Using a Feature Selection Ensemble on DNA Microarray Datasets

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Abstract. DNA microarray has brought a difficult challenge for researchers due to the high number of gene expression contained and the small samples size. Therefore, feature selection has become an indispensable preprocessing step. In this paper we propose an ensemble for feature selection based on combining rankings of features. The individual rankings are combined with different aggregation methods, and a practical subset of features is selected according to a data complexity measure –the inverse of Fisher discriminant ratio–. The proposed ensemble, tested on seven different DNA microarray datasets using a Support Vector Machine as classifier, was able to obtain the best results in different scenarios.

1 Introduction

In recent years, the real-world scenarios have incremented considerably their dimension and size. Specifically, DNA microarray experiments generate a lot of gene expressions for a low number of patients. In DNA microarray data, every sample corresponds with a patient and each feature represents a gene expression coefficient corresponding to the abundance of *Messenger Ribonucleic Acid* (mRNA) in a concrete sample. An important application of DNA microarray data is to separate healthy patients from cancer patients based on their gene expression "profile". The DNA microarray datasets pose an enormous challenge for feature selection researchers due to the high number of gene expression contained and the small samples size.

Several studies have shown that most genes in a DNA microarray experiment are not relevant for an accurate classification among different classes of the problem [1]. To reduce the data dimensionality, feature selection preprocess plays a crucial role in DNA microarray analysis. This process obtains an ideal subset of relevant features of the data, eliminating irrelevant and redundant information. As a result, the dimension of the datasets is reduced, allowing a lower use of the storage size and an improvement of the computational time of machine learning algorithms. In feature selection, there are two different approaches when conducting the evaluation of the features of a dataset: (i) *individual evaluation* and (ii) *subset evaluation* [2]. In the first case, a ranking

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of features is returned by assigning a level of relevance to each of these features. The feature selection methods with this evaluation approach are also known as *ranker methods*. In the second case, successive subsets of features are generated and evaluated iteratively, according to an optimality criterion, until reaching the final subset of selected features.

Machine learning methods have traditionally used a single learning model to solve a given problem. However, it has been recently observed that by combining different learning models, better results could be obtained (called *ensemble learning*). Ensemble learning has been normally applied to classification process but can also be thought as a means of improving other machine learning disciplines such as feature selection.

The idea of this paper is to use an ensemble of ranker methods instead of a single method. Seven different ensemble configurations will be presented, depending on the combination method used to generate the final ranking. Since the ensemble is formed by ranker methods, a threshold is necessary to obtain a practical subset of features. The novelty of the approach presented in this work is that we use complexity measures to automatically establish this threshold, in particular we employ the inverse of Fisher discriminant ratio. The results obtained in a previous work on ensembles for feature selection [3] are taken as baseline to compare the results of the proposed approach. An extensive experimentation on different DNA microarray datasets, using a Support Vector Machine (SVM) as a classifier, shows the adequacy of the proposed ensemble.

2 Ensemble feature selection

Datasets can be very large, both in number of samples or in number of features, and can also be redundant, noisy, multivariate and nonlinear. In order to make a correct choice, a user not only needs to know the domain and the characteristics of each dataset well, but also is expected to understand technical details of available algorithms [4]. In this sense, a possible way to confront this situation is to use an ensemble of feature selection algorithms. Specifically, in this study we use ranker methods, i.e. they return an ordered ranking of all the features. Thus, a threshold is necessary in order to obtain a practical subset of features. In this work, instead of choosing an arbitrary threshold, we have opted for the use of a data complexity measure to establish the threshold value automatically and tailored for the dataset, since they are a recent proposal to represent characteristics of the data which are considered difficult in classification tasks beyond estimates of error rates [5]. We assume that good candidate features would contribute to decrease the complexity and must be maintained. Figure 1 shows the proposed approach, and the pseudo-code can be seen in Algorithm 1.

Among the broad suite of feature selection methods available in the literature, four well-known filter methods were chosen to conform the final ensemble (*Chi-Square* [6], *Information Gain* [7], *mRMR* [8] and *ReliefF* [9]). The A_r rankings generated using the different feature selection methods, all of them using the same training data, must be combined in order to produce a unique final output ESANN 2016 proceedings, European Symposium on Artificial Neural Networks, Computational Intelligence and Machine Learning. Bruges (Belgium), 27-29 April 2016, i6doc.com publ., ISBN 978-287587027-8. Available from http://www.i6doc.com/en/.



Fig. 1: Diagram of the proposed ensemble method.

Al	gorithm	1:	Pseud	lo-code	e of	the	proposed	ensemb	le met	hod
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Data: R — number of ranker methods **Data**: T — number of features to be selected **Data**: V — vector of complexity measure values **Result**: E — classification test error 1 Calculate and store in V the complexity measure value of each feature. ² for each r from 1 to R do Obtaining ranking A_r using feature selection method r3 4 end A =combining rankings A_r with a ranking combination method 5 6 Use V on A to obtain the optimal T value. $\tau A_t =$ Select T top attributes from ABuild classifier SVM-RBF with the selected attributes A_t 8 Obtain test error E

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A. The combination methods, also known as 'aggregators', are responsible for conducting the fusion of several rankings in order to obtain a final ranking. To perform the combination of the input rankings several different measures are used, ranging from simple calculation to more sophisticated measures. The different ranking combination functions selected to combine the several rankings in this study were: SVM-Rank [10] (a SVM-based method for learning of ranking functions), the Min, Median, Mean and GeomMean methods [11] (based on simple arithmetic operations) and the Stuart [12] and RRA [13] methods (based on statistical sorting distributions).

Finally, we have opted for the use of a data complexity measure to establish the threshold value T to obtain a practical subset of features A_t . In this paper, the measure selected was the Fisher discriminant ratio f [5], defined as:

$$f = \frac{\sum_{i=1, j=1, i \neq j}^{c} p_i p_j (\mu_i - \mu_j)^2}{\sum_{i=1}^{c} p_i \sigma_i^2},$$
(1)

where μ_i , σ_i^2 , and p_i are the mean, variance, and proportion of the *i*th class *C*, respectively. The Fisher discriminant ratio values are calculated for each feature of the dataset individually. The formula applied to establish the threshold and obtain the final subset of features is defined as:

$$e[v] = \alpha \times 1/f + (1 - \alpha) \times \rho \tag{2}$$

where α is a parameter with value in the interval [0, 1] ($\alpha = 0.75$ for this work), ρ is the percentage of features retained (ranging from one to the total number of features of the dataset) and 1/f is the inverse of the Fisher discriminant ratio. A small complexity value e[v] represents an easier problem.

3 Experimental study

The experiments performed on seven DNA binary microarray datasets, which are listed in Table 1, consisted of a comparison between the different ensemble configurations described in the Section 2, taking as baseline the methods used in a previous study [3]. The datasets originally divided into training and test sets were maintained, whilst, those with only training set were randomly divided using the common rule 2/3 for training and 1/3 for testing for the sake of comparison. This division introduces a more challenging scenario since, in some datasets, the distribution of the classes in the training set differs from the one in the test set. Table 1 shows the number of attributes and samples and also the distribution of the binary classes, indicating if the data is unbalanced.

Dataset	Features	Samples		Train distribution $(\%)$	Test distribution (%)	
		Train	Test			
Colon	2000	42	20	67 - 33	60 - 40	
DLBCL	4026	32	15	50 - 50	53 - 47	
CNS	7129	40	20	65 - 35	65 - 35	
Leukemia	7129	38	34	71 - 29	59 - 41	
Lung	12533	32	149	50 - 50	90 - 10	
Prostate	12600	102	34	49 - 51	26 - 74	
Ovarian	15154	169	84	35 - 65	38 - 62	

Table 1: Binary microarray datasets employed in the experimental study.

A Support Vector Machine (SVM) with a Radial Basis Function (RBF) kernel has been chosen for comparing the proposed ensemble configurations with the baseline methods in terms of classification error. The results obtained by the different approaches are displayed in Table 2. This table shows the test classification error of different methods and is divided in two parts: (i) the first seven rows represent the different ensemble configurations (E-AggregatorMehod) described in this paper (Section 2), and (ii) the last six rows correspond with the baseline methods (obtained from [3]) consisting of an ensemble method (E2) and five individual filter methods.

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Ranker	Colon	DLBCL	CNS	Leukemia	Lung	Prostate	Ovarian
E-SVMRank	20.00 (5)	6.67 (5)	35.00 (5)	20.59 (5)	8.05 (5)	26.47 (10)	14.29 (5)
E-Min	15.00 (10)	13.33 (5)	40.00 (5)	8.82 (5)	0.67 (5)	26.47(5)	0.00 (5)
E-Median	15.00 (5)	13.33(5)	35.00 (5)	11.77 (5)	3.36(5)	26.47 (5)	0.00 (5)
E-Mean	20.00 (5)	13.33(5)	35.00 (5)	8.82 (5)	3.37(5)	26.47 (5)	0.00 (5)
E-GeomMean	20.00 (5)	13.33(5)	25.00 (5)	8.82 (5)	3.36(5)	26.47 (5)	0.00 (5)
E-Stuart	20.00(5)	13.33(5)	30.00 (5)	11.77(5)	5.37(5)	26.47(5)	0.00 (5)
E-RRA	20.00 (5)	13.33(5)	35.00(5)	14.71(5)	5.37(5)	26.47 (5)	1.19(5)
E2	20.00 (34)	6.67 (47)	35.00 (60)	11.76 (36)	1.34 (40)	2.94 (89)	0.00 (37)
CFS	15.00 (19)	6.67 (47)	35.00 (60)	11.76(36)	1.34(40)	2.94 (89)	0.00 (37)
Cons	20.00(3)	13.33(2)	35.00 (3)	20.59(1)	18.12(1)	73.53 (4)	0.00 (3)
INT	15.00 (16)	13.33(36)	40.00 (47)	11.76(36)	1.34(40)	29.41(73)	0.00 (27)
InfoGain	15.00 (25)	6.67 (25)	35.00(25)	11.76(25)	0.67 (25)	2.94 (25)	1.19(25)
ReliefF	15.00 (25)	6.67 (25)	30.00(25)	17.65(25)	2.01 (25)	5.88(25)	0.00 (25)

Table 2: Test classification error for DNA microarray datasets. The value shown in parenthesis represents the number of relevant features selected by each method. The best error for each dataset is highlighted in bold face.

The experimental results demonstrate the adequacy of the proposed ensemble, since they match or improve upon the results obtained in [3]. In fact, it can be seen that some of the proposed ensemble configurations (E-Min, E-Mean and *E*-GeomMean) improve the baseline error results on the *Leukemia* dataset, and the E-GeomMean configuration improves the baseline error results on the CNS dataset. In addition to this, at least one ensemble configuration matches the best baseline error when it is applied on the datasets Colon, DLBCL, Lung or Ovarian. In both cases, the proposed ensemble has the added benefit of reducing significantly the dataset dimension (using only five features, in most configurations, for the classification process). The *Prostate* dataset is the only one that does not improve their results when the proposed ensemble is applied. Notice that the *Prostate* dataset has been a big challenge for machine learning algorithms since the training and test sets were extracted from different experiments. Thus, the common assumption that the training and test data follow the same distribution is, in this case, violated. In fact, in a previous work [14] we have pointed out that some classifiers just assign all the samples to the minority class, leading to a poor classification performance around 26% global accuracy, as happens with our proposed ensemble.

4 Conclusions and discusion

DNA microarray data has brought a difficult challenge for researchers due to the high number of gene expression contained and the small samples size. For these reason, a feature selection preprocess is essential to confront the dimensionality problem. In this paper, an ensemble for feature selection was presented. The idea is to use an ensemble of methods rather than a single method, in order to take advantage of their individual strengths and overcome their weak points at the same time. This will have the added benefits of releasing the user from the task of knowing technical details about the DNA microarray scenario.

The particularity of the proposed ensemble is that it works with ordered

rankings of features, and therefore, a threshold is necessary in order to obtain a practical subset of features. Four well-known algorithms were chosen to form part of the ensemble and the individual rankings were combined with different combination methods. Since we have an ordered ranking of all the features we have opted for the use of data complexity measures to establish the threshold value, releasing the user from the task of choosing it in advance. In this study, the inverse of Fisher discriminant ratio was selected as data complexity measure. The experiments on seven DNA microarray datasets showed that our proposal was able to obtain competitive results when compared with results achieved in a previous work, with the added benefit of selecting automatically the threshold to establish the final number of features to consider in the classification stage.

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