Cancer Prognosis Prediction Using SVM for Hybrid Type and Imbalanced Data Sets

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Abstract: Cancer prognosis is one of the hot spots in the study of biological information. There have been many studies to cancer prognosis prediction using machine learning methods, which have achieved better results. Among them, the support vector machine (SVM) gets extensive attention as it is suitable to apply in small-size, high-dimensional data classification questions. However, SVM only performs well in the case where the class distribution is balanced and the input variables are numerical which are unlikely occurred in the medical domain. So in this study, we introduce a new prognosis prediction method based on SVM, which modify the standard SVM models to fit imbalanced class distribution and hybrid type of features. In details, firstly the similarity of features with nominal and numerical type is redefined in kernel function. Secondly synthetic minority oversampling technique (SMOTE) method is adopted to balance class distribution. Lastly the wrapper method SVM-RFE is introduced to select the useful features to improve the prediction performance. A series of experiments are designed and launched to validate the performance. The results have proved the effectiveness of the proposed methods.

1 INTRODUCTION

Cancer is a major disease threatening human health. Traditionally there are three common treatments, including surgery, chemotherapy and radiation therapy. So far the selection of different treatment is based on the TNM stage of the patient, that is using tumor size, location etc. to select the appropriate treatment. However, there may be a big difference that even the patients with the same TNM stage choose the same treatment. For example, the current standard of treatment for patients with early-stage Non-Small-Cell Lung Cancer(NSCLC) is surgical resection. According to the statistical data, the 5-year survival rate is only 65-70% even in those patients whose tumors have undergone complete resection(Zhu, 2009). So it is very significant to develop an accurate means of prognosis which can predict the survival time of each individual patient after surgical operation based on the characteristics of patient beside TNM stage information. With the development of molecular biological medicine and application of molecular biological diagnostic techniques, the study of tumor pathology from the molecular level has become a research hot spot, such as the use of molecular characteristics to make a prognosis.(Xu, 2016) Due to the complexity of the molecular characteristics, machine learning have absolute advantages compared with traditional statistical methods.

Machine learning methods have been successfully applied to the analysis of many different complex problems in recent years, including many bio-medical applications.(Jayasurya, 2010) The work of (Street, 1995) used a self-organizing neural network to find classes of cases with similar expected recurrence times. D. Chakraborty, U. Maulik (2014) approach prognosis as separation problem. The work of (Zhu, 2009) used SVM for prognosis in Early-Stage Non-Small-Cell Lung Cancer(NSCLC), which plotted Kaplan-Meier survival curve to prove the effectiveness of its method.

Although many different prognosis prediction methods have been proposed, these methods have poor performance in case of hybrid type and imbalanced data sets. For several cancer datasets, on the one hand, are consist of feature attributes which are likely prepossessed to hybrid type, numerical and nominal respectively ; on the other hand, the class distribution is imbalanced, which will be confronted with great difficulties in classical machine learning method. To solve this problem, we propose a novel...
prognosis prediction method based on nonlinear classification, which can be implemented using SVM. We modified the standard SVM module to fit the test data, which have the imbalanced data set and hybrid types of feature. We believe that better individual predictions may be obtained.

The rest of the article is organized as follows. In Section 2, we firstly give a brief introduction to the data for Analysis and the method adopted in the following experiments. Section 3 describes the experiments and discussion followed by conclusion in Section 4.

2 MATERIAL AND METHOD

2.1 Data for Analysis

- **NSCLC**

  The data from a series of 196 patients with early stage NSCLC was adopted in this study (Zhu, 2009). The prognosis data set is consisted of three parts, Tissue microarrays (TMA), clinical and pathologic variables. The process of construction of TMA was described in (Rosenwald, 2002). Thirty three molecular markers were chosen for investigation in the study. All of these markers have been identified in previous studies with the best candidates for the prognostic prediction (Duan, 2005). All marks of each patient are assigned to a score by three pathologists based on the number and the intensity of stained tumors cells. Furthermore, four clinical and pathologic variables are also used, including sex (male vs. female), age, cancer-cell type and tumor diameter. Thus each patient is represented by a vector of 37 dimension and the outcome is described as survival time after surgery. When patients are classified as two classes, “died before 3 years” or “not died in 3 years” after therapy, the classification distribution is faced with imbalanced data set with 150 patients died before 3 years against 46 not died in 3 years.

- **WPBC**

  We also use Wisconsin Prognostic Breast Cancer (WPBC) in which has 198 instances with 34 attributes (ID, outcome, 32 real-valued input features) in each instances. The first 30 features are computed from a digitized image of a fine needle aspirate (FNA) of a breast mass, they are described as characteristics of the cell nuclei present in the image. The last 2 features, which are tumor size and lymph node status, are observed at time of surgery, the values for features are re-coded with four significant digits. At last the outcome is described as recurrence time and disease-free time if outcome is recurrence or it is no-recurrence. The class distribution is faced with imbalanced data set with 151 non-recurrences against 47 recurrence in (Street, 1995).

  The two data sets above have the following characteristics:

  1) The training sets are small-size with redundant features. Specifically, in NSCLC, there are only 196 samples with a vector of 37 dimensions in each sample, while in WPBC, there are 198 samples with a vector of 34 dimensions in each sample. In one hand, some features may be irrelevant to the prognosis of the patient, in the other hand, the statistics show a high correlation between some features. Existence of redundant features will decrease the performance of methods.

  2) Hybrid type is existed in each sample expressed by a vector, with nominal and numerical type respectively. Therefore traditional classification method is hard to work perfectly.

  3) The class distribution is imbalanced, with the ratio of 3:1 roughly. The traditional data mining algorithm behaves undesirable in the instance of imbalanced data sets. The class boundary can be severely skewed towards the majority class which means nothing to the application.

  4) They are more likely to be non-linear classification problems. It is difficult to find the relationship between attributes and the outcome.

  From the characteristics mentioned above, the following method and strategies will be adopted in improving SVM approach to fit the datasets.

2.2 Support Vector Machine

SVM is a useful technique for data classification. A classification task usually involves separating data into training and testing sets. Each instance in the training set contains one “target value” (usually called class labels) and several “attributes”. The goal of SVM is to produce a model (based on the training data) which predicts the target values of the test data given only the test data attributes.

Given a training set of instance-label pairs \((x_i, y_i)\), \(i = 1, ..., l\) where \(x_i \in R^n\) and \(y \in \{1, -1\}\), the SVM requires the solution of the following optimization problem:

\[
\min_{w, \xi, \xi^*} \frac{1}{2} w^T w + C \sum_{i=1}^{l} \xi_i
\]

subject to:

\[
y_i (w^T x_i + b) \geq 1 - \xi_i, \quad i = 1, ..., l
\]

\[
\xi_i \geq 0, \quad i = 1, ..., l
\]
Subject to \[ y_i (w^T \phi(x_i) + b) > 1 - \varepsilon_i , \quad \varepsilon_i \geq 0 \] (1)

Function \( \phi(x) \) is used to map the training vectors into a higher dimensional space. SVM finds a linear separating hyper-plane with the maximal margin in this higher dimensional space. \( C > 0 \) is the penalty parameter of the error term.

The above optimization problem (1) can be used in a linear classification problem. But classification problem is non-linear generally. In order to solve a nonlinear classification problem, SVM first maps the training data to another dot product space (called the feature space) \( \mathcal{F} \) via a nonlinear map, thus converting the non-linear problem in the original space to linear problems in the \( \mathcal{F} \)-space. Gaussian RBF kernel function is among the most commonly used kernels, which is defined as follows:

\[ K(x_i, x_j) = \exp(-g \| x_i - x_j \|^2), \quad g > 0 \] (2)

After the kernel function is applied, the above optimization problem can be expressed by kernel function, which is defined as follows:

\[
\max_{\alpha} \quad w(\alpha) = \sum_{i=1}^{m} \alpha_i - \frac{1}{2} \sum_{i=1}^{m} \sum_{j, i}^{m} y_i y_j \alpha_i \alpha_j K(x_i, x_j) \\
\text{Subject to} \quad \sum_{i=1}^{m} \alpha_i y_i = 0 \quad \text{and} \quad 0 \leq \alpha_i \leq C \] (3)

2.3 Modified Kernel function for hybrid types

The kernel function in (2) is based on the Euclidean distance between two pairwise samples. However, when nominal features are faced, this kind of kernel cannot be worked directly. The similarity of two vector with nominal type data cannot defined as Euclidean distance. For example, if one part of the vector means the location of the cancer, such as 1,2,3, then we cannot find that 3 is far away from 1, and it is near 2. So a new definition of distance between two vectors should be given. In details, the distance between 1 and 3 is equal to the distance between 2 and 3. In conclusion, if the part of a vector is nominal, then the distance is a constant which can be predefined in the beginning when the number is not equal, while the distance is zero when the number is equal.

So the distance of two samples with nominal feature can be redefined as follows:

\[
\| x_i - x_j \|^2 = \sum_{k=1}^{n} (x_i^k - x_j^k)^2 \\
\begin{cases} x_i^k - x_j^k = 1 & \text{if } k \text{ is nominal} \\ x_i^k - x_j^k = 0 & \text{otherwise} \end{cases} \] (4)

where \( x^k \) is nominal.

2.4 Feature selection

In our case of NSCLC, 37 features are used where some features may be irrelevant to the prognosis of the patient. Accordingly, we should find ways to select the useful features to improve the prediction performance as well as reduce the computational complexity.

The existing feature selection algorithms can be generally categorized as wrapper or filter methods based on criterion functions used in searching for informative features. In wrapper methods, the performance of a learning algorithm is employed to evaluate the goodness of selected feature subsets, whereas in filter methods, criterion functions evaluate feature subsets by their information content, typically measure the correlation of feature subsets and outcome, having nothing to do with learning algorithms. In most case, filter methods are computationally much more efficient but perform worse than wrapper methods (Xu, 2012).

In this paper, the wrapper method recursive feature elimination algorithm (RFE) is adopted and SVM classifier is used as a criterion function. The SVM-RFE method is a sequential backward selection process. It starts with a set of all features, and the least important feature for the classification is removed from the whole feature set iteratively according to criteria function which is the sum square of the weight vector \( W \) of all features in support vectors in SVM model (Chakraborty, 2014).

It should be noted that the combination of the features ranked ahead can obtain better performance than the single one. Therefore, SVM-RFE algorithm can select the combination of identification features.

2.5 Strategies for imbalanced data set

Considering the training instances of one class are significantly outnumbered by those of the other class, the class boundary learned by standard SVM can be severely skewed towards the majority class. As a result, the false-negative rate can be excessively high, which may cause the classifier ineffective. Taking account of the collection of each sample is not easy to get, in order to get a balanced samples, the synthetic minority oversampling technique (SMOTE) is used to address this problem instead of under-sampling majority class samples.

SMOTE method is a kind of oversampling technique proposed by Chawla (Wu, 2003). The main idea of SMOTE is artificially generate virtual samples to minority class to over-sample the minority class. For each minority sample, its k (which is set to 5 in SMOTE) nearest neighbors of the same class are calculated, then some examples are randomly selected from them according to the over-sampling rate. After that, new synthetic samples are generated.
along the line between the minority samples and its selected nearest neighbors. The process can be described in formula (5):

$$x_{\text{new}} = x + rand(0,1) \times (\bar{x} - x)$$  \hspace{1cm} (5)

Where \( rand(0,1) \) denotes random numbers ranging from 0 to 1, \( \bar{x} \) is one of the k-nearest.

SMOTE is suitable for samples with numerical attributes. However, there is a hybrid type in each sample expressed by a vector in the data set mentioned above, with nominal and numerical type respectively. Therefore, we need to modify the formula (5) to fit hybrid types of feature. Suppose each feature \( x \) is consist of two parts \( x = \{x^1, x^2\} \), where \( x^1 \) are of nominal form and \( x^2 \) are of numerical form, the formula (5) can be turned as follows:

$$x_{\text{new}} = rand(x^1) \cup [x^2 + rand(0,1) \times (\bar{x}^2 - x^2)]$$  \hspace{1cm} (6)

Where \( rand(x^1) \) denotes the attribute value which is randomly selected from all of the specific attribute field.

3 EXPERIMENT RESULTS AND ANALYSIS

We approach the prognostic prediction as a classification problem. The following procedure with proposed method is applied to NSCLC data set and WPBC data set mentioned in section 2.

3.1 Experimental procedure

Figure 1 shows the whole experimental procedure and the detail is as following:

5) preprocessing

The standard simple imputation method (mean among the values in the training set) is used to the presence of missing values. In order to avoid attributes in greater numeric ranges dominating those in smaller numeric ranges, each numerical attribute is scaled to range \([0,1]\) when \( y_{\text{max}} = 1 \) and \( y_{\text{min}} = 0 \) by formula (7).

$$f : x \rightarrow y = (y_{\text{max}} - y_{\text{min}}) \times \frac{x - x_{\text{min}}}{x_{\text{max}} - x_{\text{min}}} + y_{\text{min}}$$  \hspace{1cm} (7)

6) Classification

The RBF kernel is used as the number of features is not very large. We use (9) and (10) to validate the performance of classification instead of (8) because (9) and (10) are more suitable for imbalanced data problems.

$$\text{Acc} = \frac{TN + TP}{TN + TP + FN + FP}$$  \hspace{1cm} (8)

$$Sn = \frac{TP}{TP + FN}$$  \hspace{1cm} (9)

$$Sp = \frac{TN}{TN + FP}$$  \hspace{1cm} (10)

In the SVM model building phase, SVM-RFE method is implemented to order the importance of each attribute. We test all subset of collection \( F \) computed by SVM-RFE, and search for the best parameter pair \((C, g)\) with grid-search method iteratively. The best SVM model can be reached with the best accuracy.

Finally, testing dataset is input into the model, after computing, confusion matrix is obtained which can be used to validate the performance of classification.

3.2 Application to NSCLC and WPBC

We approach the prognostic prediction of NSCLC and WPBC data sets as a classification problem with two classes. The main objective is to obtain the best predictive model to separate the two classes with high performance, which can help doctors and patients to make a treatment decision.

In NSCLC data set, the target value means whether a patient can survive more than 3 years or not, that is to say, the category can be defined as true if the patient survives more than 3 years in the last check-up, whereas false if the patient have died in 3 years after surgery. After sorting the patients according to their onset time in NSCLC, front 70% patients (137 patients) are used as training data and remaining 30% patients (59 patients) as testing data.
While in WPBC, we separated “patients who recurred in less than three years” from “patients who were disease free more than three years”, the target value means whether a patient would recurrent in 3 years or not. Front 70% patients are used as training data and the remaining 30% as testing data.

### 3.3 Results and analysis

The first set of experiments is to evaluate the performance of our proposed SMOTE. Table I and II show the average accuracy, sensitivity and specificity whether SMOTE is used or not.

As can be seen from the tables, we get relatively higher training accuracy but lower testing accuracy when SMOTE method is not used. Results significantly biased majority class. The Sn parameter is only 12.50% in NSCLC, 20% in WPBC, which makes the classification of minority class samples meaningless. All features in SVM model are support vectors, which illustrate that over-fitting is very serious.

Two sets of results using SMOTE method demonstrate that when using testing samples, Sn indicators has been greatly improved, and classification accuracy is more balanced than no use of SMOTE, which is displayed with Sn and Sp indicators , in NSCLC tests, are 65.22% and 61.11%, while in WPBC tests, are 74.62% and 86.57% respectively. The experiments validate the effectiveness of SMOTE strategy when faced with imbalanced data set.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Average accuracy whether use SMOTE on NSCLC</th>
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</thead>
<tbody>
<tr>
<td><strong>Strategy</strong></td>
<td>Training set as test samples</td>
</tr>
<tr>
<td></td>
<td>Acc</td>
</tr>
<tr>
<td>No use SMOTE</td>
<td>99.40%</td>
</tr>
<tr>
<td>use SMOTE</td>
<td>77.38%</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Average accuracy whether use SMOTE on WPBC</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Strategy</strong></td>
<td>Training set</td>
</tr>
<tr>
<td></td>
<td>Acc</td>
</tr>
<tr>
<td>No use SMOTE</td>
<td>92.39%</td>
</tr>
<tr>
<td>use SMOTE</td>
<td>94.54%</td>
</tr>
</tbody>
</table>

The second set of experiments is launched to evaluate the performance of attribute selection. Training datasets are used to evaluate the importance of attributes by using SVM-RFE. Grid-search is employed to find out the best parameter pair(C,g) so as to obtain the best subset of features with the highest accuracy. Then we can use the best parameter pair(C,g) and the best subset of features to train the SVM classification model. The accuracy in each iteration is calculated and shown in Fig.2.

![Fig 2: The relationship between a subset of features and prediction accuracy of WBPC and NSCLC](image)

From this figure it can be seen that we obtained the optimal attribute subset of WBPC, the front 17 attributes of collection F, with the best accuracy of 80.43%. While worked on NSCLC, we obtained the optimal attribute subset, the front 28 attributes of collection F, with the best accuracy of 68.02%. These results can demonstrate that the feature attribute subset contains more discriminating information that can greatly help prognosis prediction.

### 4 CONCLUSION

Prognosis prediction gives the clinician an unbiased method to predict treatment effect instead of traditional methods based on TNM staging. Machine learning methods have been successfully applied to the field of prognosis prediction. Its potential appears to be extremely promising and is worthy of further research.
In this work, we have proposed a novel prognosis prediction method based on SVM to create personalized predictive models. Two datasets, NSCLC and WPBC, was selected which had small size, high dimensional characteristics.

The novelty of this work is three-fold. Firstly, we have modified the standard RBF kernel function in SVM model to fit the test data, which have hybrid types of feature. This modification makes the model meet the needs of practical application. Secondly, we propose the SMOTE strategy to deal with imbalanced training-data problems. A series of experiments have demonstrated the effectiveness of SMOTE strategy when faced with imbalanced data set. Thirdly, SVM-RFE is employed to extract features collection of greatest impact on outcome. The results demonstrate that with the help of SVM-RFE, 17 out of 34 attributes of WBPC have been selected, and 28 out of 37 attributes of NSCLC have been selected which outperforms the over all attribute collection.

So far, only SVM models have been employed. In the future, we are preparing an extensive set of tests by using other machine learning method, such as random forest, deep learning, in the same manner as the SVM procedure.

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