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A Fuzzy Association Rule-Based Classification Model for High-Dimensional Problems with Genetic Rule Selection and Lateral Tuning

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Abstract—The inductive learning of fuzzy rule based classification systems suffers from exponential growth of the fuzzy rule search space when the number of patterns and/or variables becomes high. This growth makes the learning process more difficult and, in most cases, it leads to problems of scalability (in terms of the time and memory consumed) and/or complexity (with respect to the number of rules obtained and the number of variables included in each rule).

In this work, we propose a fuzzy association rule-based classification method for high-dimensional problems based on three stages to obtain an accurate and compact fuzzy rule based classifier with a low computational cost. This method limits the order of the associations in the association rule extraction and considers the use of subgroup discovery based on an Improved Weighted Relative Accuracy measure to preselect the most interesting rules before a genetic post-processing process for rule selection and parameter tuning. The results obtained over twenty-six real-world datasets of different sizes and with different numbers of variables demonstrate the effectiveness of the proposed approach.

Index Terms—Data mining, associative classification, classification, fuzzy association rules, genetic algorithms, genetic fuzzy rule selection, high-dimensional problems.

I. INTRODUCTION

F UZZY Rule Based Classification Systems (FRBCSs) [1], [2] are useful and well-known tools in the machine learning framework, since they can provide an interpretable model for the end user [3]–[6]. There are many real applications in which FRBCSs have been employed, including anomaly intrusion detection [7], image processing [8], among others. In most of these areas the available or useful data consists of a high number of patterns (instances or examples) and/or variables. In this situation, the inductive learning of FRBCSs suffers from exponential growth of the fuzzy rule search space. This growth makes the learning process more difficult and, in most cases, it leads to problems of scalability (in terms of the time and memory consumed) and/or complexity (with respect to the number of rules obtained and the number of variables included in each rule) [9], [10].

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Association discovery is one of the most common Data Mining techniques used to extract interesting knowledge from large datasets [11]. Many efforts have been made to use its advantages for classification under the name of associative classification [12]–[19]. Association discovery aims to find interesting relationships between the different items in a database [20], while classification aims to discover a model from training data that can be used to predict the class of test patterns [21]. Both association discovery and classification rules mining are essential in practical Data Mining applications [11], [22] and their integration could result in greater savings and convenience for the user.

A typical associative classification system is constructed in two stages:

- 1) discovering the association rules inherent in a database;
- 2) selecting a small set of relevant association rules to construct a classifier.

In order to enhance the interpretability of the obtained classification rules and to avoid unnatural boundaries in the partitioning of the attributes, different studies have been presented to obtain classification systems based on fuzzy association rules [23]-[28]. For instance, in [24] the authors have made use of a Genetic Algorithm (GA) [29], [30] to automatically determine minimum support and confidence thresholds, mining for each chromosome a fuzzy rule set for classification by means of an algorithm based on the Apriori algorithm [31] and adjusting the fuzzy confidence of these rules with the approach proposed by Nozaki et al in [32]. Consequently, this approach can only be used for small problems since its computational cost is very high when we consider problems that consist of a high number of patterns and/or variables. On the other hand, in [25] the authors used an algorithm based on the Apriori algorithm to mine association rules only up to a certain level and to select the K most confident ones for each class among them, in order to finally employ a genetic rule selection method that obtains a classifier from them. However, many patterns may be uncovered if we only consider the confidence measure to select the candidate rules.

In this paper we present a Fuzzy Association Rule-based Classification method for High-Dimensional problems (FARC-HD) to obtain an accurate and compact fuzzy rule-based classifier with a low computational cost. This method is based on three stages:

- Fuzzy association rule extraction for classification: A search tree is employed to list all possible frequent fuzzy itemsets and to generate fuzzy association rules for classification, limiting the depth of the branches in order to find a small number of short (i.e., simple) fuzzy rules.
- 2) Candidate rule prescreening: Even though the order of the associations is limited in the association rule extraction, the number of rules generated can be very large. In order to decrease the computational cost of the genetic post-processing stage we consider the use of subgroup discovery based on an improved Weighted Relative Accuracy measure (wWRAcc') to preselect the most interesting rules by means of a pattern weighting scheme [33].
- 3) Genetic rule selection and lateral tuning: Finally, we make use of GAs to select and tune a compact set of fuzzy association rules with high classification accuracy in order to consider the known positive synergy that both techniques present (selection and tuning). Several works have successfully combined the selection of rules with the tuning of membership functions (MFs) within the same process [34], [35], taking advantage of the possibility of different coding schemes that GAs provide. The successful application of GAs to identify fuzzy systems has led to the so-called Genetic Fuzzy Systems (GFSs) [36]–[38].

In order to assess the performance of the proposed approach, we have used twenty-six real-world datasets with a number of variables ranging from 4 to 90 and a number of patterns ranging from 150 to 19020. We have developed the following studies; first, we have shown the results obtained from comparison with three other GFSs [38]. Second, we have compared the performance of our approach with two approaches to obtain fuzzy associative classifiers. Third, we have shown the results obtained from the comparison with four other classical approaches for associative classification and with the C4.5 decision tree [39]. Furthermore, in these studies we have made use of some non-parametric statistical tests for pair-wise and multiple comparison [40]-[43] of the performance of these classifiers. Then, we have shown a study on the influence of the depth of the trees and the number of evaluations in the genetic selection and tuning process. Finally, we have analyzed the scalability of the proposed approach.

This paper is arranged as follows. The next section introduces the type of rules, rule weights and inference model used, and the basic definitions for fuzzy association rules and associative classification. Section III describes in detail each stage of the proposed approach. Section IV presents the experimental set-up. Section V shows and discusses the results obtained on the twenty-six real-world datasets. Finally, in Section VI some concluding remarks are made.

II. PRELIMINARIES

In this section, we first describe FRBCSs. Then, we introduce the basic definitions for fuzzy association rules. Finally, we present fuzzy association rules for classification.

A. Fuzzy rule based classification systems

Any classification problem consists of N training patterns $x_p = (x_{p1}, ..., x_{pm}), p = 1, 2, ..., N$ from S classes where x_{pi} is the *i*-th attribute value (i = 1, 2, ..., m) of the *p*-th training pattern. In this work, we use fuzzy rules of the following form for our classifier:

Rule
$$R_j$$
: If x_1 is A_{j1} and ... and x_m is A_{jm}
then Class = C_j with RW_j ;

where R_j is the label of the *j*-th rule, $x = (x_1, ..., x_m)$ is an m-dimensional pattern vector, A_{ji} is an antecedent fuzzy set, C_j is a class label and RW_j is the rule weight.

The rule weight of each fuzzy rule R_j has a great effect on the performance of fuzzy rule-based classifiers [44]. Different specifications of the rule weight have been proposed and examined in the literature. In [45] we can find some heuristic methods for rule weight specification. In this work, we employ the most common one, the fuzzy confidence value or Certainty Factor (CF) [46].

$$RW_{j} = CF_{j} = \frac{\sum_{x_{p} \in ClassC_{j}} \mu_{A_{j}}(x_{p})}{\sum_{p=1}^{N} \mu_{A_{j}}(x_{p})}$$
(1)

where $\mu_{A_j}(x_p)$ is the matching degree of the pattern x_p with the antecedent part of the fuzzy rule R_j . We use the fuzzy reasoning method of the weighted vote or additive combination [46] for classifying new patterns by the Rule Base (RB). With this method, each fuzzy rule casts a vote for its consequent class. The total strength of the vote for each class is computed as follows:

$$V_{Class_{h}}(x_{p}) = \sum_{R_{j} \in RB; \ C_{j} = h} \mu_{A_{j}}(x_{p}) \cdot CF_{j}$$

$$h = 1, 2, ..., S; \ R_{j} \in RB$$
(2)

The new pattern x_p is classified as the class with the maximum total strength of the vote. If multiple class labels have the same maximum value for x_p or no fuzzy rule is compatible with x_p , this pattern is classified as the class with most patterns in the training data.

B. Fuzzy association rules

Association rules are used to represent and identify dependencies between items in a database [11], [20]. They are expressions of the type $A \rightarrow B$, where A and B are sets of items and $A \cap B = \emptyset$. This means that, if all the items in A exist in a transaction then all the items in B with a high probability are also in the transaction, and A and B should not have any common items [31]. There are many previous studies for mining association rules focused on databases with binary or discrete values, however the data in real-world applications usually consist of quantitative values. Designing Data Mining algorithms, able to deal with various types of data, presents a challenge to workers in this research field.

Fuzzy set theory has been used more and more frequently in Data Mining because of its simplicity and similarity to human reasoning [1]. The use of fuzzy sets to describe associations



Fig. 1. Attributes and linguistic terms for the attributes X_1 and X_2

between data extends the types of relationships that may be represented, facilitates the interpretation of rules in linguistic terms, and avoids unnatural boundaries in the partitioning of the attribute domains. For this reason, in recent years different studies have proposed methods for mining fuzzy association rules from quantitative data [47]–[54].

Let us consider a simple database T with two attributes $(X_1$ and $X_2)$ and three linguistic terms with their associated MFs (see Fig. 1). Based on this definition, a simple example of fuzzy association rule is X_1 is $Middle \rightarrow X_2$ is High.

Support and confidence are the most common measures of interest of an association rule. These measures can be defined for fuzzy association rules as follows:

$$Support(A \to B) = \frac{\sum_{x_p \in T} \mu_{AB}(x_p)}{\mid N \mid}$$
(3)

$$Confidence(A \to B) = \frac{\sum_{x_p \in T} \mu_{AB}(x_p)}{\sum_{x_p \in T} \mu_A(x_p)}$$
(4)

where |N| is the number of transactions in T, $\mu_A(x_p)$ is the matching degree of the transaction x_p with the antecedent part of the rule and $\mu_{AB}(x_p)$ is the matching degree of the transaction x_p with the antecedent and consequent of the rule.

C. Fuzzy association rules for classification

In the last few years, different studies have proposed methods to obtain fuzzy association rule-based classifiers [23]– [28]. The task of classification is to find a set of rules in order to identify the classes of undetermined patterns. A fuzzy association rule can be considered to be a classification rule if the antecedent contains fuzzy item sets and the consequent part contains only one class label ($C = \{C_1, ..., C_j, ..., C_S\}$). A fuzzy associative classification rule $A \rightarrow C_j$ could be measured directly in terms of support and confidence as follows:

$$Support(A \to C_j) = \frac{\sum_{x_p \in ClassC_j} \mu_A(x_p)}{|N|}$$
(5)

$$Confidence(A \to C_j) = \frac{\sum_{x_p \in ClassC_j} \mu_A(x_p)}{\sum_{x_p \in T} \mu_A(x_p)}$$
(6)

III. FARC-HD: FUZZY ASSOCIATION RULE-BASED CLASSIFIER FOR HIGH-DIMENSIONAL PROBLEMS

In this section, we will describe our proposal to obtain a fuzzy association rule-based classifier for high dimensional problems. This method is based on three stages:

- Fuzzy association rule extraction for classification: A search tree is employed to list all the possible frequent fuzzy itemsets and to generate fuzzy association rules for classification.
- 2) Candidate rule prescreening: A rule evaluation criterion is used to preselect candidate fuzzy association rules.
- Genetic rule selection and lateral tuning: The best cooperative rules are selected and tuned by means of a GA, considering the positive synergy between both techniques within the same process.

Finally, we add a default rule considering the class with the most patterns in the training data. In the following subsections we will introduce the three mentioned stages, explaining in detail all their characteristics (subsections III-A, III-B and III-C) and presenting a flowchart of the algorithm (subsection III-D).

A. Stage 1. Fuzzy association rule extraction for classification

To generate the RB we employ a search tree to list all the possible fuzzy itemsets of a class. The root or level 0 of a search tree is an empty set. All attributes are assumed to have an order (in our case, the order of appearance in the training data), and the one-itemsets corresponding to the attributes are listed in the first level of the search tree according to their order. If an attribute has j possible outcomes (q_i linguistic terms for each quantitative attribute), it will have j one-item sets listed in the first level. The children of a one-item node for an attribute A are the two-item sets that include the oneitem set of attribute A and a one-item set for another attribute behind attribute A in the order, and so on. If an attribute has i > 2 possible outcomes, it can be replaced by j binary variables to ensure that no more than one of these j binary attributes can appear in the same node in a search tree. An example with two attributes $(V_1 \text{ and } V_2)$ with two linguistic terms (L and H) is detailed in Figure 2. An itemset with a support higher than the minimum support is a frequent itemset. If the support of an n-item set in a node J is less than the minimum support, it does not need to be extended more because the support of any item set in a node in the subtree led by node J will also be less than the minimum support. Likewise, if a candidate item set generates a classification rule with confidence higher than the maximum confidence, this rule has reached the quality level demanded by the user and it is again unnecessary to extend it further. These properties greatly reduce the number of nodes needed for searching. The fuzzy support of an itemset can be calculated as follows:

$$Support(A) = \frac{\sum_{x_p \in T} \mu_A(x_p)}{|N|}$$
(7)

where $\mu_A(x_p)$ is the matching degree of the pattern x_p with the itemset. The matching degree $\mu_A(x_p)$ of x_p to the different fuzzy regions is computed using a conjunction operator, in our case, the product T-norm.

Once all frequent fuzzy itemsets have been obtained, the candidate fuzzy association rules for classification can be generated, setting the frequent fuzzy itemsets in the antecedent of the rules and the corresponding class in the consequent. This process is repeated for each class.



Fig. 2. The search tree for two quantitative attributes V_1 and V_2 with two linguistic terms L and H

The number of frequent fuzzy itemsets extracted depends directly on the minimum support. The minimum support is usually calculated considering the total number of patterns in the dataset, however the number of patterns for each class in a dataset can be different. For this reason, our algorithm determines the minimum support of each class by the distributions of the classes over the dataset. Thus, the minimum support for class C_j is defined as:

$$MinimumSupport_{C_i} = minSup * f_{C_i}$$
(8)

where minSup is the minimum support determined by the expert and f_{C_i} is the pattern ratio of the class C_i .

In this stage we can generate a large number of candidate fuzzy association rules for classification. It is, however, very difficult for human users to handle such a large number of generated fuzzy rules and to intuitively understand long fuzzy rules with many antecedent conditions. For this reason, we only generate short fuzzy rules and with only a small number of antecedent conditions. Thus, the depth of the trees is limited to a fixed value $(Depth_{max})$ determined by an expert.

B. Stage 2. Candidate rule prescreening

In the previous stage, we can generate a large number of candidate rules. In order to decrease the computational costs of stage 3, we consider the use of subgroup discovery to preselect the most interesting rules from RB obtained in the previous stage by means of a pattern weighting scheme [33]. This scheme treats the patterns in such a way that covered positive patterns are not deleted when the current best rule is selected. Instead, each time a rule is selected, the algorithm stores a count i for each pattern of how many times (with how many of the selected rules) the pattern has been covered.

Weights of positive patterns covered by the selected rule decrease according to the formula $w(e_j, i) = \frac{1}{i+1}$. In the first iteration all target class patterns are assigned the same weight $w(e_j, 0) = 1$, while in the following iterations the contributions of patterns are inversely proportional to their coverage by previously selected rules. In this way the patterns already covered by one or more selected rules decrease their weights while uncovered target class patterns whose weights have not been decreased will have a greater chance of being covered in the following iterations. Covered patterns are completely eliminated when they have been covered more than k_t times.

TABLE I												
THE FIVE PATTERNS IN THIS EXAMPLE												
\overline{ID}	X_1	X_2	Class	Weight								
ID1	0.0	10.0	C_1	1.0								
ID2	2.5	4.0	C_2	1.0								
ID3	3.2	1.0	C_2	0.0								
ID4	9.0	5.0	C_2	1.0								
ID5	2.5	10	C_1	0.5								

Thus, in each iteration of the process the rules are ordered according to a rule evaluation criteria from best to worst. The best rule is selected, covered patterns are reweighted, and the procedure repeats these steps until one of the stopping criteria is satisfied: either all patterns have been covered more than k_t times, or there are no more rules in the RB. This process is to be repeated for each class.

wWRAcc' was used to evaluate the quality of intervalar rules in APRIORI-SD [33]. This measure was defined as follows:

$$wWRAcc'(A \to C_j) = \frac{n'(A)}{N'} \cdot \left(\frac{n'(A \cdot C_j)}{n'(A)} - \frac{n(C_j)}{N}\right)$$
(9)

where N' is the sum of the weights of all patterns, n'(A) is the sum of the weights of all covered patterns, $n'(A \cdot C_j)$ is the sum of the weights of all correctly covered patterns, $n(C_j)$ the number of patterns of class C_j , and N is the number of all patterns. For instance, let us consider a simple database with two attributes $(X_1 \text{ and } X_2)$, two classes $(C_1 \text{ and } C_2)$ and five training patterns. Table I shows the five training patterns and their weights in the *p*-th iteration of the process. In this iteration, the wWRAcc' value of a simple rule $R = If X_1 is [0.0, 5.0[and X_2 is [5.0, 10.0] \rightarrow C_1$ is calculated as follows:

$$wWRAcc'(R) = \frac{1.0 + 0.5}{1.0 + 1.0 + 0.0 + 1.0 + 0.5} \cdot (\frac{1.0 + 0.5}{1.0 + 0.5} - \frac{2}{5})$$

= 0.257

We have modified this measure to enable the handling of fuzzy rules. The new measure is defined as follows:

$$wWRAcc''(A \to C_j) = \frac{n''(A \cdot C_j)}{n'(C_j)} \cdot \left(\frac{n''(A \cdot C_j)}{n''(A)} - \frac{n(C_j)}{N}\right)$$
(10)

where n''(A) is the sum of the products of the weights of all covered patterns by their matching degrees with the antecedent part of the rule, $n''(A \cdot C_j)$ is the sum of the products of the weights of all correctly covered patterns by their matching degrees with the antecedent part of the rules, and $n'(C_j)$ is the sum of the weights of patterns of class C_j . Moreover, the first term in the definition of wWRAcc' has been replaced by $\frac{n''(A \cdot C_j)}{n(C_j)}$ to reward rules which cover uneliminated patterns of the class C_j .

Let us consider three linguistic terms for the attributes X_1 and X_2 (see Fig. 1). Based on this definition, a simple example of fuzzy association rule for classification is $R = If X_1$ is Low and X_2 is $High \rightarrow C_1$. This rule covers the training patterns in Table I with degrees: (ID1, 1.0), (ID2, 0.0), (ID3, 0.0), (ID4, 0.0) and (ID5, 0.5). In this situation, the wWRAcc" value of this rule is calculated as follows:

$$wWRAcc''(R) = \frac{1.0 * 1.0 + 0.5 * 0.5}{1.0 + 0.5} \cdot \left(\frac{1.0 * 1.0 + 0.5 * 0.5}{1.0 * 1.0 + 0.5 * 0.5} - \frac{2}{5}\right) = 0.5$$

This measure can obtain positive or negative values in the interval [-1.0, 1.0]. A rule with a wWRAcc" value near to one may be more useful for the classification.

C. Stage 3. Rule Selection and Lateral Tuning

We consider the use of GAs to select and tune a compact set of fuzzy association rules with high classification accuracy from the RB obtained in the previous stage. We consider the approach proposed in [35] where rules are based on the linguistic 2-tuples representation [55]. This representation allows the lateral displacement of the labels considering only one parameter (symbolic translation parameter), which involves a simplification of the tuning search space that eases the derivation of optimal models, particularly when it is combined with a rule selection within the same process enabling it to take advantage of the positive synergy that both techniques present. In this way, this process for contextualizing the MFs enables them to achieve a better covering degree while maintaining the original shapes, which results in accuracy improvements without a significant loss in the interpretability of the fuzzy labels. The symbolic translation parameter of a linguistic term is a number within the interval [-0.5, 0.5) that expresses the domain of a label when it is moving between its two lateral labels. Let us consider a set of labels S representing a fuzzy partition. Formally, we have the pair, $(S_i, \alpha_i), S_i \in S, \alpha_i \in$ [-0.5, 0.5). An example is illustrated in Fig. 3 where we show the symbolic translation of a label represented by the pair $(S_2,$ -0.3).

Let us consider the simple problem presented in the previous section. Based on this definition, an example of classic rule and linguistic 2-tuples represented rule is:

Classic Rule:

If X_1 is Low and X_2 is Middle then *Class* is C_1 .

2-Tuples Representation:



a) Simbolic Translation of a Linguistic Term



b) Lateral Displacement of a Membership Function

Fig. 3. Symbolic Translation of a Linguistic Label and Lateral Displacement of the involved MF

If X_1 is (Low, 0.1) and X_2 is (Middle, -0.3) then *Class* is C_1

In [35], two different rule representation approaches were proposed, a global approach and a local approach. In our particular case, the tuning is applied to the level of linguistic partitions (global approach). In this way, the pair (X_i , label) takes the same α value in all the rules where it is considered, i.e., a global collection of 2-tuples is considered by all the fuzzy rules. For example, X_1 is (High, 0.3) will present the same value for those rules in which the pair " X_1 is High" was initially considered. This proposal decreases the tuning problem complexity, greatly easing the derivation of optimal models. Another important issue is that, from the parameters α applied to each label, we could obtain the equivalent triangular MFs, by which a FRBCS based on linguistic 2-tuples could be represented as a classical Mamdani FRBCS. Notice that the class label and RW of the rule are not modified.

In the following, the main characteristics of the genetic approach that combines rule selection and lateral tuning are presented: genetic model, codification and initial gene pool, chromosome evaluation, crossover operator and restarting approach.

1) CHC Genetic Model: The approach proposed in [35] considers the use of a specific GA, the CHC algorithm [56].



Fig. 4. Scheme of CHC

The CHC algorithm is a GA that presents a good trade-off between exploration and exploitation, making it a good choice for problems with complex search spaces [57]. This genetic model makes use of a mechanism of Selection of Populations in order to perform an adequate global search. P parents and their corresponding offspring compete to select the best P individuals to take part in the next population. The CHC approach makes use of an incest prevention mechanism and a restarting process to encourage diversity in the population, instead of the well known mutation operator.

This incest prevention mechanism will be considered in order to apply the crossover operator, i.e., two parents are crossed if their hamming distance divided by 2 is over a predetermined threshold, L. This threshold value is initialized as the maximum possible distance between two individuals (the number of genes in the chromosome) divided by four. Following the original CHC scheme, L is decremented by one when there are no new individuals in the population in one generation. In order to make this procedure independent of the number of genes in the chromosome, in our case L will be decremented by a φ % of its initial value (with φ determined by the user, usually 10%). When L is below zero the algorithm restarts the population (for more information, see [58]).

A scheme of this algorithm is shown in Fig. 4.

2) Codification and Initial Gene Pool: To combine the rule selection with the global lateral tuning, a double coding scheme for both rule selection (C_S) and lateral tuning (C_T) is used:

• For the C_S part, each chromosome is a binary vector that determines when a rule is selected or not (alleles '1' and '0' respectively). Considering the M rules contained in the candidate rule set, the corresponding part $C_S = \{c_1, ..., c_M\}$ represents a subset of rules composing the final rule base, so that, $If c_i = 1$ then $(R_i \in RB)$ else $(R_i \notin RB)$,

with R_i being the corresponding *i*-th rule in the candidate rule set and RB being the final RB.

• For the C_T part, a real coding is considered. This part is the joint of the α parameters of each fuzzy partition. Let us consider the following number of labels per variable: (m^1, m^2, \ldots, m^n) , with *n* being the number of system variables. Then, this part has the following form (where each gene is associated to the tuning value of the corresponding label): $C_T = (c_{11}, \ldots, c_{1m^1}, c_{21}, \ldots, c_{2m^2}, \ldots, c_{n1}, \ldots, c_{nm^n})$.

Finally, a chromosome C is coded in the following way: $C = C_S C_T$. To make use of the available information, all the candidate rules are included in the population as an initial solution. To do this, the initial pool is obtained with the first individual having all genes with value '1' in the C_S part and all genes with value '0.0' in the C_T part. The remaining individuals are generated at random.

3) Chromosome Evaluation: To evaluate a determined chromosome penalizing a large number of rules, we compute the classification rate and the following function is maximized:

$$Fitness(C) = \frac{\#Hits}{N} - \delta \cdot \frac{NR_{initial}}{NR_{initial} - NR + 1.0}, \quad (11)$$

where #Hits is the number of patterns correctly classified (see subsection II-C), $NR_{initial}$ is the number of candidate rules, NR is the number of selected rules and δ is a weighting percentage given by the system expert that determines the tradeoff between accuracy and complexity. If there is at least one class without selected rules or if there are no covered patterns the fitness value of a chromosome will be penalized with the number of classes without selected rules and the number of uncovered patterns.

4) Crossover Operator: The crossover operator will depend on the chromosome part where it is applied:

- For the C_T part, we consider the Parent Centric BLX (PCBLX) operator [59] (an operator based on BLX-α). This operator is based on the concept of neighbourhood, which allows the offspring genes to be around the genes of one parent or around a wide zone determined by both parent genes. Let us assume that X = (x₁ ··· x_n) and Y = (y₁ ··· y_n), (x_i, y_i ∈ [a_i, b_i] ⊂ ℜ, i = 1 ··· n), are two real-coded chromosomes that are going to be crossed. We generate the two following offspring:
 - $O_1 = (o_{11} \cdots o_{1n})$, where o_{1i} is a randomly (uniformly) chosen number from the interval $[l_i^1, u_i^1]$, with $l_i^1 = max\{a_i, x_i I_i \cdot \alpha\}$, $u_i^1 = min\{b_i, x_i + I_i \cdot \alpha\}$, and $I_i = |x_i y_i|$.
 - $O_2 = (o_{21} \cdots o_{2n})$, where o_{2i} is a randomly (uniformly) chosen number from the interval $[l_i^2, u_i^2]$, with $l_i^2 = max\{a_i, y_i I_i \cdot \alpha\}$ and $u_i^2 = min\{b_i, y_i + I_i \cdot \alpha\}$.
- In the C_S part, the half uniform crossover scheme (HUX) is employed [58]. The HUX crossover exactly interchanges the mid of the alleles that are different in the parents (the genes to be crossed are randomly selected from among those that are different in the parents). This operator ensures the maximum distance of the offspring to their parents (exploration).

In this case, four offspring are generated by combining the two from the part C_T with the two from the part C_S . The two best offspring obtained in this way are considered as the two corresponding descendents. Notice that since we consider

a real coding scheme for the C_T part, the incest prevention mechanism has to transform each gene considering a Gray Code (binary code) with a fixed number of bits per gene (BITSGENE) that is determined by the expert to calculate the



Fig. 5. Scheme of FARC-HD method

hamming distance between two individuals in order to apply the crossover operators.

5) Restarting Approach: To get away from local optima, this algorithm uses a restart approach. In this case, the best chromosome is maintained and the remaining are generated at random. The restart procedure is applied when the threshold value L is below zero, which means that all the individuals coexisting in the population are very similar.

D. Flowchart

In accordance with the above description, the proposed algorithm to obtain a fuzzy association rule-based classifier is described below.

INPUT: A dataset with size T and m attributes, each with q_j predefined linguistic terms.

OUTPUT: A fuzzy associative classifier.

Stage 1. Fuzzy Association Rule Extraction for Classification. For each class C_i :

Step 1: Calculate the minimum support of the class C_j according to eq. (8).

Step 2: Create the levels 0 and 1 of the tree.

Step 3: Create a new level in the tree.

Step 4: Prune nodes.

Step 5: If there are more than 2 nodes in the new level and the depth of the tree is less than $Depth_{max}$, go to Step 3.

Step 6: Generate the rules with the class C_j on the right-hand side.

Stage 2. Candidate Rule Prescreening.

For each class C_j :

Step 7: Set the weight of the patterns as 1.

Step 8: Calculate the wWRAcc" value for each rule.

Step 9: Select the best rule as a part of the initial RB for Stage 3 and remove it from the candidate rule set.

Step 10: Decrease the weight of the patterns covered by the selected rule.

Step 11: If any pattern has been covered less than k_t times and there are more rules in the candidate rule set, go to Step 8.

Stage 3. Rule Selection and Lateral Tuning.

Step 12: Generate the initial population with *P* chromosomes. *Step 13:* Evaluate the population.

Step 14: Initialize the threshold value taking into account Gray codings, $L = L_{initial}$.

Step 15: Generate the next population:

- Shuffle the population.
- Select the parents two by two. Each pair is crossed if the hamming distance between the parent Gray codings divided by 2 is over L.
- Evaluate the new individuals.
- Join the parents with their offspring and select the best *P* individuals to take part in the next population.

Step 16: If the best chromosome does not change or there are no new individuals in the population, $L = L - (L_{initial} * 0.1)$. Step 17: If L < 0, restart the population and initialize L. Step 18: If the maximum number of evaluations is not reached,

go to Step 15.

A scheme of this algorithm is shown in Fig. 5.

IV. EXPERIMENTAL SET-UP

Several experiments have been carried out in this paper to evaluate the usefulness of our proposal. In the following subsections, firstly we describe the real-world databases used in these experiments, secondly we introduce a brief description of the methods considered for comparison, thirdly we show the configuration of the methods (determining all the parameters used), and finally we describe the statistical analysis adopted in this study.

A. Data sets

In order to analyze the performance of the proposed approach, we have considered 26 real-world datasets. Table II summarizes the main characteristics of the 26 datasets and shows the link to the Knowledge Extraction based on Evolutionary Learning (KEEL)-dataset repository [60] from which they can be downloaded, where Attributes(R/I/N)is the number of (Real/Integer/Nominal) attributes in the data, *Patterns* is the number of patterns and *Classes* is the number of classes. Notice that we have removed the instances with any missing value in the datasets (Cleveland and Crx) and 12 datasets have a number of variables greater than or equal to 15.

To develop the different experiments we consider a *10-fold* cross-validation model, i.e., we randomly split the data set into 10 folds, each containing 10% of the patterns of the data set, and used nine folds for training and one for testing ¹. For each of the ten partitions, we executed three trials of the algorithms.

¹The corresponding data partitions (10-fold) for these data sets are available at the KEEL-dataset repository [60]: http://sci2s.ugr.es/keel/datasets.php

Name	Attributes (R/I/N)	Patterns	Classes	Name	Attributes (R/I/N)	Patterns	Classes
Iris	4 (4/0/0)	150	3	Vowel	13 (10/3/0)	990	11
Phoneme	5 (5/0/0)	5404	2	Crx	15 (3/3/9)	653(690)	2
Monks	6 (0/6/0)	432	2	Pen-based	16 (0/16/0)	10992	10
Appendicitis	5 7 (7/0/0)	106	2	German	20 (0/7/13)	1000	2
Ecoli	7 (7/0/0)	336	8	Twonorm	20 (20/0/0)	7400	2
Pima	8 (8/0/0)	768	2	Ringnorm	20 (20/0/0)	7400	2
Yeast	8 (8/0/0)	1484	10	Wdbc	30 (30/0/0)	569	2
Glass	9 (9/0/0)	214	7	SatImage	36 (0/36/0)	6435	6
Page-blocks	10 (4/6/0)	5472	5	Texture	40 (40/0/0)	5500	11
Magic	10 (10/0/0)	19020	2	Spectfheart	44 (0/44/0)	267	2
Wine	13 (13/0/0)	178	3	Spambase	57 (57/0/0)	4597	2
Heart	13 (1/12/0)	270	2	Sonar	60 (60/0/0)	208	2
Cleveland	13 (13/0/0)	297(303)	5	Movementlibra	s 90 (90/0/0)	360	15

 TABLE II

 DATA SETS CONSIDERED FOR THE EXPERIMENTAL STUDY

Available at http://sci2s.ugr.es/keel/datasets.php

For each data set, we therefore consider the average results of 30 runs.

B. Methods considered for comparison

In these experiments, we compare the proposed approach with another ten methods which are available in the KEEL software tool [61]. What follows is a brief description of these methods:

- C4.5 [39]: This is a well-known algorithm used to generate a decision tree from a set of training data in the same way as the ID3 algorithm [62]. The extensions or improvements with respect to ID3 are that it accounts for unavailable or missing values in data, it handles continuous attribute value ranges, it chooses an appropriate attribute selection measure (maximizing gain-ratio) and it prunes the resulting decision trees.
- Classification Based on Associations (CBA) [12]: This method consists of two parts. In the first part, an algorithm based on Apriori algorithm [31] is used to mine the interval association rules for classification. In the second part, this sorts the generated rules according to their precedence relation and chooses a set of high precedence rules to cover the training data.
- CBA2 [13]: This method is the second version of the CBA algorithm which improves the previous system by using multiple class minimum support in rule generation.
- Classification based on Multiple Association Rules (CMAR) [14]: This method extends an efficient frequent (FP) pattern mining method, FP-Growth [63], constructs a class distribution-associated FP-tree, and mines large databases efficiently. Moreover, it applies a CR-tree structure to store and retrieve mined interval association rules efficiently, and it prunes rules effectively based on confidence, correlation (by using a weighted Chi-Square method) and database coverage. The classification is performed based on a weighted Chi-Square analysis using multiple strong association rules.

- Structural Learning Algorithm on Vague Environment (2SLAVE) [64]: This method is a modification of the GA of the SLAVE algorithm [65] in order to include a feature selection process. This is an inductive learning algorithm based on the Iterative Rule Learning approach, in which each chromosome represents a rule, to obtain a set of Disjunctive Normal Form (DNF) fuzzy rules. Chromosomes compete in every GA run, choosing the best rule per run. The global solution is formed by the best rules obtained when the algorithm is run multiple times.
- Learning Algorithm to discover Fuzzy Association Rules for classification (LAFAR) [24]: This method uses a GA to automatically determine the minimum fuzzy support and the minimum fuzzy confidence. To evaluate a determined chromosome this method finds frequent fuzzy grids and generates fuzzy classification rules from them. Once the whole classifier is obtained, the fitness value can be calculated, which maximizes the classification accuracy rate and minimizes the number of fuzzy rules. When reaching the termination condition, the chromosome with the maximum fitness value is used to test the performance of the proposed method.
- Classification based on Predictive Association Rules (CPAR) [15]: This method adopts a greedy algorithm to generate interval association rules directly from training data. In this process, this algorithm selects multiple literals with similar gains to build multiple rules simultaneously in order to avoid missing important rules. To perform the classification, this uses expected accuracy to evaluate each rule and uses the best k rules in prediction.
- Fuzzy Hybrid Genetic Based Machine Learning algorithm (FH-GBML) [66]: This method follows a Genetic Cooperative-Competitive Learning (GCCL) approach and consists of two processes. The first process is used for generating good fuzzy rules while the second one is used for finding good combinations of generated fuzzy rules. This method simultaneously uses multiple fuzzy partitions with different granularities for fuzzy rule extraction,

Method	Parameters
C4.5	Pruned = yes, Confidence = 0.25, InstancesPerLeaf = 2
CBA	Minsup = 0.01, Minconf = 0.5, Pruned = yes, RuleLimit = 80,000
CBA2	Minsup = 0.01, Minconf = 0.5, Pruned = yes, RuleLimit = 80,000
CMAR	$Minsup = 0.01, Minconf = 0.5, \delta = 4, DifferenceThreshold = 20\%$
2SLAVE	$Pop = 100, Iter_{change} = 500, P_c = 0.6, P_m = 0.01, \kappa_1 = 0, \kappa_2 = 1, Op_{or} = 0.1,$
	$Op_{and} = 0.1, P_{gen} = 0.1, P_{rot} = 0.1, \alpha = 0.15$
LAFAR	$t_{max} = 50, N_{pop} = 30, s = c = 10, W_{CAR} = 10, W_V = 1, P_c = 1.0, P_m = 0.01,$
	$J_{max} = 100, \eta_1 = 0.001, \eta_2 = 0.1$
CPAR	$\delta = 0.05, min_gain = 0.7, \alpha = 0.66, k = 5$
FH-GBML	$N_{rules} = 20, F_{sets} = 200, Gen = 1000, P_c = 0.9, P_{dcare} = \{0.5, 0.8, 0.95\},\$
	$P_{michigan} = 0.5$
SGERD	Q = heuristics
CFAR	$Minpsup = \{0.05, 0.1\}, Minpconf = 0.85, MS = 0.15$
FARC-HD	$Minsup = 0.05, Maxconf = 0.8, Depth_{max} = 3, k_t = 2,$
	$Pop = 50, Evaluations = 15,000, BITSGENE = 30, \delta = 0.15$

TABLE III PARAMETERS CONSIDERED FOR COMPARISON

using four homogeneous fuzzy partitions with triangular fuzzy sets and a don't care condition.

- Steady-State Genetic Algorithm for Extracting Fuzzy Classification Rules From Data (SGERD) [67]: It is a steady-state GA to generate a prespecified number of Q rules per class following a GCCL approach. In each iteration, parents and their corresponding offspring compete to select the best Q rules for each class. This method also simultaneously uses multiple fuzzy partitions with different granularities and a don't care condition for fuzzy rule extraction.
- Classification with Fuzzy Association Rules (CFAR) [27]: This method uses the Apriori algorithm to mine all the fuzzy association rules for classification and remove the conflicting and redundant rules to generate a compact set of rules denoted as CompSet. Then, this method selects the best rules to build the classifier by means of two processes. In the first process; for each pattern, CompSet is sorted by matching and confidence degree, rewarding the best rule that classify this pattern and punishing the rules that don't classify it. In the second process, the worst rules from CompSet are removed. These processes are iterated until the error rate in the training set increases.

C. Parameters of the methods

The parameters of the analyzed methods are shown in Table III ². Notice that only the rules with a number of antecedent conditions less than or equal to 3 are examined for our proposal. This restriction is intended to facilitate the discovery of a small number of short (i.e., simple) fuzzy rules. The parameters of the remaining methods were selected according to the recommendation of the corresponding authors within each proposal, which are the default parameter settings included

in the KEEL software tool [61]. Notice that in the FH-GBML algorithm the authors used three different probabilities of don't care (0.5, 0.8 and 0.95 depending on the size of the dataset) to obtain fuzzy rules with a few antecedent fuzzy sets. In these experiments, we have used these three probabilities of don't care in each dataset and have shown in the tables the best average result obtained for each one. Furthermore, in the CFAR algorithm the authors used 0.1 as the minimim support and this could be very high for some datasets (we are using 0.05 as the minimim support in our proposal). Likewise, we have used these two minimum supports in each dataset (0.1 and 0.05) and we have shown in the tables the best average result obtained in each one.

The initial linguistic partitions for our proposal and the fuzzy methods analyzed are comprised of *five linguistic terms* with uniformly distributed triangular MFs giving meaning to them, except in the FH-GBML and SGERD algorithms where the partitions are comprised of *two, three, four and five linguistic terms* for each attribute. The discretization of the continuous attributes for the CBA, CBA2, CMAR and CPAR algorithms is done using the Entropy method [68]. Notice that we use a crisp label for each value of the nominal variables.

D. Statistical analysis

In order to assess whether significant differences exist among the results, we adopt statistical analysis [41]–[43] and in particular non-parametric tests, according to the recommendations made in [40], where a set of simple, safe and robust non-parametric tests for statistical comparisons of classifiers has been introduced.

For pair-wise comparison we use Wilcoxon's Signed-Ranks test [69], [70], and for multiple comparison we employ Friedman's test [71], Iman and Davenport's test [72] and Holm's method [73]. In order to perform multiple comparisons, it is necessary to check whether all the results obtained by the algorithms present any significant difference (Friedman's test and Iman-Davenport's test) and, in the case of finding one,

²With these values for our proposal we have tried to facilitate comparisons, selecting standard common parameters that work well in most cases instead of searching for very specific values.

	2SLAVE			2	FH-GBML			SGERD				FARC-HD				
Dataset	# R	#C	Tra	Tst	# R	#C	Tra	Tst	# R	#C	Tra	Tst	# R	# C	Tra	Tst
Iris	4.0	3.2	94.32	94.44	14.9	3.3	98.89	94.00	3.4	2.0	95.14	94.89	4.0	1.1	98.59	96.00
Phoneme	11.5	24.2	77.52	76.41	17.4	4.5	79.57	79.66	3.6	1.9	75.74	75.55	17.8	2.2	83.52	82.14
Monks	3.0	1.3	97.22	97.26	14.7	2.1	98.36	98.18	2.2	1.4	80.56	80.65	14.2	2.0	99.92	99.77
Appendicitis	4.4	7.5	91.20	82.91	13.8	7.0	93.19	86.00	2.5	2.0	87.88	84.48	6.8	1.8	93.82	84.18
Ecoli	12.6	9.6	89.51	84.53	10.3	4.2	75.83	69.38	9.4	1.6	76.53	74.05	33.8	2.4	92.33	82.19
Pima	7.8	8.8	76.35	73.71	10.6	6.0	77.18	75.26	3.1	2.0	74.01	73.37	22.7	2.4	82.90	75.66
Yeast	23.6	9.8	55.54	51.27	7.5	5.9	52.31	51.42	11.3	1.5	39.83	38.77	35.2	2.6	63.81	58.50
Glass	15.1	9.3	74.25	58.05	9.4	5.0	64.85	57.99	6.9	2.0	61.31	58.49	22.7	2.5	81.10	70.24
Page-blocks	7.5	10.3	91.39	91.39	7.4	8.1	94.37	94.21	6.5	2.0	90.83	90.72	19.1	2.3	95.62	95.01
Magic	4.1	10.5	73.97	73.96	9.9	8.2	81.23	81.30	3.1	2.0	72.17	72.06	43.3	2.5	85.36	84.51
Wine	5.5	10.3	92.52	89.47	9.2	4.7	95.51	92.61	4.2	2.0	93.67	91.88	8.7	1.6	99.94	94.35
Heart	4.3	10.7	75.35	71.36	12.7	3.2	84.65	75.93	2.7	1.9	74.83	73.21	27.0	2.6	93.91	84.44
Cleveland	11.9	12.8	54.24	48.82	6.9	4.5	58.29	53.51	6.4	2.0	56.55	51.59	61.3	2.9	88.18	55.24
Vowel	63.1	15.6	82.06	71.11	9.2	13.0	67.41	67.07	18.0	1.9	72.99	65.83	72.3	2.9	80.48	71.82
Crx	2.4	6.7	74.36	74.06	11.6	6.2	86.32	86.60	2.1	1.9	85.04	85.03	25.4	2.6	91.17	86.03
Pen-based	40.0	18.9	81.32	81.16	18.4	8.0	50.69	50.45	15.9	2.0	68.17	67.93	152.8	2.8	97.04	96.04
German	6.5	8.4	72.44	70.53	5.1	4.0	87.11	87.01	3.4	2.0	68.54	67.97	85.7	2.8	86.81	72.80
Twonorm	26.5	15.5	87.45	86.99	12.0	7.6	86.26	85.97	3.1	2.0	74.49	73.98	60.9	2.6	96.64	95.28
Ringnorm	4.6	23.7	80.12	79.63	6.9	11.3	87.34	86.92	6.8	2.0	73.21	72.63	24.0	1.9	95.13	94.08
Wdbc	5.2	8.1	92.43	92.33	7.2	4.9	95.12	92.26	3.7	2.0	91.79	90.68	10.4	1.7	98.57	95.25
SatImage	57.9	25.1	84.03	81.69	16.5	36.0	74.90	74.72	12.2	2.0	77.15	77.10	76.1	2.7	88.68	87.32
Texture	34.9	23.9	82.87	81.57	14.6	40.0	69.91	70.15	18.6	2.0	72.12	71.66	54.5	2.7	93.71	92.89
Spectfheart	6.1	21.7	80.71	79.17	10.8	44.0	79.28	72.36	2.1	1.9	79.05	78.16	12.9	1.8	91.43	79.83
Spambase	7.9	11.4	69.87	70.14	3.9	18.5	77.86	77.22	3.7	2.0	72.90	72.98	29.8	2.4	92.37	91.93
Sonar	9.6	17.5	77.92	71.42	10.3	4.7	80.56	68.24	3.2	2.0	74.22	71.90	18.0	2.3	98.77	80.19
Movementlibras	47.4	26.5	90.13	67.04	12.1	90.0	77.87	68.89	22.9	2.0	72.37	68.09	83.1	2.9	95.52	76.67
Mean	16.4	13.5	80.73	76.94	10.9	13.6	79.80	76.82	6.9	1.9	75.43	73.99	39.3	2.3	90.97	83.94

 TABLE IV

 Results obtained by the analyzed methods

then we can find out by using a post-hoc test for comparing the control algorithm with the remaining algorithms (Holm's test). We use $\alpha = 0.05$ as the level of confidence in all cases. A wider description of these tests, together with software for their use, can be also found on the web site available at: http://sci2s.ugr.es/sicidm/.

V. EXPERIMENTAL RESULTS

In this section we analyze the results obtained in the different experiments. This section is organized as follows:

- In subsection V-A, we show a statistical study obtained from the comparison with another three GFSs including: FH-GBML [66], 2SLAVE [64] and SGERD [67].
- In subsection V-B, we compare the performance of our approach with two other approaches to obtain a fuzzy associative classifier: LAFAR algorithm [24] and CFAR algorithm [27].
- In subsection V-C, we compare the performance of our approach with the C4.5 decision tree [39] and four classical approaches for associative classification: CBA algorithm [12], CBA2 algorithm [13], CMAR algorithm [14] and CPAR algorithm [15].
- In subsection V-D, we show an analysis of the performance of our approach depending on the depth of the trees and the number of evaluations in the genetic process.

• In subsection V-E, we analyze the scalability of our proposal.

A. Comparison with other GFSs

This section analyzes the performance of our model against three recognized GFSs. The results obtained by the analyzed methods are shown in Table IV, where

- #R stands for the average number of rules,
- #C stands for the average number of conditions in the antecedent of the rules,
- *Tra* for the average classification percentage obtained over the training data,
- and *Tst* for the average classification percentage obtained over the test data.

The best global result for each one is stressed in boldface in each case.

In order to compare the results, we have used nonparametric tests for multiple comparison to find the best approach (see subsection IV-D), considering the average results obtained in test (Tst). First of all, we have used the Friedman and ImanDavenport tests in order to find out whether significant differences exist among all the mean values. Table V shows the Friedman and ImanDavenport statistics and it relates them to the corresponding critical values for each distribution

TABLE V Results of the Friedman and ImanDavenport tests ($\alpha=0.05)$

Friedman Test								
Statistic (\mathcal{X}_F^2)	p value							
40.823	7.814	< 0.0001						
Iman	Davenport Test							
Statistic (\mathcal{F}_F)	Critical Value	p value						
27.452	2.7386	< 0.0001						

TABLE VI Average Rankings of the methods

Method	Ranking
SGERD	3.2884
2SLAVE	2.9808
FH-GBML	2.5577
FARC-HD	1.1731

by using a level of significance $\alpha = 0.05$. The *p* value obtained is also reported for each test. Given that the statistics of Friedman and ImanDavenport are clearly greater than their associated critical values, there are significant differences among the observed results with a level of significance $\alpha \leq 0.05$. Table VI shows the rankings (computed using a Friedman test) of the different methods considered in this study.

We now apply Holm's test to compare the best ranking method (FARC-HD) with the remaining methods. Table VII presents these results. In this table, the methods are ordered with respect to the z-value obtained. Holm's test rejects the hypothesis of equality with the rest of the methods $(p < \alpha/i)$. Therefore, analyzing the statistical study shown in Tables VI and VII we conclude that our model is a solid approach to deal with high dimensional datasets, as it has shown itself to be the best accuracy method when compared with the remaining fuzzy GFSs applied in this study.

Finally, the results presented in Table IV show that our proposal obtains a higher average number of rules (39.2 rules on average) than all the GFSs (good approaches for obtaining very compact models), showing a good trade-off closer to the accuracy with rules involving no more than three attributes in their antecedent, giving the advantage of easier understanding with respect to 2SLAVE and FH-GBML.

B. Comparison with other fuzzy associative classifiers

This section compares the performance of our model with two other approaches to obtain a fuzzy associative classifier, LAFAR algorithm [24] and CFAR algorithm [27]. The results obtained by these methods are shown in Table VIII (this kind of table was described in the previous subsection). Notice that we show less datasets; this is due to scalability problems in the LAFAR and CFAR algorithms which cannot run in all datasets.

In order to compare the two algorithms, we use a Wilcoxon's test, which is shown in Table IX. We can observe that the null hypothesis for the Wilcoxon's test has been rejected (*p*-value $\langle = \alpha \rangle$) and our proposal has achieved a

TABLE VII Holm Table for the selection methods with $\alpha=0.05$

i	Method	z	p	lpha/i	Hypothesis
3	SGERD	5.01	3.46E-9	0.0166	Rejected
2	2SLAVE	5.05	4.45E-7	0.0125	Rejected
1	FH-GBML	3.87	1.10E-4	0.05	Rejected

TABLE IX WILCOXON'S TEST ($\alpha = 0.05$)

Comparison	R^+	R^{-}	Hypothesis	p-value
FARC-HD vs. CFAR	161	10	Rejected	0.001
FARC-HD vs. LAFAR	34	2	Rejected	0.025
LAFAR vs. CFAR	32	4	Rejected	0.05

TABLE XI Results of the Friedman and ImanDavenport tests ($\alpha=0.05)$

Friedman Test									
Statistic (\mathcal{X}_F^2)	p value								
15.84	11.0705	< 0.01							
Iman	Davenport Test								
Statistic (\mathcal{F}_F)	Critical Value	p value							
3.4703	2.2885	< 0.01							

TABLE XII Average Rankings of the methods

Method	Ranking
CBA	3.807
CMAR	3.775
CBA2	3.730
C45	3.576
CPAR	3.346
FARC-HD	2.307

higher ranking. We may conclude that our proposal also presents the best performance in this case.

On the other hand, the results presented in Table VIII show that our approach obtains an average number of rules lower than the LAFAR and CFAR algorithms. However, the CFAR algorithm obtains less rules than our approach in 11 of the 18 datasets.

C. Comparison with classical approaches

This section analyzes the performance of our model against five classical approaches. The results obtained by the analyzed methods are shown in Table X.

In order to compare the results, we have applied the nonparametric tests described in subsection V-A. Table XI shows that the statistics of Friedman and ImanDavenport are clearly greater than their associated critical values and there are significant differences among the observed results with a level of significance $\alpha \leq 0.05$. Table XII shows the rankings (computed using a Friedman test) of the different methods considered in this study.

Table XIII shows that Holm's test rejects the hypothesis of equality with the rest of the methods ($p < \alpha/i$). Therefore, analyzing the statistical study shown in Tables XII and XIII, we

	LAFAR	CFAR	FARC-HD					
Dataset	#R #C Tra Tst	#R #C Tra Tst	#R #C Tra Tst					
Iris	17.4 1.0 98.74 92.67	9.1 1.5 91.33 90.67	4.0 1.1 98.59 96.00					
Phoneme	16.9 1.1 78.49 77.87	2.0 1.0 70.65 70.65	17.8 2.2 83.52 82.14					
Monks	126.4 3.9 94.64 93.36	12.1 1.8 100.00 99.55	14.2 2.0 99.92 99.77					
Appendicitis	39.1 1.1 95.91 86.00	14.8 1.4 88.47 87.82	6.8 1.8 93.82 84.18					
Ecoli	68.4 2.1 81.58 72.36	4.0 3.1 48.45 47.61	33.8 2.4 92.33 82.19					
Pima	32.3 1.1 77.36 75.40	2.0 1.9 65.10 65.11	22.7 2.4 82.90 75.66					
Yeast	40.2 2.0 54.53 50.90	3.0 1.0 36.44 36.44	35.2 2.6 63.81 58.50					
Glass	36.0 2.5 68.59 52.17	3.4 2.1 44.19 44.66	22.7 2.5 81.10 70.24					
Page-blocks		2.0 1.0 89.78 89.78	19.1 2.3 95.62 95.01					
Magic		2.0 1.5 64.84 64.84	43.3 2.5 85.36 84.51					
Wine		115.7 3.0 99.63 93.24	8.7 1.6 99.94 94.35					
Heart		45.3 3.3 86.01 82.22	27.0 2.6 93.91 84.44					
Cleveland		2.0 2.1 53.87 53.88	61.3 2.9 88.18 55.24					
Crx		89.9 5.1 89.45 86.79	25.4 2.6 91.17 86.03					
Pen-based		6.9 2.9 36.54 36.43	152.8 2.8 97.04 96.04					
German		2.0 2.7 70.00 70.00	85.7 2.8 86.81 72.80					
Twonorm		498.7 3.7 91.96 91.66	60.9 2.6 96.64 95.28					
Sonar		37.3 3.0 83.01 72.48	18.0 2.3 98.77 80.19					
Mean	47.1 1.8 81.23 75.09	47.3 2.3 72.76 71.32	36.6 2.3 90.52 82.92					

TABLE VIII Results obtained by the analyzed methods

TABLE X Results obtained by the analyzed methods

СВА		CBA CBA2				CMAR			CPAR			C4:	FARC-HD					
Data	# R	Tra	Tst	# R	Tra	Tst	# R	Tra	Tst	# R	Tra	Tst	#R	Tra	Tst	#R	Tra	Tst
Iris	4	96.7	93.3	4	96.7	93.3	41	96.4	94.0	34	96.4	96.0	5	98.0	96.0	4	98.6	96.0
Phoneme	81	81.4	80.5	116	82.2	81.2	358	79.3	78.6	1109	84.0	81.9	129	91.9	86.8	18	83.5	82.1
Monks	11	100.0	100.0	11	100.0	100.0	126	100.0	100.0	79	100.0	100.0	5	100.0	100.0	14	99.9	99.8
Appendicitis	6	91.5	89.6	6	91.5	89.6	42	90.8	89. 7	25	89.5	87.8	3	91.0	83.3	7	93.8	84.2
Ecoli	29	87.3	78.0	35	89.2	77.1	178	82.5	77.7	158	82.6	76.2	20	91.7	79.5	34	92.3	82.2
Pima	42	79.6	72.7	44	79.9	72.5	237	79.4	75.1	166	79.1	74.5	18	83.2	74.0	23	82.9	75.7
Yeast	48	59.4	54.7	67	61.8	55.9	262	57.2	53.6	427	59.4	56.3	169	81.5	55.6	35	63.8	58.5
Glass	25	82.3	70.8	28	83.1	71.3	174	78.9	70.3	121	77.2	68.9	26	93.7	67.4	23	81.1	70.2
Page-blocks	172	94.6	94.0	256	98.2	95.9	1348	94.7	94.4	464	97.4	96.1	43	98.5	97.1	19	95.6	95.0
Magic	728	82.7	81.5	1015	84.8	83.7	4200	79.3	78.8	5422	87.9	84.9	321	90.9	85.1	43	85.4	84.5
Wine	9	99.9	93.8	9	99.9	93.8	73	99.9	96.7	44	99.7	95.6	5	98.9	93.3	9	99.9	94.3
Heart	40	93.8	83.0	40	94.2	81.5	306	91.5	82.2	93	90.1	80.7	17	91.7	78.5	27	93.9	84.4
Cleveland	30	64.3	56.9	47	70.0	54.9	230	54.1	53.9	107	61.8	54.9	40	83.1	54.5	61	88.2	55.2
Vowel	130	77.8	63.6	215	92.6	74.9	945	71.2	60.4	888	76.1	63.0	95	97.1	81.5	72	80.5	71.8
Crx	97	95.2	83.6	97	94.9	85.0	728	92.0	85.0	204	91.1	87.3	21	90.8	85.3	25	91.2	86.0
Pen-based	653	85.3	83.1	592	90.3	88.1	4328	72.0	71.7	2536	97.9	93.1	191	99.3	96.5	153	97.0	96.0
German	177	94.5	75.3	227	92.9	73.7	1984	84.4	71.9	666	88.1	73.3	83	84.9	72.5	86	86.8	72.8
Twonorm	708	97.7	91.6	724	97.0	91.7	2163	97.7	95.5	1646	99.0	89.4	286	98.7	84.5	61	96.6	95.3
Ringnorm	295	96.3	94.1	184	97.2	93.7	2788	87.2	83.7	1209	99.1	92.2	203	98.7	90.2	24	95.1	94.1
Wdbc	50	100.0	94.7	51	99.9	95.1	309	99.9	94.9	66	99.0	95.1	12	99.1	95.2	10	98.6	95.3
SatImage	722	88.5	85.2	587	87.6	83.6	7083	86.6	84.9	2032	97.3	85.8	287	97.7	85.8	76	88.7	87.3
Texture	377	88.3	84.5	50	53.5	52.5	5616	76.3	73.8	1492	98.5	90.7	152	99.0	92.6	55	93.7	92.9
Spectfheart	31	92.0	79.8	33	91.8	79.8	305	85.5	79.4	87	87.3	78.3	19	98.3	76.5	13	91.4	79.8
Spambase	278	95.0	93.2	302	95.2	93.0	2317	92.9	92.0	686	97.9	93.5	109	97.4	92.7	30	92.4	91.9
Sonar	27	92.3	75.4	39	97.7	77.9	234	98.3	78.8	92	95.8	75.0	14	97.7	70.5	18	98.8	80.2
Movementlibras	41	54.8	36.1	4	11.6	7.2	510	51.7	39.2	636	85.3	63.6	47	94.4	69.4	83	95.5	76.7
Mean	185	87.4	80.3	184	85.9	78.7	1419	83.8	79.1	788	89.1	82.1	89	94.1	82.5	39	91.0	83.9

TABLE XIII Holm Table for the selection methods with $\alpha=0.05$

i	Method	z	p	lpha/i	Hypothesis
5	CMAR	3.71	2.1035E-4	0.01	Rejected
4	CBA	2.89	0.0038	0.0125	Rejected
3	CBA2	2.74	0.0060	0.0166	Rejected
2	C45	2.45	0.0144	0.025	Rejected
1	CPAR	2.01	0.0453	0.05	Rejected

conclude that our model is the best performing method when compared with the remaining classical approaches applied in this study. Finally, the results presented in Table X show that our proposal obtains an smaller average number of rules than the remaining approaches.

D. Analysis of the influence of $Depth_{max}$ and the number of evaluations

In this subsection several experiments have been carried out to analyse the performance of our approach depending on $Depth_{max}$ and the number of evaluations in the genetic selection and tuning process (using the experimental setting described in subsection IV). In order to make this analysis easier to interpret, we have used four representative datasets in this experiments: Yeast, Vowel, Ringnorm and Spectfheart (8, 13, 20 and 44 variables respectively). Table XIV shows the results obtained with three different values for $Depth_{max}$ (2, 3 and 4), where #R1 stands for the average number of rules obtained at the end of Stage 1, #R2 stands for the average number of rules obtained at the end of Stage 2, #R3 stands for the average number of rules obtained at the end of Stage 3, and *time* stands for the average runtime (in format hh : mm : ss).

Analysing the results presented in Table XIV, we can highlight the following facts:

- Candidate rule prescreening allows the selection of a reduced number of interesting rules with the three values for $Depth_{max}$, decreasing the computational cost in the genetic selection and tuning process. Notice that the number of rules obtained in the stage 1 for the dataset Spectfheart is higher than 300,000 rules.
- When we use $Depth_{max} = 4$ we can see how the proposed approach does not obtain important improvements in training for 3 of the 4 datasets and only improves the results obtained in test for 2 of the 4 datasets. Moreover, the increase of the computational cost is high in all datasets, being $Depth_{max} = 3$ a value with a good compromise between both properties.

On the other hand, Fig. 6 depicts the accuracy obtained over the training data along with different numbers of evaluations in the genetic process with $Depth_{max} = 4$. In this figure we can highlight how this process obtains the best solution in less than 14,000 evaluations in all datasets because the initial RBs consist of a reduced number of rules.

Taking into account both studies, a good neutral choice ensuring the convergence may be to use 3 for $Depth_{max}$ and 15,000 for the number of evaluations in the genetic process,



Fig. 6. The accuracy obtained over the training data with different numbers of evaluations in the genetic process with $Depth_{max} = 4$

obtaining a good trade-off between accuracy and computational cost (good accuracy and not too great a computational cost).

E. Analysis of Scalability

Table XV shows the average runtime of the analyzed methods in the previous subsections on the 26 real-world problems (with a number of variables ranging from 4 to 90 and a number of patterns ranging from 150 to 19020) and using the 10fold cross-validation model. The methods were implemented using Java and all of the experiments were performed using a Pentium Corel 2 Quad, 2.5GHz CPU with 4Gb of memory and running Linux.

Analyzing the results presented in Table XV we can draw the following conclusions:

- The SGERD algorithm presents a very low average runtime in all datasets, obtaining a good scalability when we increase the size of the problem. This method, however, should be the worst in Friedman's test when we compare the results obtained in the test data (see table VI).
- The 2SLAVE, FH-GBML, LAFAR and CFAR algorithms expend a large amount of time when the number of attributes and patterns in the dataset is high. Notice that CFAR and LAFAR can not run in 7 and 18 of the 26 datasets, respectively.
- The remainder of the methods obtain low computational costs in all datasets and present good results in accuracy. However, our proposal obtains the best ranking in Friedman's test when we compare the results obtained in the test partitions.
- Notice that the CBA and CBA2 algorithms present similar runtimes to CMAR and CPAR algorithms because they limit the total number of candidates rules generated in datasets with more than 15 variables since they cannot be completed within this limit.
- The FARC-HD approach presents a good computational cost in all datasets, obtaining a good scalability and the best performance in accuracy.

	$Depth_{max} = 2$					$Depth_{max} = 3$						$Depth_{max} = 4$						
Data	#R1	#R2	#R3	Tra	Tst	Time	#R1	#R2	#R3	Tra	Tst	Time	#R1	#R2	#R3	Tra	Tst	Time
Yeast	96.9	38.9	27.9	62.2	57.7	0:00:51	373.2	56.3	35.2	63.8	58.5	0:01:00	845.5	68.1	43.7	64.6	57.6	0:02:03
Vowel	211.9	48.3	35.2	60.4	52.3	0:00:58	1482.3	97.5	72.3	80.4	71.8	0:02:16	6423	111.1	90.5	89.8	78.4	0:07:10
Ringnorm	1460.4	36.7	25.9	94.8	93.8	0:05:44	9927.7	41.4	24.0	95.1	94.1	0:07:28	35201.1	45.4	28.0	95.9	94.6	0:19:42
Spectfheart	2870.0	37.6	13.8	91.3	79.1	0:00:19	45508.1	34.7	12.9	91.4	79.8	0:01:59	389983	32.8	15.4	92.6	77.5	1:10:21

 TABLE XIV

 Analysis of the performance depending on $Depth_{max}$

TABLE XV AVERAGE RUNTIME OF THE ANALYZED METHODS (HH:MM:SS)

Dataset	Var	Patt	2SLAVE	FH-GBML	SGERD	CFAR	LAFAR	CBA	CBA2	CMAR	CPAR	C4.5	FARC-HD
Iris	4	150	00:00:05	00:12:57	00:00:00	00:00:00	00:09:39	00:00:00	00:00:00	00:00:00	00:00:00	00:00:00	00:00:01
Phoneme	5	5404	00:03:01	01:39:48	00:00:01	00:00:05	01:08:27	00:00:01	00:00:01	00:00:01	00:00:05	00:00:00	00:01:44
Monks	6	432	00:00:07	00:10:57	00:00:00	00:00:01	07:39:18	00:00:00	00:00:00	00:00:00	00:00:00	00:00:00	00:00:09
Appendicitis	7	106	00:00:03	00:02:05	00:00:00	00:00:05	01:00:44	00:00:00	00:00:00	00:00:00	00:00:00	00:00:00	00:00:02
Ecoli	7	336	00:00:50	00:11:05	00:00:00	00:00:01	01:20:10	00:00:00	00:00:00	00:00:00	00:00:01	00:00:00	00:00:15
Pima	8	768	00:00:41	00:23:31	00:00:00	00:00:06	04:18:18	00:00:00	00:00:00	00:00:00	00:00:01	00:00:00	00:00:22
Yeast	8	1484	00:07:04	00:48:51	00:00:00	00:00:01	15:27:38	00:00:00	00:00:00	00:00:00	00:00:01	00:00:01	00:01:00
Glass	9	214	00:00:38	00:08:05	00:00:00	00:00:04	07:58:01	00:00:00	00:00:00	00:00:00	00:00:00	00:00:00	00:00:18
Page-blocks	10	5472	00:09:59	03:09:10	00:00:02	00:01:08	-	00:00:06	00:01:09	00:00:10	00:00:04	00:00:07	00:02:15
Magic	10	19020	01:03:25	13:04:02	00:00:21	00:16:30	-	00:00:28	00:00:50	00:01:18	00:01:20	00:00:52	00:17:52
Wine	13	178	00:00:12	00:10:16	00:00:00	00:00:45	-	00:04:45	00:43:04	00:15:58	00:00:00	00:00:00	00:00:09
Heart	13	270	00:00:15	00:12:51	00:00:00	00:00:12	-	00:00:00	00:00:00	00:00:00	00:00:00	00:00:00	00:00:14
Cleveland	13	303	00:01:29	00:14:36	00:00:00	00:00:09	-	00:00:00	00:00:00	00:00:00	00:00:00	00:00:00	00:00:37
Vowel	13	990	00:13:20	00:13:40	00:00:00	00:00:10	-	00:01:12	00:02:13	00:00:05	00:00:03	00:00:00	00:02:16
Crx	15	690	00:00:35	00:35:37	00:00:00	00:02:02	-	00:02:06	00:00:18	00:06:19	00:00:01	00:00:00	00:00:27
Pen-based	16	10992	02:32:58	10:29:28	00:00:13	00:00:07	-	00:00:29	00:00:52	00:01:24	00:00:36	00:00:27	00:45:08
German	20	1000	00:02:26	01:03:06	00:00:00	00:02:17	-	00:01:16	00:00:15	00:04:46	00:00:03	00:00:00	00:02:20
Twonorm	20	7400	00:39:37	09:29:46	00:00:07	05:03:33	-	00:00:49	00:00:39	00:00:13	00:00:14	00:00:10	00:15:46
Ringnorm	20	7400	00:08:04	10:05:13	00:00:07	-	-	00:02:12	00:01:14	00:13:33	00:00:16	00:00:31	00:07:28
Wdbc	30	569	00:00:30	00:50:01	00:00:00	-	-	00:00:09	00:00:05	01:01:00	00:00:01	00:00:01	00:00:51
SatImage	36	6435	01:57:23	04:07:42	00:00:13	-	-	00:00:43	00:00:24	00:04:50	00:00:54	00:00:18	00:35:08
Texture	40	5500	01:29:43	03:40:14	00:00:12	-	-	00:00:28	00:00:15	00:03:50	00:00:40	00:00:15	00:36:42
Spectfheart	44	267	00:00:31	00:14:30	00:00:00	-	-	00:00:09	00:00:05	00:02:39	00:00:00	00:00:10	00:01:59
Spambase	57	4597	00:42:58	13:01:47	00:00:10	-	-	00:00:15	00:00:12	00:24:55	00:00:13	00:00:27	00:14:36
Sonar	60	208	00:01:04	00:21:02	00:00:01	00:00:20	-	00:00:03	00:00:08	00:10:26	00:00:01	00:00:00	00:17:27
Movementlibras	90	360	00:10:45	05:17:58	00:00:01	-	-	00:00:01	00:00:01	00:00:01	00:00:06	00:00:27	01:20:10

VI. CONCLUDING REMARKS

In this paper we have proposed a new fuzzy associative classification method for high-dimensional datasets, named FARC-HD. Our aim is to obtain accurate and compact fuzzy associative classifiers with a low computational cost. To do this, we mine fuzzy association rules limiting the order of the associations in order to obtain a reduced set of candidate rules with less attributes in the antecedent. We have made use of a pattern weighting scheme in order to reduce the number of candidate rules, preselecting the rules with the best quality. A genetic rule selection and lateral tuning is applied to select a small set of fuzzy association rules with a high classification accuracy.

Taking into account the results obtained, we can conclude that our model is a solid approach to deal with high dimensional datasets, as it obtains the best accuracy in the experimental study. Moreover, FARC-HD obtains models with a reduced number of rules (39,2 rules on average) and particularly with few attributes in the antecedent. Finally, the limit in the depth of the trees, along with candidate rule prescreening using the fuzzy measure wWRACC", allow us to reduce the search space considerably. Thus, the genetic process for selection and tuning does not introduce an excessive computational cost in to the whole process.

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