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Improving Naive Bayes Classifier Using Conditional Probabilities

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Abstract

Naive Bayes classifier is the simplest among Bayesian Network classifiers. It has shown to be very efficient on a variety of data classification problems. However, the strong assumption that all features are conditionally independent given the class is often violated on many real world applications. Therefore, improvement of the Naive Bayes classifier by alleviating the feature independence assumption has attracted much attention. In this paper, we develop a new version of the Naive Bayes classifier without assuming independence of features. The proposed algorithm approximates the interactions between features by using conditional probabilities. We present results of numerical experiments on several real world data sets, where continuous features are discretized by applying two different methods. These results demonstrate that the proposed algorithm significantly improve the performance of the Naive Bayes classifier, yet at the same time maintains its robustness.

Keywords: Bayesian Networks, Naive Bayes, Semi Naive Bayes, Correlation

1 Introduction

Classification is the task to identify the class labels for instances based on a set of features, that is, a function that assigns a class label to instances described by a set of features. Learning accurate classifiers from pre classified data is an important research topic in machine learning and data mining. One of the most effective classifiers is Bayesian Networks (Shafer 1990, Heckerman 1995, Jensen 1996, Pearl 1996, Castillo 1997). A Bayesian Network (BN) is composed of a network structure and its conditional probabilities. The structure is a directed acyclic graph where the nodes correspond to domain variables and the arcs between nodes represent direct dependencies between the variables. Considering an instance $X = (X_1, X_2, ..., X_n)$ and a class C, the classifier represented by BN is defined as

 $\arg \max_{c \in C} P(c|x_1, x_2, ..., x_n) \propto \arg \max_{c \in C} P(c) P(x_1, x_2, ..., x_n | c),$ (1)

where x_i, c are the values of X_i, C respectively.

However, accurate estimation of $P(x_1, x_2, ..., x_n | c)$ is non trivial. It has been proved that learning an optimal BN is NP-hard problem (Chickering 1996, Heckerman 2004). In order to avoid the intractable complexity for learning BN, the Naive Bayes classifier has been used. In the Naive Bayes (NB) (Langley 1992, Domingos 1997), features are conditionally independent given the class. The simplicity of the NB has led to its wide use, and to many attempts to extend it (Domingos 1997). Since NB assumes the strong assumption of independency between features, learning semi Naive Bayes has attracted much attention from researchers (Langley 1994, Kohavi 1996, Paz-zani 1996, Friedman 1997, Kittler 1986, Zheng 2000, Webb 2005). The semi Naive Bayes classifiers are based on the structure of NB, requiring that the class variable be a parent of every feature. However, they allow additional edges between features that capture correlation among them. The main aim in this area of research has involved maximizing the accuracy of classifier predictions.

In this paper, we propose a new version of the Naive Bayes classifier (semi Naive Bayes) without assuming independence of features. The proposed algorithm finds dependencies between features using conditional probabilities. This algorithm is a new algorithm and different from the existing semi Naive Bayes methods (Langley 1994, Kohavi 1996, Pazzani 1996, Friedman 1997, Kittler 1986, Zheng 2000, Webb 2005).

Most of data sets in real world applications often involve continuous features. Therefore, continuous features are usually discretized (Lu 2006, Wang 2009, Ying 2009, Yatsko 2010). The main reason is that the classification with discretization tend to achieve lower error than the original one (Dougherty 1995). We apply two different methods to discretize continuous features. The first one, which is also the simplest one, transforms the values of features to $\{0, 1\}$ using their mean values. We also apply the discretization algorithm using sub-optimal agglomerative clustering algorithm from (Yatsko 2010) which allows us to get more than two values for discretized features. This leads to the design of a classifier with higher testing accuracy in most data sets used in this paper.

We organize the rest of the paper as follows. We give a brief review to the Naive Bayes and some semi Naive Bayes classifiers in Section 2. In Section 3, we present the proposed algorithm. Section 4 presents an overview of the discretization algorithm using sub-optimal agglomerative clustering. The numerical experiments are given in Section 5. Section 6 concludes the paper.

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2 Naive Bayes and Semi Naive Bayes Classifiers

The Naive Bayes (NB) assumes that the features are independent given the class, it means that all features have only the class as a parent (Kononenko 1990, Langley 1992, Domingos 1997, Mitchell 1997). A sample of the NB with *n* features is depicted in Figure 1. The NB, classifies an instance $X = (X_1, X_2, ..., X_n)$ using Bayes rule, by selecting

$$\arg\max_{c\in C} P(c) \prod_{i=1}^{n} P(x_i|c).$$
⁽²⁾

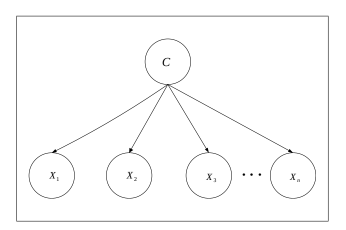


Figure 1: Naive Bayes

NB has been used as an effective classifier for many years. Unlike many other classifiers, it is easy to construct, as the structure is given a priori. Although the independence assumption is obviously problematic, NB has surprisingly outperformed many sophisticated classifiers, especially where the features are not strongly correlated (Domingos 1997). In spite of NB's simplicity, the strong independency assumption harms the classification performance of NB when it is violated. On the other hand, learning BN requires searching the space of all possible combinations of edges which is NP-hard problem (Chickering 1996, Heckerman 2004).

In order to relax the independence assumption of NB, a lot of effort has focussed on improving NB. The improved NB classifiers use exhaustive search to join features based on statistical methods. There are some improved algorithms of the NB. Langley and Sage (Langley 1994) considered Backwards Sequential Elimination (BSE) and Forward Sequential Selection (FSS) in which their methods select a subset of features using leave-one-out cross validation error as a selection criterion and establish a NB with these features. Starting from the full set of features, BSE successively eliminates the features whose elimination most improves accuracy, until there is no further accuracy improvement. FSS uses the reverse search direction, that is iteratively adding the features whose addition most improves accuracy, starting with the empty set of features. The work of Pazzani (Pazzani 1996) introduces Backward Sequential Elimination and Joining (BSEJ). It uses predictive accuracy as a merging criterion to create new Cartesian prod-uct features. The value set of a new compound fea-tures is the Cartesian product of the value sets of the two original features. As well as joining features, BSEJ also considers deleting features. BSEJ repeatedly joins the pair of features or deletes the features

that most improves predictive accuracy using leaveone-out cross validation. This process terminates if there is no accuracy improvement. Kohavi (Kohavi 1996) proposed the NB Tree, a strategy that is a hybrid approach combining NB and decision tree learning. It partitions the training data using a tree struc-ture and establishes a local NB in each leaf. It uses 5-fold cross validation accuracy estimate as the splitting criterion. A split is defined to be significant if the relative error reduction is greater than 5 percent and the splitting node has at least 30 instances. When there is no significant improvement, NB Tree stops the growth of the tree. As the number of splitting features is greater than or equals one, NB Tree is an x-dependence classifier. The classical decision tree predicts the same class for all the instances that reach a leaf. In NB Tree, these instances are classified using a local NB in the leaf, which only considers those non tested features. Friedman et al. (Friedman 1997) introduced Tree Augment Naive Bayes (TAN) based on tree structure. It approximates the interactions between features by using a tree structure imposed on the NB structure. In TAN, each feature has the class and at most one other feature as parents. Super Parent algorithm is proposed by Keogh and Pazzani (Keogh 1999). This algorithm uses the same representation as the Tree Augment Naive Bayes, but utilizes leave-one-out cross validation error as a criterion to add a link. The Super Parent is the feature that is the parent of all the other orphans, the features without a non-class parent. There are two steps to add a link: first selecting the best Super Parent that improves accuracy the most, and then selecting the best child of the Super Parent from orphans. This method stops adding links when there is no accuracy improvement. Zheng and Webb (Zheng 2000) developed Lazy Bayesian Rules (LBR), which adopts a lazy approach, and generates a new Bayesian rule for each test example. The antecedent of a Bayesian rule is a conjunction of feature-value pairs, and the consequent of the rule is a local NB, which uses those features that do not appear in the antecedent to classify. LBR stops adding feature value pairs into the antecedent if the outcome of a one tailed pairwise sign test of error difference is not better than 0.05. As the number of the feature value pairs in the antecedent is greater than or equals one, LBR is anx-dependence classifier. Webb et al. (Webb 2005) proposed Averaged One Dependence Estimators (AODE), which averages the predictions of all qualified 1-dependence classifiers. In each 1-dependence classifier, all features depend on the class and a single feature.

In the next section, we introduce a new version of the Naive Bayes classifier (semi Naive Bayes) without assuming independence of features. The proposed algorithm approximates the interactions between features by using conditional probabilities.

3 The Proposed Algorithm

In this section, we present a new algorithm that maintains the basic structure of the NB, and thus ensure that the class C is the parent of all features. The proposed algorithm, however, removes the strong assumption of independence in the NB by finding correlation between features, while also capturing much of the computational efficiency of the NB. In this algorithm, the class has no parents and each feature has the class and at most one other feature as parents. Therefore, each feature can have one augmenting edge pointing to it. The procedure for learning these edges is based on the Pearson's correlation and conditional

probabilities. First, we construct a basic structure of the NB with n features $X_1, X_2, ..., X_n$ from the set X and the class C. After that, we find the Pearson's A and the class C. After that, we find the Featson's correlations between each feature X_i and the class C using the formula (3), $Corr(X_i, C)$. Then we reorder the set X as a set X^* in a descending order of $|Corr(X_i, C)|$. In the ordered set X^* , an arc from the first feature is added to the second one. Finally, for all remain features, we find the conditional probabilities of each feature with the previous features given the class values in the ordered set X^* , formula (4). The highest value of these conditional probabilities between features is used to recognize the parent of each feature. The conditional probabilities described in (4), first introduced by Quinn et al. (Quinn 2009) and called influence weights, have been used directly for data classification. However, here, we used them for finding the dependencies between features.

The correlation coefficient (Graham 2008) between two random variables X_i and X_j is defined as :

$$Corr(X_i, X_j) = \frac{N \sum_{i,j=1}^{N} X_i X_j - \sum_{i=1}^{N} X_i \sum_{j=1}^{N} X_j}{\sqrt{(N \sum_{i=1}^{N} X_i^2 - (\sum_{i=1}^{N} X_i)^2)(N \sum_{j=1}^{N} X_j^2 - (\sum_{j=1}^{N} X_j)^2)}}$$
(3)

where N is the number of data points. This measure has the property of $|Corr(X_i, X_j)| \leq 1$. When this value is close to 1, it denotes the perfect linear cor-relation between X_i and X_j , and $Corr(X_i, X_j) = 0$ stands for no linear correlation.

The proposed algorithm consists of six main steps:

Algorithm. Proposed Algorithm

Step 1. Construct a basic structure of the Naive Bayes with n features, $X = \{X_1, X_2, ..., X_n\}, \text{ and the class } C.$

Step 2. Compute the correlation between each feature X_i , i = 1, ..., n and the class C using the formula (3), $Corr(X_i, C)$.

Step 3. Reorder X as a set $X^* = \{X_1^*, X_2^*, ..., X_n^*\}$ in a descending order of $|Corr(X_i, C)|, i = 1, ..., n.$

Step 4. Add an arc from X_1^* to X_2^* .

Step 5. For j = 3, ..., n: 5.1 Find X_i^* that has the highest value of

$$\sum_{k=1}^{N} |P(X_{ki}^*, X_{kj}^*|C) - P(X_{ki}^*, X_{kj}^*|\overline{C})|, \ i < j, \quad (4)$$

where $X_i^* = (X_{1i}^*, X_{2i}^*, ..., X_{Ni}^*)^T$, N is the number of instances and $\overline{C} = -C$. 5.2 Add an arc from X_i^* to X_j^* .

Step 6. Compute the conditional probability tables inferred by the new structure.

Figure 2 shows the structure of Symguide1 data set, taken from LIBSVM, with four features (see Table 1) using the proposed algorithm. The solid lines are those edges required by the Naive Bayes classifier. The dashed lines are correlation edges between features found by our algorithm.

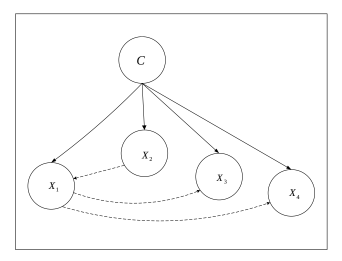


Figure 2: Proposed algorithm, Svmguide1

$\mathbf{4}$ Discretization Algorithm Using Sub-Optimal Agglomerative Clustering (\overline{SOAC})

Discretization is a process which transform continuous numeric values into discrete ones. In this paper, we apply two different methods to discretize continuous features. The first one, which is also the simplest one, transforms the values of features to 0,1 using their mean values. We also apply the discretization algorithm using sub-optimal agglomerative clustering algorithm which allows us to get more than two values for discretized features. In this section, we introduce discretization algorithm SOAC which is an efficient discretization method for the NB learning. Details of this algorithm can be found in (Yatsko 2010).

Consider a finite set of points A in the n dimensional space R^n , that is $A = \{a^1, ..., a^m\}$, where $a^i \in R^n$, i = 1, ..., m. Assume that the sets $A^{j}, j = 1, ..., k$ be clusters, and each cluster A^{j} can be identified by its centroid $x^j \in \mathbb{R}^n$, j = 1, ..., k. The discretization algorithm SOAC proceeds as follows.

Algorithm. Discretization Algorithm SOAC

Step 1. Set k = m, and a small value of parameter θ , $0 < \theta < 1$. Sort values of the current feature in the ascending order. Each feature requiring discretization is treated in turn. Step 2. Calculate the center of each cluster:

$$x^{j} = \sum_{a \in A^{j}} \frac{a}{|A^{j}|}, \ j = 1, ..., k$$

and the error E_k of the cluster system approximating set A:

$$E_k = \sum_{j=1}^k \sum_{a \in A^j} \|x^j - a\|^2$$

Step 3. Merge in turn each cluster with the next tentatively. Calculate the error increase after each merge $E_{k-1} - E_k$ and choose the pair of clusters giving the least increase. Merge these two clusters permanently. Set k = k - 1.

Step 4. Once the error of the current cluster system is over the set fraction of the maximum error corresponding to the single cluster $E_k \geq \theta E_1$ stop, otherwise go to Step 2.

5 Numerical Experiments

To verify the efficiency of the proposed algorithm, numerical experiments with a number of real world data sets have been carried out. We use 10 real world data sets. The detailed description of the data sets used in this experiments can be found in the UCI machine learning repository, with the exception of "Fourclass", "Svmguide1" and "Svmguide3". These three data sets are downloadable on tools page of LIBSVM. A brief description of data sets is given in Table 1. We discritize the values of features in data sets using two different methods. In the first one, we apply a mean value of each feature variable to discritize the values to $\{0, 1\}$. The second one is the discrization algorithm SOAC (Yatsko 2010) which is presented in Section 4.

We conduct empirical comparison for the NB and the proposed algorithm in terms of test set accuracy using two different discritization methods. The results of the NB and the new algorithm on each data set were obtained via 1 run of 10-fold cross validation. Runs were carried out on the same training sets and evaluated on the same test sets. In particular, the cross validation folds were the same for all experiments on each data set.

The test set accuracy obtained by the NB and the proposed algorithm on 10 data sets using mean values for discretization summarized in Table 2. The results presented in this table demonstrate that the test set accuracy of the new algorithm is much better than that of the NB. The proposed algorithm works well in that it yields good classifier compared to the NB. Its performance was further improved by introducing some additional edges in the NB, using conditional probabilities. Improvement is noticed mainly in large data sets. In 8 cases out of 10, the new algorithm has higher accuracy than the NB. The accuracy of this algorithm is same with the NB in data sets Fourclass and Svmguide1.

Table 3 presents the test set accuracy obtained by the NB and the proposed algorithm on 10 data sets using discretization algorithm SOAC. The results from this table show that the accuracy obtained by the new algorithm in all data sets are higher than those obtained by the NB.

Figures 3 to 4 show the scatter plot comparing the proposed algorithm with the NB, using two different discritization methods. In these plots, each point represents a data set, where the x coordinate of a point is the percentage of miss classifications according to the NB and the y coordinate is the percentage of miss classification according the proposed algorithm. Therefore, points above the diagonal line correspond to data sets where the NB performs better, and points below the diagonal line correspond to data sets where the proposed algorithm performs better.

According to the results explained above, the proposed algorithm outperforms the NB, yet at the same time maintains its robustness. However, the proposed algorithm requires more computational effort than the NB since we need to compute conditional probabilities between features to recognize the parent of each feature in our algorithm.

Table 1: A brief description of data sets

Data sets	# Features	# Instances
Congres Voting Records	16	435
Credit Approval	14	690
Diabetes	8	768
Fourclass	2	862
Haberman Survival	3	306
Heart Disease	13	270
Phoneme CR	5	5404
Spambase	57	4601
Svmguide1	4	7089
Svmguide3	21	1284

Table 2: Test set accuracy of NB and the proposed algorithm using mean value for discretization

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Data Sets	Naive Bayes	Proposed Algorithm
Congres Voting Records	90.11	91.47
Credit Approval	84.85	86.85
Diabetes	75.78	77.68
Fourclass	76.82	76.82
Haberman Survival	74.51	75.66
Heart Disease	84.14	85.18
Phoneme CR	75.96	78.30
Spambase	90.13	93.45
Svmguide1	92.17	92.17
Svmguide3	80.61	87.18

Table 3: Test set accuracy of NB and the proposed algorithm using discretization algorithm SOAC

Data Sets	Naive Bayes	Proposed Algorithm
Congres Voting Records	90.11	91.47
Credit Approval	84.85	86.85
Diabetes	75.78	77.68
Fourclass	78.58	79.70
Haberman Survival	74.66	75.33
Heart Disease	78.62	79.31
Phoneme CR	77.01	79.36
Spambase	89.30	92.30
Svmguide1	95.61	97.54
Svmguide3	77.25	80.85

6 Conclusion

In this paper, we have developed the new version of the Naive Bayes classifier without assuming independence of features. An important step in this algorithm is adding edges between features that capture correlation among them. The proposed algorithm finds dependencies between features using conditional probabilities. We have presented the results of numerical experiments on 10 data sets from UCI machine learning repository and LIBSVM. The values of features in data sets are discritized by using mean value of each feature and applying discretization algorithm SOAC. We have presented results of numerical experiments. These results clearly demonstrate that the proposed algorithm significantly improve the performance of the Naive Bayes classifier, yet at the same time maintains its robustness. Furthermore, this improvement becomes even more substantial as the size of the data sets increases.

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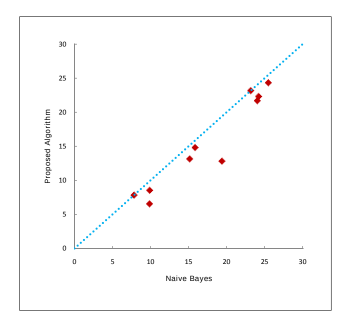


Figure 3: Scatter plot comparing miss classifications of the proposed algorithm (y coordinate) with Naive Bayes (x coordinate); using mean value for discritization

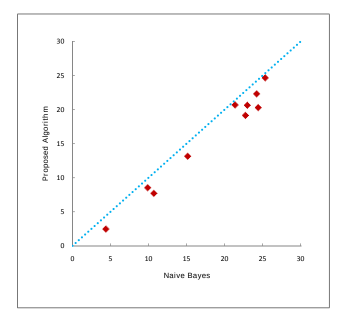


Figure 4: Scatter plot comparing miss classifications of the proposed algorithm (y coordinate) with Naive Bayes (x coordinate); using Algorithm SOAC for discritization