On the Selection of the Globally Optimal Prototype Subset for Nearest-Neighbor Classification

Emilio Carrizosa
Facultad de Matemáticas, Universidad de Sevilla, 41012 Sevilla, Spain, ecarrizosa@us.es

Belén Martín-Barragán
Departamento de Estadística, Universidad Carlos III de Madrid, 28903 Getafe, Madrid, Spain, belen.martin@uc3m.es

Frank Plastria
MOSI–Department of Mathematics, Operational Research, Statistics and Information Systems for Management, Vrije Universiteit Brussel, B-1050 Brussel, Belgium, frank.plastria@vub.ac.be

Dolores Romero Morales
Saïd Business School, University of Oxford, Oxford OX1 1HP, United Kingdom, dolores.romero-morales@sbs.ox.ac.uk

The nearest-neighbor classifier has been shown to be a powerful tool for multiclass classification. We explore both theoretical properties and empirical behavior of a variant method, in which the nearest-neighbor rule is applied to a reduced set of prototypes. This set is selected a priori by fixing its cardinality and minimizing the empirical misclassification cost. In this way we alleviate the two serious drawbacks of the nearest-neighbor method: high storage requirements and time-consuming queries. Finding this reduced set is shown to be NP-hard. We provide mixed integer programming (MIP) formulations, which are theoretically compared and solved by a standard MIP solver for small problem instances. We show that the classifiers derived from these formulations are comparable to benchmark procedures. We solve large problem instances by a metaheuristic that yields good classification rules in reasonable time. Additional experiments indicate that prototype-based nearest-neighbor classifiers remain quite stable in the presence of missing values.

Key words: classification; optimal prototype subset; nearest neighbor; dissimilarities; integer programming; variable neighborhood search; missing values

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1. Introduction

A classification problem has a database with objects belonging to \(|C|\) different classes, and one wants to derive a classification rule, i.e., a procedure that labels every future entry as a member of one of the \(|C|\) existing classes. Classification procedures are of two types: parametric and nonparametric. Parametric procedures assume that each object from class \(c \in C\) is associated with a random vector with known distribution, perhaps up to some parameters to be estimated, (e.g., data are multivariate normal vectors with unknown mean \(\mu_c\) and covariance matrix \(\Sigma_c\)), and primarily use of statistics (see e.g. McLachlan 1992). For complex databases with no evident distributional assumptions on the data (typical for databases with both quantitative and qualitative variables), nonparametric methods, such as the one described in this paper, are needed.

There has been increasing interest in deriving (nonparametric) classification rules via mathematical programming, requiring, for each object \(i\), a vector \(v^i\) of \(n\) numerical variables. This assumes variables to be ratio-scaled, rather than nominal or ordinal. Moreover, no null values are allowed, which excludes its direct use for cases in which some measures are missing or simply do not apply; see Cristianini and Shawe-Taylor (2000), Freed and Glover (1981), Gehrlein (1986), Gochet et al. (1997), and Mangasarian (1994).

A more flexible methodology, which requires only knowledge of a metric (or, as discussed in Section 2.1, a dissimilarity), is the nearest-neighbor (NN) method (Cover and Hart 1967, Dasarathy 1991, Devroye et al. 1996, Hastie et al. 2001), which provides excellent results (King et al. 1995, Pekalska et al. 2006). In NN methods, for each new entry \(i\) the distances (or dissimilarities) \(d(i, j)\) to some objects \(j\) in the database (called prototypes) are computed, and \(i\) is classified according to these distances. In particular, in the classical NN method, (Cover and Hart 1967), all objects are prototypes, and \(i\) is classified as member of the class \(c^*\) to which its closest prototype \(j^*\) (satisfying \(d(i, j^*) \leq d(i, j) \ \forall j\)) belongs.
A generalization of the NN method is the $k$-NN method (Devroye et al. 1996), which classifies each $i$ to the class most frequently found among the $k$ prototypes closest to $i$. In particular, NN is the special case of $k$-NN for $k = 1$.

These classification rules, however, require distances to be calculated to all data in the database for each new entry, involving high storage and time resources, making it impractical to perform online queries, so several variants have been proposed (Bennett and Willeman 2004, Bezdek and Kuncheva 2001, Dasarathy 1991, Devroye et al. 1996, Geva and Sitte 1991, Hart 1968, Kuncheva 1997, Kuncheva and Bezdek 1998). Most proposals differ in the way they attempt to provide heuristically a set of prototypes of small size and low misclassification cost (Bezdek and Kuncheva 2001). An extreme case is the condensed nearest-neighbor (CNN) rule (Hart 1968), in which the full database $I$ is replaced by a so-called minimal consistent subset, a smallest subset $S$ of records such that, if the NN classifier is used with $S$ (instead of $I$) as the set of prototypes, all objects in $I$ are correctly classified.

Since the size of a minimal consistent subset is unpredictable and might still be too large, several procedures have been suggested to reduce its size. Although such procedures do not necessarily correctly classify all items in the database (i.e., they are not consistent), they may have similar or even better behavior to predict class membership on future entries because they may reduce the possible overfitting suffered by CNN (Brighton and Mellish 2002, Lipowezky 1998).

In Bezdek and Kuncheva (2001) a number of procedures of this type are classified according to three different issues:

- The prototype design: the prototypes can either be selected from $I$ or be constructed (e.g. by considering centroids for numerical variables)
- The use of labels, i.e., whether the class labels of the sample data are used or not to select/construct the prototypes
- The control on the size of the set of prototypes, i.e., whether the number of prototypes is specified in advance or is automatically determined by the algorithm.

We propose a method in which a set of prototypes of pre-specified cardinality $p$ is sought, minimizing an empirical misclassification cost. As prototype design, we assume that prototypes are to be chosen from a given set, not necessarily equal to the set of available data. Hence, we give the highest freedom in this issue. Labels from sample data are used, so all existing information is taken into account. Finally, the user chooses the number of prototypes, and therefore fully controls the query times, which are critical when the computation of dissimilarities is costly. Indeed, the effort needed to classify a new entry is directly proportional to $p$ and may therefore guide the choice of $p$.

We restrict ourselves to the classification rule based on the closest distance, so our method is as a variant of NN. However, our results may directly be extended to considering the $k$ closest distances, $k \geq 1$, in the classification procedure, leading to a variant of the $k$-NN method.

The model is introduced in Section 2 and shown to be $NP$-hard. In Section 3, two mixed integer programming (MIP) formulations are proposed and theoretically compared. Numerical results are given in Section 4. When the optimization problems are solved exactly (with a standard MIP solver) the behavior of the classification rule is promising, but at the price of enormous preprocessing times. For this reason, a heuristic procedure is also proposed, and its quality and speed is explored. The rules obtained with this heuristic procedure have similar behavior on testing samples as the optimal ones. The method remains quite stable in the presence of missing values. Some concluding remarks and possible extensions are in Section 5.

### 2. The Model

#### 2.1. Classification Rules

A key concept in NN-based classification methods is the concept of distance, or, more generally, dissimilarity. A dissimilarity on a set $J$ is a function $d: J \times J \to \mathbb{R} \cup \{+\infty\}$, satisfying

$$d(u, v) \geq 0, \quad \forall u, v \in J$$

$$d(u, u) = 0, \quad \forall u \in J.$$  

When the set $J$ is (a subset of) the $n$-dimensional space $\mathbb{R}^n$, the most popular dissimilarities are those derived from metrics, such as the (weighted) Euclidean or the Mahalanobis distance. See Plastria (1995, 2001) for further details, extensions, and modelling aspects.

However, not all interesting dissimilarity measures correspond to metrics. In a typical example (Kaufman and Rousseeuw 1990) $J$ is a finite set of the $n$-dimensional space, but for some objects some of its coordinates cannot be used (because they are missing, or strongly suspected to be wrong). In that case, denoting, for each $u \in \mathbb{R}^n$, by $D(u)$ the set of coordinates of $u$ that can be used, we can extend the definition of Euclidean distance to the dissimilarity (not necessarily metric)

$$d(u, v) = \begin{cases} 
\left( \frac{\sum_{j \in D(u) \cap D(v)} \omega_j}{|D(u) \cap D(v)|} (u_j - v_j)^2 \right)^{1/2} 
& \text{if } D(u) \cap D(v) \neq \emptyset \\
+\infty & \text{otherwise,}
\end{cases}$$

for given weights $\omega_1, \ldots, \omega_n$. 

Natural definitions abound of dissimilarities for sets $J$ for which a metric is neither feasible nor recommended; see Kaufman and Rousseeuw (1990), where some dissimilarities are defined for sets $J$ of objects for which $n$ variables are measured, some quantitative, some ordinal or nominal, and nulls, as in (3), exist. Other methods for deriving dissimilarities can be found in protein/amino-acid alignment in bioinformatics (Altschul et al. 1994, 1990; Pearson and Lipman 1998), or fuzzy analysis, usually as the complement to 1 of a fuzzy similarity relation (Zimmermann 1991). In Yang and Shih (2001) set $J$ is a set of portraits of people from three different families.

Let $J$ be a finite set of objects with a dissimilarity $d$ defined on it, partitioned into $|C|$ classes $J_c$ ($c \in C$). A classification rule is a function $\varphi: J \rightarrow C$ that associates with each object $s \in J$ a class that might be its correct class or not. We consider only classification rules based on selecting prototypes for the different classes in $C$ as follows.

For each class $c \in C$, we are given a nonempty set $R_c \subset J$ of prototype candidates of class $c$. We denote the full set of candidates as $R$ and assume that the sets $R_c$ produce a partition of $R$, i.e., $\bigcup_{c \in C} R_c = R$ and $R_c \cap R_{c'} = \emptyset$, $c \neq c'$.

Some nonempty $S \subset R$, the set of prototypes, is to be chosen. For a given $S$, let $\varphi_S$ be the $S$-based NN classification rule, namely the classification rule that labels each $i \in J$ with the (known) label of the prototype in $S$ closest (i.e., least dissimiliar) to $i$. In other words, if $d(i, S \cap R_c)$ denotes $d(i, S \cap R_c) = \min_{j \in S \cap R_c} d(i, j)$, then let

$$\varphi_S(i) = \arg \min_{c \in C} d(i, S \cap R_c),$$

(4)

if this minimum is attained at a single $c$. In case of ties, a least-dissimiliar $c$ must be chosen as $\varphi_S(i)$, as detailed in the next section.

### 2.2. Performance Measure

For each $i \in J$ let $c(i) \in C$ denote the class to which $i$ belongs. In general, misclassification errors, i.e., objects $i$ with $\varphi_S(i) \neq c(i)$, will exist. Since not all misclassification errors are equally important, we assume we know for each $c, c' \in C$, the misclassification penalty $r(c' | c) \geq 0$, associated with each object of class $c$ labelled as a member of class $c'$. Let $r(c | c) = 0$, i.e., the cost of correct classification is zero.

A particular but important case is when all wrong classifications contribute the same cost,

$$r(c' | c) = \begin{cases} r_c & \text{if } c' \neq c \\ 0 & \text{otherwise} \end{cases}$$

(5)

where, for each $c \in C$, $r_c > 0$. We will call this the uniform case. Moreover, when all $r_c$ are equal, say, to unity, one obtains the binary case

$$r(c' | c) = \begin{cases} 1 & \text{if } c' \neq c \\ 0 & \text{otherwise} \end{cases}$$

(6)

which counts the number of misclassified objects (Mangasarian 1994). Our method accommodates cost structures more general than (6), e.g., as needed in Carrizosa et al. (2005), who construct a classifier for a cancer-diagnosis problem where different misclassification types naturally imply different misclassification costs.

Assignment rules should also be defined in case of ties: assignment will be done to the least dissimilar prototype, and ties will be broken randomly or by some user-defined procedure.

To compute the cost associated with a classifier, we use a worst-case approach. Since $R$ is a finite set, we can arbitrarily sort its labels, giving a strict total order $\prec$ on $R$. For each $i \in J$, let $\prec$ be the strict total order on $R$ yielding the assignment for $i$: For any $j_1, j_2 \in R$, $j_1 \prec j_2$ iff one of the following holds:

- $d(i, j_1) < d(i, j_2)$
- $d(i, j_1) = d(i, j_2)$, $r(c(j_1) | c(i)) > r(c(j_2) | c(i))$
- $d(i, j_1) = d(i, j_2)$, $r(c(j_1) | c(i)) = r(c(j_2) | c(i))$, $j_1 \neq j_2$.

Since $\prec$ is a strict total order on $R$, it is also a strict total order on any nonempty $S \subset R$. Hence, $\{j \in S: \text{no } j' \in S \text{ satisfies } j' \prec j\}$ is a singleton and its class defines $\varphi_S(i)$. This definition is consistent with (4) and extends it to the case of ties in dissimilarities.

With this cost structure, the total cost $\tau_i(S)$ within $I$ of the $S$-based NN classification rule $\varphi_S$ is $\tau_i(S) = \sum_{i \in I} r(\varphi_S(i) | c(i))$, which is a performance measure of the classification rule.

Evaluation of $\tau_i(\cdot)$ implicitly requires complete knowledge of the member class of each object in $I$. In practice, $c(i)$ will be known only for objects in a set $I \subset J$, called the training set. Hence, the definition of $\tau_i$ is of limited use since it cannot be calculated. However, if we assume that the training set $I$ has been obtained by a sampling in $J$, (unbiased) estimators $\delta_i$ of $\tau_i$ can be used as surrogates; see Cochran (1977) for an introduction to sampling and statistical estimation strategies.

Indeed, suppose that $I$ has been obtained after sampling in $J$, using a sampling design such that, for any $c \in C$, any object $i$ in class $c$ is included in the sample with probability $r(c) > 0$. For instance, if $I$ is obtained by random sampling without replacement of size $s$, then

$$r(c) = \frac{s}{|I|} \quad \forall c \in C.$$ 

(7)

With stratified random sampling, a random sample without replacement $I$ of size $s(c)$ is drawn from each $c \in C$, yielding (Cochran 1977) $r(c) = s(c)/|c|$. In any
case, an unbiased estimator for $\pi_I(S)$ is the Horwitz-Thompson estimator (Thompson 2002)

$$\hat{\pi}_I(S) = \sum_{i \in I} r(\varphi_S(i) \mid c(i)) / pr(c(i)).$$

We assume that, as in (7), all probabilities $pr(c)$ are equal, so $\pi_I(S)$ is proportional to the total cost $\pi_I(S)$ within the training sample $I$,

$$\pi_I(S) = \sum_{i \in I} r(\varphi_S(i) \mid c(i)),$$  \hspace{1cm} (8)

called the empirical classification cost. Thus, for the particular cost structure (6), the empirical classification cost is simply the number of misclassified objects in the training sample $I$.

Given an integer $p$, $|S| \leq p \leq |R|$, we determine the classification rule $\varphi_S$ with minimal empirical cost, measured as (8), such that $S$ is a subset of $R$, with cardinality $p$, and at least one prototype from each class $c$ is included, i.e. $S \cap R \neq \emptyset$, $\forall c$. We call this the optimal $p$-prototypes nearest-neighbor (p-PNN) model. If $R = I$, i.e., admitting the full training set as candidates to prototypes, then $|I|$-PNN is NN.

2.3. Complexity

In this section we prove that finding a p-PNN rule is $\mathcal{NP}$-hard and that the problem remains $\mathcal{NP}$-hard even when restricted to two-class case, the set of candidates to prototypes coinciding with the training set, the dissimilarity is a metric and the misclassification costs are uniformly equal to one. We formalize this in the following decision problem

PERFECT CLASSIFICATION: Given a number $p$ and a finite set $I$, partitioned into two subsets $I_1$, $I_2$ and equipped with a metric $d$, does there exist a subset $S$ of $I$ of cardinality $p$ such that the corresponding classification rule $\varphi_S$ classifies all elements of $I$ correctly?

PROPOSITION 1. PERFECT CLASSIFICATION is an $\mathcal{NP}$-complete problem.

PROOF. Our starting point is the following $\mathcal{NP}$-complete problem (Garey and Johnson 1979):

DOMINATING SET: Given a graph $(V, E)$ and a positive number $l \leq |V|$, does there exist an $l$-dominating vertex set? An $l$-dominating vertex set is a subset $V' \subseteq V$ with $|V'| \leq l$ and such that all vertices in $V \setminus V'$ are adjacent to $V'$.

For instance, consider the graph $(V, E)$ in Figure 1. $\{v_4, v_5, v_6, v_8\}$ is a $4$-dominating vertex set for $(V, S)$, whereas $\{v_4, v_5, v_6\}$ is not a $3$-dominating vertex set, since it contains no vertex adjacent to $v_9$.

Given an instance of DOMINATING SET, we construct an instance of PERFECT CLASSIFICATION as follows. Let $I_1 = V$ and $I_2 = \{w\}$ where $w$ is an arbitrary object not in $V$ and $p = l + 1$. The dissimilarities are defined by the shortest-path distances in the extended graph $(V \cup \{w\}, E \cup (V \times \{w\}))$ with edge lengths 2 on $E$ and 3 on all new edges. See in Figure 2 the extended graph for the graph $(V, E)$ of Figure 1, where edges of length 2 are plotted as continuous lines, and the edges of length 3 (those adjacent to $w$) are dashed.

Then for any set of prototypes $S \subseteq R$,

- $w$ is always correctly classified because $I_2$ is a singleton.
- For any $v \in I_1 = V$ only one of the following three possibilities for assignment may arise:
  - if $v \in S$, it is assigned to itself since dissimilarity is then 0, while $d(v, w) > 0 \quad \forall v \neq w$
  - if $v \notin S$ but adjacent to some $v' \in S \cap V$, it will be assigned to $v'$ since $d(v, v') = 2$, which is the minimal nonzero possibility for the dissimilarity
  - if $v \notin S$ and not adjacent to $S \cap V$, it will be assigned to $w$ since $d(v, w) = 3$, while the dissimilarity to any prototype in $V$ is at least 4.

Thus, misclassification by $S$ of some object is equivalent to being vertex-dominated by $\{w\}$.

Figure 1 An Example of Graph $(V, E)$

Figure 2 Extended Graph for the Graph $(V, E)$ of Figure 1
For instance, see Figure 1 with \( S = \{ v_0, v_4, v_5, v_6 \} \) as set of prototypes. Since \( v_5 \in S \), \( d(v_5, v_0) = 0 < 3 = d(v_5, w) \); hence, \( v_5 \) is assigned to class \( I_1 \) so is correctly classified. On the other hand, since \( v_1 \) is adjacent to \( v_5 \in S \), \( d(v_1, v_5) = 2 < 3 = d(v_1, w) \) and \( v_1 \) is also correctly classified. However, \( v_0 \) is misclassified since \( d(v_0, v_5) = 6 > 3 = d(v_0, w) \), \( d(v_0, v_6) = 6 > 3 = d(v_0, w) \), and \( d(v_0, v_5) = 4 > 3 = d(v_0, w) \), so \( v_0 \) is assigned to \( I_2 \). Hence, such \( S \) does not classify correctly all objects, or, equivalently, it is not a 4-dominating vertex set for the extended graph.

Hence, \( S \cap V \) being an \( l \)-dominating set is equivalent to all objects being correctly classified by \( S \) with \( |S| \leq l + 1 \).

**Corollary 2.** Finding a \( p \)-PNN rule is \( \mathcal{NP} \)-hard.

In Section 3 we formulate as integer programs the problem of determining the classification rule \( \varphi_s \) with minimal empirical misclassification cost, measured as (8), both for general misclassification penalties and also for the uniform case (5).

### 3. Integer-Programming Formulations

#### 3.1. General Costs

For each \( i \in I \) and \( s \in R \), let \( R_{i,s} \) denote the set of prototypes that are more preferred by \( i \) (according to \( \prec_i \)) than \( s \), \( R_{i,s} = \{ t \in R : t \prec_i s \} \). Let

- \( x_i \in \{0, 1\} \), \( \forall s \in R \), answering the question “Is candidate \( s \) chosen to be a prototype?” Hence, a classification rule \( \varphi_s \), as defined in (4), is identified by the vector \( x \in \{0, 1\}^{|R|} \) defined by

\[
x_s = \begin{cases} 1 & \text{if } s \in S \\ 0 & \text{otherwise.} \end{cases}
\]

- \( y_{is} \in \{0, 1\} \), \( \forall i \in I \) and \( s \in R \), answering the assignment question “Is \( s \) the prototype least dissimilar to \( i \)?” Although, in principle, these variables should be binary, the model allows us to consider them as continuously relaxed to lie between 0 and 1.

The misclassification cost for object \( i \in I \) belonging to the class \( c(i) \) is then \( \sum_{s \in R} r(c(s) \mid c(i)) y_{is} \), so the empirical misclassification cost \( \pi(S) \) of classification rule \( \varphi_S \), as defined in (4), is \( \sum_{c \in C} \sum_{i \in I} r(c(s) \mid c(i)) y_{is} \).

The problem is to find the set \( S \subset R \) with cardinality \( |S| = p \) (with \( p \) given) containing at least one element in each class \( c \), such that the empirical misclassification cost is minimized. This yields the following (mixed) integer program:

\[
\begin{align*}
\text{(P1)} \quad & \min \sum_{i \in I} \sum_{s \in R} r(c(s) \mid c(i)) y_{is} \\
\text{subject to } & \sum_{s \in R} x_s \geq 1 \quad \forall c \in C \quad (9) \\
& \sum_{s \in R} x_s = p \\
& \sum_{s \in R} y_{is} = 1 \quad \forall i \in I \\
& x_s - y_{is} \leq \sum_{t \in R_{i,s}} x_t \quad \forall (i, s) \in I \times R \\
& y_{is} \leq x_s \quad \forall (i, s) \in I \times R \\
& x_i \in \{0, 1\} \quad \forall s \in R \\
& y_{is} \in \{0, 1\} \quad \forall (i, s) \in I \times R.
\end{align*}
\]

Constraints (9) force each class to have at least one prototype and (10) says that we choose \( p \) prototypes in total. (11) and (13) ensure that each object has a prototype and that an object can only be assigned to a candidate chosen as prototype. Following Plastria (2002), (12) ensure that an object is assigned to the least dissimilar of the prototypes in the following way:

If \( s \) is a prototype and there is no prototype preferred by \( i \) then \( i \) must be assigned to \( s \) or equivalently

If \( x_i = 1 \) and \( x_i = 0 \), \( \forall t \in R_{i,s} \), then \( y_{is} = 1 \), which, from Plastria (2002), is expressed by \( 1 - y_{is} \leq \sum_{t \in R_{i,s}} x_t + (1 - x_i) \quad \forall (i, s) \in I \times R \), yielding (12).

#### 3.2. Uniform Costs

Now we discuss in some detail the particular case in which the cost structure is given by (5), because, as shown below, we can derive an alternative formulation with fewer variables and constraints than (P1). The additional advantage of this formulation is, remarkably, that its LP relaxation is at least as tight as the LP relaxation of (P1).

Indeed, in this case, the objective of (P1) is

\[
\sum_{i \in I} \sum_{s \in R} r(c(s) \mid c(i)) y_{is}
\]

Define, for each \( i \in I \), \( z_i = \sum_{s \in R_{i,s}} y_{is} \) which answers the (fuzzy) question “Is object \( i \) correctly classified?” With this, the objective at any feasible solution is \( \sum_{i \in I} r_{i} - \sum_{i \in I} r_{i} z_i \).

To guarantee that the variables \( z_i \) answer the question above, the only thing that must be specified is when \( z_i \) must be 0: when the choice of \( z_i \)'s value is left
free there will always be an optimal solution where \( z_i = 1 \). The former is obtained by stating

If \( t \not\in R_{c(i)} \) is chosen as a prototype and no prototype in \( R_{c(i)} \) is preferred (according to \( \prec_i \)) than \( t \), then \( i \in I \) will not be correctly classified or, for each \( t \not\in R_{c(i)} \).

If \( x_i = 1 \) and \( x_t = 0 \) (\( \forall s \in R_{c(i)} \cap R_{c(t)} \)), then \( z_i = 0 \), which is expressed by the constraint

\[
z_i \leq (1 - x_i) + \sum_{s \in R_{c(i)} \cap R_{c(t)}} x_s. \tag{14}
\]

Note that this constraint indeed expresses exactly the desired property even when \( z_i \) is continuously relaxed (Plastria 2002). Thus, we may rewrite (P1) as

\[
\begin{align*}
(P2) \quad & \min \sum_{i \in I} r_{c(i)} - \sum_{i \in I} r_{c(i)} z_i \\
& \text{subject to } \sum_{s \in R_c} x_s = p \\
& \quad \sum_{s \in S} x_s \geq 1 \quad \forall c \in C \\
& \quad z_i \leq (1 - x_i) + \sum_{s \in R_{c(i)} \cap R_{c(t)}} x_s \quad \forall i \in I, t \not\in R_{c(i)} \\
& \quad x_s \in [0, 1] \quad \forall s \in R \\
& \quad z_i \in [0, 1] \quad \forall i \in I.
\end{align*}
\]

Calling (LP1) and (LP2) the LP relaxations of (P1) and (P2), we may state

**Proposition 3.** (LP2) is at least as tight as (LP1).

**Proof.** See Carrizosa et al. (2005). □

Since the integer variables are the same in both models, the subproblems generated when fixing some of these variables satisfy the same property. Therefore, if available, (P2) is preferred when solving the problem with a branch-and-bound algorithm.

Note that \( \sum_{i \in I} r_{c(i)} \) in (P2)’s objective is constant, and, since we are only interested in optimal solutions, and not in the optimal objective value, it may be dropped. Sign inversion then leads to the simpler objective \( \max \sum_{i \in I} r_{c(i)} z_i \) defining our model (P2) subject to the same constraints as (P2) above.

The variables above can also be used to model as IPs other variants of NN. For instance, finding the consistent subset of minimal cardinality, i.e., Hart’s CNN-rule (Hart 1968), amounts to solving

\[
\begin{align*}
(PCNN) \quad & \min \sum_{s \in S} x_s \\
& \text{subject to } \sum_{s \in R_c} x_s \geq 1 \quad \forall c \in C \\
& \quad 1 \leq (1 - x_i) + \sum_{s \in R_{c(i)} \cap R_{c(t)}} x_s \quad \forall i \in I, t \not\in R_{c(i)} \\
& \quad x_s \in [0, 1] \quad \forall s \in R.
\end{align*}
\]

### 4. Computational Experience

#### 4.1. Aims

The storage requirements and processing time of the \( p \)-PNN rule are smaller than those of NN. Our aim here is to compare empirically the classification power of the \( p \)-PNN rule, for different values of \( p \), against NN. For completeness, we also compare \( p \)-PNN with other benchmark methods. In particular, we tested the performance of \( p \)-PNN against

- \( k \)-NN (\( k \)-NN in the tables) for different values of \( k \).
- Support vector machines (SVM), Cristianini and Shawe-Taylor (2000), with linear kernel (Lin), polynomial kernel (Pol), and radial bases function kernel (Rbf).
- Classification trees, denoted here by Trees, (Breiman et al. 1984), with and without pruning (Pruned and Crude respectively), as implemented in Matlab 6.5 Statistics Toolbox.

We performed numerical tests on different standard databases from the UCI Machine Learning Repository (Blake and Merz 1998). The details are in Section 4.2. Since some of the benchmark methods do not accommodate in a simple way different misclassification costs, we used the test set error rate.

#### Table 1: Parameters of the Databases

| Database | \( J \) | \( |C| \) | \( n \) |
|----------|------|------|------|
| glass    | 214  | 6    | 9    |
| glassw   | 163  | 3    | 9    |
| wine     | 178  | 3    | 13   |
| yeastME  | 258  | 3    | 8    |
| abalone  | 4,177| 3    | 7    |
| spam     | 4,601| 2    | 57   |

#### Table 2: Results with \( k \)-NN, SVM, and Classification Trees

<table>
<thead>
<tr>
<th>Database</th>
<th>1-NN</th>
<th>2-NN</th>
<th>3-NN</th>
<th>4-NN</th>
<th>5-NN</th>
<th>Lin</th>
<th>Pol</th>
<th>Rbf</th>
<th>Pruned</th>
<th>Crude</th>
</tr>
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<tbody>
<tr>
<td>glass</td>
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<td>test</td>
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<td>100</td>
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<td>100</td>
<td>100</td>
<td>60</td>
<td>72</td>
<td>67</td>
<td>78</td>
<td>89</td>
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<td>glassw</td>
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<td>62</td>
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<tr>
<td>test</td>
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<td>63</td>
<td>67</td>
<td>70</td>
<td>71</td>
<td>56</td>
<td>67</td>
<td>68</td>
<td>71</td>
<td>69</td>
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<tr>
<td>yeastME</td>
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<td>84</td>
<td>86</td>
<td>86</td>
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<tr>
<td>spam</td>
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<td>90</td>
<td>99</td>
<td>92</td>
<td>92</td>
</tr>
</tbody>
</table>
costs, the comparisons were made using the binary cost structure (6). Hence, both (P1) and (P2) apply. Since the LP bound of (P2) is at least as good as that from (P1), by Proposition 3, all our results refer to the simpler variant (P2′) of (P2).

4.2. The Databases
The UCI-Repository databases we used are of different sizes. A first group of databases consists of the Glass Identification Database (called here glass), a subset of the glass database (glassw) consisting only of the “window glass” classes, the Wine Recognition Database (wine), and Yeast Database (yeastME), from which only the three “membrane protein” classes (denoted as ME1, ME2, ME3 in the UCI Repository) are used. Moreover, two bigger databases are also considered: the Abalone Database (abalone), and the Spambase Database (spam). In abalone, three classes (grouping classes 1–8, 9–10, and 11 on) are considered, as cited in Blake and Merz (1998), and the qualitative variable was excluded.

For each database \( J \), the total number of objects \( |J| \), the number of classes \( |C| \), and the number of variables (all quantitative) \( n \) are in Table 1.

The databases contain only continuous variables, and one can thus calculate dissimilarities according to the weighted Euclidean distance, defined for \( u = (u_1, \ldots, u_n) \in \mathbb{R}^n \), \( v = (v_1, \ldots, v_n) \in \mathbb{R}^n \) as

\[
d(u, v) = \left( \sum_{1 \leq j \leq n} \omega_j (u_j - v_j)^2 \right)^{1/2},
\]

where \( \omega_j \) is the weight of the \( j \)th variable in the database, respectively. This model is equivalent to considering the unweighted Euclidean distance after rescaling each variable to the interval [0, 1].

All results presented were obtained by ten-fold cross-validation (Kohavi 1995).

The averaged percentages of correctly classified objects in both the training samples (tr) and testing samples (test) obtained using the benchmark methods are in Table 2.

4.3. Solving to Optimality
The worst-case complexity of the problem was addressed in Section 2.3, and now we discuss the empirical behavior. To compare the running times and the classification power of \( \tilde{p} \)-PNN, the IPs were solved on a 2.86 GHz Pentium 4 and 256 MB RAM. CPLEX 8.1.0 was used as the MIP solver. Due to the hardness of these MIP formulations we imposed an upper bound (MAXT) on the computing time of 10,800 seconds. In most instances, running times exceeded MAXT and the optimization process stopped prematurely.

Tables 3–6 display for different values of the number \( p \) of prototypes (first column), the percentage of

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Results of glass</th>
</tr>
</thead>
<tbody>
<tr>
<td>( p )</td>
<td>( p \text{PNN} ) (%)</td>
</tr>
<tr>
<td>6</td>
<td>66</td>
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<tr>
<td>10</td>
<td>70</td>
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<tr>
<td>15</td>
<td>76</td>
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<td>20</td>
<td>80</td>
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</table>

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Results of glassw</th>
</tr>
</thead>
<tbody>
<tr>
<td>( p )</td>
<td>( p \text{PNN} ) (%)</td>
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<td>3</td>
<td>67</td>
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<tr>
<td>5</td>
<td>72</td>
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<td>10</td>
<td>80</td>
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<td>15</td>
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<tr>
<td>30</td>
<td>92</td>
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<tr>
<td>40</td>
<td>96</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 5</th>
<th>Results of wine</th>
</tr>
</thead>
<tbody>
<tr>
<td>( p )</td>
<td>( p \text{PNN} ) (%)</td>
</tr>
<tr>
<td>3</td>
<td>99</td>
</tr>
<tr>
<td>5</td>
<td>95</td>
</tr>
<tr>
<td>10</td>
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<td>30</td>
<td>100</td>
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<tr>
<td>40</td>
<td>100</td>
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</table>

with each weight \( \omega_j \) given by

\[
\omega_j = \frac{1}{(\hat{\Delta}_j - \tilde{\Delta}_j)^2}
\]

where \( \hat{\Delta}_j \) and \( \tilde{\Delta}_j \) represent the highest and the lowest value for the \( j \)th variable in the database, respectively.

<table>
<thead>
<tr>
<th>Table 6</th>
<th>Results of yeastME</th>
</tr>
</thead>
<tbody>
<tr>
<td>( p )</td>
<td>( p \text{PNN} ) (%)</td>
</tr>
<tr>
<td>3</td>
<td>97</td>
</tr>
<tr>
<td>5</td>
<td>95</td>
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<tr>
<td>10</td>
<td>100</td>
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<td>30</td>
<td>100</td>
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<tr>
<td>40</td>
<td>100</td>
</tr>
</tbody>
</table>
correctly classified objects in the training sample (second column) and in the testing sample (third column) for the four smallest databases. (The other columns are for later use in Section 4.4.) Further computational results can be found in Carrizosa et al. (2005).

Comparing with the benchmark results in Table 2, several conclusions can be drawn. First, no method systematically outperforms the others. In particular, the p-PNN shows to be comparable against the remaining methods. Moreover, an adequate choice of the parameter \( p \) makes our method among the best classifiers. However, how to choose \( p \) is not evident to us, and a crossvalidation process seems to be needed unless the choice of \( p \) is guided by the query times requirements.

On the other hand, the computing times are, in all cases, extremely large, suggesting heuristic procedures for solving (P2); see Section 4.4. Moreover, from the columns giving the proportion of correctly classified objects in the training and the testing samples, the former strongly overestimates the latter. Hence, overfitting happens.

### 4.4. Heuristic Approach

The \( NP \)-hardness of the problem as well as the empirical results of Section 4.3 suggest heuristic procedures to speed up computing times. A first and simple choice, yielding promising results (Pekalska et al. 2006), might consist of randomly selecting \( p \) prototypes. The results (random choice in Tables 3–8) are discouraging, particularly when \( p \) is small. Hence, more sophisticated heuristics are needed.

The structure of the problem is such that several existing (meta) heuristic procedures can be easily adapted to our problem. Multistart, genetic algorithms or tabu search have been proposed for prototype selection, with encouraging results (Bezdek and Kuncheva 2001).

VNS combines local search with redefinitions of the neighborhood structure. We use the same neighborhoods as Hansen and Mladenović (1998, 2001a) for the \( p \)-median problem. Given a feasible solution, i.e. a set of \( p \) prototypes including at least one for each class, its neighborhood of order \( \ell \) consists of all feasible solutions that differ from it in at most \( \ell \) prototypes. The procedure works as described in Figure 3.

In our experiments, the procedure stops after 5,000 calls to step 2(b) in Figure 3. The results for the small data sets for which the exact solution was also sought with CPLEX are under Heuristic in Tables 3–6. Moreover, much larger data sets, such as abalone or spam, can be handled (Tables 7 and 8).

A very simple heuristic yields, with very low computing times, rather sharp solutions on the training samples. However the quality of the procedures should not be measured on the training sample (which overestimates the quality results) but on the testing sample. On testing samples, the heuristic yields (at much lower computing times) solutions with comparable quality than those obtained with the exact method.

We selected variable neighborhood search (VNS), proposed by Hansen and Mladenović (2001b), both

<table>
<thead>
<tr>
<th>Table 7</th>
<th>Results for abalone</th>
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<tbody>
<tr>
<td>( p )</td>
<td>Random choice (%)</td>
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<td></td>
<td>tr</td>
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<tr>
<td>3</td>
<td>61</td>
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<tr>
<td>5</td>
<td>68</td>
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<tr>
<td>10</td>
<td>69</td>
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<td>25</td>
<td>72</td>
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<td>50</td>
<td>75</td>
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<tr>
<td>100</td>
<td>78</td>
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<tr>
<td>500</td>
<td>85</td>
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</table>

<table>
<thead>
<tr>
<th>Table 8</th>
<th>Results for spam</th>
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<tbody>
<tr>
<td>( p )</td>
<td>Random choice (%)</td>
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<tr>
<td></td>
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<tr>
<td>2</td>
<td>61</td>
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<td>5</td>
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<td>500</td>
<td>85</td>
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<table>
<thead>
<tr>
<th>Table 9</th>
<th>Results with Missing Values for glass</th>
</tr>
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<tbody>
<tr>
<td>( p )</td>
<td>Fraction of missing values</td>
</tr>
<tr>
<td></td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>0.051</td>
</tr>
<tr>
<td>8</td>
<td>0.063</td>
</tr>
<tr>
<td>10</td>
<td>0.075</td>
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<tr>
<td>12</td>
<td>0.087</td>
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<tr>
<td>14</td>
<td>0.099</td>
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<tr>
<td>16</td>
<td>0.111</td>
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<td>18</td>
<td>0.123</td>
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Figure 3 VNS Heuristic

1. **Initialization**. Randomly choose an initial solution \( x \).
   Choose a stopping criterion.
2. Repeat until the stopping condition:
   (a) Set \( \ell \leftarrow 1 \).
   (b) Repeat until \( \ell = p \):
      i. Generate randomly a new solution \( x' \) differing in
         at most \( \ell \) prototypes with the current solution \( x \).
      ii. If \( x' \) is better than \( x \), set \( x \leftarrow x' \) and go to 2(a);
         otherwise, set \( \ell \leftarrow \ell + 1 \).
3. Return the best solution found so far.
for simplicity of its implementation and the excellent results obtained for related problems, such as the $p$-median problem. Other metaheuristics could be used to tackle the MIP (P1), but an empirical comparison of such methods is beyond the scope of this paper.

4.5. Missing Values
As mentioned in Section 2.1, dissimilarities can also be constructed for databases with missing values. We performed experiments to explore the stability of the classification rule with respect to the existence of (many) missing values. For different values of $\vartheta$, a fraction $\vartheta$ of data were randomly chosen and replaced by nulls.

We considered the dissimilarity described in (3), with each $\omega$, defined by (16). The optimization was performed using the VNS heuristic from Section 4.4 with its stopping rule. To reduce the random effects due to the inclusion of nulls, we ran each test 100 times. The average proportion of correctly classified objects in the testing sample, for different values of $p$ and $\vartheta$, together with the slopes of the regression lines linking percentage of correctly classified objects with fraction of missing data, are shown in Tables 9–12 for the small databases; see Carrizosa et al. (2005) for further details.

As expected, the quality of the classification rule deteriorates as the number of nulls increases. However, the correct classification rates decrease slowly since, for instance, in yeastME, the rules still classify correctly more than 70% of the objects when 30% of the values are missing. It is not evident how the degradation in classification is affected by the number of prototypes, since no trend is found in the slopes of the corresponding regression lines.

Table 11  Results with Missing Values for wine
\begin{tabular}{ccccccc|c}
\hline
\text{Fraction of missing values} & $p$ & 0 & 0.025 & 0.05 & 0.075 & 0.1 & 0.15 & 0.175 & 0.2 & 0.25 & 0.3 & Slope \\
\hline
3 & 92 & 94 & 93 & 94 & 93 & 92 & 91 & 85 & 92 & 90 & -14.58 \\
5 & 96 & 95 & 90 & 95 & 93 & 93 & 92 & 93 & 89 & 88 & -20.61 \\
7 & 96 & 94 & 95 & 94 & 93 & 94 & 93 & 92 & 90 & 89 & -20.27 \\
9 & 94 & 94 & 93 & 93 & 93 & 92 & 90 & 88 & 90 & 88 & -21.53 \\
\hline
\end{tabular}

Table 12  Results with Missing Values for yeastME
\begin{tabular}{ccccccc|c}
\hline
\text{Fraction of missing values} & $p$ & 0 & 0.025 & 0.05 & 0.075 & 0.1 & 0.15 & 0.175 & 0.2 & 0.25 & 0.3 & Slope \\
\hline
3 & 82 & 78 & 76 & 74 & 79 & 77 & 76 & 74 & 73 & 71 & -25.90 \\
5 & 77 & 79 & 81 & 79 & 75 & 75 & 74 & 70 & 71 & 71 & -30.81 \\
7 & 84 & 79 & 79 & 75 & 78 & 75 & 76 & 75 & 73 & 71 & -32.79 \\
9 & 78 & 74 & 79 & 76 & 76 & 74 & 74 & 71 & 72 & 73 & -19.45 \\
\hline
\end{tabular}

5. Conclusion and Further Research
We have introduced an optimization-based method for multiclass classification problems. Since the only requirement for the data is knowledge of a dissimilarity between entries, no statistical assumptions on data are needed, and qualitative variables, as well as missing values, are easily handled. Contrary to other competitive procedures, differences in misclassification costs are naturally accommodated within the model.

MIP formulations have been developed yielding classifiers with performance comparable to benchmark procedures. However, the computational effort required makes them prohibitive for databases of moderate size. Stronger MIP formulations, using valid inequalities, as well as more sophisticated bounding strategies, should be investigated since they might allow solving larger instances of this $\mathcal{NP}$-hard problem in reasonable time.

For large databases heuristics seem to be the only feasible approach. The results obtained using VNS are encouraging. An empirical comparison with other (meta)heuristics remains.

The dissimilarity $d$ has been considered given. However, the dissimilarity itself can be seen as a (modelling) decision variable. Hints to choose an appropriate $d$, e.g. by choosing appropriate weights $\omega$, in (15), are now under research.

Further study is also needed for the choice of the parameter $p$. Indeed, our numerical results do not lead to clear guidelines for choosing $p$. We propose crossvalidation, but in applications in which querying time is a critical issue, $p$ will have to be fixed exogenously.

Acknowledgments
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