo	2	0.88
CNS	MGRFE	4	0.921
	McTwo	4	0.85
Stroke	MGRFE	1	0.825
	McTwo	1	0.75

### 3.4 Comparison with other methods on Dataset Two

The performance comparison based on *Acc* and the gene number with other state-of-the-art algorithms of feature selection on the three benchmark datasets are presented in Tables 4, 5, and 6, respectively.

For the SRBCT dataset, MGRFE selected five genes and achieved 100% *Acc* in both 5-fold and 10-time 10-fold CV. In our computational experiments, combinations of four genes can reach 100% train and test *Acc* in 5-fold CV, but these gene combinations did not show classification stability in 10-time 10-fold CV. On the SRBCT dataset, Khan *et al.* [44] have applied an artificial neural network (ANN) and selected 96 genes to achieve 100% *Acc*. Tibshirani *et al.* [57] have used the nearest shrunken centroid-based method (NSC) and achieved 100% *Acc* by means of 43 genes. Fu and Fu-Liu [48] employed support vector machine (SVM)-RFE and achieved 100% *Acc* by means of 19 genes. Pal *et al.* [54] have applied feature selection multilayered perceptron (FSMLP) and non-Euclidean relational fuzzy c-means clustering (NERFCM) and found seven genes important for 100% *Acc*. Mohamad *et al.* [53] carried out improved binary PSO, and six genes were selected. Kar *et al.* [23] applied PSO and KNN and six genes were selected too. Moosa *et al.* [24] have achieved 100% *Acc* with the modified artificial bee colony algorithm (ABC) by means of five genes. Sharma *et al.* [55] have applied successive feature selection (SFS) with linear discriminant analysis (LDA) and nearest centroid classifier (NCC) and achieved 100% train and test *Acc* using four genes.

For the ALL\_AML dataset, MGRFE selected two

genes and achieved 100% 5-fold *Acc* and 98.2% 10-time 10-fold CV *Acc*. On this dataset, Fu and Fu-Liu [48] have achieved 100% train *Acc* by means of 19 genes via SVM-RFE. Yang *et al.* [59] have employed a gene-scoring technique and SVM, and four genes were selected to achieve 98.6% *Acc* in leave one out cross-validation (LOOCV). Mohamad *et al.* [53] have selected two genes to reach 100% CV *Acc* based on improved binary PSO. Dashtban and Balafar [27] have applied integer-encoding GA and SVM and selected 15 genes with 100% *Acc*. Ge *et al.* [16] have designed a two-step MIC-based method, and two genes were selected to reach 100% *Acc*.

For the MLL dataset, MGRFE selected three genes and achieved 100% 5-fold *Acc* and 99.7% *Acc* for 10-time 10-fold CV. On this dataset, Sharma *et al.* [55] have selected four genes with 100% train and test *Acc* based on SFS, LDA, and NCC. Mohamad *et al.* [53] have selected four genes with 100% CV *Acc* based on improved binary PSO. Dashtban and Balafar [27] have applied integer-encoding GA and SVM and selected 15 genes with 100% *Acc*. Kar *et al.* [23] have employed PSO and KNN to select four genes with 100% train and test *Acc* and 92.5% CV *Acc*.

### 3.5 Independent validation of selected features

We performed totally 10-group validation experiments on independent datasets to verify the generalization ability of selected gene subsets by MGRFE. For each experiment, firstly, the selected gene probe features from the first dataset were transformed into the official gene symbols; secondly, the obtained gene symbols were transformed into corresponding gene probe Ids in the second dataset; thirdly, three kinds of classifier were used to perform 10 times *k*-fold cross validation using the samples and selected gene probe features on the second dataset. Particularly, no feature mapping between Gas1 and Gas2 for they are generated simultaneously and have identical feature set.

On tested datasets GSE56315 and GSE2604 with gene features from DLBCL and ALL1 respectively, NB, SVM and RF (Random Forest) classifiers all achieved 1.0 cross validation accuracy in each test. In particular, there are only 14 samples totally on GSE2604, which means the classifiers were trained on merely about 10 samples in each 5-fold cross validation. Thus, the selected unique gene *CD3D* is one ideal discrimination

**TABLE 4:** Performance comparison among the methods on the SRBCT dataset

Experiments	Methods	Genes				CV Acc(%)				Train Acc(%)	Test Acc(%)
Khan <i>et al.</i> (2001) [44] Tibshirani <i>et al.</i> (2002) [57] Fu and Fu-Liu (2005) [48] Yang <i>et al.</i> (2006) [59]	ANN	96				-				100	100
	NSC	43				-				100	100
	SVM-RFE	19				-				100	100
		5CV				5CV					
		KNN	SVM	KNN	SVM	KNN	SVM	KNN	SVM		
	GS1	88	93	57	34	98	97.9	98.8	98.8	-	-
	GS2	90	99	77	96	98.1	99	98.8	100	-	-
	Cho's	98	98	82	80	90.2	94.3	92.8	98.8	-	-
	F-test	90	95	89	78	98	99.2	98.8	100	-	-
Pal <i>et al.</i> (2007) [54]	FSMLP+NERFCM	7				-				100	100
Li and Shu (2009) [52]	KLLE+LLE+PCA	20				-				100	100
Ji <i>et al.</i> (2011) [50]	PLSVIP	24				-				100	100
	PLSVEG	15				-				100	100
Mohamad <i>et al.</i> (2011) [53]	IBPSO	6				100				-	-
Zainuddin and Ong (2011) [60]	MSFCM+WNN	10				10CV				-	-
						100					
Lee <i>et al.</i> (2011) [51]	AGA+KNN	14				-				100	100
Sharma <i>et al.</i> (2012) [55]	SFS+LDA with NCC	4				-				100	100
	SFS+Bayes classifier	4				-				100	90
	SFS+NNC	4				-				100	95
Chen <i>et al.</i> (2014) [63]	PSODT	-				5CV				-	-
						92.94					
Kar <i>et al.</i> (2015) [23]	PSO+KNN	6				98.0159				100	100
Moosa <i>et al.</i> (2016) [24]	ABC	5				-				100	100
Dashthan and Balafar (2017) [27]	GA+SVM	18				-				100	100
<b>This paper</b>	<b>MGRFE</b>	5				5CV 10-10CV				<b>100</b>	<b>100</b>
						100 100					

In Tables 4, 5, and 6, 5CV means 5-fold cross validation; 10CV means 10-fold cross validation; 10-10CV represents 10-time 10-fold cross validation; and LOOCV represents leave one out cross validation.

**TABLE 5:** Performance comparison among the methods on the ALL\_AML (Leukaemia) dataset

Experiments	Methods	Genes				CV Acc(%)				Train Acc(%)	Test Acc(%)
Fu and Fu-Liu (2005) [48] Yang <i>et al.</i> (2006) [59]	SVM-RFE	4				-				100	97.06
		5CV				5CV					
		KNN	SVM	KNN	SVM	KNN	SVM	KNN	SVM		
	GS1	100	93	60	4	97.9	97.9	98.6	98.6	-	-
	GS2	85	98	10	25	97.1	97.4	98.6	98.6	-	-
	Cho's	100	98	9	80	96.8	97	97.2	98.6	-	-
	F-test	96	99	25	33	97.4	97.5	98.6	98.6	-	-
Shen <i>et al.</i> (2008) [56]	Stepwise	3				-				90.83	88.14
	Pure TS	5				-				95.83	94.24
	Pure PSO	7				-				94.75	94.19
	HPSOTS	7				-				98.08	95.81
Wong and Liu (2010) [58]	Probabilistic mechanism	-				SVM KNN				-	-
						97.38 98.21					
Ji <i>et al.</i> (2011) [50]	PLSVIP	9				-				100	100
		8				-				100	100
Mohamad <i>et al.</i> (2011) [53]	IBPSO	2				100				-	-
Zainuddin and Ong (2011) [60]	MSFCM+WNN	10				10CV				-	-
						98.61					
Chandra and Gupta (2011) [47]	RNBC	-				10CV				-	-
						RNBC NBC KNN					
						94.29 84.29 85.71					
Kumar <i>et al.</i> (2012) [49]	CSA	10				100				-	-
Kar <i>et al.</i> (2015) [23]	PSO+KNN	3				95.8868				100	97.0588
Ge <i>et al.</i> (2016) [16]	McTwo	2				-				100	100
Dashthan and Balafar (2017) [27]	GA+SVM	15				-				100	100
<b>This paper</b>	<b>MGRFE</b>	2				5CV 10-10CV				<b>100</b>	<b>100</b>
						100 98.2					

**TABLE 6:** Performance comparison among the methods on the MLL dataset

Experiments	Methods	Genes				CV Acc(%)				Train Acc(%)	Test Acc(%)
Yang <i>et al.</i> (2006) [59]		5CV				5CV					
		KNN	SVM	KNN	SVM	KNN	SVM	KNN	SVM		
	GS1	29	99	97	56	94.8	95.2	97.2	97.2	-	-
	GS2	91	87	90	91	94.9	94.7	97.2	97.2	-	-
	Cho's	93	89	23	44	96	95.5	97.2	95.8	-	-
	F-test	99	100	65	31	95.4	94.8	95.8	95.8	-	-
Mohamad <i>et al.</i> (2011) [53]	IBPSO	4				100				-	-
Chandra and Gupta (2011) [47]	RNBC	-				10CV				-	-
						RNBC NBC KNN					
						87.14 80 68.57					
Sharma <i>et al.</i> (2012) [55]	SFS+LDA with NCC	4				-				100	100
	SFS+Bayes classifier	4				-				100	100
	SFS+NNC	4				-				100	93
Chen <i>et al.</i> (2014) [63]	PSODT	-				5CV				-	-
						100					
Kar <i>et al.</i> (2015) [23]	PSO+KNN	4				92.5439				100	100
<b>This paper</b>	<b>MGRFE</b>	3				5CV 10-10CV				<b>100</b>	<b>100</b>
						100 99.7					

for B-cell acute lymphoblastic leukemia (ALL) and T-cell ALL. On GSE2685, the sample number is merely 30, and only contains one gene probe Id mapped from the *LIFR*, one of three selected genes on Gastric dataset. But NB and SVM still achieved acceptable cross validation accuracies over 0.8. Except three tested datasets of GSE8511, GSE2685 and GSE8514 with samples less than 50, the prediction accuracies of three classifiers are above 0.9 on all other seven datasets in cross validation. The independent validation results proved that the selected genes features by MGRFE in each dataset have strong association with the disease phenotype and can be selected as biomarker candidates.

### 3.6 Biological inferences of the genes selected by MGRFE

The sizes of gene subsets selected by MGRFE with the 100% 5-fold CV *Acc* on datasets Leuk, Gas, and ALL1 are only two, three, and one, respectively. For each gene selected by MGRFE on these three datasets, we surveyed the number of published literatures involving the gene of interest and the related cancer phenotype in PubMed on July 9, 2018. The literature-mining results on these three datasets are shown in Table 8. Moreover, the gene probes finally selected by MGRFE on all 19 datasets are provided in the "S5" section of the Supplementary Material. In the Leukaemia dataset, our selected genes are *CD33* and *TCF3*. In PubMed, there are 3001 published literatures about *CD33*, among which 1753 (58.94%) papers discuss the relevance of *CD33* to leukemia. And there are 569 publications about *TCF3* in PubMed, among which 115 (20.21%) papers confirming the association between *TCF3* and leukemia. According to GeneCards, the E protein encoded by *TCF3* performs a critical function in lymphopoiesis and is necessary for B and T lymphocytes. This gene is related to cancers including ALL (t(1;19), with *PBX1*), childhood leukemia (t(19;19), with *TFPT*), and acute leukemia (t(12;19)). In the Gastric dataset, genes *COL8A1*, *SEMA6D*, and *LIFR* are selected by MGRFE, and there are 187 publications in PubMed confirming their relevance to cancer, but only five papers reveal their relations with gastric cancer. According to the excellent classification performance of these three genes on gastric cancer, they could be novel biomarker candidates for gastric cancer. In the ALL1 dataset, only one gene, *CD3D*, is selected by MGRFE. There are 84 publications in PubMed about *CD3D*, among which 13 (15.48%) papers revealing the relevance of *CD3D* to leukemia. In [64], it has also been pointed out that gene *CD3D* is one ideally discriminatory feature and gave a diagnostic rule when the expression of *CD3D* is below a certain

cutoff limit. Regarding *CD3D*, GeneCards explains that this gene is involved in T-cell development and signal transduction, whereas defects in this gene will lead to severe combined immunodeficiency.

## 4 DISCUSSION

The proposed MGRFE is a novel multilayer recursive feature elimination algorithm based on an embedded integer-coded genetic algorithm. MGRFE is aimed at selecting minimal discriminatory gene feature subset associated closely with the disease phenotype. MGRFE could be regarded as a complementary feature selection algorithm for high-dimensional data especially for gene expression data analysis.

The main innovation of MGRFE is effectively combining the advantages of evolution calculation of the embedded GA with the explicit feature reduction manner of the RFE process in GA-RFE. Therefore, our developed MGRFE can perform explicit feature elimination along with the evolution optimization search and achieve relatively quick convergence speed. First, compared with other evolutionary-computation-based feature selection algorithms, our proposed MGRFE has shown higher convergence speed and obtained a slightly smaller discriminatory gene subset. For selecting informative gene features in a microarray, the state-of-the-art methods are commonly evolutionary-computation based. Meanwhile, almost all the existing evolution-based gene selection methods mainly rely on binary encoding and none of them take advantage of the RFE technique [14], [23], [24], [27], [65], [66], [67]. Nonetheless, the binary encoding has the shortcomings of the probable existing irrelevant features in a selected feature subset and high time cost to converge because there are thousands of genes in a microarray. Meanwhile, the fixed coding length of binary encoding leads to impossibility of explicit recursive feature reduction. Instead, MGRFE utilizes a variable-length integer-encoding technique in embedded GA and cuts down the encoding length recursively in an RFE process, which can quickly remove the irrelevant and redundant features and converge to a minimal informative feature combination. In 2017, Dashtban and Balafar also proposed an integer-coded GA with dynamic coding length for gene selection [27], but they did not employ the recursive feature reduction technique. In fact, their method selected 18 and 15 genes with *Acc* 100% on the SRBCT and ALL\_AML datasets, respectively. But MGRFE only needs five and two genes to accomplish the same performance. Second, compared with the original SVM-RFE [17], MGRFE is more flexible and robust. SVM-RFE ranks all gene features by the weight vector from SVM with a linear kernel and removes the feature with

**TABLE 7:** Independent validation of selected gene features by MGRFE with 10-time  $k$ -fold cross validation.

Feature From / #Features	Feature Tested / #Features	#Samples	Classifier	$S_n$	$S_p$	$Acc$	$Ave$	$MCC$	$AUC$
Leuk / 2	MLL / 4	52	NB	<b>0.963</b>	<b>0.955</b>	<b>0.961</b>	<b>0.959</b>	<b>0.929</b>	<b>0.993</b>
			SVM	0.935	0.887	0.913	0.911	0.844	0.975
			RF	0.960	<b>0.955</b>	0.959	0.958	0.925	0.977
Gas1 / 2	Gas2 / 3	124	NB	<b>0.968</b>	<b>0.966</b>	<b>0.967</b>	<b>0.967</b>	<b>0.937</b>	<b>0.993</b>
			SVM	0.952	<b>0.982</b>	<b>0.967</b>	<b>0.967</b>	<b>0.937</b>	0.992
			RF	0.957	0.931	0.944	0.944	0.895	0.987
Gas2 / 3	Gas1 / 2	144	NB	<b>0.949</b>	0.968	<b>0.958</b>	<b>0.958</b>	<b>0.920</b>	<b>0.975</b>
			SVM	0.941	<b>0.972</b>	0.956	0.956	0.916	0.970
			RF	0.936	0.958	0.947	0.947	0.900	0.974
DLBCL / 3	GSE56315 / 7	88	NB	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
			SVM	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
			RF	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
Prostate / 4	GSE8511 / 5	41	NB	<b>0.884</b>	<b>0.852</b>	<b>0.870</b>	<b>0.868</b>	<b>0.753</b>	<b>0.935</b>
			SVM	<b>0.900</b>	0.665	0.806	0.783	0.582	0.900
			RF	0.868	0.752	0.822	0.810	0.646	0.917
Gastric / 3	GSE2685 / 1	30	NB	0.919	<b>0.650</b>	<b>0.846</b>	<b>0.785</b>	<b>0.584</b>	0.861
			SVM	<b>0.990</b>	0.440	0.843	0.715	0.464	<b>0.865</b>
			RF	0.862	0.500	0.765	0.681	0.365	0.686
Gastric / 3	GSE66229 / 7	400	NB	0.903	<b>0.896</b>	0.902	0.900	0.764	0.961
			SVM	<b>0.955</b>	0.864	0.932	0.909	0.823	<b>0.971</b>
			RF	0.950	0.894	<b>0.936</b>	<b>0.922</b>	<b>0.835</b>	<b>0.971</b>
Adenoma / 1	GSE8514 / 3	15	NB	0.900	0.800	0.867	0.850	0.700	<b>0.960</b>
			SVM	0.900	0.500	0.767	0.700	0.400	0.920
			RF	<b>0.910</b>	<b>0.820</b>	<b>0.880</b>	<b>0.865</b>	<b>0.730</b>	0.950
Colon / 6	GSE44076 / 23	148	NB	<b>0.988</b>	0.950	<b>0.976</b>	<b>0.969</b>	<b>0.948</b>	0.996
			SVM	0.969	0.952	0.963	0.961	0.924	0.995
			RF	0.977	<b>0.960</b>	0.972	<b>0.969</b>	0.940	<b>0.998</b>
ALL1 / 1	GSE2604 / 4	14	NB	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
			SVM	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>
			RF	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>	<b>1.000</b>

On the tested datasets with sample number greater than 50, 10-time 10-fold CV were performed with different random seeds. Meanwhile, 10-time 5-fold CV were performed on datasets with samples less than 50.

The later seven validation datasets are retrieved from GEO and named as their GEO accessions.

On MLL, only "ALL" and "AML" samples are used. On GSE8511, the "Local Prostate Cancer" and "Metastatic Prostate Cancer" samples are combined together. On GSE44076, the "Mucosa sample from healthy Normal donor", "Normal paired sample from patient" samples are combined together. On GSE8511, the features containing Null values have been abandoned. On GSE2604, the samples with Null values have been removed.

NB, SVM and RF represent Gaussian Naive Bayes, Support Vector Machine and Random Forest classifiers, respectively. The **bold** face values denote the highest performance achieved by classifiers.

**TABLE 8:** Literature mining in PubMed for the selected genes on Leukaemia, ALL1 and Gastric datasets

Dataset	Probe ID	Gene	PubMed hits for gene of interest	PubMed hits for gene of interest and leukemia <sup>(1)</sup> (Ratio1)	
Leukaemia	M23197_at	CD33 Molecule(CD33)	3001	1753(58.41%)	
	M31523_at	Transcription Factor 3(TCF3)	569	115(20.21%)	
ALL1	38319_at	CD3d molecule(CD3D)	84	13(15.48%)	
Dataset	Probe ID	Gene	PubMed hits for gene of interest	PubMed hits for gene of interest and cancer <sup>(2)</sup> (Ratio2)	PubMed hits for gene of interest and gastric cancer <sup>(3)</sup> (Ratio3)
Gastric	226237_at	collagen type VIII alpha 1 chain(COL8A1)	66	15(22.73%)	2(13.33%)
	226492_at	semaphorin 6D(SEMA6D)	41	13(31.71%)	1(7.69%)
	227771_at	leukemia inhibitory factor receptor alpha(LIFR)	463	159(34.34%)	2(1.26%)

<sup>1</sup> gene of interest [All Fields] AND ("leukemia"[All Fields]).

<sup>2</sup> gene of interest [All Fields] AND ("tumour"[All Fields] OR "neoplasms"[MeSH Terms] OR "neoplasms"[All Fields] OR "tumor"[All Fields] OR "cancer"[All Fields] OR "carcinoma"[All Fields]).

<sup>3</sup> gene of interest [All Fields] AND ("stomach"[All Fields] OR "gastric"[All Fields]) AND ("tumour"[All Fields] OR "neoplasms"[MeSH Terms] OR "neoplasms"[All Fields] OR "tumor"[All Fields] OR "cancer"[All Fields] OR "tumor"[All Fields] OR "carcinoma"[All Fields]).

<sup>4</sup> Ratio1 = # (gene of interest-leukemia related literatures) / # (gene of interest literatures).

<sup>5</sup> Ratio2 = # (gene of interest-cancer related literatures) / # (gene of interest literatures).

<sup>6</sup> Ratio3 = # (gene of interest-gastric cancer related literatures) / # (gene of interest-cancer related literatures).

the smallest weight recursively. Nonetheless, SVM-RFE has the following potential limitations: 1) the greedy feature elimination manner makes it not very robust in achieving the minimal discriminatory feature subset; 2) features are only selected on training dataset, no independent test or validation involved; 3) other classifiers that can be used to replace the linear kernel SVM are limited. In GA-RFE, the population evolution of GA is more robust and fault-tolerant than the feature refining process on a single feature subset in original SVM-RFE. Meanwhile, the generation and evaluation processes of feature subsets are separated

in GA-RFE. Thus, independent validation for feature subset could be done and all kinds of classifiers could be embedded in MGRFE to obtain their most suitable feature subsets. In 2005, Fu and Fu-Liu evaluated SVM-RFE on datasets SRBCT and ALL\_AML and finally selected 19 and four genes to achieve 100% and 97.6% test *Accs*, respectively [48]. But MGRFE selected only five and two genes to attain 100% *Accs* in 5-fold CV for the same datasets. As for the selection operator of our embedded GA, we find that the widely used roulette wheel selection [61] is inferior to simple truncation selection in this gene selection problem.

The fitness gaps between different GA individuals are usually slight and account for only a tiny proportion of a fitness value. This situation provides all individuals with nearly the same area occupation in the roulette wheel and leads to the inefficiency of roulette wheel selection.

The 19 popular benchmark microarray datasets including multiclass and imbalanced datasets are employed to validate MGRFE. According to the performance comparison with other various algorithms, our proposed MGRFE is proved to be superior to most of the current state-of-the-art feature selection methods. MGRFE offers smaller informative gene subsets but the same or higher phenotype diagnosis accuracies. Many promising results are obtained by MGRFE on these datasets. MGRFE can reach *Acc* 100% within only five genes for 10 (52.6%) of 19 datasets, and *Acc* higher than 90% within 10 genes for all 19 datasets, in 5-fold CV. MGRFE also possesses strong robustness for multiclass datasets and imbalanced datasets according to metrics *Sn*, *Sp*, *Av*, *MCC*, and *AUC*.

To conclude, the chief research contribution in theory is providing a novel feature selection method which combines embedded genetic algorithm with recursive feature elimination process, working as a creative thought for future research. To the best of our knowledge, none previous studies have designed an evolutionary algorithm using variable length integer encoding approach in a recursive manner to select minimal discriminatory feature subset in high-dimension data, which is described in this paper. Meanwhile, through theoretical and experimental comparisons, our proposed MGRFE could outperform mostly other state-of-the-art algorithms for gene selection on microarray. Therefore, the proposed method MGRFE is worthy to be generalized to more feature selection problems on high-dimensional data characterized by the "large  $p$  small  $n$ " paradigm and applied in several practical fields.

Furthermore, our presented MGRFE would be useful in medical diagnosis as well as further biomedical research. The biological associations with phenotypes using literature mining in PubMed for the selected genes confirmed that the genes selected by MGRFE are biologically relevant to cancer phenotypes. Therefore, the informative genes selected by MGRFE could be novel biomarker candidates that are useful for better understanding the molecule mechanism related to the disease phenotypes and developing potential early detection and molecularly-targeted therapies for cancer diseases. For clinical applications involving microarrays, MGRFE can contribute to the development of a potential simplified procedure for diagnosis of cancer subgroups by selecting the minimal discriminatory

gene subsets, which will cut down the cost of medical diagnoses.

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# Supplementary For MGRFE: multilayer recursive feature elimination based on an embedded genetic algorithm for cancer classification

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## S1. THE DATASETS USED IN THIS STUDY

This study used total 19 benchmark microarrays subdivided into two large Datasets to validate the performance of our proposed MGRFE. In Tables 1 and 2, we provide the brief description of each dataset in Dataset One and Dataset Two.

## S2. PSEUDOCODE OF THE PROPOSED MGRFE

In the main manuscript, we provided the flow chart of the proposed MGRFE in Fig. 2. Here, we supplement the pseudocodes of our methodology. Pseudocode 1 describes the complete procedure of MGRFE. Pseudocodes 2 and 3 explain the two key processes of MGRFE: GA-RFE and embedded GA.

## S3. IMPLEMENTATION NOTES AND COMPUTATION TIME

We implemented the proposed MGRFE in Python version 3.6.0 environment (<https://www.python.org/>) on a common laptop computer with Intel(R) Core(TM) i5-4210U CPU and 8G memory. The Python SciPy package version 0.19.0 [3] was involved in the  $t$ -test process, and minepy package version 1.2.0 [4] was used to perform the MIC calculation. Some parameter settings about MGRFE: 1) the evolution iteration

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### Pseudocode 1: MGRFE: multilayer iterative feature selection using GA-RFE

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**Input :** A microarray gene expression data  
**Output:** The optimal gene feature combination for phenotype classification  
 The  $t$ -test-based gene ranking to generate the candidate gene set  $G$ ;  
 The MIC-based gene ranking to narrow  $G$ ;  
 Set  $GC$ , the list of optimal gene combinations in MGRFE, to empty;  
**while** the maximal iterative layer number not reached **do**  
   Initialize and run a layer of GA-RFE (Pseudocode 2) based on  $G$ ;  
   **for each** GA-RFE **do**  
     Add the returned optimal gene combinations to  $GC$ ;  
   Sort the optimal gene combinations in  $GC$  and only preserve the top ranked ones;  
   Use the genes in the top ranked gene combinations in  $GC$  to form a reduced  $G$ ;  
 Multiple k-fold CV on the gene combinations in  $GC$ ;  
 Return the final selected gene combination;

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number of embedded GA was dynamically set to 1 to 3 (smaller iteration number used for larger chromosome length to save time); 2) the reduced feature number between two GA runs, the RFE step, was dynamically set to 1 to 3 (larger reduction step used for larger chromosome length to save time); and 3) the iterative layer number of MGRFE being 3 with three, two and

TABLE 1  
Summary of the 17 binary classification datasets in Dataset One from ref. [1]

ID	Dataset	Samples	Features	Summary
1	DLBCL <sup>1</sup>	77	7 129	DLBCL patients (58) and follicular lymphoma (19)
2	Pros(Prostate) <sup>1</sup>	102	12 625	prostate (52) and non-prostate (50)
3	Colon <sup>2</sup>	62	2 000	tumour (40) and normal (22)
4	Leuk(Leukaemia) <sup>2</sup>	72	7 129	ALL (47) and AML (25)
5	Mye(Myeloma) <sup>3</sup>	173	12 625	presence (137) and absence (36) of focallesions of bone
6	ALL1 <sup>1</sup>	128	12 625	B-cell (95) and T-cell (33)
7	ALL2 <sup>1</sup>	100	12 625	patients that did (65) and did not (35) relapse
8	ALL3 <sup>1</sup>	125	12 625	with (24) and without (101) multidrug resistance
9	ALL4 <sup>1</sup>	93	12 625	with (26) and without (67) the t(9;22) chromosome translocation
10	CNS <sup>1</sup>	60	7 129	medulloblastoma survivors (39) and treatment failures (21)
11	Lym(Lymphoma) <sup>1</sup>	45	4 026	germinalcentre (22) and activated B-like DLBCL (23)
12	Adeno(Adenoma) <sup>1</sup>	36	7 457	colon adenocarcinoma (18) and normal (18)
13	Gas(Gastric) <sup>3</sup>	65	22 645	tumors (29) and non-malignants (36)
14	Gas1(Gastric1) <sup>3</sup>	144	22 283	non-cardia (72) of gastric and normal (72)
15	Gas2(Gastric2) <sup>3</sup>	124	22 283	cardia (62) of gastric and normal (62)
16	T1D <sup>3</sup>	101	54 675	T1D (57) and healthy control (44)
17	Stroke <sup>3</sup>	40	54 675	ischemic stroke (20) and control (20)

In Tables 1 and 2, "Samples" and "Features" indicate the total sample number and feature number of each dataset, and "Summary" column describes the sample classes and the related sample numbers in parenthesis.

<sup>1</sup> These datasets were retrieved from <http://www.broadinstitute.org/cgi-bin/cancer/datasets.cgi>.

<sup>2</sup> Colon and Leuk datasets were downloaded from the R/Bioconductor packages *colonCA* and *golubEsets*, respectively.

<sup>3</sup> These datasets were downloaded from <https://www.ncbi.nlm.nih.gov/geo/>.

TABLE 2  
Summary of the 3 classification datasets in Dataset Two from ref. [2]

ID	Dataset	Classes	Samples	Features	Summary
1	SRBCT <sup>1</sup>	4	88	2 308	EWS (29), NHL (11), NB (18) and RMS (25)
2	ALL_AML <sup>2</sup>	2	72	7 129	ALL (47) and AML (25)
3	MLL <sup>3</sup>	3	72	12 582	ALL (24), MLL (20) and AML (28)

<sup>1</sup> SRBCT dataset was downloaded from <http://research.nhgri.nih.gov/microarray/Supplement/>. This dataset includes 88 samples totally, but five of them are irrelevant and thus only 83 samples were used.

<sup>2</sup> ALL\_AML in Dataset Two and Leuk in Dataset One are the same dataset in actual.

<sup>3</sup> MLL dataset was retrieved from [http://portals.broadinstitute.org/cgi-bin/cancer/publications/pub\\_paper.cgi?paper\\_id=63](http://portals.broadinstitute.org/cgi-bin/cancer/publications/pub_paper.cgi?paper_id=63).

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**Pseudocode 2:** GA-RFE: recursive feature elimination with embedded GA

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**Input :** Candidate gene set  $G$ , Maximal chromosome length  $L$   
**Output:** The optimal gene feature combinations in GA-RFE  
Randomly generate the first GA population  $P$  from  $G$  with chromosome length equal to  $L$ ;  
Set  $GC$ , the list of optimal gene combinations in GA-RFE, to empty;  
**do**  
    Execute embedded GA (Pseudocode 3) using the population  $P$ ;  
    Add the returned gene combinations by GA to  $GC$ ;  
    **if** the current chromosome length  $> 1$  **then**  
        Chromosome length drop: each individual in  $P$  randomly discard several genes with the number equal to the RFE step;  
    **while** the current chromosome length  $\geq 1$ ;  
    Sort the optimal gene combinations in  $GC$  and only preserve the top ranked ones;  
    Return the optimal gene combinations in  $GC$ ;

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**Pseudocode 3:** Embedded GA

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**Input :** GA population  $P$ , Maximal evolution times  $T$   
**Output:** Updated  $P$ , The optimal gene feature combinations in GA  
Set  $GC$ , the list of optimal gene combinations in GA, to empty;  
**while** the maximal evolution times  $T$  not reached  
    **do**  
        Perform mutation operator;  
        Perform crossover operator;  
        Fitness calculation of each GA individual by k-fold CV;  
        Truncation selection to form the updated  $P$ ;  
    Sort the GA individuals in  $P$  and select the top ones to form  $GC$ ;  
    Return the updated population  $P$  and the optimal gene combinations in  $GC$ ;

---

one GA-RFE processes at each layer is usually enough. In the experiments, we limited the size of the final selected gene combination in each dataset to below 10 genes. According to the experiment records, in each of the 19 microarray datasets, the running time of MGRFE is commonly between 500 seconds (8.33 minutes) and 900 seconds (15 minutes). The running time has included the whole filter screen and later wrapper search processes. Additionally, it is well worthy to mention that the final chosen gene subset in each dataset might be already found by the first GA-RFE process in the first layer of MGRFE, which just costs 2~3 minutes. More implementation details and experiment results of MGRFE in the 19 datasets are available at <https://github.com/Pengeace/MGRFE-GaRFE>.

Because Kar *et al.* also employed an evolutionary-computation method PSO, which is similar to GA, to select minimal informative genes in microarray and provided their program running time records on three datasets SRBCT, ALL\_AML, and MLL [2], here, we offer a simple running time comparison between their method and MGRFE. Their PSO-based method cost 2.7956, 2.7906 and 7.1488 hours on SRBCT, ALL\_AML and MLL respectively to find their optimal gene subsets. In contrast, MGRFE merely used 10.8230, 9.0108 and 8.8739 minutes respectively in the same three datasets and thus showed much higher converge speed. Moreover, according to Tables 4, 5 and 6 in the main manuscript, the gene subsets selected by MGRFE had smaller sizes but higher classification accuracies compared with Kar *et al.*'s method. We noted that Kar *et al.* didn't employ the filter techniques to cut down the feature search space and their binary-coded PSO don't has any explicit feature decline mechanism like RFE.

#### S4. PERFORMANCE OF MGRFE IN 10-TIME 10-FOLD CV

In the main manuscript, the performance of MGRFE on the two large Datasets in 10-time 10-fold cross validation (CV) are shown in the box-plot form. Here, we supplement the detailed mean accuracies (*Mean Accs*) and standard deviations (*S.D.s*) of MGRFE on all the 19 datasets. In each dataset, 10-fold CV is repeated 10 times based on different random seeds. In each 10-fold CV, the mean accuracy value in 10-fold is calculated and recorded. Then after 10 repetitions of the 10-fold CV, the *Mean Acc* and *S.D.* of MGRFE in a dataset is calculated from the recorded total 10 mean accuracy values.

#### S5. THE GENE PROBES SELECTED BY MGRFE

The gene probes finally selected by MGRFE on all the 19 datasets are listed in Table 4. These differentially

TABLE 3  
Mean Acc and S.D. of MGRFE on 19 benchmark datasets in 10-time 10-fold CV

Dataset	DLBCL	Pros	Colon	Leuk	Mye	ALL1	ALL2	ALL3	ALL4	CNS
Mean Acc	0.987	0.979	0.971	0.982	0.933	0.998	0.880	0.920	0.963	0.980
S.D.	0.007	0.003	0.012	0.007	0.011	0.004	0.013	0.007	0.010	0.007

Dataset	Lym	Adeno	Gas	Gas1	Gas2	T1D	Stroke	SRBCT	MLL
Mean Acc	1.000	1.000	1.000	0.974	1.000	0.897	1.000	1.000	0.997
S.D.	0.000	0.000	0.000	0.004	0.000	0.014	0.000	0.000	0.006

Note that the Mean Acc and S.D. of MGRFE in dataset ALL\_AML are same as the records in Leuk for these two are the same dataset in actual.

TABLE 4  
The gene probes finally selected by MGRFE on the 19 microarray datasets

Dataset	Probe number	Gene probes
DLBCL	3	[X69433_at, Z84497_s_at, M15205_at]
Pros	4	[37639_at, 38634_at, 1909_at, 37537_at]
Colon	6	[Hsa.36952, Hsa.36696, Hsa.94, Hsa.442, Hsa.5226, Hsa.5756]
Leuk	2	[M23197_at, M31523_at]
Mye	7	[35977_at, 33130_at, 31366_at, 34571_at, 38013_at, 1368_at, 41150_r_at]
ALL1	1	[38319_at]
ALL2	8	[37502_at, 39885_at, 1291_s_at, 39408_at, 1838_g_at, 819_at, 31331_at, 39336_at]
ALL3	8	[38907_at, 38478_at, 34284_at, 37693_at, 201_s_at, 34497_at, 37809_at, 41259_at]
ALL4	6	[39631_at, 38119_at, 36795_at, 36873_at, 39905_i_at, 1265_g_at]
CNS	7	[S76475_at, M96739_at, X64624_s_at, X93511_s_at, K01911_at, S78693_f_at, X78565_at]
Lym	3	[GENE3332X, GENE3261X, GENE1191X]
Adeno	1	[D43636]
Gas	3	[225571_at, 236118_at, 237466_s_at]
Gas1	3	[213125_at, 41037_at, 208897_s_at]
Gas2	2	[212344_at, 210766_s_at]
T1D	7	[1566232_at, 215728_s_at, 215612_at, 226585_at, 239474_at, 219870_at, 244223_at]
Stroke	4	[1567009_at, 240084_at, 239389_at, 233835_at]
SRBCT	5	[245330.0, 784257.0, 43733.0, 784224.0, 295985.0]
MLL	3	[38242_at, 37710_at, 1389_at]

expressed genes could be potential biomarker candidates that are useful to related phenotype researches.

## S6. THE STATISTICALLY SIGNIFICANT GENES IN *t*-TEST

In *t*-test, this study adopt the widely used  $p = 0.05$  significance threshold to select the differentially expressed genes with  $p$ -values lower than the threshold. The chosen of  $p=0.05$  has also been experimentally validated by the sample distribution condition of 17 binary classification datasets and our final experiment results on these datasets. Table 5 illustrates the number of significant genes with  $p$ -value less than 0.05 in the *t*-test. From Table 5 we can note that,  $p=0.05$  is

a relatively accommodative condition on 17 binary-class datasets, which can not only identify the most differentially expressed genes, but also avoid the inappropriate exclusion of too many genes.

## S7. COMPARE OTHER FILTER METHODS WITH THE *t*-TEST AND MIC COMBINATION

In the feature space reduction stage, the study used *t*-test and MIC for their efficiency and convenience in gene filtering process. The *t*-test has been widely used and validated for detecting differentially expressed genes in microarray [5], [6], [7]. But *t*-test has limitation in dealing with multi-class dataset for multivariate *t*-test can't be performed directly. The recently

TABLE 5  
Number of statistically significant features with  $t$ -test-based  $p$ -values less than 0.05 on 17 binary classification datasets.

Dataset	DLBCL	Pros	Colon	Leuk	Mye	ALL1	ALL2	ALL3	ALL4
Significant features	2632	5061	594	2449	1720	4387	644	571	1279
Total features	7129	12625	2000	7129	12625	12625	12625	12625	12625
Dataset	CNS	Lym	Adeno	Gas	Gas1	Gas2	T1D	Stroke	
Significant features	334	804	1799	8260	16454	15601	10159	5569	
Total features	7129	4026	7457	22645	22283	22283	54675	54675	

TABLE 6  
The performance comparison among different filter method combinations by 5-fold cross validation

Filter Methods	Dataset	Genes	$S_n$	$S_p$	$Acc$	$Avc$	$MCC$	$AUC$
$t$ -test+MIC	Adeno	1	1.0	1.0	1.0	1.0	1.0	1.0
	Gas1	3	0.986	0.973	0.980	0.980	0.961	0.99
	Pros	4	0.980	0.982	0.981	0.981	0.963	0.98
	DLBCL	3	1.0	1.0	1.0	1.0	1.0	1.0
	Leuk	2	1.0	1.0	1.0	1.0	1.0	1.0
	CNS	7	1.0	1.0	1.0	1.0	1.0	1.0
Anova+FC	Adeno	2	1.0	1.0	1.0	1.0	1.0	1.0
	Gas1	3	0.987	0.973	0.980	0.980	0.960	0.979
	Pros	4	0.980	0.982	0.981	0.981	0.963	0.982
	DLBCL	3	1.0	1.0	1.0	1.0	1.0	1.0
	Leuk	4	1.0	1.0	1.0	1.0	1.0	1.0
	CNS	8	1.0	1.0	1.0	1.0	1.0	1.0
Volcano plot+MIC	Adeno	1	1.0	1.0	1.0	1.0	1.0	1.0
	Pros	4	0.980	0.982	0.980	0.981	0.963	0.968
	DLBCL	3	1.0	1.0	1.0	1.0	1.0	1.0
	Leuk	2	1.0	1.0	1.0	1.0	1.0	1.0

proposed MIC shows excellent performance in detecting a wide range of associations in large datasets including microarray [1], [8], and MIC can cope with multi-class dataset. Thus, we combined  $t$ -test and MIC to complete the feature screen task.

For the execution order, we perform the  $t$ -test first and then MIC. By the  $p=0.05$  significance threshold in the  $t$ -test, we can quickly find the statistically significant genes, which could notably reduce the gene feature range. Besides, it has been noticed that the MIC calculation is kind of time-consuming compared with  $t$ -test, thus it is suitable to perform  $t$ -test first to decrease the gene number.

We also compared the performance of other filter method combinations with the  $t$ -test+MIC combination, the result are shown in Table 6.

Firstly, the combination of first Anova then Fold change (FC), Anova+FC. The experiment was carried on 3 balanced datasets (Adeno, Gas1 and Pros) and 3 imbalanced datasets (DLBCL, Leuk, and CNS) using 5-fold cross validation. For Anova, the  $p$ -value threshold

was also set as 0.05 as in  $t$ -test. From Table 6, it can notice that with the combination of Anova+FC, the sizes of finally selected genes are 2, 4, and 8 on datasets Adeno, Leuk and CNS, respectively. But by the  $t$ -test+MIC combination, simply 1, 2 and 7 genes are needed to achieve the same performance on the 3 datasets. On the rest of 3 datasets, the two filter method combinations have similar performance. Thus, the filter method combination of Anova+FC is little inferior to the combination of  $t$ -test+MIC in finding the minimal discriminative gene subset.

Secondly, the combination of first "volcano plot" then MIC, Volcano plot+MIC. The "volcano plot" can combine the advantages of  $t$ -test and fold-change, thus we use the "volcano plot" to replace the  $t$ -test. The experiments were performed on 2 balanced datasets (Adeno and Pros) and 2 imbalanced datasets (DLBCL and Leuk) by 5-fold cross validation. According to experiment results in Table 6, these two methods select same size of genes and achieve similar performance on all the tested 4 datasets. In the experiments, we

noted one defect of “volcano plot” for gene selection in a range of different microarray datasets. When we use the “volcano plot” to selected informative genes, in each microarray dataset, we need to hand-tune the  $p$ -value threshold in  $t$ -test and fold-change threshold value in FC to obtain the satisfactory result. For example, on dataset Adeno, there are 894 informative genes have FC value larger or equal to 2; but on dataset Pros, the highest FC value in all genes is just 1.46. Thus, for different datasets, the threshold values in “volcano plot” should be different. In fact, for each of the tested 4 datasets, the threshold values in “volcano plot” have been hand-tuned individually and the finally assigned threshold values vary among different datasets. This situation pose difficulty for building automatically application on a wide range of microarray datasets. In contrast, the  $t$ -test has the consistent  $p$ -value setting in all the 17 datasets in experiments and shows more convenience.

To conclude, the filter method combination of  $t$ -test+MIC are more efficient and convenient than the Anova+FC or Volcano plot+MIC combinations for the feature range reduction task in our study.

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