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LODE: A distance-based classifier built on ensembles of positive and negative observations

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Abstract

Current work on assembling a set of local patterns such as rules and class association rules into a global model for the prediction of a target usually focuses on the identification of the minimal set of patterns that cover the training data. In this paper we present a different point of view: the model of a class has been built with the purpose to emphasise the typical features of the examples of the class. Typical features are modelled by frequent itemsets extracted from the examples and constitute a new representation space of the examples of the class. Prediction of the target class of test examples occurs by computation of the distance between the vector representing the example in the space of the itemsets of each class and the vectors representing the classes.

It is interesting to observe that in the distance computation the critical contribution to the discrimination between classes is given not only by the itemsets of the class model that match the example but also by itemsets that do not match the example. These absent features constitute some pieces of information on the examples that can be considered for the prediction and should not be disregarded. Second, absent features are more abundant in the wrong classes than in the correct ones and their number increases the distance between the example vector and the negative class vectors. Furthermore, since absent features are frequent features in their respective classes, they make the prediction more robust against over-fitting and noise. The usage of

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features absent in the test example is a novel issue in classification: existing learners usually tend to select the best local pattern that matches the example - and do not consider the abundance of other patterns that do not match it. We demonstrate the validity of our observations and the effectiveness of LODE, our learner, by means of extensive empirical experiments in which we compare the prediction accuracy of LODE with a consistent set of classifiers of the state of the art. In this paper we also report the methodology that we adopted in order to determine automatically the setting of the learner and of its parameters.

**Keywords:** Data Mining, Supervised Learning, Concept Learning, Associative Classifier

1. **Introduction**

In the last years novel approaches to classification emerged joining two distinct research trends in data mining and machine learning: from one side the rich area of itemsets and class association rules algorithms, which studied in depth efficient algorithms on very large data-sets and on high dimensional data being able to extract large volumes of patterns from data and to enforce on them a rich set of evaluation constraints [1–8]. On the other side the even richer field of machine learning discoveries, whose algorithms and theories were able to combine in powerful classifiers an ensemble of already powerful elementary learners built on a wide spectrum of inductive learning techniques [9–14].

The framework From Local Patterns to Global Models (LEGO) approach to learning showed how it is possible to join these two worlds trying to combine the benefits of the respective fields: manage an increased volume of predictive patterns and at the same time evaluate and assemble them into powerful learners. The resulting model constitutes a general model for learning [15].

Current work [16] on assembling a set of local patterns such as rules and class association rules into a global model for the prediction of a target usually focuses on the identification of the minimal set of patterns that is able to cover the training data. In this paper we present a new learner named LODE (Learning On Distance between Ensembles) which takes a different viewpoint. We are convinced that a good model of a class should emphasise the typical features of the examples of the class and that for effective results
in classification all of them should be used *at the same time* at prediction time.

In this approach to induction of classification patterns and global model construction we abandon the usual learning strategy based on covering. Covering algorithms often remove the training examples covered by the local patterns already discovered. The removal of examples is motivated by the aim of identifying a global model constituted by a minimal set of as much as possible diverse patterns.

Consider the example shown in Figure 1 in which the examples of the positive class are denoted by the symbol ‘+’. By a classical covering algorithm, like Ripper [9] and the rule induction systems from propositional learning [17], each new learnt classification rule covers the most numerous set of examples that have not been covered yet by the already learnt rules. Suppose the first classification rule discovered is $ABC \Rightarrow +$. After this, all the positive examples matching $ABC$ are discarded from further considerations. In Figure 1 these examples are included in the square labelled by $ABC$. After the elimination of these examples, another rule like $DEF \Rightarrow +$ has no possibility to be extracted because without the discarded examples, it does not cover a sufficiently large set of examples. The unfortunate consequence of this covering strategy is that the algorithm misses some rules that have a large support in terms of covered examples and represent a distinctive
set of characteristics of the positive class that would give a more complete characterization to it. As a conclusion, the examples removal by the covering strategy introduces a distortion in the training set which prevents the induction process to learn a sufficient number of characteristics from the removed examples. In practice, it diverts the learning towards the construction of a class model composed of a comprehensive set of characteristics that describes satisfactorily the examples of a class. Instead, by LODE, all the classification rules that cover a sufficient number of examples are considered. LODE is characterized by the following distinguishing issues:

1. the model of each class includes those classification patterns that are frequent w.r.t. the training examples of the class. The frequency constraint is a guarantee that the pattern has a high coverage and represents recurrent features in the class. Though, we do not require that any two local patterns included in the class model should cover distinct examples. We believe that we should create a global model that emphasises the class characteristics and therefore could contain more patterns that represent the same examples.

2. The selected patterns generate a probabilistic class model that consists of a different representation space of the examples. The model of each class is a vector whose components are the selected patterns observed in the training examples of the class. Each vector component is represented with a magnitude equal to the probability with which the corresponding pattern occurs in a random example of the class.

3. We used as classification patterns the frequent set of items (called itemsets). Itemsets are introduced in Subsection 4.1 [18]. In LODE itemsets are selected by the innovative criterion of $\Delta$ [19]. $\Delta$ is the departure of the observed frequency of a pattern w.r.t. an estimated frequency of that pattern computed on the basis of the observed frequency of its subsets in the condition of maximum entropy. As a result, a high $\Delta$ occurs in itemsets whose occurrence could not be determined from the observation of their subsets. This fact is an indication that the itemset is not redundant with respect to its subsets in terms of information quantity.

The adoption of $\Delta$ has the aim to control the volume of patterns in the class model. In fact, an increase in the volume of patterns could occur, due to the combinatorial explosion of the items and the combination of unrelated, independent elements into the itemsets.
In a related paper [20] we have validated the use of $\Delta$ for itemsets selection in classification. As in LODE, in [20] the class models are based on ensemble of itemsets as well. We have compared $\Delta$ with different alternative measures such as accuracy, KL divergence [21], strong jumping emerging patterns [7]. Experiments showed that $\Delta$ allows to identify the itemsets that make effective the classifiers.

4. Prediction of a test instance occurs by distance computation between the vectors representing the class models and the projection of the test instance into each class space. In this projection, the role played by the typical characteristics of class examples is crucial: those characteristics that are absent in a test instance but are present in the class model will make the difference between the classes. The predicted class will be the one in which the typical characteristics of the class are absent in the instance as little as possible.

Thus, one of the fundamental differences between our learner and existing prediction techniques based on rules is that our learner uses for the prediction all the patterns of the class model at the same time - also the patterns that are absent in the test example. Instead, other pattern-based classifiers, like Class Based Association Rules (CBA), RIPPER, decision lists or decision trees, choose a single rule for the prediction; whereas instance-based classifiers, like k-nn, rely too much on the single examples, that come from local portions of the data-set and could be affected by noise or result in over-fitting.

In the new classifier that we present in this paper, LODE, prediction of a test instance occurs by distance computation between vectors representing ensembles of local patterns. Our learner is at some extent, similar to ensemble learners, since both combine by an operation of weighed aggregation the contributions of a large number of elementary predictions. Though, ensemble learners like RandomForest or AdaBoost [22] still base the single predictions on patterns which have been recognized present and not absent from the test instance.

We do not compare in this paper LODE with ensemble learners but only with elementary classification techniques since our purpose here is to show that our learning technique in itself is already ready to be employed as a basic learning technique. The local patterns employed in LODE, a combination both of present and absent patterns from test instances, could still be further combined into a more sophisticated global model, having performed an
evaluation and aggregation of local patterns typical of boosting or bagging’s combination of elementary classifiers. This is a matter for future work.

The contributions of the paper are the following. We describe and formalize a new classifier in the context of the LEGO framework. Differently from many approaches based on local patterns, in LODE the generation of the class model by local patterns does not occur by a covering strategy but by frequent pattern search. Frequent patterns produce class descriptive models. The class models are vectors composed of the itemsets occurring in the training class examples and are weighed by their probability in the class. By the experimental results of Section 7, we make evident that a descriptive model of the classes, after a suitable simplification and evaluation by a wrapper approach, can be used effectively with a predictive purpose.

The descriptive model is turned into a predictive model by an optimization, randomized algorithm (Simulated Annealing) whose working points represent the possible cuts in the ranking of the class itemsets. Ranking has been performed by $\Delta$, proposed in our previous works [19, 20]. $\Delta$ is an uncommon criterion for the determination of the importance of the itemsets and is related to a non redundancy test between the itemsets and their subsets.

Prediction by local patterns occurs by an ensemble in which all the patterns of the class model are used at the same time. Furthermore, prediction makes use not only of the patterns present in the example but also of the absent patterns. We show that the adoption of a multitude of patterns selected from the original class description models makes the prediction more robust against noise and the risk of over-fitting. In this paper we have performed extensive experiments on many data-sets and have conducted a comparison with a consistent set of classifiers of the state of the art. We show that LODE is specially suitable to the prediction in noisy environments since its characteristics and its probabilistic nature make it robust against noise. As a conclusion we are able to show its excellent performances.

The last contribution is to show the methodology for the tuning of the parameters in LODE. Tuning exploits the memory resources of the system and occurs without the intervention of the user. The contributions of this paper with respect to previous work [20] consist in the formalization of the overall LODE model, the wrapper approach based on Simulated Annealing, a more extensive set of experiments that consider both the real and the noisy data and the tuning strategy of the parameters.

This paper is organized as follows. Section 2 provides an overview of the related work while Section 3 puts LODE in the context of the research
works that combine local patterns into a global model for learning. Section 4 presents the details of the inductive learning technique. Section 5 discusses one of the positive issues of this type of classifier: it has at the same time descriptive and predictive capabilities. Section 6 presents the methodology we adopted in order to automatically set the learner parameters. Section 7 presents the experimental results. Finally Section 8 draws the conclusions.

2. Related work

Associative classification is a popular classification technique which combines association mining with classification. CBA [1] is one of the earliest associative classifier which uses association mining to extract association rules from the training data-set. It then ranks the rules based on their confidence. Later it builds a global model from these rules by using a wrapper approach. HPWR [23] is an associative classifier which uses statistical residual analysis [24] for the selection of the best associative patterns in the data-set and then uses the weight of evidence [25] in making predictions. IGLUE [26] uses the concept lattice in order to re-describe each instance of the dataset. In the new description of instances the attributes are represented by numerical values according to the number of their occurrences in the concept lattice. It later uses K-nn for prediction. Classifiers based on Emerging Patterns [6–8] choose the associative patterns for each class that are able to discriminate between classes on the basis of their support ratio in different classes. The patterns with larger support ratio are given priority. L3 [27] is another associative classifier which uses a compression method for maintaining more associative rules compared to other associative classifiers. It divides these rule sets into two parts where the first part contains all the specific rules and the second part contains spare rules. During prediction if a matching rule is not found in the first part then the spare rules are used.

Instance based classifiers like K-nn [10, 28, 29] use distance calculation similar to ours. The main difference between our approach and K-nn is that K-nn calculates the distance between each test instance and a (possibly large) set of training instances for making prediction. In our approach we calculate the distance between a test instance and a whole global model for each class. Decision tree classifiers like J48 [30] use decision trees where the internal nodes represent tests on attributes and the leaf nodes represent classes. The best matching single path from root to the leaf node is chosen for making any prediction. Rule based classifiers are the closest cousins of
associative classifiers. Typical rule based classifiers like Ripper [30] apply the if – then paradigm for building rules and uses the best rule for any prediction. Probabilistic classifiers like Naive Bayes [31] use prior probabilities for calculating all the class conditional probabilities of each attribute. For making a prediction they calculate the posterior probability of the attributes in a test instance. The class with higher posterior probability is chosen as the predicted class. SVM [32] uses the concept of maximum margin hyperplane to find a decision boundary which maximizes the distance between the examples of distinct classes. The decision boundary is the global model which has been built by giving a special relevancy to some local observations in the training set (closest observations coming from different classes). In this sense it is sensitive to the presence of noise in those local portions of the dataset.

Almost all the above mentioned classifiers use local information for making a prediction, in the sense that they use only those patterns or attributes which are present in both of the test instance and the model constructed from training instances. Differently, in LODE we use the global information of a prototypical model of the classes for making any prediction, in the sense that we use all the information contained in each class model as we calculate the distance between each class model and the test instance. According to our knowledge this is a new approach.

Our learner is at some extent, similar to ensemble learners because it combines by an operation of weighted aggregation the contributions of a set of elementary predictions. Though, ensemble learners that make use of bagging, like RandomForest, or boosting, like AdaBoost [22] still base the single predictions on patterns which have been recognized present and not absent from the test instance. Moreover, the basic mechanism by which they learn is different to ours: they generate randomly the basic learners and during the learning adjust their weights. Instead, in LODE we generate first a descriptive model of the class (that acts as a sort of prototype) and learn how to simplify the model by elimination of some of the features. [33] shares the same idea that an ensemble must be composed of millions of patterns but presents important differences w.r.t. our approach: it randomly generates patterns which have a uniform (also low) coverage in terms of the number of examples they match. Patterns are formulated by checking that they do not cover negative examples and are weighted in a uniform way.
3. From Local Patterns To Global Models

LODE is an instantiation of the LEGO approach to classification [15]. LEGO requires an initial extraction of local patterns from the training data, possibly selected by means of some constraints. Resulting patterns could still be redundant or loosely correlated to the target. Thus local patterns are selected by means of some measure of subsets evaluation. Resulting patterns can be thought as the features on which the classification is based upon. Final patterns are in turn aggregated into an ensemble that constitutes the final global model which is used for prediction of test data.

In Figure 2 we present LODE work-flow according to this approach. In LODE the terms (local) patterns and (frequent) itemsets are meant as interchangeable. Furthermore, each itemset is a feature in the new representation space of the class examples and each of the features corresponds to a vector component in the class model.

1. From the training examples of source data, frequent itemsets are extracted as the local patterns and separately from each class. Any algorithm for frequent itemsets extraction would be valid. The step of frequent itemsets mining can be considered as a black-box from the viewpoint of the final LODE classifier whose need is to collect the largest
possible set of frequent itemsets that main memory allows\(^1\). Resulting frequent itemsets generate a lattice. The itemsets with the same cardinality are at the nodes of the same level in the lattice and have edges to their subsets and their supersets at the closer lattice levels nodes.

2. Then the itemsets in the lattice of each class \(C_i\) are ranked into ranking \(R_i\) according to some evaluation measure. The purpose of this step is to raise at the top of the ranking the patterns to be included in the model of class \(C_i\). In the case of LODE, as we will see in Section 3.1, the evaluation measure is the normalized \(\Delta\). Normalized \(\Delta\) allows to compare itemsets of different cardinality and, as \(\Delta\), allows the identification of which itemsets are not redundant w.r.t. their subsets.

3. The correct top portion of the ranking must be determined so that the ensemble can be simplified, the classifier can be made more robust and efficient and the risk of over-fitting is reduced. As a result of this simplification, each itemset that is in the top portion of the ranking is placed in the ensemble.

4. The ensemble model of the class \(i\) is a vector \((M_i)\). Any vector component \((p_{ij})\) represents an itemset in the ranking \(R_i\) with a weight equal to its probability \((p_{ij})\) to occur in an example of the class \(i\).

5. The global model is composed of all the class vectors.

   The prediction of a test instance is made by projection of the instance in the feature space of each class. The proximity of the projected vector to any class vector is then evaluated for the different classes: the class which is closest to the test instance is predicted.

3.1. \(\Delta\) as the Measure to Select the Relevant Itemsets for the Characterization of the Classes

We use the criterion of \(\Delta\) \([19]\) for the selection of local patterns that will be included in the global model. \(\Delta\) is the departure of the observed frequency of a pattern w.r.t. an estimated frequency of that pattern:

\[
\Delta = P_{\text{observed}} - P_{\text{estimated}} \tag{1}
\]

\(P_{\text{observed}}\) and \(P_{\text{estimated}}\) are the relative frequencies of a pattern in each class.

\(^1\)In the context of our implementation we adopted LCM \([34]\), the winner of FIMI-2004, a competition on Frequent Itemset Mining algorithms on very dense datasets.
The referential estimation is computed in the condition of maximum entropy. It represents the frequency that the itemset would have in the case it was maximally difficult to predict its presence when the presence of its subsets was known. In other terms, it is the condition in which the presence of the itemset gives the maximum information when we know the presence of its subsets.

Notice that the estimated probability is a generalized version of the condition of independence of the itemset with respect to its subsets: for a pair of items it corresponds to the product of their individual probabilities. For a number of itemsets higher than two there is not a closed formula: the solution is found by an iterative, numerical procedure that finds the zero of the derivative of the itemset entropy function [35]. ∆ is a departure from a generalized condition of independence between \( n \) items. Thus, it is able to determine a existing dependency between all the items in the n-itemset and to distinguish when a dependency, though present in the itemset, is present because it has been “inherited” by dependencies already existing in the subsets. Summarising, it gives us an indication necessary to distinguish the intrinsic utility of the itemset w.r.t its subsets.

**Normalised \( \Delta \).** In this paper we adopted a normalized version of \( \Delta \) as follows:

\[
\frac{\Delta}{P_{\text{observed}}} = \frac{P_{\text{observed}} - P_{\text{estimated}}}{P_{\text{observed}}} \quad (2)
\]

This normalisation is necessary in order to compare itemsets with different cardinality. In fact, it is well-known that itemsets with a higher cardinality tend to have a lower value of probability. As a consequence, the expected values of \( \Delta \) for higher cardinality itemsets are lower.

Itemsets with an absolute value of \( \Delta \) close to zero are considered redundant w.r.t. their subsets. If two independent subsets are merged to form a new itemset, the contribution of the new itemset to the model would be low because the new itemset does not add new information w.r.t. the subsets. In case of independent subsets, the probability of the supersets corresponds to the estimated probability, obtained in the condition of maximum entropy. Therefore, a \( \Delta \) close to zero identifies an itemset that has been formed merely by the combinatorial process of union of items in the itemset formation but do not constitute any specific information, per se. This is an indication that the itemset can be eliminated. Conversely, a high \( \Delta \) occurs in an itemset if its subsets are dependent. In that case, the superset would be interest-
ing because identifies related pieces of information that combined appear as non casual. Itemsets that result from the selection based on a high value of normalized \( \Delta \) represent the features of interest.

The resulting class model is probabilistic and consists in a characterization of the class based on frequent itemsets composed by non independent subsets. This is motivated by the need of eliminating irrelevant itemsets from the multitude of frequent itemsets. Indeed, some frequent itemsets, though might occur in the class with a high probability, could be constituted by independent elements; in this case they should not be kept since they do not convey significant additional information for the class with respect to their subsets.

3.2. Selection of the abstraction level

In LODE, ranking is used also to determine unknown characteristics of the local patterns. The main unknown characteristics of the patterns is the itemset cardinality.

Any lattice level can be considered as an abstraction level at which the class model can be constructed. It is the purpose of the inductive algorithm to learn the correct abstraction level at which the itemsets must be selected in the global model. In fact, itemsets coming from different abstraction levels should not be kept in the model: there would be a duplication of information in the two itemsets when a relationship of specialization (or set containment) exists between them.

In LODE we used ranking by normalized \( \Delta \) also to determine the abstraction level of the itemsets in the lattice. Ranking the itemsets makes emerge the patterns that are more relevant for the class model. We determined from the top portion of the itemset ranking which are the most recurrent values of the itemset cardinality and produced a rank of the cardinality values denoted by \( R_{\text{levels}} \). The best value of the cardinality \( j \) will be selected from \( R_{\text{levels}} \) by a wrapper approach. The wrapper is based on the accuracy obtained by the classifier LODE with the class models built on the portion of the itemset rankings with cardinality \( j \). In particular, the portion \( R_{ij} \) of the ranking \( R_i \) is composed of the itemsets extracted from class \( i \) with cardinality \( j \). The method is an optimization algorithm based on Simulated Annealing that is responsible also for the ranking reduction (feature selection) and is described in Section 6.5.
4. The Distance-Based Learner on Ensembles of Itemsets

In this Section we describe how our distance-based learner works.

4.1. Preliminary definitions

Before entering into the details of the prediction, let us introduce some preliminary definitions. Each test and training example is described by some attributes whose values characterize the instance itself. The values of each attribute belong to a certain domain that could be continuous or discrete. Continuous attribute values are not suitable to be employed in classification by means of class association rules, since they do not often occur frequently in the data. The search for recurrent and frequent itemsets usually works on discrete (categorical or numerical) values in the class examples. Thus continuous values are usually discretized in a preprocessing step, by a supervised algorithm like [36].

- Any example in the data-set is represented by a set of attribute-value pairs. We call item an attribute-value pair. For each item a binary variable is associated. For each example of the data-set the binary variable associated to an item has a true value if the example contains that attribute with that value, false otherwise. In this way, the data-set conceptually can be represented as a binary matrix with a row for any example and a column for any item. In each cell of the matrix there is a true or a false value according to the value of the item variable for that example.

- Similarly, from any example we can generate itemsets, as those sets of item variables assuming true values for the example.

- Since we are interested only in recurrent characteristics observed in the examples of a class, we recall only frequent itemsets that occur with a certain, minimum frequency in the examples of the class.

- Let be \( \mathcal{C} = \{C_1, C_2, ..., C_L\} \) the set of classes.

- \( M_k \) denotes the model of the class \( C_k \), with \( k = 1..L \). \( M_k \) is represented as a vector of \( n_k \) components. \( M_k \) constitutes a new feature space of representation of examples. Each component of the feature space of class \( C_k \) is one of the frequent itemsets extracted from the training examples of class \( C_k \).
• $p_{ki}$ denotes the probability with which $i$-th itemset (or $i$-th component) of the feature space of the class $C_k$ occurs in a random example of class $C_k$.

• $M_k \equiv \langle p_{k1}, p_{k2}, ..., p_{kn_k} \rangle$

4.2. Projection of a test instance in the class space

Each test instance is described by some attributes whose values characterize the instance itself. Similarly, as we represented training instances, by means of itemsets variables with true or false values, we represent test instances, adopting the same representation.

• In order to predict the class of a test instance $t$ it must be represented in the feature space of each class $C_k$. This is a projection operation.

• The projection works as follows. We check the presence of each component (an itemset) of the class model $M_k$ in the instance $t$. Let us denote this generic component as the $i$-th component. If it is present in $t$ (that is, if every item in the itemset $i$ is present in $t$), we set the value of the $i$-th component of $t$ vector to 1; 0 otherwise. In fact, not all the itemsets observed frequently in the objects of a certain class will be present in every test instance, even if the instance comes from that class.

• In the projection we discard those components of $t$ that are not present in class $C_k$. In fact, not all the itemsets that are present in the instance $t$ could be present in a certain class, even if that instance comes from that class.

As a conclusion, the vector of instance $t$ projected into the feature space $M_k$ of class $C_k$ is denoted by $\overrightarrow{t} \perp \overrightarrow{M_k}$ whose $i$-th component is:

$$Ind(i, t)$$

where the itemset $i$ is the $i$-th component of the feature space $M_k$ and $Ind(i, t)$ is the indicator function whose value is 1 if itemset $i$ belongs to test instance $t$, 0 otherwise.

Vectors are later used for class prediction in a distance computation. Since, class vectors could have a different number of components (features in the feature space) according to the number of frequent itemsets that could
be extracted from the examples of that class, we do not want to favour in the distance computation those classes whose feature spaces have a low number of features since in those spaces data is less sparse and therefore distances result reduced. To solve the problem we normalized the vectors such that their length is 1 and as a consequence distances are not influenced by the number of features in the space. In the following we will use the normalized version of the vectors, indicated by the $u$ operator. The generic $i$-th component of the normalized vectors is:

$$u(t_i) = \frac{\text{Ind}(i, t)}{\sqrt{\sum_{j \in M_k} \text{Ind}(j, t)}}$$

(4)

$$u(M_{ki}) = \frac{p_{ki}}{\sqrt{\sum_{j \in M_k} p_{kj}^2}}$$

(5)

where $t_i$ and $M_{ki}$ represent respectively the $i$-th component of $\vec{t}$ and $\vec{M_k}$.

4.3. Distance computation between the test vector and the class vectors

The proximity between the two vectors $\vec{M_k}$ and $\vec{t}$ can be computed in many ways, either as a measure of distance (like the Euclidean distance), or as a measure of similarity (like the cosine similarity), Jaccard or extended Jaccard, etc. Here, we report results obtained by application of the Euclidean distance and the cosine similarity (the results do not differ significantly). Notice, however, that they have an opposite behavior: the former increases with the dissimilarity of the instance w.r.t. the class, while the latter decreases.

$$\text{Euclidean distance}(\vec{u}(M_k), \vec{u}(t)) = \sqrt{\sum_{i \in M_k} (u(M_{ki}) - u(t_i))^2}$$

(6)

$$\text{cosine similarity}(\vec{u}(M_k), \vec{u}(t)) = \frac{\vec{u}(M_k) \cdot \vec{u}(t)}{|\vec{u}(M_k)| \cdot |\vec{u}(t)|}$$

(7)

Justification of the proximity formulas. It is clear from the formulas 6 and 7 that when a feature (itemset) is present in the class model $M_k$ but it is absent in the test instance $t$, its contribution does not increase the value of cosine similarity and increases the Euclidean distance. On the other side, when the total number of features in the class is high, the contribution to
each of them is lower (because of the normalisation factor). However, in this
normalisation factor even absent features have their weight and contribute
to decrease the weight of all the other features (also the present ones).

4.4. Formalization of the problem with an objective function

LODE framework can be formalized as a problem of learning the com-
position of the class models in terms of the frequent itemsets extracted from
the training examples. The learning makes use of an objective function 8 that
guides the tuning of the models and can be formalized as follows.

Let \( \mathcal{C} = \{ C_1, C_2, \ldots, C_L \} \) be the set of classes and \( \mathbf{Tr} = \bigcup_{i=1}^{L} \mathbf{Tr}_i \) the training
set where \( \mathbf{Tr}_i \) is the subset of the examples of class \( C_i \). The goal of the
learning task is to find the set of class models \( \{ M_1, M_2, \ldots, M_L \} \) in which each
\( M_i \) is composed of frequent itemsets occurring in the examples of \( \mathbf{Tr}_i \). Each
itemset \( j \) describes a set of characteristics occurring in the examples of \( \mathbf{Tr}_i \)
with frequency \( p_{ij} \). Each \( M_i \) maximizes an objective function:

\[
\sum_{t \in \mathbf{Tr}_i} \text{Ind}\{ \text{proximity}(\mathbf{u}(M_i), \mathbf{u}(t \perp M_i)) > \text{proximity}(\mathbf{u}(M_j), \mathbf{u}(t \perp M_j)) \}, \text{ with } j \neq i
\]

where the proximity function is equation (6) or equation (7), \( t \) is an example
of class \( C_i \) in the training set \( \mathbf{Tr}_i \). The unary indicator function \( \text{Ind}\{ \cdot \} \) is 1
if its parameter is true, 0 otherwise. The parameter:

\[
\text{proximity}(\mathbf{u}(M_i), \mathbf{u}(t \perp M_i)) > \text{proximity}(\mathbf{u}(M_j), \mathbf{u}(t \perp M_j))
\]
is the condition for the true class prediction. The proximity is computed
between two vectors: the unit vector representing the class model and the
unit vector representing a single training instance projected in the class space.
The objective function increases for the class model \( M_i \) when the proximity
between the class model vector and the training instance vector is higher
then for the other class models \( M_j \). In other terms, the objective function
represents the number of correct predictions for class \( C_i \).

The tuning step of LODE for the determination of the composition of
\( M_i \) selects the itemsets from a ranking \( R_i \). The itemsets in \( R_i \) have been
ranked by an evaluation measure (that in our case is the normalised \( \Delta \)). Let
denote by \( R_{il} \) the set of itemsets in \( R_i \) of cardinality \( l \). The selection of the
itemsets in \( M_i \) is made by determination of the itemset cardinality value \( l \)
(abstraction level) and of a cut of the ranking \( R_{il} \), such that the top portion
of the ranking is retained in \( M_i \) and the portion below the cut is discarded.
The cut has been formalised by the determination of a percentage \( r_i \) of the
itemsets to be retained in \( M_i \).
4.5. LODE on a toy example

In Figure 3 we show a toy sample from the Play Tennis data-set, a well-known example of a prediction problem of the suitable weather conditions for playing tennis. From the examples of the two classes (Yes and No) separately, LODE extracts the frequent itemsets. The two diamonds represent the itemsets search space that has the form of a lattice. LODE ranks the itemsets according to the value of the normalized $\Delta$ (NDelta). Then, the itemsets cardinality equal to a value of three is selected (see the method described in Subsection 3.2). For each itemset, the value of the relative frequency in the class is reported. In the picture, due to lack of space, only the first six itemsets are shown in the ranking, out of the total of eleven extracted itemsets. These ensembles constitute a descriptive representation of the classes in which the order of the features is informative of its importance in the description.

In Figure 4 we show instead the test phase of LODE. On the left, the vector model of class Yes is composed by the eleven itemsets that were in the corresponding ensemble. Each itemset is represented in the vector by its relative frequency in class Yes. Analogously for the other class whose vector is shown on the right. Below, we show both the vector models with numerical values, after normalization.

In the center of the picture a test example is taken. At the bottom, the picture reports the vectors representing the test example under the viewpoint of the classes. They have been obtained by means of projection of the original test vector in the feature space of each class and normalized. Finally, the distance between the unit test projected vector and the class vector is computed. In this example, class No is predicted. In fact, class Yes has only one present feature (which in addition occurs in the class with a low frequency) while class No has four features (and the first one is very frequent). In this example, both the number and the frequency of the present features make the difference in the computation of the two distances which depend also on the number of absent features in the test vector projected in the class space.

5. Descriptive and predictive capabilities of LODE

The distinctive issue in LODE is the adoption of a descriptive model for a final predictive goal. The question that we want to answer with this work is: does a descriptive model support a predictive task? We will ask with the empirical evidence of the extensive experiments performed at Section 7.
The benefit is that it is possible to obtain both a descriptive model of the classes and a predictive model, at the same time. Furthermore, the test instance vector projected onto the class models constitutes a descriptive representation of the test instance itself. This model is interpretable: it is composed by an ensemble of itemsets where each itemset is in practice
a sentence composed by a conjunction of predicates that are true for the instance and are frequent in the class. Furthermore, each predicate sentence is equipped with a weight equal to the probability of its occurrence in the class examples. This weight is then reduced by the total number of sentences (if many sentences are applicable each of them weighs less).

The number of class features that apply to the test instance tells us how much that instance is similar to the class. Finally, the evaluation measure of the proximity between the test vector and the class vector quantifies the similarity between the test instance and the class.

As regards the descriptive capability, we believe that using Δ - this novel
concept, unfortunately rarely adopted in data mining - it is possible to distin-
tinguish which of the itemsets are really relevant to determine a complete,
but non redundant, set of characteristics describing a class of examples.

Certainly related to this concept are the concepts of miki, maximally
informative k-itemsets [37] and pattern teams [38]. Both make use of the
concept of maximum entropy but they do not make use of the departure
with respect to it. Indeed, their aim is orthogonal to ours. Miki’s aim is
to identify the minimal set of items that is able to distinguish between the
examples in the data and search for the set of items that are distributed in
the most uniform way in the data-set. Similarly pattern teams’ aim is to
select from a set of local patterns the minimal set that is non redundant and
that allows a maximal covering in terms of the number of the examples they
represent. Instead, our aim is descriptive: in the class model we want to
identify the most complete set of patterns that catch all the class features
observed in the training set. This ensemble of patterns collectively provides
a probabilistic description of the class. It is composed of itemsets that have
a good coverage each (because each of them is frequent) and such that each
of them is not redundant.

Related to our goal is the field of subgroup discovery [39, 40]. In subgroup
discovery, the set of patterns selected for the representation of a subgroup of
the population does not need to provide a complete description of the target
but needs to represent a set of interesting characteristics. The difference of
our approach w.r.t. subgroup discovery is that our model aims to provide a
complete description of the class while subgroup discovery accepts to describe
only a portion of the target but it achieves a statistically sound description of
a subgroup of examples. It would be possible to adopt a subgroup discovery
model and adapt it to a predictive goal by means of the distance-based ap-
proach on positive and negative descriptions as described in this paper. This
will be the theme of a future research work. Currently, the authors of [40]
made the reverse: they adapt a rule induction algorithm with a predictive
goal to the subgroup discovery task.

On itemset redundancy. Itemsets redundancy is tested in our model by $\Delta$
with respect to the itemsets subsets and not with respect to their peer-
itemsets. This operation by $\Delta$ is simpler than other standard statistical
methods like the test of independence by $\chi^2$ or on correlation by Pearson
coefficient. Both these tests work on pairs of patterns. Instead, $\Delta$ does not
involve the peer-itemsets but their subsets (whose number is much lower than
the number of peer-itemsets). Furthermore, the results of the experiment we are describing in the following do not justify the search of redundancies between the peer-itemsets. We checked the number of dependent peer-itemsets by \( \chi^2 \) test. The test is done at different cardinality levels of the itemsets. In Figure 5 we report the percentages of redundant itemsets at the cardinality level 2 and 3 detected by \( \chi^2 \) in a few UCI data-sets. We observe that the percentage of redundant itemsets reduces dramatically as soon as the itemsets cardinality increases. This is due to the following fact: as a consequence of the construction of itemsets, the space gets more sparse and the number of redundancies between itemsets results reduced.

In agreement to these observations, many algorithms of rule learning try to reduce redundancies between the rules. Very often they work on redundancies between the original features (in our case, the items). One example is [41] which learns class association rules by an Apriori-style algorithm [18]. During rule generation Apriori-C checks that for each rule there is not already one of its generalisations that has a better coverage. This test aims to reduce the number of rules and is based (as \( \Delta \)) on the comparison between an itemset and its subsets. Rules are ordered according to their quality measure but, in contrast to LODE, one single rule is applied to each test example.

6. Methodology of Learner Setting

In this Section we present the methodology we adopted to generate the local patterns, and later tune the model: select a subset of the local patterns for the creation of a global model.
6.1. Motivations for the ensemble reduction

In Figure 6 we present a study on the effect of the dimension of the ranking of local patterns $R_k$ on the misclassification rate. As we explained, we ranked the local patterns by the evaluation measure (normalized $\Delta$). We reduced the dimension of the class models by elimination of the bottom part of the rankings. Figure 6 shows an initial gradual improvement of the classification accuracy which is due to a simplification of the models that leads to a better generalization capability and a reduction of over-fitting. After the optimum point, the model worsens because too many itemsets have been eliminated from the model and the error rate starts to increase.

![Figure 6: Error rate on Wisconsin-Breast-cancer with class models built on different number of itemsets](image1)

![Figure 7: Misclassification on Wisconsin-Breast-cancer by class models on frequent itemsets at different minimum support](image2)

6.2. Effect of the minimum support threshold on classification

In Figure 7 we observe the relationship between the minimum support threshold that governs the algorithms of frequent itemsets mining and the misclassification rate of the ensemble of local patterns. We notice an improvement in classification accuracy by decreasing the support threshold. This tells us that even itemsets with a not very high frequency could be useful for the classification since their normalized $\Delta$ value could be high.

6.3. Determination of the value of minimum support threshold

How many itemsets do we have to collect? We believe that we could try to collect the highest possible number of itemsets that our computing system can allow: it will be responsibility of the normalized $\Delta$ their final selection in the global model. Thus in Figure 8 we plot the total number of itemsets
having a frequency in the interval indicated at the $x$ axis. Knowing this histogram we can decide a suitable value of the minsup parameter (i.e., the minimum itemsets frequency allowed) given the total amount of memory in our system: it is sufficient to sum up the total number of itemsets taken from the histogram starting from the highest support value and going towards the left until the maximum memory size is reached or the support reached is 0. (In other words we consider the histogram as a probability density function of the itemsets, and compute the area under it from the right).

The construction of the exact histogram is of high computational cost if it is constructed by running an algorithm of frequent itemset mining on a large data-set. Thus we suggest the following strategy.

1. We simply count the frequency of singletons (items) in the data.
2. We compute the frequency of other itemsets under the hypothesis of statistical independence of items and generate the histogram of the number of itemsets with a frequency equal to a certain value determined by the interval indicated at $x$ axis (see Figure 8). You can notice that the real and the estimated histograms differ only at low frequency thresholds.
3. We select the support threshold directly on the estimated histogram and this choice is conservative.

If we wished to establish instead the correct number of itemsets at lower support values, we could collect a sample of the data and test in the sample the total number of itemsets without the hypothesis of statistical independence. In Figure 9 we show that this approach is
practical since the total number of itemsets collected does not differ significantly with the dimension of the sample (the percentages of the sampled examples are indicated at the $x$ axis). Thus this strategy is computationally feasible, since it requires to collect only a little portion of the data-set. In this initial sample we lower the minimum support threshold and observe the total number of itemsets extracted. The right threshold is the one that gives the maximum number of itemsets that the system memory allows.

![Figure 9: Total number of itemsets from Breast data-set with progressive sampling](image)

6.4. Initialisation of LODE

Figure 10 shows the diagram of the initialization of LODE. After the first step of determination of minimum support threshold (according to the algorithm described in Section 6.3) the algorithm of frequent itemsets mining is launched on the training-set and produces a set of frequent itemsets that are ranked according to their normalized $\Delta$ value (ranking $R_{all}$). This initial, unique ranking tells us some things regarding some unknown but characteristic parameters of the itemsets, as already explained in Subsection 3.2. First of all the most recurrent value of cardinality of the itemsets at the top portion of the ranking (top portion is set to 2/3). The most recurrent value of itemsets cardinality at the top of the ranking is the first value that will be tested in the following tuning process of LODE in order to select itemsets for the class models (see Section 6.5). Then, the other cardinality levels are ordered in a rank ($R_{levels}$) according to the frequency of the cardinality value
of the itemsets at the top portion of the ranking $R_{all}$. This is done because it is necessary to determine a unique value of cardinality for the itemsets in the class models. Indeed, it makes no sense to include itemsets with a different cardinality value as components of the class vectors since some dependencies exist between itemsets and their subsets.

6.5. Tuning step of LODE

The pseudo-code of the algorithm that performs the tuning process of the class models is Algorithm 1. This tuning process has been performed in all the experiments that will be described in Section 7 within a 10-fold cross-validation (from line 6 to 37): we repeated this process 10 times on different training sets and with a different validation set and test set. For each training set there is a separate validation set. On each training set the learning algorithm produces a model (lines 14-19) with parameters which are optimised by the tuning algorithm on the validation set (lines 22-33). As can be observed from Algorithm 1, since the discretization performed by Fayyad and Irani’s method [36] is a supervised step, we have to guarantee that no information on the class outcomes towards the test sets in an hidden way through the information on the discretization. Thus, the discretization has been applied ten times as a pre-processing step to each different training-set (see lines 10-11) and re-applied ten times to the corresponding test set (line 34). Each test-set is then used to evaluate the classification accuracy of the generated models (lines 35-36) and gives in output the average of the values of misclassification for the ten test folds (line 38).

1. At lines 7-11 the cross-validation procedure is prepared; discretization is applied as a pre-processing step to each different training-set of the cross-validation loop.
Algorithm 1 Feature Reduction Tuning Process of LODE within Cross-Validation

1: Input: \( D \) data-set
2: Input: MaxMemorySize
3: Output: TotErr LODE classification error
4: - - set small and big perturbation (percentage reduction) in rankings - -
5: \( s = 0.0002, b = 0.05, \) TotErr = 0, Nfolds = 10
6: for testFolds = 1 to Nfolds - - 10-folds cross-validation - -
7: divide \( D \) into disjoint portions: training-set \( Tr \), validation-set \( V \), test-set \( T \)
8: - - apply Fayyad Irani supervised discretization on training-set and validation-set - -
9: set of attribute value intervals \( S = \text{FindSupDiscr}(Tr \cup V) \)
10: \( \text{discTr} = \text{applyDiscretization}(Tr, S) \)
11: \( \text{discV} = \text{applyDiscretization}(V, S) \)
12: - - Initialisation of minsup and ranking of cardinality levels by Algo. in Fig. 9
13: initialise(\( \text{discTr}, \) MaxMemorySize, minsup, \( R_{levels} \))
14: - - extract itemsets from discretised training-set and rank them by Norm. Delta
15: \( R_{all} = \text{generate-itemsets-and-rankings}(% \text{discTr}, \) minsup\)
16: - - \( R_{ij} \) is the portion of \( R_{all} \) with freq. itemsets at cardinality \( j \) for class \( i \)
17: iteration \( it = 1 \)
18: \( l = \) first cardinality level from \( R_{levels} \)
19: Optimal\( R_i = R_{il} \) - - initialisation step - -
20: BestErr = HIGHEST ERR VALUE - - initialise with worst possible error
21: stop = FALSE
22: repeat
23: - - Tune class models by Simulated Annealing on LODE classification error - -
24: CurrentErr = SA (\( \text{discV}, R_{il}, s, b \)) - - call Algo. 2
25: \( \Delta \text{Err} = \text{BestError}-\text{CurrentErr} \)
26: IF (\( \Delta \text{Err} < 0 \))
27: BestErr = CurrentErr
28: Optimal\( R_i = R_{il} \)
29: END IF
30: IF (\( e(\Delta \text{Err})/it \leq \text{rand}(0,1) \)) stop = TRUE
31: \( it++ \)
32: \( l = \) next cardinality level in \( R_{levels} \)
33: until stop
34: \( \text{discT} = \text{applyDiscretization}(T, S) \)
35: errorOnTestFold = LODE(\( \text{discT}, \) Optimal\( R_i \))
36: TotErr = TotErr + errorOnTestFold
37: end for
38: return TotErr/Nfolds

2. At line 13 the initialization phase determines the minimum support value (by algorithm described at Figure 10) and generates the rank of
itemsets cardinality levels ($R_{\text{levels}}$).

3. At the lines 16-20, during the first iteration ($\text{it}=1$), the sets of frequent itemsets are extracted separately from the classes of the training set (with the same minimum support proposed by the initialisation step); the itemsets for class $i$ are ranked in ranking $R_i$. They are given in input to the phase of itemset cardinality selection (lines 22-33) for the construction of the class vectors. The level is taken from the rank of levels ($R_{\text{levels}}$) obtained at initialisation.

4. From the collections of itemsets at the selected level of cardinality $l$ separate rankings are obtained for the various classes ($R_{il}$). We denote by $R_{il}$ the different portions of the ranking $R_i$, where $l$ represents the cardinality level of the itemsets.

5. This is the phase of feature reduction on the rankings $R_{il}$ (at line 24). Since from Figure 6 we saw that the feature selection on the rankings (ranking reduction from the bottom) is beneficial for classification, but we do not know how many minima of the classification error are present, we adopt an algorithm of global optimization ($\text{Simulated Annealing, SA}$) in order to search for the optimum reduction. The pseudo-code of the algorithm of SA is shown in Algorithm 2. We note that SA is executed on the validation set on the rankings obtained on the training set.

6. After SA algorithm stops, a minimum of classification error is obtained in correspondence to the rankings of itemsets at the cardinality level $l$. Delta error ($\Delta Err$) is obtained w.r.t. the best error obtained at previous iterations (which worked on rankings of itemsets with a different value of cardinality $l$). Stop condition (line 33) is probabilistic. It is set to true when $e^{(\Delta Err/\text{it})} \leq \text{rand}(0,1)$. It is similar to the condition seen for entering into a big perturbation in SA: it depends on the reduction in error computed in the current iteration w.r.t. the best error obtained so far, the number of iterations already computed and by a random value. According to this exponential formula, stop is encouraged if the number of iterations is high and if the error is not improving ($\Delta Err$ is negative).

The approach to feature selection operated by SA in Algorithm 2 is a wrapper approach. The rankings are progressively reduced and the decrease in classification error on validation data by the classification algorithm LODE is monitored so that an optimum is found. If we are in search for the next
local minimum (line 12 of Algorithm 2) small perturbations are sufficient (small ranking reduction). Otherwise, if we need to escape from a local minimum (line 17), in order to find a global minimum in the error, we have the chance to escape if we adopt a big perturbation (percent big reduction of the rankings). In the experiments we set the big reduction to 5% while the small reduction consists in 0.0002 (corresponding to a single itemset with a ranking of 5 thousands itemsets). Our implementation of SA has been adjusted by us to the particular contest: the direction of movement on the ranking is unique (bottom-up) when a small reduction is involved. Instead, when a big reduction takes place, it searches the closer minimum in both the directions (top-down and bottom-up).

The condition for the adoption of a big perturbation is again probabilistic: it is governed by a random function \( \text{rand} \), decreases with the number of iterations computed (parameter \( I \)) and increases with the obtained improvement in the error (\( \Delta \text{Err} \)).

**Algorithm 2** Itemsets Selection Algorithm by *Simulated Annealing*

1: Input: \( V \) validation-set
2: Input: \( R_i \) ranking of frequent itemsets.
3: Input: \( s \) small ranking reduction (perturbation)
4: Input: \( b \) big ranking reduction (perturbation)
5: Output: Optimal reduction of \( R_i \) based on classification error.
6: – Tune class models by calling LODE algorithm –
7: \( \text{InitialError} = \text{LODE} (V, R_i) \)
8: \( \text{CurrentError} = \text{InitialError} \)
9: \( \text{MinError} = \text{InitialError} \)
10: \( I = 1 \)
11: repeat
12: \( \text{perturb} (R_i, s) \)
13: \( \text{CurrentError} = \text{LODE} (V, R_i) \)
14: \( \Delta \text{Error} = \text{CurrentError} - \text{MinError} \)
15: IF ( \( \Delta \text{Error} < 0 \))
16: \( \text{MinError} = \text{CurrentError} \)
17: \( \text{OptimalR}_i = R_i \)
18: ELSE IF ( \( \Delta \text{Error} >= 0 \) )
19: \( \text{IF} (e^{(\Delta \text{Error})/I} > \text{rand}(0,1)) \)
20: \( \text{perturb} (R_i, b) \)
21: \( I++ \)
22: until \( I > \) upper limit of \( I \)
23: return \( \text{MinError} \)
7. Experimental Results

We have implemented LODE in C++. \( \Delta \) computation and ranking generation is instead in java. We implemented some scripts in python and perl for the automatising of the whole procedure. In future work we are going to implement all the modules in a unique programming language and equip the system with a web-based graphical user interface. Current version of the software can be down-loaded from:

\texttt{http://www.di.unito.it/~meo/Algo/LODE.zip}.

We have performed classification experiments with LODE on several datasets from the Machine Learning Repository, maintained by UCI as a service to the machine learning community (\texttt{http://archive.ics.uci.edu/ml/}). Table 1 reports the various characteristics of the datasets, chosen for their wide variability in terms of the type of data, number of classes, number of examples, number of attributes, total number of instances per class (indicated in table separated by commas), availability in competitive algorithms, etc. Experiments were run on a Pentium Intel core Duo processor (P9500), running at 2.53Ghz with a RAM of 1.95 GB.

We compared the classification performances of our classifier with many well known classification algorithms: \textit{Knn} [10], \textit{J48} [30] (an implementation of Decision Trees), \textit{Naive Bayes} [42], Support Vector Machines [43] and decision table [44], CBA [1] and RIPPER [9] as representative learners for conjunctive rules-based classifiers. We included also L3 [27] as a representative learner from the set of classifiers that learn by a multitude of rules. We used the implemented version of these classifiers that is available in \textit{Weka} (\texttt{http://www.cs.waikato.ac.nz/ml/weka/}), a collection of machine learning algorithms for data mining tasks.

Notice that some datasets contain only categorical attributes, while others contain also continuous ones. Itemsets are usually extracted from categorical attributes: in fact, from continuous attributes itemsets would have hardly a sufficient frequency. In order to be able to run experiments with LODE and the associative classifiers, we performed a preprocessing step consisting in a supervised discretization of continuous attributes by means of a method based on entropy [36] that minimizes the entropy of the class given the interval of discretization. We used a unique data-set discretization method (by Fayyad and Irani) for all the classifiers that require a discretized data-set and that do not perform discretization by themselves. Instead, for those methods that have their own discretization embedded in the learning algorithm (like
Table 1: Description of datasets

<table>
<thead>
<tr>
<th>Data-set Name</th>
<th>Number of attributes</th>
<th>Number of classes</th>
<th>Number of instances per class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analcatdata-Bankruptcy</td>
<td>7</td>
<td>2</td>
<td>25,25</td>
</tr>
<tr>
<td>Analdata-cyyoung8092</td>
<td>11</td>
<td>2</td>
<td>24,200</td>
</tr>
<tr>
<td>Analcatdata-Creditscore</td>
<td>7</td>
<td>2</td>
<td>27,73</td>
</tr>
<tr>
<td>Analcatdata-Lawsuit</td>
<td>5</td>
<td>2</td>
<td>245,19</td>
</tr>
<tr>
<td>BioMed</td>
<td>9</td>
<td>2</td>
<td>75,134</td>
</tr>
<tr>
<td>Bupa</td>
<td>7</td>
<td>2</td>
<td>145,200</td>
</tr>
<tr>
<td>Credit</td>
<td>16</td>
<td>2</td>
<td>397,383</td>
</tr>
<tr>
<td>Diabetes</td>
<td>9</td>
<td>2</td>
<td>500,268</td>
</tr>
<tr>
<td>Haberman</td>
<td>4</td>
<td>2</td>
<td>225,81</td>
</tr>
<tr>
<td>Horse</td>
<td>28</td>
<td>2</td>
<td>99,201</td>
</tr>
<tr>
<td>HD</td>
<td>14</td>
<td>2</td>
<td>165,138</td>
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<tr>
<td>Hepatitis</td>
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<td>2</td>
<td>32,123</td>
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<td>Heartstatlog</td>
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<td>2</td>
<td>156,120</td>
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<td>2</td>
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<td>Wisconsin-Breast-Cancer</td>
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<td>258,241</td>
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<td>3</td>
<td>73,79,254</td>
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<tr>
<td>Cmc</td>
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<td>3</td>
<td>629,335,911</td>
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<tr>
<td>Iris</td>
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<td>3</td>
<td>50,50,50</td>
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<tr>
<td>Tae</td>
<td>6</td>
<td>4</td>
<td>49,50,52</td>
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<tr>
<td>Grubsdamage</td>
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<td>4</td>
<td>49,41,46,49</td>
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<tr>
<td>Vehicle</td>
<td>19</td>
<td>4</td>
<td>212,217,218,199</td>
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<td>Analdata-Dmft</td>
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<td>6</td>
<td>127,132,124,155,136,123</td>
</tr>
<tr>
<td>Glassp</td>
<td>10</td>
<td>6</td>
<td>70,76,17,13,9,29</td>
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<tr>
<td>Ecoli</td>
<td>8</td>
<td>8</td>
<td>143,77,32,35,20,5,22</td>
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<tr>
<td>Yeast</td>
<td>9</td>
<td>10</td>
<td>463,129,244,163,31,44,35,30,20,15</td>
</tr>
</tbody>
</table>

decision trees) or that do not require a discretized data-set (like K-nn, SVM, etc) we let them the original data-set.

We did discretization and parameters tuning within a 10-fold cross-validation. The division in folds is the same for all the learning algorithms. We performed cross-validation with (disjoint) training set, validation set and test set. Validation-set and test-set are folds from the 10-fold-cross-validation. Training set instead is made by the remaining part of the data-set. We never used the test folds for parameter tuning or discretization: we used test folds only for accuracy testing of the algorithms. For each training-set in the cross-validation, we performed learning of the model and used the validation set to estimate the suitability of the parameters values.

We carefully tuned the parameters values of the competitors as Figure 2 shows. We tested all the combinations of values of their parameters taken from a wide spectrum of possible values with the reported variation step.

In Figure 3 we report the results (mean values and standard deviation)
Table 2: Parameters and their range of values used in the tuning process of learning competitors

<table>
<thead>
<tr>
<th>learner</th>
<th>parameter</th>
<th>range of values</th>
<th>step</th>
</tr>
</thead>
<tbody>
<tr>
<td>J48</td>
<td>pruning confidence</td>
<td>0.05 – 0.5</td>
<td>0.05</td>
</tr>
<tr>
<td>J48</td>
<td>instances per leaf</td>
<td>2 – 10</td>
<td>1</td>
</tr>
<tr>
<td>DTABLE</td>
<td>n. folds cross-valid.</td>
<td>1 – 10</td>
<td>1</td>
</tr>
<tr>
<td>DTABLE</td>
<td>perf. eval. measure</td>
<td>[acc, rmse, mae]</td>
<td></td>
</tr>
<tr>
<td>NB</td>
<td>no parameters needed</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>SVM (SMO)</td>
<td>complexity C</td>
<td>-3 – 3</td>
<td>0.25</td>
</tr>
<tr>
<td>SVM (SMO)</td>
<td>polykernel exp.</td>
<td>1 – 8</td>
<td>0.5</td>
</tr>
<tr>
<td>RIPPER</td>
<td>n. folds for REP</td>
<td>1 – 10</td>
<td>1</td>
</tr>
<tr>
<td>RIPPER</td>
<td>min inst. weight in split</td>
<td>0.5 – 5</td>
<td>0.5</td>
</tr>
<tr>
<td>KNN</td>
<td>k</td>
<td>1 – 10</td>
<td>1</td>
</tr>
<tr>
<td>CBA</td>
<td>minsup</td>
<td>1%</td>
<td>–</td>
</tr>
<tr>
<td>CBA</td>
<td>minconf</td>
<td>50%</td>
<td>–</td>
</tr>
</tbody>
</table>

of the optimal parameter values that we obtained by the parameter tuning process of LODE. In particular, for LODE, we applied the tuning process described by Algorithm 1.

The overall results of our experiments in comparison with other classifiers are presented in the series of tables Figure 11–Figure 5. Each Figure shows the results obtained for the comparison of LODE with the other competitors on many viewpoints: classification accuracy, training time, test time. In order to compare the statistical significance of the observed difference we adopted the approach overviewed in [45]. It presents the statistical test proposed by Friedman on average rankings applied to classifiers performance results. We briefly summarize the test here.

1. The performance of each classifier on a certain issue (accuracy, training times, etc) is determined on each data-set.
2. The classifiers are ranked on each data-set according to the results.
3. For each classifier, its position in the various rankings is recorded and its average position w.r.t. the data-sets is computed. This is the resulting value that we give in output in the tables. The advantage is that it allows to present a comparison of multiple classifiers on multiple data-sets.
4. The observed differences between the average rankings are compared with the critical difference CD which establishes whether the differences are statistically significant: $CD = q_{\alpha} \sqrt{\frac{k(k+1)}{6N}}$ where $N$ is the number of datasets, $k$ is the total number of classifiers, $\alpha$ is the significance level and $q_{\alpha}$ is the critical value for $\frac{\alpha}{(k-1)}$ based on the Studentized range statistic.
Table 3: Optimal parameters values resulting from tuning process in LODE

<table>
<thead>
<tr>
<th>Data-set name</th>
<th>Cardinality level</th>
<th>Minimum support threshold</th>
<th>Percentage of features retained (mean of ten folds)</th>
<th>Standard deviation of retained features (ten folds)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analcatdata-Bankruptcy</td>
<td>3</td>
<td>0.01</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>Analdata-cyyoung8092</td>
<td>3</td>
<td>0.05</td>
<td>85.36</td>
<td>1.7</td>
</tr>
<tr>
<td>Analcatdata-Creditscore</td>
<td>4</td>
<td>0.01</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>Analcatdata-Lawsuit</td>
<td>3</td>
<td>0.01</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>German</td>
<td>4</td>
<td>0.3</td>
<td>75.37</td>
<td>2.36</td>
</tr>
<tr>
<td>BioMed</td>
<td>5</td>
<td>0.01</td>
<td>76.89</td>
<td>3.7</td>
</tr>
<tr>
<td>Bupa</td>
<td>3</td>
<td>0.05</td>
<td>87.47</td>
<td>1.73</td>
</tr>
<tr>
<td>Australia</td>
<td>3</td>
<td>0.3</td>
<td>84.3</td>
<td>1.9</td>
</tr>
<tr>
<td>Labor</td>
<td>4</td>
<td>0.1</td>
<td>96.9</td>
<td>1.13</td>
</tr>
<tr>
<td>Sonar</td>
<td>4</td>
<td>0.3</td>
<td>72.23</td>
<td>0.7</td>
</tr>
<tr>
<td>Wave</td>
<td>4</td>
<td>0.2</td>
<td>75.39</td>
<td>2.33</td>
</tr>
<tr>
<td>Lymph</td>
<td>4</td>
<td>0.3</td>
<td>89.9</td>
<td>7.81</td>
</tr>
<tr>
<td>Diabetes</td>
<td>3</td>
<td>0.05</td>
<td>78.37</td>
<td>1.3</td>
</tr>
<tr>
<td>Haberman</td>
<td>2</td>
<td>0.01</td>
<td>93.13</td>
<td>2.37</td>
</tr>
<tr>
<td>Horse</td>
<td>2</td>
<td>0.15</td>
<td>90.71</td>
<td>1.29</td>
</tr>
<tr>
<td>HD</td>
<td>3</td>
<td>0.2</td>
<td>86.43</td>
<td>5.18</td>
</tr>
<tr>
<td>Hepatitis</td>
<td>4</td>
<td>0.2</td>
<td>89.5</td>
<td>1.3</td>
</tr>
<tr>
<td>Heartstatlog</td>
<td>3</td>
<td>0.1</td>
<td>81.13</td>
<td>3.29</td>
</tr>
<tr>
<td>Monkst</td>
<td>5</td>
<td>0.01</td>
<td>85.16</td>
<td>4.33</td>
</tr>
<tr>
<td>Prnsynth</td>
<td>2</td>
<td>0.01</td>
<td>87.31</td>
<td>1.36</td>
</tr>
<tr>
<td>Titanic</td>
<td>2</td>
<td>0.01</td>
<td>85.3</td>
<td>2.19</td>
</tr>
<tr>
<td>Wisconsin-Breast-Cancer</td>
<td>4</td>
<td>0.01</td>
<td>92.27</td>
<td>1.39</td>
</tr>
<tr>
<td>Cars</td>
<td>5</td>
<td>0.01</td>
<td>93.14</td>
<td>2.79</td>
</tr>
<tr>
<td>Cinc</td>
<td>4</td>
<td>0.05</td>
<td>90.12</td>
<td>1.36</td>
</tr>
<tr>
<td>Iris</td>
<td>4</td>
<td>0.05</td>
<td>97.1</td>
<td>1.7</td>
</tr>
<tr>
<td>Tae</td>
<td>2</td>
<td>0.05</td>
<td>94.13</td>
<td>3.69</td>
</tr>
<tr>
<td>Grubsdamage</td>
<td>4</td>
<td>0.05</td>
<td>95.1</td>
<td>4.1</td>
</tr>
<tr>
<td>Vehicle</td>
<td>4</td>
<td>0.2</td>
<td>93.24</td>
<td>1.3</td>
</tr>
<tr>
<td>Analdata-Drift</td>
<td>3</td>
<td>0.1</td>
<td>97.4</td>
<td>1.2</td>
</tr>
<tr>
<td>Glassp</td>
<td>3</td>
<td>0.01</td>
<td>91.39</td>
<td>2.1</td>
</tr>
<tr>
<td>Ecoli</td>
<td>5</td>
<td>0.05</td>
<td>96.27</td>
<td>1.7</td>
</tr>
<tr>
<td>Yeast</td>
<td>3</td>
<td>0.1</td>
<td>98.1</td>
<td>1.3</td>
</tr>
</tbody>
</table>
In our tests we used the value of $\alpha$ equal to 0.05 and the corresponding value of $q_\alpha$ equal to 2.394.

The first important observation that comes out from the Table in Figure 11 is that LODE outperforms the other learners as regards the classification accuracy (on test set, with 10-folds cross-validation). LODE mean ranking is equal to 1.46875 and since the second ranked classifier is SMO with a mean rank of 4.15625 and the value of $CD$ is equal to 1.864995 the observed differences are statistically significant. The graphical representation of the Friedman test on the differences in classification accuracy is presented in Figure 12.

<table>
<thead>
<tr>
<th>Classifier Name</th>
<th>Mean Rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>LODE</td>
<td>1.46875</td>
</tr>
<tr>
<td>SMO</td>
<td>4.15625</td>
</tr>
<tr>
<td>CBA</td>
<td>4.203125</td>
</tr>
<tr>
<td>KNN</td>
<td>5.03125</td>
</tr>
<tr>
<td>NB</td>
<td>5.515625</td>
</tr>
<tr>
<td>J48</td>
<td>5.578125</td>
</tr>
<tr>
<td>L3</td>
<td>6.078125</td>
</tr>
<tr>
<td>RIPPER</td>
<td>6.171875</td>
</tr>
<tr>
<td>DTABLE</td>
<td>6.796875</td>
</tr>
</tbody>
</table>

Figure 11: Mean ranking on classification accuracy of classifiers on original datasets (test set)

In Figure 13 we show the analogous statistical test on the difference between the accuracy on training-set and the accuracy on test-set. This experiment aims at putting in evidence the capability of a classifier to escape from over-fitting. We can see that NB is ranked first, CBA second and LODE third but these differences are not statistically significant.

In Table 4 we report the details of the classification error (and of the difference between the error on the training-set and on the test-set) of LODE and of SMO, which is among the learners the one whose performance gets closer to LODE. We reported also the details regarding the kind of datasets (number of classes, balanced classes, number of features) in order to ascertain if the obtained good results could be also related to the typology of data or not. From the results it is clear that good results could be obtained both in binary and in multi-class data, balanced or unbalanced, high-dimensional or not.

In Figure 15 and in Figure 17 we show the results on execution times. NB
Table 4: Comparison with SVM (SMO)

<table>
<thead>
<tr>
<th>Dataset Name</th>
<th>N. classes</th>
<th>High dimens. (N. features)</th>
<th>Balanced</th>
<th>LODE</th>
<th>SMO</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Test err</td>
<td>Train-Test err</td>
</tr>
<tr>
<td>Anal.-Bank.</td>
<td>2</td>
<td>N (7)</td>
<td>Y</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Anal.-young8092</td>
<td>2</td>
<td>Y (11)</td>
<td>N</td>
<td>13.36</td>
<td>0.66</td>
</tr>
<tr>
<td>Anal.-Credit.</td>
<td>2</td>
<td>N (7)</td>
<td>N</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Anal.-Lawsuit</td>
<td>2</td>
<td>N (5)</td>
<td>N</td>
<td>1.05</td>
<td>0.05</td>
</tr>
<tr>
<td>BioMed</td>
<td>2</td>
<td>N (9)</td>
<td>N</td>
<td>2.89</td>
<td>1.19</td>
</tr>
<tr>
<td>Bupa</td>
<td>2</td>
<td>N (7)</td>
<td>Y</td>
<td>31.05</td>
<td>0.96</td>
</tr>
<tr>
<td>Credit-a</td>
<td>2</td>
<td>Y (16)</td>
<td>Y</td>
<td>12</td>
<td>1.1</td>
</tr>
<tr>
<td>Diabetes</td>
<td>2</td>
<td>N (9)</td>
<td>N</td>
<td>17.66</td>
<td>1.36</td>
</tr>
<tr>
<td>Haberman</td>
<td>2</td>
<td>N (4)</td>
<td>N</td>
<td>22.7</td>
<td>2.23</td>
</tr>
<tr>
<td>Horse</td>
<td>2</td>
<td>Y (28)</td>
<td>N</td>
<td>17.7</td>
<td>2</td>
</tr>
<tr>
<td>HD</td>
<td>2</td>
<td>Y (14)</td>
<td>Y</td>
<td>12.43</td>
<td>2.05</td>
</tr>
<tr>
<td>Hepatitis</td>
<td>2</td>
<td>Y (20)</td>
<td>N</td>
<td>11.35</td>
<td>1.05</td>
</tr>
<tr>
<td>Heartstatlog</td>
<td>2</td>
<td>Y (14)</td>
<td>Y</td>
<td>12.37</td>
<td>0.98</td>
</tr>
<tr>
<td>MonkSl</td>
<td>2</td>
<td>N (5)</td>
<td>Y</td>
<td>44.67</td>
<td>0</td>
</tr>
<tr>
<td>Prnsynth</td>
<td>2</td>
<td>N (3)</td>
<td>Y</td>
<td>12.4</td>
<td>0.71</td>
</tr>
<tr>
<td>Titanic</td>
<td>2</td>
<td>N (4)</td>
<td>N</td>
<td>20.1</td>
<td>0.04</td>
</tr>
<tr>
<td>Wiscon.-B.</td>
<td>2</td>
<td>Y (10)</td>
<td>N</td>
<td>2.27</td>
<td>0.88</td>
</tr>
<tr>
<td>Cars</td>
<td>3</td>
<td>N (9)</td>
<td>N</td>
<td>20.3</td>
<td>1.2</td>
</tr>
<tr>
<td>Cmc</td>
<td>3</td>
<td>Y (10)</td>
<td>N</td>
<td>40.3</td>
<td>2.7</td>
</tr>
<tr>
<td>Iris</td>
<td>3</td>
<td>N (5)</td>
<td>Y</td>
<td>4</td>
<td>2.25</td>
</tr>
<tr>
<td>Tae</td>
<td>3</td>
<td>N (6)</td>
<td>Y</td>
<td>42.3</td>
<td>1.61</td>
</tr>
<tr>
<td>Grubdamage</td>
<td>4</td>
<td>N (9)</td>
<td>N</td>
<td>48.19</td>
<td>3.49</td>
</tr>
<tr>
<td>Vehicle</td>
<td>4</td>
<td>Y (19)</td>
<td>Y</td>
<td>22.97</td>
<td>3.27</td>
</tr>
<tr>
<td>Anal.-Dmft</td>
<td>6</td>
<td>N (5)</td>
<td>Y</td>
<td>74.2</td>
<td>7.1</td>
</tr>
<tr>
<td>Glassp</td>
<td>6</td>
<td>Y (10)</td>
<td>N</td>
<td>23.6</td>
<td>1.6</td>
</tr>
<tr>
<td>Ecoli</td>
<td>8</td>
<td>N (8)</td>
<td>N</td>
<td>12.3</td>
<td>0.6</td>
</tr>
<tr>
<td>Yeast</td>
<td>10</td>
<td>N (9)</td>
<td>N</td>
<td>40.96</td>
<td>3.8</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td></td>
<td></td>
<td>21.1133</td>
<td>22.6442</td>
</tr>
</tbody>
</table>

is first, KNN is second and Ripper is third. LODE ranks only 7th. However, we must say that the results of these experiments favour those learners (like NB and KNN) which do not have parameters or have a few parameters only. In these cases, the parameter tuning time is totally eliminated or strongly reduced. In fact, tuning time depends both on the number of parameters and on the range of values tested for each of them. In Section 7.1 we report also more studies on the family of LODE classifiers with alternative feature selection methods. Further experiments are conducted in order to test also if the Simulated Annealing approach to tuning allows to get significant improvements in accuracy and training times. Our answers will be positive.

In Figure 17 we report results on Friedman test on the mean execution time needed by each classifier to predict a single test instance. In this experiment LODE ranks 6th. We can notice that all the classifiers that have a class model composed by a multitude of patterns, like L3, DTABLE and
LODE, are slow in testing, while decision trees, NB and RIPPER, that have mechanisms of strong pruning of rules, are faster.

<table>
<thead>
<tr>
<th>Classifier Name</th>
<th>Mean Rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>threshold (CD)</td>
<td>1.864995</td>
</tr>
<tr>
<td>J48</td>
<td>2.109375</td>
</tr>
<tr>
<td>NB</td>
<td>2.219785</td>
</tr>
<tr>
<td>RIPPER</td>
<td>3.734375</td>
</tr>
<tr>
<td>KNN</td>
<td>4.796875</td>
</tr>
<tr>
<td>CBA</td>
<td>5.5625</td>
</tr>
<tr>
<td>LODE</td>
<td>5.578125</td>
</tr>
<tr>
<td>DTABLE</td>
<td>5.96875</td>
</tr>
<tr>
<td>SMO</td>
<td>6.5625</td>
</tr>
<tr>
<td>L3</td>
<td>8.46875</td>
</tr>
</tbody>
</table>

Figure 17: Mean ranking on unit test time on original datasets

Experiments on noisy datasets. One of the believed benefits of our model-based classifier is that it is supposed to be more robust to the presence of noise in data. This occurs in virtue of the prototypical class models of LODE that represent the frequent characteristics of examples on which noise super-imposition should not have many effects. In order to establish concretely this claim we experimented with the datasets with a variable amount of noise. We added noise to the datasets in the form of a random change of the class. We varied the percentage of noise from 5% until 20% of the instances.

Figure 19 and Figure 21 show classification results on the datasets affected by noise at the extreme values of this range (5% and 20%). These experiments clearly show that LODE outperforms the other classifiers, and the improvement increases especially with an increasing amount of noise. When comparing the results in Figure 19 with Figure 21, some algorithms such as NB or J48 appear to improve their classification accuracy. This is due to the fact that NB and J48 predictive mechanisms rely on a global model that is able to generalize better and results more robust w.r.t. a marked presence of noise. The other learners (like KNN, SMO, RIPPER, CBA) instead, can sustain only lower levels of noise because they make predictions that are based on a restricted number of local patterns. With a marked level of noise they lost at a greater extent their accuracy, due to the noisy modification of the data values that results detrimental for their final prediction.

As regards the effect of over-fitting, we monitored the differences between accuracy in training and test set. In Figure 23 we can notice that LODE is
ranked second when the level of noise in the datasets is set to the level of 5%; LODE improved its position w.r.t. the analogous experiment on the original datasets (it was third). Furthermore, when the level of noise is increased to 20% LODE reaches the first position in the ranking (see Figure 25).

7.1. Impact on LODE of Feature Selection and Composite Features

In this new set of experiments we want to determine the relative impact that the different settings have to the performance of LODE. In other terms, we want to establish if the positive results observed in terms of classification accuracy and execution times are due to the feature selection results and the corresponding wrapper (in particular, to the quite sophisticated tuning process based on Simulated Annealing) or to the fact that composite features (itemsets) are adopted instead of simple ones (items). We therefore carry out a performance study on a family of learners based on LODE, in which these
issues are taken one by one or in combinations. We furthermore consider two types of wrappers: a Simple Backward Elimination (SBE) in which itemsets are eliminated from the bottom of the rankings - one by one - until the first minimum is found and a Backward Elimination by Simulated Annealing (SA). In this experiment, we wish to establish if SA is relevant or not.

Results on classification error are reported in Figure 27 while results on execution times for the classification model generation are reported in Figure 5.

We can notice two issues. First, adopting composite features (itemsets) is extremely beneficial. All the results of LODE with composite features are significantly improved with respect to the experiments in which simple features (items) are adopted. Second, using a feature reduction method is always beneficial, both when composite features (itemsets) are used and
when simple features (items only) are adopted. Furthermore, the observed differences are statistically significant when the feature reduction method is based on SA.

From Figure 5, in which results on training times are reported, we can notice another main benefit of SA: not only it improves accuracy results but also it is useful to speed-up training execution times. This result is due to its faster achievement of the optimal working point because of the “jumps” it performs based on the big reduction rate.

8. Conclusions

In this paper we have proposed LODE, a new classifier whose class models are composed of frequent itemsets extracted from the instances of each class in the training set. Class models have been generated originally for a descriptive purpose but can be employed for prediction if we the number of the itemsets is reduced by a wrapper approach. Prediction occurs by computation of the distances between two vectors that represent respectively a class model and a test example.

The adoption of a model-based distance is an advantage w.r.t. learners based on local patterns that apply the best single local pattern for each example. Indeed, using an ensemble of local patterns for prediction reduces
the chance of model overfitting and is more robust w.r.t. the presence of noise in the data.

We have validated this new approach to classification by several experiments on many available UCI datasets, with and without noise. We have shown that LODE outperforms traditional IBL such as Knn and other classifiers in the state of the art, like decision trees, SVM, NB and rule-based classifiers. In a related paper [20] we have experimented with different techniques of itemsets ranking employed in order to simplify class models: accuracy, KL divergence, strong jumping emerging patterns and an entropy based measure (normalized ∆). Experiments showed that the observed good performances of our classifier are due not only to the mechanism of model-based distance computation - that works on itemsets features and considers both the present and the absent features from the test instances - but also to the effectiveness of ∆.

In future work, we plan to investigate on learning the features weights in the ensembles and in employing further mechanisms of feature generation and selection based on principles of subgroup discovery and on predictive and discriminative capabilities.

References


[35] R. Meo, Maximum independence and mutual information, IEEE TRANSACTIONS ON INFORMATION THEORY 48 (1).


