COMBINING NEURAL NETWORK AND SUPPORT VECTOR MACHINE INTO INTEGRATED APPROACH FOR BIODATA MINING

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Abstract: Bioinformatics is the science of managing, mining, and interpreting information from biological sequences and structures. In this paper, we discuss two data mining techniques that can be applied in bioinformatics: namely, Neural Networks (NN) and Support Vector Machines (SVM), and their application in gene expression classification. First, we provide description of the two techniques. Then we propose a new method that combines both SVM and NN. Finally, we present the results obtained from our method and the results obtained from SVM alone on a sample dataset.

1 INTRODUCTION

Data mining is a process that uses a variety of data analysis tools to discover patterns and relationships in data that may be used to make valid predictions. The continuous progress in data mining research has led to developing various efficient methods for mining patterns in large databases. Data mining approaches include: Neural Networks, Support Vector Machines, Evolutionary Programming, Memory Based Reasoning, Decision Trees, Genetic Algorithms, and Nonlinear Regression Methods.

Data classification is a process that groups data in categories possessing similar characteristics. The classification process involves refining each group by defining its shared characteristics. Data analysis is becoming the bottleneck in gene expression classification. Data integration is necessary to cope with an ever increasing amount of data, to cross-validate noisy data sets, and to gain broad interdisciplinary views of large biological data sets. Noise and disparities in experimental protocols strongly limit data integration. Noise can be caused by systematic variation, experimental variation, human error, and variation of scanner technology, variation in which biologists are not interested.

Another issue with gene classification is, currently available databases typically contain low number of instances, though each instance quantifies the expression levels of several thousands of genes. Due to the high dimensionality and the small sample size of the experimental data, it is often possible to find a large number of classifiers that can separate the training data perfectly, but their diagnostic accuracy on unseen test samples is quite poor and different.

According to the mentioned problems we may conclude that the choice of machine learning technique selection is the most important aspect in classifying gene expression. Based on this, in this paper we study and compare two approaches to deal with this problem: namely NN and SVM. We proposed a novel approach which integrates the advantages of both for better biodata mining. Experimental results reported on a sample dataset demonstrate the effectiveness and applicability of our approach.

The rest of this paper is organized as follows. Section 2 is a brief overview of neural networks. Section 3 presents a short coverage of SVM. Section 4 is dedicated to feature extraction. Section 5 includes experimental results. Section 6 discusses the results. Section 7 is the conclusions.
2 NEURAL NETWORKS

A Neural Network (NN) is an information-processing paradigm inspired by the way biological nervous systems, such as the brain, process information. Neural networks are made up of a number of artificial neurons. An artificial neuron is simply an electronically modeled biological neuron. How many neurons are used depends on the problem we are trying to solve. Figure 1 represents a picture of a neuron in a neural network. Each neuron accepts a weighted set of inputs and responds with an output.

The real power of neural networks comes when we combine neurons in multi-layer structures. Figure 2 represents a sample neural network. The number of nodes in the input layer corresponds to the number of inputs and the number of nodes in the output layer corresponds to the number of outputs produced by the neural network. When the network is used, the input variable values are placed in the input units, and then the hidden and output layer units are progressively executed. Each of them calculates its activation value by taking the weighted sum of the outputs of the units in the preceding layer, and subtracting the threshold. The activation value is passed through the activation function to produce the output of the neuron. When the entire network has been executed, the outputs of the output layer act as the output of the entire network.

Once the number of layers and number of units in each layer has been selected, the network's weights and thresholds must be set so as to minimize the prediction error made by the network. This is the role of the training algorithms. The error of a particular configuration of the network can be determined by running all the training cases through the network, comparing the actual output generated with the desired or target outputs. The differences are combined together by an error function to give the network error.

3 SUPPORT VECTOR MACHINES

The support vector machine (SVM) algorithm (Boser et al., 1992; Vapnik, 1998) is a classification algorithm that has received a great consideration because of its astonishing performance in a wide variety of application domains such as handwriting recognition, object recognition, speaker identification, face detection and text categorization (Cristianini and Shawe-Taylor, 2000). Generally, SVM is useful for pattern recognition in complex datasets. It usually solves the classification problem by learning from examples.

During the past few years, the support vector machine-learning algorithm has been broadly applied within the area of bioinformatics. The algorithm has been used to detect new unknown patterns within and among biological sequences, which help to classify genes and patients based on gene expression, and has recently been used in several advance biological problems. There are two main motivations that suggest the use of SVM in bioinformatics. First, many biological problems involve high-dimensional, noisy data, and the difficulty of a learning problem increases exponentially with dimension. It has been a common practice to use dimensionality reduction to relief this problem. SVMs use a different technique, based on margin maximization, to cope with high dimensional problems. Empirically, they have been shown to work in high dimensional spaces with remarkable performance. In fact, rather than reducing dimensionality as suggested by Duda and Hart, the SVM increases the dimension of the feature space. The SVM computes a simple linear classifier, after mapping the original problem into a much higher dimension space using a non-linear kernel function. In order to control over fitting in this extremely high-dimensional space, the SVM attempts to maximize the margin characterized by the distance between the nearest training point and the separating discriminant.

Second, in contrast to most machine learning methods, SVMs can easily handle non-vector inputs,
such as variable length sequences or graphs. These types of data are common in biology applications, and often require the engineering of knowledge-based kernel functions.

![Input Space Feature Space](image)

Figure 3: Support vector machine: mapping non-separable data from input space to higher-dimensional feature space, where a separating hyper-plane can be constructed.

![Hyper-plane and margin](image)

Figure 4: Hyper-plane and margin, circular dots and square dots represent samples of class -1 and class +1.

### 4 FEATURE SELECTION

A feature is a meaningful and distinguishing characteristic of a data sample used by a classifier to associate it with a particular data category. The process of selecting or extracting features involves mathematically manipulating the data sample, and producing a useful pattern. The process of selecting or extracting several features to form a feature set is known as feature selection or feature extraction.

In classification tasks, feature selection is often used to remove irrelevant and noisy features as well as producing useful features. The selected feature set can be refined until the desired classification performance is achieved. Thus, manually developing a feature set can be a very time consuming and costly endeavor. In the area of gene classification by feature selection, we are interested in identifying the subset of genes whose expression levels are most relevant for classification or diagnosis.

In the current bioinformatics research, the following three approaches are mostly considered in order to do feature selection in gene expression datasets:

The guiding principle of the first approach is that the features, which can best be used for classification of the tissue sample, should be chosen (Xiong et al, 2001). A consequence of this principle is that one must know exactly how tissue samples will be classified before feature selection can be done. The process of feature selection wraps around the classifier in the following procedure:

1. A candidate set of features is considered.
   a. Tissue samples are divided into training and test sets.
      i. The classifier is trained on the training set of tissue samples.
      ii. The classifier is used on the test set of tissue samples.
   b. Step 1(a) is repeated with alternative divisions into training and test sets.
   c. The candidate feature set is evaluated using all classifications from 1(a)(ii).

2. Step 1 is repeated with another candidate feature set.

In this way, many candidate feature sets are evaluated using the training set of tissue samples, and the feature set that performs best is chosen. A major advantage of the feature wrapper approach is accuracy, because the feature selection is “tuned” for the classification method. Another advantage is that the approach provides some protection against overfitting because of the internal cross validation. Yet another advantage will become apparent when classifiers are employed to distinguish between more than two tissue types, because most feature selection methods used to date have been specific to binary classification. One drawback of feature wrapper methods is that the methods can be computationally intensive.

Filter type methods are essentially data preprocessing or data filtering methods. Features are selected based on the intrinsic characteristics, which determine their relevance or discriminant powers with regard to the targeted classes. In filters, the characteristics in the feature selection are uncorrelated to that of the learning methods; thus, filter methods are independent of the technique for classifier design; they may be used in conjunction with any such algorithm, and they have better generalization property.

In embedded methods, feature selection and classifier design are accomplished jointly. Embedded methods incorporate variable selection as part of the training process and may be more efficient in several respects: they make better use of the available data by not needing to split the training data into a training and validation set; they reach a solution faster by avoiding retraining a predictor from scratch for every variable subset investigated.

SVM RFE improves feature selection based on feature ranking by eliminating the orthogonally assumptions of correlation methods. This method...
allows us to find nested subsets of genes that lend themselves well to a model selection technique that finds an optimum number of genes. RFE method uses the following iterative procedure to eliminate the features:

1. Initialize the data set to contain all features.
2. Train SVM on the data set.
3. Rank features according to criterion c.
4. Eliminate the lowest-ranked feature.
5. If more than one feature remains, return to step 2.

In practice, removing half of the features in step 4 speeds up the algorithm. RFE ranks the features based on their weight learned by SVM. Features are removed one by one (or by chunks); SVM is re-run at each iteration.

Among all minimization algorithms for feature selection using SVMs, RFE has empirically been observed to achieve the best results on classification tasks using gene expression data.

5 INTEGRATED APPROACH AND EXPERIMENTS

For our experiments we used PC with AMD 1900+ CPU, 1GB memory. The experiments were performed on Matlab 6.5 using Neural Network and Spider toolboxes.

In our study, we propose and apply the SVM method of Recursive Feature Elimination (RFE) to gene selection. By using RFE, we eliminated chunks of genes at a time. At the first iteration, we reached the number of genes, which is the closest power of 2. At subsequent iterations, we eliminated half of the remaining genes. We thus obtained nested subsets of genes of increasing informative density. Using Neural Network as a classifier then assessed the quality of these subsets of genes. We propose this method based on the following arguments:

• If it is possible to identify a small set of genes that is indeed capable of providing complete discriminatory information, inexpensive diagnostic assays for only a few genes might be developed and be widely deployed in clinical settings.
• Knowledge of a small set of diagnostically relevant genes may provide important insights into the mechanisms responsible for the disease itself.
• In the cases in which datasets have large number of features, Neural Networks show their limits on running time and space complexity. They require polynomial time and storage space with higher degree comparing to Support Vector Machines. And in practice, even 3000-feature-dataset exceeds the limit of computational power for matrix in Matlab. Therefore, some approach with lower time and space complexity like SVM is introduced in the preprocessing of classification as feature reduction tool. As a result, we are able to use Neural Networks to classify datasets that are generally too big, and cannot be handled by the Neural Network in their original form.

In the experiment, the general three-layer model of Neural Network is used as the main classifier. There are sixty neurons in each layer with their weights randomly initialized. For the input layer and hidden layer, the Tan-Sigmoid is chosen as transfer function, which redistributes input from previous layer to next layer at the range of [-1, 1]. And the Log-Sigmoid is used as transfer function on the final output layer and determines the result -- positive/negative, true/false or class#1/class#0. Conjugate Gradient Back propagation with Powell-Beale Restarts (CGB) is the training algorithm for the network because of its decent performance and short running-time.

To test our approach we downloaded several published biological datasets. The datasets used in the experiments consist of matrix of gene expression vectors obtained from DNA micro-arrays. We will provide short description for each dataset.

The first dataset was obtained from cancer patients with two different types of leukemia. The problem is to distinguish between two variants of leukemia (ALL and AML). Distinguishing between ALL and AML is critical because the two types of leukemia require different treatment. The dataset consists of 72 samples (47 ALL vs. 25 AML) over 7129 probes from 6817 human genes.

Table 1: Results obtained by using 50/50 ratio for training data

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Highest Accuracy</th>
<th>Lowest Accuracy</th>
<th>Num Features</th>
<th>Test vs Train (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALL/AML</td>
<td>0.9722</td>
<td>0.9444</td>
<td>60</td>
<td>50/50</td>
</tr>
<tr>
<td>1</td>
<td>0.9722</td>
<td>50</td>
<td>50/50</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.9722</td>
<td>40</td>
<td>50/50</td>
<td></td>
</tr>
<tr>
<td>Central Nervous System</td>
<td>0.83</td>
<td>0.60</td>
<td>60</td>
<td>50/50</td>
</tr>
<tr>
<td>0.83</td>
<td>0.67</td>
<td>50</td>
<td>50/50</td>
<td></td>
</tr>
<tr>
<td>0.83</td>
<td>0.72</td>
<td>40</td>
<td>50/50</td>
<td></td>
</tr>
<tr>
<td>Colon Tumor</td>
<td>0.9677</td>
<td>0.8710</td>
<td>60</td>
<td>50/50</td>
</tr>
<tr>
<td>0.9032</td>
<td>0.7419</td>
<td>50</td>
<td>50/50</td>
<td></td>
</tr>
<tr>
<td>0.871</td>
<td>0.7097</td>
<td>40</td>
<td>50/50</td>
<td></td>
</tr>
<tr>
<td>Lung Cancer</td>
<td>0.989</td>
<td>0.956</td>
<td>60</td>
<td>50/50</td>
</tr>
<tr>
<td>1</td>
<td>0.989</td>
<td>50</td>
<td>50/50</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.9231</td>
<td>40</td>
<td>50/50</td>
<td></td>
</tr>
</tbody>
</table>

Note: Number of Neurons for each NN is 60
The purpose of the second dataset is to analyze the outcome of the treatment. It contains a total of 60 instances. The samples are classified in two classes. Survivors - patients that responded to the treatment and Failures – patients that did not benefit from the treatment. The dataset consists of 21 survivors and 39 failures samples. There are 7129 genes in the dataset.

The third data set contains 62 samples collected from colon-cancer patients. Among them, 40 tumor biopsies are from tumors and 22 biopsies are from healthy parts of the colons of the same patients. Two thousand out of around 6500 genes were selected based on the confidence in the measured expression levels.

Our fourth data set is used for classification between malignant pleural mesothelioma (MPM) and adenocarcinoma (ADCA) of the lung. It consists of 181 tissue samples (31 MPM vs. 150 ADCA) and each sample is described by 12533 genes.

### Table 2: Comparison between our method and SVM

<table>
<thead>
<tr>
<th></th>
<th>Average Rate SVM+NN</th>
<th>Average Rate SVM</th>
<th>Num Features for NN</th>
<th>Train vs. Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALL/AML</td>
<td>0.9353</td>
<td>0.9361</td>
<td>60</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>0.9861</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central Nervous System</td>
<td>0.645</td>
<td>0.6681</td>
<td>60</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>0.65</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Central Nervous System</td>
<td>0.22</td>
<td>0.6166</td>
<td>60</td>
<td>70</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Colon Tumor</td>
<td>0.80645</td>
<td>0.8080</td>
<td>60</td>
<td>70</td>
</tr>
<tr>
<td></td>
<td>0.75</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lung Cancer</td>
<td>0.96155</td>
<td>0.9922</td>
<td>60</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>0.96155</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: Number of Neurons for each NN is 60

### Table 4: Results obtained from increasing the training data set size to 60% of the data

<table>
<thead>
<tr>
<th></th>
<th>Highest Accuracy Rate</th>
<th>Lowest Accuracy Rate</th>
<th>Num Features</th>
<th>Ratio Test vs Train (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALL/AML</td>
<td>1</td>
<td>0.9310</td>
<td>60</td>
<td>60/40</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>0.9655</td>
<td>50</td>
<td>60/40</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>0.9655</td>
<td>40</td>
<td>60/40</td>
</tr>
<tr>
<td>Central Nervous System</td>
<td>0.7083</td>
<td>0.6667</td>
<td>60</td>
<td>60/40</td>
</tr>
<tr>
<td></td>
<td>0.7917</td>
<td>0.6250</td>
<td>50</td>
<td>60/40</td>
</tr>
<tr>
<td>Central Nervous System</td>
<td>0.72</td>
<td>0.6250</td>
<td>40</td>
<td>60/40</td>
</tr>
<tr>
<td>Colon Tumor</td>
<td>0.9231</td>
<td>0.8077</td>
<td>60</td>
<td>60/40</td>
</tr>
<tr>
<td></td>
<td>0.9231</td>
<td>0.8077</td>
<td>50</td>
<td>60/40</td>
</tr>
<tr>
<td>Lung Cancer</td>
<td>0.9863</td>
<td>0.9231</td>
<td>60</td>
<td>60/40</td>
</tr>
<tr>
<td></td>
<td>0.9863</td>
<td>0.9231</td>
<td>50</td>
<td>60/40</td>
</tr>
</tbody>
</table>

Note: Number of Neurons for each NN is 60

### 6 THE RESULTS

To normalize the Neural Network training set we used the formula:

$$\text{normData} = \frac{2 \times (\text{data} - \min(\text{data}))}{(\max(\text{data}) - \min(\text{data}))} - 1$$

where $\text{normData}$ – normalized value, $\text{data}$ – current value, $\min(\text{data})$ – minimum value in the corresponding column, $\max(\text{data})$ – maximum value in the corresponding column.

The formula normalizes dataset entries in the range $[-1, 1]$. Then we mapped the two class instances to 0 and 1 and then to 0.05/0.95 in order to use the Log sigmoid transfer function in Neural Network for the output layer. The initial number of features for all datasets was too big for the Neural Network. As a result, we applied the proposed method and reduced the number of features to 60, 50, and 40 for the different trials.

In addition, we applied a permutation on the samples in our datasets. The motivation for such step was to obtain an even distribution of the two classes that are going to be classified in both training and testing sets. Our experiments have shown that if that condition is not satisfied the performance of the Neural Network degrades. This was due to the fact that during the training process the Neural Network has not seen enough samples from both classes.

To test our network, we divided the datasets in the training and testing part. The ratios used in different trials are specified in the results. Finally, we implemented a program in Matlab that creates 5 different Neural Networks in each trial and outputs the best results. The accuracy rate that we are providing is computed as the ratio of the correct predictions over the total predictions made by the Neural Network. For example, for ALL/AML
dataset using 50/50 ratio for test and train set, the accuracy rate of 0.9722 that out of 36 samples 35 were correctly classified and only one was predicted wrong.

7 CONCLUSIONS

As we can see in the results the method we have proposed in this paper achieves high success rate. In some trials we obtained a rate as high as 100%. Our method allows using Neural Networks for datasets that are too large in their original form and the Neural Network is not able to handle the input data. As a result, we can apply this approach for problems where it is important to minimize the empirical risk and the use of a Neural Network is desirable over SVM classifier.

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