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IPADE: Iterative Prototype Adjustment for Nearest Neighbor Classification

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Abstract—Nearest prototype methods are a successful trend of many pattern classification tasks. However, they present several shortcomings such as time response, noise sensitivity, and

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storage requirements. Data reduction techniques are suitable to alleviate these drawbacks. Prototype generation is an appropriate process for data reduction, which allows the fitting of a dataset for nearest neighbor (NN) classification. This brief presents a methodology to learn iteratively the positioning of prototypes using real parameter optimization procedures. Concretely, we propose an iterative prototype adjustment technique based on differential evolution. The results obtained are contrasted with nonparametric statistical tests and show that our proposal consistently outperforms previously proposed methods, thus becoming a suitable tool in the task of enhancing the performance of the NN classifier.

Index Terms—Classification, differential evolution, nearest neighbor, prototype generation.

I. INTRODUCTION

Classification is one of the most important tasks in machine learning and data mining [1], [2]. Most machine learning methods build a model during the learning process, known as eager learning methods [3], but there are some approaches where the algorithm does not need a model. These algorithms are known as lazy learning methods [4].

The nearest neighbor (NN) algorithm [5] and its derivatives belong to the family of lazy learning. It has proved itself to perform well for classification problems in many domains [2], [6] and is considered one of the top ten methods in data mining [7]. NN is a nonparametric classifier, which requires the storage of the entire training set and the classification of unseen cases, finding the class labels of the closest instances to them. In order to determine how close two instances are, several distances or similarity measures have been proposed [8]–[10]. The effectiveness and simplicity of the NN may be affected by several weaknesses such as high computational cost, high storage requirement, and sensitivity to noise. Furthermore, NN makes predictions over existing data and assumes that input data perfectly delimits the decision boundaries among classes.

Several approaches have been suggested and studied to tackle the drawbacks mentioned above, for instance, weighting schemes [11], [12] have been widely used to improve the results of the NN classifier.

A successful technique that simultaneously tackles the computational complexity, storage requirements, and sensitivity to noise of NN is based on data reduction. These techniques aim to obtain a representative training set with a lower size compared to the original one and with similar or even higher classification accuracy for new incoming data. Apart from feature selection [13], data reduction can be divided into two different approaches, known as prototype selection [14], [15] and prototype generation (PG) or abstraction [16], [17]. The former process consists of choosing a subset of the original training data, while PG can also build new artificial prototypes to better adjust the decision boundaries between classes in NN classification.

In the specialized literature, a great number of PG techniques have been proposed. Since the first approach to PNN based on merging prototypes [18] and divide-and-conquerbased schemes [19], many other proposals of PG were considered, for instance, Mixt_Gauss [20], ICPL [17], and RSP [21].

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Positioning adjustment of prototypes is another perspective within the PG methodology. It aims to correct the position of a subset of prototypes from the initial set by using an optimization procedure. Many proposals belong to this family, such as learning vector quantization (LVQ) [22] and its successive improvements [23], [24], genetic algorithms [25], and particle swarm optimization (PSO) [26], [27].

Many existing positioning adjustment of prototype techniques start with an initial set of prototypes and try to improve the classification accuracy by adjusting it. Two schemes of initialization are commonly used.

- 1) The number of representative instances for each class is proportional to their number in the input data.
- All the classes are represented by the same number of prototypes.

This initialization process is their main drawback due to the fact that this parameter can be very dependent on the problem tackled. Some PG approaches [23], [25] compute the number of needed prototypes to be retained automatically, but in complex domains they require to retain many prototypes. We propose a novel procedure to automatically find the smallest reduced set that achieves suitable classification accuracy over different types of problems. This method follows an iterative prototype adjustment scheme with an incremental approach. At each step, an optimization procedure is used to adjust the position of the prototypes, and the method adds new prototypes if needed. As a second contribution of this brief, we will adopt the differential evolution (DE) [28], [29] technique as optimizer. Our proposal will be denoted "iterative prototype adjustment based on differential evolution" (IPADE).

In experiments on 50 real-world benchmark datasets, the classification accuracy and reduction rate of our approach are investigated and its performance is compared with classical and recent PG models.

The rest of this brief is organized as follows. Section II describes the background of PG and DE. Section III explains the proposed algorithm IPADE. Section IV discusses the experimental framework and presents the analysis of results. Finally, in Section V we summarize our conclusions.

II. BACKGROUND

This section covers the background information necessary to define and describe our proposal. Section II-A presents the background on PG. Section II-B shows the main characteristics of DE.

A. PG

PG is an important technique in data reduction. It has been widely applied to instance-based classifiers and can be defined as the application of instance construction algorithms over a dataset to improve the classification accuracy of a NN classifier.

More specifically, PG can be defined as follows. Let \mathbf{x}_p be an instance where $\mathbf{x}_p = (\mathbf{x}_{p1}, \mathbf{x}_{p2}, \dots, \mathbf{x}_{pm}, \mathbf{x}_{p\omega})$, with \mathbf{x}_p belonging to a class ω of Ω possible classes given by $\mathbf{x}_{p\omega}$ and an *m*-dimensional space in which \mathbf{x}_{pi} is the value of the *i*th feature of the *p*th sample. Furthermore, let \mathbf{x}_t

be an instance where $\mathbf{x}_t = (\mathbf{x}_{t1}, \mathbf{x}_{t2}, \dots, \mathbf{x}_{tm}, \mathbf{x}_{t\psi})$, with \mathbf{x}_t belonging to a class ψ , which is unknown, of Ω possible classes. Then, let us assume that there is a training set *TR* which consists of *n* instances \mathbf{x}_p and a test set *TS* composed of *s* instances \mathbf{x}_t . The purpose of PG is to obtain a prototype generated set *GS* that consists of *r*, r < n, prototypes \mathbf{p}_u where $\mathbf{p}_u = (\mathbf{p}_{u1}, \mathbf{p}_{u2}, \dots, \mathbf{p}_{um}, \mathbf{p}_{u\omega})$, which are generated from the examples of *TR*. The prototypes of the generated set are determined to represent efficiently the distributions of the classes and to discriminate well when used to classify the training objects. Their cardinality should be sufficiently small to reduce both the storage and evaluation time spent by an NN classifier.

The PG approaches can be divided into several families depending on the main heuristic operation followed. The first approach that we can find in the literature, called PNN [18], belongs to the family of methods that carry out a merging of prototypes of the same class in successive iterations, generating centroids. Other well-known methods are those based on a divide-and-conquer scheme, by separating the *m*-dimensional space into two or more subspaces with the purpose of simplifying the problem at each step [19]. Recent advances that follow a similar operation include Mixt_Gauss [20], which is an adaptive PG algorithm considered in the framework of mixture modeling by Gaussian distributions while assuming a statistical independence of features, and the RSP3 technique [21] which tries to avoid drastic changes in the form of decision boundaries associated with TR, which is the main shortcoming observed in the classical approach [19].

One of the most important families of methods is based on adjusting the position of the prototypes that can be viewed as an optimization process. The main algorithm belonging to this family is LVQ [22]. LVQ can be understood as an artificial neural network in which a neuron corresponds to a prototype and a competition weight based is carried out in order to locate each neuron in a concrete place of the *m*-dimensional space to increase the classification accuracy. The third version of this algorithm, LVQ3, reported the best results. Several approaches have been proposed that modify the basic LVQ, for instance LVQPRU [23] which extends LVQ by using a pruning step to remove noisy instances, or the HYB algorithm [24] that constitutes a hybridization of several prototype reduction techniques. Specifically, HYB combines support vector machines (SVMs) with LVQ3 and executes a search in order to find the most promising parameters of LVQ3.

As a positioning adjustment of prototypes technique, a genetic algorithm called ENPC was proposed for PG in [25]. This algorithm executes different operators in order to find the most suitable position of the prototypes. PSO was proposed for PG in [26], [27], and they also belong to the positioning adjustment of prototypes category of methods. The main difference between them is the type of codification of the particles. The PSO approach proposed in [26] codifies a complete solution *GS* per particle. However, AMPSO [27] encodes each prototype of *GS* in a single particle. AMPSO has been shown to be more effective than PSO [26].

B. DE

DE follows the general procedure of an evolutionary algorithm. It starts with a population of *NP* candidate solutions, the so-called individuals. The generations in DE are denoted by $G = 0, 1, \ldots, G_{max}$. It is usual to denote each individual as a *D*-dimensional vector $X_{i,G} = \{x_{i,G}^1, \ldots, x_{i,G}^D\}$, called a "target vector".

After initialization, DE applies the mutation operator to generate a mutant vector $V_{i,G}$, with respect to each individual $X_{i,G}$, in the current population. For each target $X_{i,G}$, at the generation G, its associated mutant vector $V_{i,G} = \{V_{i,G}^1, \ldots, V_{i,G}^D\}$. The method of creating this mutant vector is that which differentiates one DE scheme from another. We focus on the *DE/Rand/1*, which generates the mutant vector as follows:

$$V_{i,G} = X_{r_1,G} + F \cdot (X_{r_2,G} - X_{r_3,G}).$$
(1)

After the mutation phase, the crossover operation is applied to each pair of the target vector $X_{i,G}$ and its corresponding mutant vector $V_{i,G}$ to generate a new trial vector which we denote $U_{i,G}$. There are three kinds of crossover operators known as "binomial," "exponential," and "arithmetic."

Specifically, we will focus on the well-known *DE/CurrentToRand/1* strategy [30], which generates the trial vector $U_{i,G}$ by linearly combining the target vector $X_{i,G}$ and the corresponding mutant vector $V_{i,G}$ as follows:

$$U_{i,G} = X_{i,G} + K \cdot (V_{i,G} - X_{i,G}).$$
(2)

Now incorporating (1) in (2) and simplifying, we obtain

$$U_{i,G} = X_{i,G} + K \cdot (X_{r_1,G} - X_{i,G}) + F \cdot (X_{r_2,G} - X_{r_3,G}).$$
(3)

The indices r_1^i , r_2^i , and r_3^i are mutually exclusive integers randomly generated within the range [1, NP], which are also different from the base index *i*. The scaling factor *F* is a positive control parameter for scaling the different vectors. *K* is a random number from [0, 1].

When the trial vector has been generated, we must decide which individual between $X_{i,G}$ and $U_{i,G}$ should survive in the population of the next generation G + 1. If the new trial vector yields an equal or better solution than the target vector, it replaces the corresponding target vector in the next generation, otherwise the target is retained in the population.

The success of the DE algorithm in solving a specific problem crucially depends on the appropriately choice of its associated control parameter values that determine the convergence speed. Hence, a fixed selection of these parameters can produce a slow and/or premature convergence depending on the problem. Thus, researchers have investigated the parameter adaptation mechanisms to improve the performance of the basic DE algorithm. One of the most successful adaptive DE algorithms is SFLSDE [31]. It uses two local search algorithms in the scale factor space to find the appropriate parameters for a given $X_{i,G}$.

III. IPADE

In this section, we present and describe the IPADE approach in depth. IPADE follows an iterative scheme in which it determines the most appropriate number of prototypes per class and

```
1: GS = Initialization(TR)
2: DE_Optimization(GS, TR)
3: Accuracy_{Global} = Evaluate(GS, TR)
4: registerClass[0..\Omega] = optimizable
5: while Accuracy<sub>Global</sub> <>1.0 or all classes are non -
    optimizables do
6:
       lessAccuracy = \infty
       for i = 1 to \Omega do
7:
8:
          if registerClass[i] == optimizable then
            AccuracyClass [i] = Evaluate (GS, Examples
9:
            of class i in TR)
10:
            if AccuracyClass [i] < lessAccuracy then
11:
               lessAccuracy = AccuracyClass[i]
12:
               targetClass = i
13:
            end if
14:
          end if
15:
       end for
       GS_{test} = GS \cup \text{RandomExampleForClass} (TR,
16:
       targetClass)
17:
       DE_Optimization(GS_{test}, TR)
18:
       accuracy_{Test} = \text{Evaluate}(GS_{test}, TR)
19:
       if accuracy_{Test} > Accuracy_{Global} then
20:
         Accuracy_{Global} = accuracy_{Test}
         GS = GS_{test}
21:
22:
       else
          registerClass[targetClass] = non-optimizable
23:
24:
       end if
25: end while
26: return GS
```

Fig. 1. IPADE algorithm-basic structure.

their best positioning. Concretely, IPADE is divided into three different stages: initialization (Section III-A), optimization (Section III-B), and addition of prototypes (Section III-C). Fig. 1 shows the pseudocode of the model proposed. In the following, we describe the most significant instructions, enumerated from 1 to 26.

A. Initialization

A random selection (stratified or not) of examples from TR may not be the most adequate procedure to initialize the GS. Instead, IPADE iteratively learns prototypes in order to find the most appropriate structure of GS. Instruction 1 generates the initial solution GS. In this step, GS must represent each class with one prototype and should cover the entire search space as much as possible. For this reason, each class distribution is represented with its respective centroid. This initialization was satisfactorily used by the approaches proposed in [16] and [20]. The centroid of the class does not completely cover the region of each class and does not avoid misclassifications. Thus, instruction 2 applies the first optimization stage using the initial GS composed of centroids for each class. The optimization stage must modify the prototypes of GS using the movement idea in the *m*-dimensional space, adding or subtracting some quantities to the attribute values of the prototypes. It is important to point out that we normalize all attributes of the dataset to the [0, 1] range.

B. DE Optimization for IPADE

In this section, we explain the proposal to apply the underlying idea of the DE algorithm to the PG problem as a position adjusting of prototypes scheme.

First of all, it is necessary to define the solution codification. In the proposed DE algorithm, each individual in the population encodes a single prototype without the class label and, as such, the dimension of the individuals is equal to the number of attributes of the specific problem. An individual classifies an example of TR when it is the closest particle (in terms of Euclidean distance) to that example.

The DE algorithm uses each prototype \mathbf{p}_u of GS, provided by the IPADE algorithm, as an initial population. Next, mutation and crossover operators guide the optimization of the positioning of each \mathbf{p}_u in the *m*-dimensional space. It is important to point out that these operators only produce modifications in the attributes of the prototypes of GS. Hence, the class value remains unchangeable throughout the evolutionary cycle. We will focus on the well-known *DE/CurrentToRand/1* strategy [30] to generate the trial prototypes \mathbf{p}'_u because it has reportedly the best behavior. It can be viewed as

$$\mathbf{p}'_{u} = \mathbf{p}_{u} + K \cdot (\mathbf{p}_{r_{1}} - \mathbf{p}_{u}) + F \cdot (\mathbf{p}_{r_{2}} - \mathbf{p}_{r_{3}}).$$
(4)

The examples \mathbf{p}_{r_1} , \mathbf{p}_{r_2} , and \mathbf{p}_{r_3} are randomly extracted from *TR* and they belong to the same class as \mathbf{p}_u . In the hypothetical case that *TR* does not contain enough prototypes of the \mathbf{p}_u class, i.e., there is not at least three prototypes of this class in *TR*, we artificially generate the necessary number of new prototypes \mathbf{p}_{r_j} , $1 \le j \le 3$, with the same class label as \mathbf{p}_u , using little random perturbations such as $\mathbf{p}_{r_j} =$ $(\mathbf{p}_{u1} + rand[-0.1, 0.1], \mathbf{p}_{u2} + rand[-0.1, 0.1], \dots, \mathbf{p}_{um} + rand[-0.1, 0.1], \mathbf{p}_{u\omega})$.

After applying this operator, we check if there have been values out of range [0, 1]. If a computed value is greater than 1, we truncate it to 1, and if it is lower than 0, we establish it at 0.

After the mutation process over all the prototypes of GS, we obtain a trial solution GS', which is constituted for each \mathbf{p}'_u . The selection operator decides which solution GS' or GS should survive for the next iteration. The 1NN rule guides this operator to obtain the corresponding fitness value. We try to maximize this value, so the selection operator can be viewed as follows:

$$GS = \begin{cases} GS' & \text{if } accuracy(GS') >= accuracy(GS) \\ GS & \text{otherwise.} \end{cases}$$
(5)

In order to guarantee a high-quality solution, we use the ideas established in [31] to obtain a self-adaptive algorithm. Instruction 3 evaluates the accuracy of the initial solution, measured by classifying the examples of TR with the prototypes of GS by using the NN rule.

C. Addition of Prototypes

After the first optimization process, IPADE enters in an iterative loop (instructions 5-25) to determine which classes need more prototypes to faithfully represent their class distribution. In order to do this, we need to define two types of classes. A class ω is said to be optimizable if it allows the addition of new prototypes to improve its local classification accuracy. The local accuracy of ω is computed by classifying the examples of *TR* whose class is ω with the prototypes kept in *GS* (using the NN rule). The *target class* will be the *optimizable* class with the least accuracy registered. From instructions 7–15, the algorithm identifies the target class in each iteration. Initially, all classes start as optimizable (instruction 4)

In order to reduce the classification error of the target class, IPADE extracts a random example of this class from *TR* and adds this to the current *GS* in a new trial set *GS*_{test} (instruction 16). This addition forces the re-positioning of the prototypes of GS_{test} by again using the optimization process (instruction 17) and its corresponding evaluation (instruction 18) of predictive accuracy.

After this process, we have to ensure that the new positioning of prototypes of GS_{test} , generated with the optimizer, has reported a successful improvement of the accuracy rate with respect to the previous GS. If the global accuracy of the GS_{test} is less than the accuracy of GS, IPADE does not add this prototype to GS and this class is registered as *non-optimizable*. Otherwise, $GS = GS_{test}$.

The stopping criterion is satisfied when the accuracy rate is 1.0 or all the classes are registered as non-optimizable. The algorithm returns GS as the smallest reduced set that is able to classify the TR appropriately.

IV. EXPERIMENTAL FRAMEWORK AND ANALYSIS OF RESULTS

This section presents the experimental framework (Section IV-A) and the comparative study between our proposal and other PG techniques (Section IV-B).

A. Experimental Framework

In this section, we show the issues related to the experimental study. In order to compare the performance of the algorithms, we use four measures, *accuracy* [1], [32], the *reduction rate* measured as

$$Reduction \ rate = 1 - size(GS)/size(TR)$$
(6)

Acc·Red measured as accuracy·reduction rate, and execution time.¹

We use 50 datasets² from the KEEL dataset repository³ [33], [34]. These datasets contain between 100 and 20 000 instances, and the number of attributes ranges from 2 to 60. The datasets considered are partitioned using the 10-fold cross-validation (10-fcv) procedure.

³Available at http://sci2s.ugr.es/keel/datasets.

¹Reduction rate and execution time information can be found in the web page.

²Datasets: abalone, appendicitis, australian, balance, banana, bands, breast, bupa, car, chess, cleveland, coil2000, contraceptive, crx, dermatology, ecoli, flare-solar, german, glass, haberman, hayes-roth, heart, hepatitis, housevotes, iris, led7digit, lymphography, magic, mammographic, marketing, monks, newthyroid, page-blocks, pima, ring, saheart, satimage, segment, sonar, spectheart, splice, tae, thyroid, tic-tac-toe, titanic, twonorm, wine, wisconsin, yeast, zoo.

TABLE I Parameter Specification for All the Methods Employed in the Experimentation

Algorithm	Parameters
IPADE	Iterations of basic $DE = 300/500/1000$, iterSFGSS = 8,
	iterSFHC = 20, $Fl = 0.1$, $Fu = 0.9$
RSP3	Subset choice = diameter
Mixt_Gauss	Reduction rate $= 0.95$
ENPC	Iterations $= 300/500/1000$
AMPSO	Iterations = $300/500/1000$, C1 = 1.0, C2 = 1.0, C3 = 0.25,
	Vmax = 1, W = 0.1, X = 0.5, Pr = 0.1, Pd = 0.1
LVQPRU	Iterations = $300/500/1000$, $\alpha = 0.1$, WindowWidth = 0.5
HYB	Search_Iter = 300/500/1000, Optimal_Iter = 1000
	$\alpha = 0.1, I_{\epsilon} = 0, F_{\epsilon} = 0.5$
	Initial_Window = 0, Final_Window = 0.5
	$\delta = 0.1, \delta$ _Window = 0.1
	Initial Selection = SVM
LVQ3	Iterations = $300/500/1000$, $\alpha = 0.1$, WindowWidth = 0.2,
	$\epsilon = 0.1$

Many different configurations are established by the authors of each paper for the different techniques. We focus this experimentation on the recommended parameters proposed by their respective authors, assuming that the choice of the values of the parameters was optimally chosen. However, we have done a previous study for each method, which depends on the number of iterations performed, with 300, 500, and 1000 iterations in all the datasets. This parameter can be very sensitive to the problem tackled. An excessive number of iterations may produce overfitting for some problems, and a lower number of iterations may not be enough to tackle other datasets. For this reason, we present the results of the best performing number of iterations in each method and dataset. The complete set of results can be found in the associated web site (http://sci2s.ugr.es/ipade/). The configuration parameters of IPADE and the methods used in the comparison are shown in Table I. In this table, the values of the parameters F_l , F_u , *iterSFGSS*, and *iterSFHC* of the IPADE algorithm are the recommended values established in [31]. Furthermore, Euclidean distance is used as a similarity function, and those which are stochastic methods have been run three times per partition.

Implementations of the algorithms can be found in the web site associated or in the KEEL software tool [33].

B. Analysis of Results

In this section, we analyze the results obtained. Specifically, we check the performance of the IPADE model and seven other PG techniques.

In the scatterplot of Fig. 2, each point compares IPADE to a second algorithm on a single dataset. The x-axis position of the point is the accuracy of IPADE, and the y-axis position is the accuracy of the comparison algorithm. Therefore, points below the y = x line correspond to datasets for which IPADE performs better than a second algorithm.

In order to test the reduction capabilities of PG methods in comparison with IPADE, Fig. 3 shows at each point the $Acc \cdot Red$ obtained on a single dataset.



Fig. 2. Accuracy results over 50 datasets.



Fig. 3. Acc-Red results over 50 datasets.



Fig. 4. Map of convergence over three different datasets.

Fig. 4 shows a graphical representation of the convergence of the IPADE model over three different datasets. The graph shows a line representing the accuracy rate in each step

TABLE II Average Rankings of the Algorithms (FA + HAPV) for the Accuracy Measure

Algorithm	Accuracy FA	Accuracy HAPV
IPADE	109.63	_
LVQPRU	199.66	6.4064×10^{-4}
LVQ3	203.22	6.4064×10^{-4}
RSP3	231.80	7.9160×10^{-6}
1NN	236.53	4.2657×10^{-6}
AMPSO	248.11	5.0703×10^{-7}
ENPC	259.13	5.4223×10^{-8}
HYB	268.14	5.6781×10^{-9}
Mixt_Gauss	272.02	3.4239×10^{-6}

TABLE III Average Rankings of the Algorithms (FA + HAPV) for the $Acc \cdot Red$ Measure

Algorithm	Acc·Red FA	Acc·Red HAPV
IPADE	53.83	_
LVQ3	125.92	0.0055
Mixt_Gauss	169.18	2.2324×10^{-5}
LVQPRU	170.38	2.2324×10^{-5}
AMPSO	182.54	2.9965×10^{-6}
ENPC	267.91	9.3280×10^{-16}
RSP3	275.32	9.9684×10^{-17}
HYB	362.57	1.5321×10^{-31}
1NN	421.83	1.5321×10^{-44}

and its corresponding reduction rate (in brackets). The *x*-axis represents the number of iterations of the main loop of IPADE, and the *y*-axis represents the accuracy rate currently achieved.

Tables II and III present the statistical analysis conducted by nonparametric multiple comparison procedures for *accuracy* and *Acc*·*Red*, respectively. More specifically, we have used the Friedman aligned (FA) procedure [35], [36] to compute the set of rankings that represent the effectiveness associated with each algorithm (second column). Both tables are ordered from the best to the worst ranking. In addition, the third column shows the adjusted *p*-value with the Holm's test (*HAPV*) [35]. Note that IPADE is established as the control algorithm because it has obtained the best FA ranking. By using a level of significance $\alpha = 0.01$, IPADE is significantly better than the rest of the methods, considering both *accuracy* and *Acc*·*Red* measures. More information about these tests and other statistical procedures can be found at http://sci2s.ugr.es/sicidm/.

For the sake of simplicity, we only include the graphical and statistical results achieved, whereas the complete results can be found at the web page associated with this brief.

Looking at Tables II and III and Figs. 2–4, we want to make some interesting comments.

 Fig. 2 shows that the proposed IPADE outperforms, on average, the rest of the PG techniques with the parameter setting established. The most competitive algorithms for IPADE, in terms of the accuracy measure, are the LVQ3 and LVQPRU algorithms. In this figure, most of the LVQ3 and LVQPRU points are close to the y = x line. However, the statistical test confirms that IPADE significantly outperforms these methods.

- 2) The tradeoff between accuracy and reduction rate is an important factor because the efficiency of the NN classifier depends on the resulting number of prototypes of the *GS*. Fig. 3 shows that achieving this balance between accuracy and reduction rate is a difficult task. IPADE is the best performing method considering the balance between accuracy and reduction rates. In Fig. 3, there are more points under the y = x line in comparison with Fig. 2. Furthermore, Table III also supports this statement, showing smaller *p*-values when the reduction rate is considered.
- 3) Observing the map of convergence of Fig. 4, we can highlight the DE algorithm as a promising optimizer because it is able to reach highly accurate results very fast. This implies that the IPADE scheme needs a small number of iterations.

V. CONCLUSION

In this brief, we have presented a new data reduction technique called IPADE which iteratively learns the most adequate number of prototypes per class and their respective positioning for the NN algorithm, acting as a PG method. This technique uses a real parameter optimization procedure based on DE in order to adjust the positioning of the prototypes at each step. The large experimental study performed allowed us to show that IPADE is a suitable method for PG in NN classification. Furthermore, due to the fact that IPADE is a heuristic optimization approach, as future work this technique could be used for building an ensemble of classifiers.

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Linear Discriminant Analysis for Signatures

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Abstract—We propose signature linear discriminant analysis (signature-LDA) as an extension of LDA that can be applied to signatures, which are known to be more informative representations of local image features than vector representations, such as visual word histograms. Based on earth mover's distances between signatures, signature-LDA does not require vectorization of local image features in contrast to LDA, which is one of the main limitations of classical LDA. Therefore, signature-LDA minimizes the loss of intrinsic information of local image features while selecting more discriminating features using label information. Empirical evidence on texture databases shows that signature-LDA improves upon state-of-the-art approaches for texture image classification and outperforms other feature selection methods for local image features.

Index Terms—Earth mover's distance, feature selection, linear discriminant analysis, signature, texture classification.

I. INTRODUCTION

Linear discriminant analysis (LDA) is one of the most popular feature selection methods for classification tasks and has been widely used in machine learning and computer vision [1]–[3]. Numerous extensions of LDA have been exploited to improve the original LDA [4]–[11]. One of the main limitations of classical LDA is that it can be applied to data only in a vector space. Therefore, the given data, each of which is not represented as a vector, must be transformed into vectors before applying LDA.

Recently, local feature approaches have been widely used for texture image classification, object recognition, and image matching tasks because these approaches are descriptive as well as robust to occlusion and background clutter [12]–[14]. After obtaining local features through detection of the region of interest [12], [15]–[17] and descriptor computation [13], [18], visual word histograms comprising the frequencies of each visual word in images are typically employed to represent a large number of local features [13].

However, visual histograms often suffer from quantization or binning problems [19]. To resolve the problems, signatures [19] (see Section II for the definition) can be utilized to represent local image features instead of visual histograms. Previous works [18], [20] have shown empirically that signature-based methods generally outperform visual histograms on texture classification and object recognition tasks. Signatures are more informative than visual word histograms in that a signature is expressed not as a vector but as a collection of cluster center vectors and their relative weights. However, in contrast to histograms, signatures are beyond the scope

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