Naive Bayes Classifier with Mixtures of Polynomials

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Abstract: We present in this paper a methodology for including continuous features in the Naive Bayes classifier by estimating the density function of the continuous variables through the Mixtures of Polynomials model. Three new issues are considered for this model: i) a classification oriented parameter estimation procedure ii) a feature selection procedure and iii) the definition of new kind of variable, to deal with those variables that are in theory continuous, but their behavior makes the estimation difficult. These methods are tested with respect to classical discrete and Gaussian Naive Bayes, as well as classification trees.

1 INTRODUCTION

Classification problems are present in all science fields, and deal with many real-world problems everyday. That is why they have caught the attention of the statistics, artificial intelligence or machine learning community. In a classification problem there is a qualitative (discrete) variable, called class, whose value we want to predict given some other variables, called features. If the class value is known for a data sample, it is called supervised classification, but if it is unknown, it is called unsupervised classification or clustering. We focus in this paper on supervised classification. There are several techniques available that try to solve this problem, as for example neural networks, classification trees, logistic regression, or probabilistic graphical models. Bayesian networks (BNs) (Pearl, 1988; Jensen and Nielsen, 2007) are a specific type of probabilistic graphical models that are widely used in classification, mainly due to their mixture of model-complexity and accuracy results (Friedman et al., 1997). In fact there is a wide variety in the BN-models for classification that can be used, ranging from the simplest one, called Naive Bayes (NB) (Minsky, 1963), to the most complex one, a BN with no restrictions. The election of the appropriate model depends on the complexity of the problem and the data available. However, the NB model is probably the most used model, because of its simplicity and good results (Domingos and Pazzani, 1997). Originally, BN models were defined only for categorical variables, however in the last decades several frameworks have emerged for dealing with hybrid BNs, in which discrete and continuous variables coexist. The main goal of this paper is to present a new NB-classifier based on the Mixtures of Polynomials (MoP) model, which we will note by NB-MoP, able to deal simultaneously with discrete and continuous features. This model was already presented in (López-Cruz et al., 2013), however it was introduced as an example of application of a more general model. We include in this paper some improvements, such as a feature selection procedure, a specific classification-oriented parameter estimation scheme, and the definition of a pseudo-continuous type of variables in order to make the most of the ability of the model to work with a hybrid set of variables.

Section 2 presents the basics of BNs in classification, Section 3 presents two preprocessing steps to apply in the learning procedure, Section 4 presents the MoP model and the parameter estimation algorithm designed specifically for classification. Section 5 shows an experimental evaluation of the proposed algorithms, and finally in Section 6 conclusions and future work are outlined.

2 BAYESIAN NETWORKS FOR CLASSIFICATION

A BN is composed by two components: a qualitative part and a quantitative part:

1. The qualitative part is a directed acyclic graph in
which every node represents a random variable, and the presence of an edge between two variables shows a dependence relation between them.

2. The quantitative part is the definition of a set of conditional probabilities, in particular one for each variable given its parents in the graph, \( p(x_i | pa(x_i)) \), \( \forall i = 1 \ldots n \).

BNs have traditionally been used only for discrete data. In the presence of continuous data, most of the researchers and practitioners discretize the data, using some common techniques such as \textit{equal width, equal frequency} or \textit{k-means} (Dougherty et al., 1995), more sophisticated algorithms that dynamically discretize the distributions according to the evidence (Kozlov and Koller, 1997), or using specific algorithms for classification problems that relate every continuous feature variable to the class variable (Fayyad and Irani, 1993). However, discretizing the data imply some loss of information and cause problems of accuracy, so alternative solutions have been proposed in the literature. The first approach is to define the continuous variables as Gaussians, which leads to the Conditional Gaussian model (Lauritzen and Vermunt, 1989; Cowell et al., 1999). It is the alternative solution most widely used, however it has some disadvantages, as for example that discrete nodes cannot have continuous nodes as parents in the graph, and that the distribution of the continuous variables must be Gaussian, which does not hold for every continuous variable. Following this idea appears the Mixtures of Truncated Exponentials model (MTE) (Morales et al., 2007), in which any probability distribution can be approximated and there is no restriction in the topology of the network. This model has been successfully applied in recent years to classification and regression problems (Aguilera et al., 2010; Fernández and Salmerón, 2008; Morales et al., 2007) and in general to hybrid BNs (Romero et al., 2006). The Mixtures of Polynomials model (MoP) (Shenoy and West, 2011; Shenoy, 2011) is similar, in the sense that it can also approximate any probability distribution, but uses polynomials as the basis functions, gaining fitting power just increasing the degree. They have mostly been applied to approximate specific known distributions, as in (Shenoy et al., 2011), but not to include them as a tool to approximate any dataset. There are some recent publications that deal with this problem using different properties of the polynomials (Langseth et al., 2013; López-Cruz et al., 2013).

For classification problems it is usual to restrict the structure of the network in order to obtain simpler but effective models. The simplest BN model for classification is the NB model, in which all the features are assumed to be independent given the class (see Fig. 1). Even though this assumption is not always true, NB is known to be an accurate classifier with a relatively small computational complexity. The lack of independence affects to the estimation of the probability of the classes, but the class with maximum probability is still correct (Domingos and Pazzani, 1996). Some other Bayesian classifiers have been proposed, such as the Tree Augmented Network model (Friedman et al., 1997), the kdB model (Sahami, 1996) or the FAN model (Lucas, 2002) but all of them increase the complexity of the model adding more links to the structure.

NB was first proposed to deal only with discrete variables, however, there are some approaches in which continuous variables are allowed as features, representing them as Gaussian distributions (Domingos and Pazzani, 1997), kernel density estimations (John and Langley, 1995; Pérez et al., 2009) or MTEs (Morales et al., 2007). Since MoPs are a novel and efficient way to represent and compute with continuous variables in BNs, the aim of this paper is to introduce this representation framework in the NB model together with some preprocessing steps in order to improve its applicability.

3 PREPROCESSING STEPS

Prior to the specific procedure of estimating the parameters of the network, we will also investigate in this paper the effect of two different pre-processing methods, a classical filter-wrapper feature selection, and some partial discretization of the variables.

3.1 Discretization

The discretization of continuous variables is usual in classification tasks, but it means that some information from the dataset will be lost.

Nevertheless, there are some variables that cannot be considered as discrete neither continuous because of the great number of factors or the difficulty for fitting a polynomial. In this paper we have considered three kinds of features:

![Figure 1: Structure of Naive Bayes classifier.](image_url)
• Discrete: Character or string features as well as numeric features that have less than 20 different values.
• Pseudo-Continuous: numeric features which have more than 20 different values but less than the 5% of the total number of observations of the dataset.
• Totally-Continuous: numeric features which are not pseudo-continuous

Pseudo-Continuous variables are mainly found in large databases (more than 400 observations). For instance, in a database composed by 1000 observations, a variable will be pseudocontinuous if it has between 20 and 50 different values (which is the 5% of 1000).

In the Adult dataset there are two examples of what we have called pseudo-continuous variables. They are capital-gain and capital-loss. The number of different values is 118 and 90, respectively, and, more importantly, 91 and 95% of those values, respectively, are zeroes, due to the fact that they represent the purchases or sales that a person does in one year, which frequently are none. Usually these variables are processed as continuous, however such a variable is difficult to represent by means of a continuous distribution.

Keeping those three kinds of variables in mind, we have designed two different alternatives for the NB-MoP algorithm and tested them in the experiments section:

• Process each variable as it originally is, discrete variables are processed as discrete and continuous variables, both pseudo-continuous and totally-continuous, as continuous.
• Process discrete and pseudo-continuous variables as discrete, i.e., discretize those variables that are not clearly neither discrete nor continuous.

In the last case the discretization procedure used is the Fayyad-Irani method (Fayyad and Irani, 1993).

3.2 Feature Selection

It is known that selecting the appropriate feature variables in a classification problem may lead to an increase of accuracy in the model, as well as to avoid overfitting and noise in the model (Dash and Liu, 1997). A filter-wrapper approach for feature selection has been implemented, in which, firstly, the mutual information between the class and each variable is computed, and different models are learned, starting with one feature variable and adding variables following the order of the previous ranking, until no improvement in terms of accuracy is achieved.

The mutual information is defined as

\[
MI(X;C) = \sum_{i=1}^{n} \sum_{j=1}^{m} p(x_i, c_j) \log \frac{p(x_i, c_j)}{p(x_i)p(c_j)}
\]

in the case of \( X \) being a discrete feature variables, and as

\[
MI(X;C) = \sum_{j=1}^{m} \int_{\Omega} f(x, c_j) \log \frac{f(x, c_j)}{f(x)p(c_j)}
\]

in the case of \( X \) being a continuous feature variable where \( f(x) \) is obtained using a MoP.

Note that, a filter-wrapper approach as this one is not properly a pre-processing step, since it is performed during the learning of the model, not before. However, we decided to include it in this Section for a better reading of the paper.

4 LEARNING THE MODEL

Given the structure of the NB, to learn this model it is needed to estimate the probability distribution of the class, and the conditional distribution of every feature variable given the class.

These conditional distributions will be a conditional probability table (CPT) in the case of discrete features, and a MoP density function for each value of the class variable, in the case of continuous features.

Once the model is learnt, it is able to predict the class value \( c_i \) of a given observation \( x = (x_1, \ldots, x_n) \) using the following formula:

\[
e^* = \arg\max_{c_i \in \Omega_c} p(c_i | x)
\]

where each \( p(c_i | x) \) is computed as

\[
p(c_i | x) = p(c_i | x_1, \ldots, x_n) \propto p(c_i) \prod_{j=1}^{n} p(x_j | c_i)
\]

In the following sections the process of estimating a MoP density from data will be deeply explained.

4.1 Mixtures of Polynomials

The Mixtures of polynomials (MoPs) framework is able to approximate any continuous distribution by a piecewise function which has in each piece a polynomial function. This allows to work directly with the continuous variables without the need of discretizing them. A MoP function is defined as follows (Shenoy and West, 2011):

Definition 1. A one-dimensional function \( f : \mathbb{R} \rightarrow \mathbb{R} \) is said to be a mixture of polynomials function if it is a piecewise function of the form

\[
f(x) = \begin{cases} a_0 + a_1 x + \cdots + a_m x^n & \text{for } x \in A_i, \ i = 1, \ldots, k \\ 0 & \text{otherwise} \end{cases}
\]
where \(A_1,\ldots,A_k\) are disjoint intervals in \(\mathbb{R}\) and \(a_0,\ldots,a_n\) are real constants for all \(i\).

There are some variants to this definition, in (Shenoy, 2011) Shenoy et al re-defines the MoP function to include multivariate functions in which \(A_1,\ldots,A_k\) may be also hyper-rhombuses. This re-definition makes MoP functions richer because now they can deal with deterministic functions, however it makes computations more complex (Shenoy et al., 2011; Rumí et al., 2012). We do not use this variant because, in the NB model, there is no need of estimating multivariate continuous MoPs.

The main reason for using MoPs is that they have a great fitting power, as we can see in Fig. 2 in which we show a Gaussian distribution and the corresponding MoP approximation, according to the parameter estimation procedure explained in Section 4.2.

### 4.2 Learning Mixtures of Polynomials from Data

In previous papers, MoPs were designed to approximate known distributions, such as Gaussian or Chi-square, taking advantage of some properties of the functions, like symmetry or inflection points (López-Cruz et al., 2013). In this paper we design a methodology for learning MoP distributions from a sample (dataset), without knowing the true underlying distribution. This approach is similar to the one shown in (Langseth et al., 2013), but different in some details and focused on a posterior application to the NB model.

To obtain a univariate MoP density from a dataset we have applied the mean squared error (MSE) parameter estimation procedure, by means of the lm method implemented in software R (R Core Team, 2013). An outline of the most important steps is:

1. Input: A sample \((x_1,\ldots,x_n)\) from variable \(X\).
2. Output: a function \(f: \mathbb{R} \rightarrow \mathbb{R}\) expressed as in Def. 1.
3. Obtain the pairs \((x_i^*,y_i^*)\), \(i = 1,\ldots,m\) where \(x_i^*\) is a value in the support set of \(X\) and \(y_i^*\) is the associated density value estimated through a kernel density estimator (Simonoff, 1996).
4. Set the degree to \(i = 1\)
5. Estimate the parameters of the polynomial of degree \(i\) \((p_i(x))\) by minimizing the MSE in the pairs \((x_i^*,y_i^*)\).
6. \(p_i(x)\) may be negative in some points of the domain. Select only the most important positive part, in terms of weight or size (detailed in Section 4.4)
7. Normalize the polynomial so that its integral is equal to the proportion of points of the sample included in the selected subinterval.
8. If a part of the domain of \(p_i(x)\) has been removed because of the negative values, add the necessary tails to extend the domain of the polynomial to include the whole domain of \(X\) (in this case, a piecewise function is obtained as seen in Section 4.4)
9. Compute the corresponding MSE
10. If \(i < 15\) then \(i := i + 1\) and repeat from step 5.
11. Select the best out of the 15 different polynomials, according to the criteria explained in Section 4.4.

Using this procedure, we end up with a MoP following Def. 1, that may be defined in pieces (at most in three), and at most with degree 15 (in the central piece, and 0 in the tails, if there are any). See Section 4.4 for more details.

We will briefly expand some of the steps of the above algorithm.

### 4.3 Kernel Density Estimator

To be able to apply the MSE procedure, we need as input data a grid of \(x^*\) and \(y^*\) points (Step 3). They are obtained through a kernel density estimator with the reflection boundary effect correction (Schuster, 1985). Initially 500 kernel points equally spaced are selected¹ and the domain is then divided in 10 subintervals of equal length. These points are filtered in such a way that more importance is given to those subintervals of the domain with more sample-frequency. In those intervals that actually contain less than 10% sample, the number of kernel points is reduced until they represent the same proportion as the sample-frequency (these points are chosen to be also equally spaced within the correspondent interval). The selection of the number of initial kernel points (500) was done after trying several other sizes.

In Fig. 3, it can be seen an example of the kernel points selection. In this case we used the variable \textit{hours-per-week} from the dataset \textit{Adult} (see Section 5 for more information). In the first plot it is shown an equally spaced griddedata representing the kernel density estimations for that variable, while in the second plot the points are selected following the method described above, that is the reason why there some subintervals of the domain with fewer data. In this way, some subintervals with small sample frequency will have less importance in the polynomial fitting than other subintervals with a great proportion of sample points.

¹The use of more points has been dismissed because its lack of influence in the results
4.4 Parameter Estimation

Once we have the points $x_1^*, \ldots, x_m^*$ and $y_1^*, \ldots, y_m^*$ obtained from $X$, we can calculate the polynomial that best fits the grid-data through minimizing the MSE (Step 5). However, this procedure has two main disadvantages: the degree of the polynomial has to be set in advance and the returned function may have negative values.

The solution for the first problem is to design an iterative procedure (see Sect. 4.2) in which several different degrees are checked.

The solution for the second problem is to keep only the positive part of the function, omitting the negative one (Step 6). Also, an increase in the degree of the polynomial implies an increase in the number of roots, which may lie within the domain of the variable. Two different approaches were observed in order to select the subinterval of the domain to keep:

1. The most distant roots, i.e., the longest interval that can be considered included in the minimum and maximum value of $X_i$ for which the function is positive.

2. The couple of roots which include the higher proportion of points $x_i^*, y_i^*$ between them, for which the function is positive.

It is usual that the longest interval also includes the major number of points, however there are some exceptions in which there are lots of points in a small interval and the rest are disperse.

(Step 7) It is necessary to normalize it so that the integral of the function between the roots is 1. If $r_{\min}$ and $r_{\max}$ are the selected roots, the polynomial $p_{\text{norm}}$ is:

$$p_{\text{norm}}(x) = \frac{p(x) \times \text{prob}}{\int_{r_{\min}}^{r_{\max}} p(x)}$$

where prob stands for the proportion of points of the sample in $(r_{\min}, r_{\max})$.

(Step 8) After defining the initial domain $(r_{\min}, r_{\max})$ where the polynomial $p_{\text{norm}}(x)$ will be defined, it is necessary to extend it to include the removed negative parts. This means to include (if necessary) one or two tails to $p_{\text{norm}}(x)$. The probability of each tail will be the proportion of sample points that lie on it ($p_j$, $j = 1, 2$). This way $p(x)$ will become an actual density function. Let $p_1$ is the probability of the left side of $r_{\min}$ and $p_2$ the probability of the right side of $r_{\max}$, and let range$1 = r_1 - \text{min}(x)$ and range$2 = \text{max}(x) - r_2$ the MoP will be:

$$p(x) = \begin{cases} p_{\text{norm}} & \text{for } x < r_{\min} \\ \frac{p_1}{\text{range}_1} & \text{for } r_{\min} < x < r_{\max} \\ \frac{p_2}{\text{range}_2} & \text{for } x > r_{\max} \end{cases}$$

In Figure 4 there is an example of a polynomial fitted to a sample from Adult dataset, in which we can
see the inclusion of a right tail to the initial MoP estimation.

4.5 Related Work

Some other approaches for estimating parameters in MoPs densities include constraints in the MSE problem in order to assure non-negative functions (López-Cruz et al., 2013; Langseth et al., 2013), however we followed a different approach due to several reasons: i) If the restrictions were included in the algorithm, the calculations would get more complex and the whole algorithm less stable due to the fact that it would be an optimization problem, possibly linked to numerical instability ii) The estimation is carried out splitting the domain and fitting a polynomial in each subinterval, leading probably to a more heavier polynomial, in terms of parameters to estimate iii) Despite the fact that the estimation of the parameters can be appropriate, it is also possible that the approximation in the tails is not so good. In a classification problem as the studied in this paper, it can be a key point, as the choice of the class may depend on a value lying on a tail. That is the reason why we decided to estimate the tails using a piecewise function with a constant for each tail instead of including some constraints in order not to have negative values. In this case, the estimation will be at least as accurate as the discretization.

A MoP parameter estimation based on splines interpolation was used in a NB classifier in (López-Cruz et al., 2013). However, they only deal with totally continuous problems. We focus more on hybrid problems. Also, the R-package provided in (López-Cruz et al., 2013) only estimates the parameters of the MoP distributions, not the NB model. For these two reasons, we do not include this model in the experimental set up.

However, in order to briefly compare this estimation procedure with the one proposed here, the same distribution mentioned in the former Section (see figure 4) was learnt using the algorithm proposed in (López-Cruz et al., 2013) and displayed together with our proposal in Figure 5 and the histogram for the original data. As we can see, the two MoP distributions are quite similar. The B-Splines based MoP is defined in 16 different intervals, with a 3-degree polynomial fitted on each one, requiring a total of 64 parameters to estimate. However, the MoP estimation procedure proposed here yields a MoP density defined on two pieces, one with a 12-degree polynomial fitted on it, and a tail in which a uniform (constant) distribution is fitted, requiring a total of 14 parameters to estimate.

In table 1 we can also observe some statistics for a more detailed comparison between both approaches for this variable.

Table 1: Statistics for comparison between the B-splines based MoP estimation (B-Spline) and our proposal (MoP).

<table>
<thead>
<tr>
<th>Statistic</th>
<th>B-Spline</th>
<th>MoP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>182989.4</td>
<td>184519.6</td>
</tr>
<tr>
<td>1st quartile</td>
<td>170859.4</td>
<td>175907.3</td>
</tr>
<tr>
<td>Median</td>
<td>211942.9</td>
<td>219630.9</td>
</tr>
<tr>
<td>3rd quartile</td>
<td>311405.6</td>
<td>322937.7</td>
</tr>
<tr>
<td>Standard deviation</td>
<td>94595.7</td>
<td>95474.4</td>
</tr>
</tbody>
</table>

As we can see in figure 5, most of the intervals defined in the B-Splines based MoP are included in the tail of the MoP proposed here, so we get a similar MoP distribution but much simpler.

However, the main goal of this paper is not a comparison between different MoP estimation procedures, but to check the validity of the proposed procedure as a competitive classifier, so the experiments we will show in the next section will be devoted to that aim.

5 EXPERIMENTS

The aim of the paper is to validate the MoP-NB model, in terms of accuracy of the predictions. This
is done through a k-fold cross validation process, in which overfitting is avoided because all the available data is used, both for learning and testing the model. The proposed model is compared to an implementation of the CART methodology, available through the package \textit{rpart} in R (R Core Team, 2013) and some Naive Bayes classifiers used to manage continuous variables (Bouckaert, 2004), such as the Gaussian, Kernel and Discrete NB models implemented in WEKA (Hall et al., 2009). The comparison of the different accuracies is carried out by means of a statistical test.

A total of 13 different datasets have been selected for the experiments from the UCI Machine Learning Repository (Bache and Lichman, 2013) and the KEEL repository (Alcalá-Fdez et al., 2011). They all have both discrete and continuous features, with a wide range of number of cases (from 61 to 30162), number of features (from 3 to 39) and number of categories for the class variable (from 2 to 5). In the case of missing values, the corresponding cases were removed from the dataset.

From the experiments several issues are to be considered i) The improvement of the results due to the Feature selection scheme ii) The improvement of the results when pseudo-continuous variables are taken into account iii) The validity of the MoP-NB model in comparison, as mentioned above, to well-known classifiers, in particular the classical discrete NB, the Gaussian NB, the Kernel NB (Pérez et al., 2009) and Classification Trees (Breiman et al., 1984).

The algorithms for learning and classification the different models proposed in this paper were implemented in R (R Core Team, 2013).

The results of the different classifiers over the selected datasets are depicted in Table 3. \textit{No discretization and Pseudo-discretization} refers to the NB models listed in Section 3.1, whilst \textit{Classification Tree} refers to the CART classification tree models, \textit{Gauss NB} refers to the Gaussian Naive-Bayes model, \textit{Kernel NB} refers to the Kernel Naive-Bayes model and \textit{Disc. NB} refers to the Multinomial version of the Naive-Bayes model.

Each method has been tested using a 10-fold cross validation method, therefore, the \textit{Accuracy} is the mean of the accuracy of every fold. \textit{# Variables} represents, for the proposed algorithms the mean number of features included in the model for each fold, and for the state-of-the-art algorithms the number of variables of the final model after performing feature selection. For Gauss, Kernel and Discrete NB models, a Wrapper Feature Selection procedure was performed to reduce the size of the model.

Table 3 also shows, in those datasets that contain pseudo-continuous variables (abbreviated as P-C features in the table), that are \textit{Adult}, \textit{Australian}, \textit{Credit} and \textit{German}, the mean number of pseudo-continuous variables included in the model for each fold, whether they have been discretized or treated as continuous features. Notice that this part does not make sense in those datasets where there are not pseudo-continuous features.

To answer hypothesis i), whether or not the feature selection makes a difference, a Wilcoxon paired rank test was performed, returning a \textit{p-value} of 0.001221, and so rejecting the hypothesis of equal performance between Feature Selection and No Feature Selection.

In order to answer ii), whether results improve when discretizing pseudo-continuous variables, no statistical test was carried out, since only four dataset contain this kind of variables. However, in 3 out of these 4 datasets the accuracy when discretizing these P-C variables is higher. The reason for this may be double i) the pseudo-continuous variable is not selected in the feature selection procedure, because its probability distribution was not properly estimated ii) it was included, but added extra-noise to the model, because of the wrong estimation of the probability
Table 2: Databases used in experiments.

<table>
<thead>
<tr>
<th>Datasets</th>
<th>Discrete Features</th>
<th>Pseudo-Continuous Features</th>
<th>Totally-Continuous Features</th>
<th>Class Categories</th>
<th>Instances</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult</td>
<td>9</td>
<td>4</td>
<td>1</td>
<td>2</td>
<td>30162</td>
</tr>
<tr>
<td>Australian</td>
<td>8</td>
<td>1</td>
<td>5</td>
<td>2</td>
<td>690</td>
</tr>
<tr>
<td>Band</td>
<td>25</td>
<td>0</td>
<td>14</td>
<td>2</td>
<td>277</td>
</tr>
<tr>
<td>Cleveland</td>
<td>8</td>
<td>0</td>
<td>5</td>
<td>5</td>
<td>297</td>
</tr>
<tr>
<td>Credit</td>
<td>9</td>
<td>1</td>
<td>5</td>
<td>2</td>
<td>653</td>
</tr>
<tr>
<td>Echocardiogram</td>
<td>4</td>
<td>0</td>
<td>8</td>
<td>2</td>
<td>61</td>
</tr>
<tr>
<td>German</td>
<td>21</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>1000</td>
</tr>
<tr>
<td>Haberman</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>2</td>
<td>306</td>
</tr>
<tr>
<td>Hepatitis</td>
<td>13</td>
<td>0</td>
<td>6</td>
<td>2</td>
<td>80</td>
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<tr>
<td>Hungarian</td>
<td>6</td>
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<td>4</td>
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<td>261</td>
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<tr>
<td>Ionosphere</td>
<td>2</td>
<td>0</td>
<td>32</td>
<td>2</td>
<td>351</td>
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<tr>
<td>Liver</td>
<td>1</td>
<td>0</td>
<td>5</td>
<td>2</td>
<td>345</td>
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<tr>
<td>Switzerland</td>
<td>7</td>
<td>0</td>
<td>3</td>
<td>5</td>
<td>105</td>
</tr>
</tbody>
</table>

Table 3: Results of the experiments. Proposed algorithms are shown firstly: without any discretization (with and without feature selection) and discretizing only pseudo-continuous variables. For each dataset the best accuracy result is boldfaced. Feature selection is included in all methods except for the first one, which is specified in the table.

<table>
<thead>
<tr>
<th>Datasets</th>
<th>Proposed algorithms</th>
<th>State of the art algorithms</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No Disc.</td>
<td>No Disc.</td>
</tr>
<tr>
<td></td>
<td>No FS</td>
<td>Pseudo-Continuous</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Totally-Continuous</td>
</tr>
<tr>
<td></td>
<td>Accuracy</td>
<td>Class. Tree</td>
</tr>
<tr>
<td>Adult</td>
<td>0.7750</td>
<td>0.7140</td>
</tr>
<tr>
<td>Australian</td>
<td>0.7898</td>
<td>0.8507</td>
</tr>
<tr>
<td>Band</td>
<td>0.4055</td>
<td>0.6462</td>
</tr>
<tr>
<td>Cleveland</td>
<td>0.5244</td>
<td>0.5420</td>
</tr>
<tr>
<td>Credit</td>
<td>0.7909</td>
<td>0.8468</td>
</tr>
<tr>
<td>Echocardiogram</td>
<td>0.7928</td>
<td>0.9344</td>
</tr>
<tr>
<td>German</td>
<td>0.7490</td>
<td>0.7380</td>
</tr>
<tr>
<td>Haberman</td>
<td>0.7400</td>
<td>0.7287</td>
</tr>
<tr>
<td>Hepatitis</td>
<td>0.6250</td>
<td>0.6500</td>
</tr>
<tr>
<td>Hungarian</td>
<td>0.6894</td>
<td>0.7509</td>
</tr>
<tr>
<td>Ionosphere</td>
<td>0.7918</td>
<td>0.8831</td>
</tr>
<tr>
<td>Liver</td>
<td>0.4299</td>
<td>0.6579</td>
</tr>
<tr>
<td>Switzerland</td>
<td>0.2433</td>
<td>0.4666</td>
</tr>
</tbody>
</table>

distribution. These two issues are partially avoided by the discretization of these variables.

So, we select the pseudo-Disc model (column #3 from table 3) as the optimal proposed one, and compare it with some state-of-the-art algorithms (Class. Tree and Gauss, Kernel and Discrete NB from Table 3) performing a Friedman’s test with significance level 0.05. The result indicated a statistically significant difference (p-value of 0.0059), but a post-hoc comparison using Wilcoxon-Nemenyi-
McDonald-Thompson’s post-hoc test (Hollander and Wolfe, 1999) yields that the only significant differences (marked in green in figure 6) is between the Kernel NB and the Classification trees. There are no differences between the MoP-NB proposed method and the rest of the algorithms.

It is also important to note that the inclusion of continuous variables leads to more compact models, according to the number of variables included. The mean number of variables included is obviously very similar for the No discretization and pseudo-discretization versions of the MoP-NB model (they only vary in 4 datasets) 3.51 and 3.46 respectively, however, this mean number of variables moves to 4.23 for the classification trees (recall that CART methodology does a feature selection internally when selecting the variables to split). The mean number of variables for the Gaussian, Kernel and Discrete NB models are 6, 7.61 and 5.46 respectively.

6 SUMMARY AND CONCLUSIONS

We have presented in this paper a novel classifier, based on Naive Bayes and modeling the distribution of the continuous variables through the MoP model. This work differs from other models using MoP in several senses, we have included a feature selection procedure, as well as a classification-oriented parameter estimation procedure in which tails are given importance. From a more general point of view, we have defined a new kind of variables, called pseudo-continuous which seem to be continuous, but have a behavior that makes them appropriate for discretizing.

The experiments designed show that this new kind of classifier is competitive with respect to other classifiers such as Gaussian, Kernel and Discrete Naive Bayes and CART classification trees. These results also indicate an improvement when the pseudo-continuous variables are differentiated from the totally-continuous and discretized.

There are more complex BN classification models, such as the TAN model, the kDb model or the AODE model. They require more computations in their learning stage, as well as a general inference algorithm to perform the final predictions. We plan to continue the application of MoPs in these more sophisticated models, as well as emphasizing the idea of the pseudo-continuous variables with a definition ad-hoc for each different problem.

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