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Fuzzy Inference System for Data Processing in Industrial Applications

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1. Introduction

In the last years Fuzzy Inference Systems (FIS) have been used in several industrial applications in the field of automatic control, data classification, decision analysis, expert systems, time series prediction, and pattern recognition.

The large use of FIS in the industrial field is mainly due to the nature of real data, that are often incomplete, noisy and inconsistent, and to the complexity of several processes, where the application of mathematical models can be impractical or even impossible, due to the lack of information on the mechanisms ruling the phenomena under consideration. Fuzzy theory is in fact essential and applicable to many complex systems and the linguistic formulation of its rule basis provides an optimal, very suitable and intuitive tool to formalise the relationships between input and output variables.

In real world database anomalous data (often called *outliers*) can be frequently found, which are due to several causes, such as erroneous measurements or anomalous process conditions. Outliers elimination is a necessary step, for instance, when building a training database for tuning a model of the process under consideration in standard operating conditions. On the other hand, in many applications, such as medical diagnosis, network intrusion or fraud detection, rare events are more interesting than the common samples. The rarity of certain patterns combined to their low separability from the rest of data makes difficult their identification. This is the case, for instance, of classification problems when the patterns are not equally distributed among the classes (the so-called *imbalanced dataset* (Vannucci et al., 2011)). In many real problems, such as document filtering and fraud detection, a binary classification problem must be faced, where the data belonging from the "most interesting" class are far less frequent than the data belonging to the second class, which corresponds to normal situations. The main problem with imbalanced dataset is that the standard learners are biased towards the common samples and tend to reduce the error rate without taking the data distribution into account.

In this chapter a preliminary brief review of traditional outlier detection techniques and classification algorithms suitable for imbalanced dataset is presented. Moreover some recent practical applications of FIS that are capable to outperform the widely adopted traditional methods for detection of rare data are presented and discussed.

2. Outlier detection

Outliers are measurements that are different from other values of the same dataset and can be due to measurement errors or to the variability of the phenomenon under consideration. Hawkins (Hawkins, 1980) defined an outlier as "*an observation that deviates so much from other observations as to arouse suspicion that it was generated by a different mechanism*".

The detection of outlier is an important step of data mining because it improves the quality of data and it represents a useful pre-processing phase in many applications, such as financial analysis, network intrusion detection and fraud detection (Hodge, 2004).

Classical outlier detection methods can be classified into four main groups: distance-based, density-based, clustering-based and statistical-based approaches. All these approaches have several advantages or limitations and in the last years a lot of contributions have been proposed on this subject. Artificial intelligence techniques have been widely applied to overcome the traditional methods and improve the cleanness of data; in particular some fuzzy logic-based approaches proved to outperform classical methodologies.

2.1 Distance-based methods

Distance-based method is based on the concept of the neighborhood of a sample and it was introduced by Knorr and Ng (Knorr & Ng, 1999). They gave the following definition: "*An object O in a dataset T is a DB(p,D)-outlier if at least fraction p of the objects in T lie at a distance greater than D from O*". The parameter p represents the minimum fraction of samples that is out of an outlier's D-neighborhood. This definition needs to fix a parameter and do not provide a degree of outlierness. Ramaswamy et al. (Ramaswamy et al., 2000) modified the definition of outlier: "*Outliers are the top n data points whose distance to the kth nearest neighbor is greatest*". Jimenez-Marquez et al. (Jimenez-Marquez et al., 2002) introduced the Mahalanobis Outlier Analysis (MOA) which uses Mahalnobis distance (Mahalanobis, 1936) as outlying degree of each point. Another outlier detection method based on Mahalanobis distance is defined as the distance between each point and the center of mass. This approach considers outliers data points that are far away from their center of mass.

2.2 Density-based methods

Density-based methods calculate the density distribution of data and classify as outliers the points lying in low-density regions. Breunig et al. (Breunig et al., 2000) allocate a local outlier factor (LOF) to each point on the basis of the local density of its neighborhood. In order to understand the formula concerning the Local Outlier Factor is necessary introduce several definitions. The **k-distance** of a point *x* is the distance between two points *x*, *y* belonging to the dataset *D* such that for at least *k* points data $d(x,y') \leq d(x,y)$; where $y' \in D-\{x\}$ and *k* an integer value. The **k distance neighborhood** of a data point *x* includes points whose distance from *x* is not greater than the *k distance*. Moreover the **reachability distance** of *a* data point *x* is defined as the inverse of the mean reachability distance based on the *MinPts*-nearest neighbors of *x*. Finally the Local Outlier Factor is defined as the mean of the ratio of the local reachability density of *x* and the cardinality of the set including the *MinPts*-nearest

neighbors of *x*. It is evident that *MinPts* is an important parameter of the proposed algorithm. Papadimitriou et al. (Papadimitriou et al., 2003) propose LOCI (Local Correlation Integral) which uses statistical values belonging to data to solve the problem of choosing values for *MinPts*.

2.3 Clustering-based methods

Clustering-based methods perform a preliminary clustering operation on the whole dataset and then classify as outliers the data which are not located in any cluster.

Fuzzy C-means algorithm (FCM) is a method of clustering developed by Dunn in 1973 (Dunn, 1973) and improved by Bezdek in 1981 (Bezdek, 1981). This approach is based on the notion of fuzzy c-partition introduced by Ruspini (Ruspini, 1969). Let us suppose $X=\{x_1, x_2, ..., x_n\}$ be a set of data where each *sample* x_h (h=1, 2, ..., n) is a vector with dimensionality p. Let U_{cn} be a set of real $c \times n$ matrices where c is an integer value which can assume values between 2 and n. The fuzzy C-partition space for x is the following set:

$$M_{cn} = \{ U \in U_{cn}; u_{ih} \in [0, 1] ; \sum_{i=1}^{c} u_{ih} = 1, 0 < \sum_{h=1}^{n} u_{ih} < n \}$$
(1)

where u_{ih} is the degree of membership of x_n in cluster i ($1 \le i \le c$). The objective of FCM approach is to provide an optimal fuzzy C-partition minimizing the following function:

$$J_m(U,V;X) = \sum_{h=1}^n \sum_{i=1}^c (u_{ih})^m \|x_k - v_i\|^2$$
(2)

where $V = (v_1, v_2, ..., v_c)$ is a matrix of cluster centres, $\|.\|$ is the Euclidean norm and *m* is a weighting exponent (*m*>1).

Many clustering-based outlier approaches have been recently developed. For instance, Jang et al. (Jang et al., 2001) proposed an outlier-finding process called OFP based on k-means algorithm. This approach considers small clusters as outliers. Yu et al. (Yu et al., 2002) proposed an outlier detection method called FindOut, which is based on removing of clusters from original data to identify outliers. Moreover He et al. (He et al., 2003) introduced the notion of cluster-based local outlier and outlier detection method (FindCBLOF), which exploits a cluster-based LOF in order to identify the outlierness of each sample. Finally Jang et al. (Jang et al., 2005) proposed a novel method in order to improve the efficiency of FindCBLOF approach.

2.4 Statistical-based methods

Statistical-based methods use standard distribution to fit the initial dataset. Outliers are defined considering the probability distribution and assuming that the data distribution is a priori known. The main limit of this approach lies in the fact that, for many applications, the prior knowledge is not always distinguishable and the cost for fitting data with standard distribution could be considerable. A widely used method belonging to distribution-based approaches has been proposed by Grubbs (Grubbs, 1969). This test is efficient if data can be approximated by a Gaussian distribution. The Grubbs test calculates the following statistics:

$$G_i = \frac{\max_i \left(x_i - \mu \right)}{\sigma} \tag{3}$$

where μ is the mean value and σ the standard deviation of data. When *G* is greater than a fixed threshold the *i*-th data-point is classified as an outlier. The critical value depends on the required significance level of test; common values are 1% and 5%. Other similar tests that assume that data are normally distributed are Rosner's test (Gibbons, 1994) and Dixon's test (Dixon, 1993).

2.5 Fuzzy Inference System based method

In the last years novel interesting FIS-based outlier detection approaches have been proposed in order to outperform the classical approaches.

Yousri et al. (Yousri et al., 2007) proposes an approach which combine an outlier detection method with a clustering algorithm. The outlier detection method is used for two objectives: to give a hard membership for outliers to decide if the considered pattern is an outlier or not and to give a degree of outlierness. A clustering algorithm is then used to allocate patterns to clusters. Let *P* indicate the dataset and *p* its generic sample and let *To* be the outlier detection technique and *Tc* the adopted clustering algorithm. The combination of outlier detection and clustering algorithms is provided by the following formula, providing the degree of outlierness of a sample *p*:

$$O_p = \frac{w_o O_{To}(p) + w_c O_{Tc}(p)}{w_o + w_c}$$
(4)

where $O_{T_o}(p)$ is the degree of outlierness resulting from T_o for each sample p while $O_{T_c}(p)$ is the degree of outlierness resulting from T_c considering an outlier the patterns allocated to tiny clusters or not assigned to any cluster. Finally w_o and w_c represent the weights given to both algorithms to determine outliers. The two weights must be not negative (w_o , $w_c \ge 0$) and their sum must be positive ($w_o+w_c > 0$). Equation (4) can be rewritten as follows:

$$O_{p} = \frac{\frac{W_{o}}{W_{c}}O_{To}(p) + O_{Tc}(p)}{\frac{W_{o}}{W_{c}} + 1}$$
(5)

The parameter w_0/w_c should be accurately fixed considering a balance between the membership degree given to the outlier cluster and the clusters in the set of initial groups.

The main advantage of this approach is that it is general, i.e. it can combine any outlier detection method with any clustering algorithm, and it is effective with low and high dimensional dataset.

Another novel fuzzy approach is proposed by Xue et al. (Xue et al., 2010). The approach is called *Fuzzy Rough Semi-Supervised Outlier Detection* (FRSSOD) and combines two methods: the *Semi-Supervised Outlier Detection method* (SSOD) (Gao et al., 2006) and the clustering method called *Fuzzy Rough C-Means clustering* (FRCM) (Hu & Yu, 2005). The aim of this approach is to establish if samples on the boundary are outliers or not, by exploiting the advantages of both approaches. In order to understand FRSSOD a brief description of SSOD and FRCM is necessary.

SSOD is a semi-supervised outlier detection method (Li et al. 2007; Zhang et al. 2005; Gao et al., 2006; Xu & Liu, 2009) main that uses both unlabeled and labelled samples in order to improve the accuracy without the need for a high amount of data. Let us suppose that X is the dataset with *n* samples forming *K* clusters. The first *l* samples are labelled as binary values: the null samples are considered outliers while the unitary value are not outliers. If we consider that outliers are not included in any of the *K* clusters, an $n \times K$ matrix must be found whose elements t_{ih} (i=1,2...,n; h=1, 2, ..., K) are unitary when x_i belongs to cluster C_h . Outliers are determined as points that do not belong to any clusters through the minimization of the following objective function:

$$Q = \sum_{i=1}^{n} \sum_{h=1}^{K} t_{ih} dist (c_h, x_i)^2 + \gamma_1 (n - \sum_{i=1}^{n} \sum_{h=1}^{K} t_{ih}) + \gamma_2 \sum_{i=1}^{l} \left| u_i - \sum_{h=1}^{K} t_{ih} \right|$$
(6)

where c_h represent the centroid of cluster C_h , *dist* is the Euclidean distant and γ_1 , γ_2 are adjusting parameters. The objective function is the sum of three parts. The first part come from k-means clustering and outliers are not considered, the second part is used to constrain the number of outliers below a certain threshold and finally the third part is used to maintain consistency of labelling introduced by authors with existing label.

FRCM is a combination between Fuzzy C-means algorithm and Rough C-means approach. Fuzzy C-means method was introduced by Dunn (Dunn, 1974) and it is an unsupervised clustering algorithm. The approach assigns a membership degree to each sample for each cluster by minimizing the following objective function:

$$J_m(U,V;X) = \sum_{k=1}^n \sum_{i=1}^c u_{ik}^m \|x_k - v_i\|^2$$
(7)

where $V = (v_1, v_2, ..., v_c)$ is the vector representing the centres of the clusters, u_{ik} is the degree of membership of the sample x_k to the cluster *i*. The iteration stops when a stable condition is reached and the sample is allocated to the cluster for which the membership value is maximum. The centres initialization is an important step affecting the final result.

The RCM approach is based on the concept of C-means clustering and on the concept of rough set. Rough set was introduced by Pawlak (Pawlak, 1982; Pawlak, 1991). In the rough set concept each observation of the universe has a specified amount of information. The objects which have the same information are indistinguishable. A rough set, unlike a precise set, is characterized by lower approximation, upper approximation and boundary region. The lower approximation includes all objects belonging to the considered notion, while the upper approximation contains objects which possibly belong to the notion. The boundary region represents the difference between the two regions. In RCM method each cluster is considered as a rough set having the three regions. A sample, unlike in the classical clustering algorithm, can be member of more than one cluster. Also, it is possible to have overlaps between clustering. Lingras and West (Lingras & West, 2004) proposed a method based on the following four properties:

- A sample can belong only to one of lower approximation regions.
- The lower approximation region of a cluster must be a subset of its upper approximation region.
- If a sample do not belong to any lower approximation regions then it is member of at least two upper approximation regions.
- Samples belonging to boundary region are undecided data and are assigned to two or more upper approximation regions.

FRCM integrates the concept of rough set with the fuzzy set theory adding a fuzzy membership value of each point to the lower approximation and boundary region of a cluster. The approach divides data into two sets, a *lower approximation region* and a *boundary region*; then the points belonging to the boundary region are fuzzified. Let us suppose that $X=\{x_1, x_2, ..., x_n\}$ is the available dataset and \underline{C}_h and \overline{C}_h are respectively the lower and upper approximation of the cluster *h*. The boundary region is calculated as the difference between the two regions. If $u=\{u_{ih}\}$ are memberships of clusters the problem of FRCM become the optimization of the following function:

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$$J_m(u,v) = \sum_{i=1}^n \sum_{h=1}^H (u_{ih})^m d_{ih}^2$$
(8)

FRSSOD combines the two methods above described, i.e. FRCM and SSOD, in order to create a novel approach. Let *X* be the set of data with *n* samples and *Y* its subset formed by the first l < n samples. The elements of *Y* are labelled as $y_i = \{1, 0\}$ where null value indicates that the considered point is an outlier. The normal points, i.e. points that are not considered outliers, form *C* clusters and each point normal point belong to each cluster with a membership value, while outliers do not belong to any cluster. The main aim of FRSSOD is to create a $n \times c$ matrix called *u*, whose generic entry u_{ik} represents the fuzzy membership degree of the i_{th} sample on the cluster. The optimization problem consists in the minimization of the following function:

$$J_m(u,v) = \sum_{i=1}^n \sum_{k=1}^C (u_{ik})^m d_{ik}^2 + \gamma_1 (n - \sum_{i=1}^n (\sum_{k=1}^C u_{ik})^m) + \gamma_2 \sum_{i=1}^l (y_i - \sum_{k=1}^C u_{ik})^2$$
(9)

where $\gamma_1 e \gamma_2$ are adjusting positive parameters in order to make the three terms compete with each other and *m* is a fuzziness weighting exponent (*m*>1). As the idea of SSOD approach only normal points are divided in two clusters and also the points considered as outliers don't compare in the first term of the equation. Then the second term of the equation is used to maintain the number of outliers under a certain limit and finally the third term maintains consistency of user labelling with existing label punishing also the mislabelled samples. The proposed method is applied on a synthetic dataset and on real data and results show that FRSSOD can be used in many fields having fuzzy information granulation. The experimental results show also that the proposed method has many advantages over SSOD improving outlier detection accuracy and reducing false alarm rate thanks to the control on labelled samples. The main disadvantages of the FRSSOD method are that the result depends on the determination of number of cluster, initialization of the centres of clusters and adjustment parameters.

Another fuzzy based method to detect outliers is proposed by Cateni et al. (Cateni et al., 2009). The proposed approach combines different classical outlier detection techniques in order to overcome their limitations and to use their advantages. An important advantage of the proposed method lies in the fact that the system is automatic and no a priori assumptions are required. The method consists in calculating four features for each pattern by using the most popular outlier detection techniques:

1. **Distance-based.** The Mahalanobis distance is calculated and normalized with respect to its maximum value. Patterns which assume value near 1 are considered outliers.

- 2. Clustering-based. The clustering algorithm used for each method is the fuzzy C-means (FCM) that has already been described. The output of the clustering represents the membership degree of the selected samples to clusters and it lies in the range [0,1]. Samples for which such features is close to 0 have to be considered outliers. The FCM approach requires the number of clusters to be a priori known. If the distribution is unknown, it is not easy to find the optimal number of clusters. In this approach a validity measure based on intra-cluster and inter-cluster distance measures (Ray & Turi, 1999) is calculated in order to determine automatically the most suitable number of clusters. This step is fundamental because the result of clustering strongly depends on this parameter.
- 3. **Density-based.** For each pattern the Local Outlier Factor (LOF) is evaluated. This feature requires that the number of the samples of nearest neighbours *K* must be known a-priori. Here *K* corresponds to the number of elements in the less populated cluster that has been previously calculated by the Fuzzy C Means algorithm. The LOF parameter lies in the range [0;1] where the unitary value means that the considered sample is an outlier.
- 4. **Distribution-based.** The Grubbs test is performed and the result is a binary value: an unitary value indicates that the selected pattern is an outlier, is null otherwise.

This four features are fed as inputs to a FIS of the Mandani type (Mandani, 1974). The output of the system, called *outlier index*, represents the degree of outlierness of each pattern. Finally a threshold, set to 0.6, is used to point out the outliers.

The proposed method has been tested in a dataset provided by a steelmaking industry in the pre-processing phase. The extracted data represent the chemical analysis and process variables associated to the liquid steel fabrication. A dataset has been analyzed in order to find factors that affect the final steel quality. In this case outliers can be caused by sensor failures, human error during registration of data of off-line analysis or abnormal conditions; this kind of outliers are the most difficult to discover because they do not differ so much from some correct data. In the considered problem the variables which are mostly affected by outliers are tapping temperature, reheating temperature and the addition of aluminium. First of all each variable has been normalised in order to obtain values in the range [0,1]. Table 1 illustrates the results that have been obtained on 1000 measurements through 6 different approaches: a distance-based approach, a clustering-based approach a densitybased approach, a distribution-based approach and two fuzzy-based approaches. The effective outliers are five and they have been pointed out by technical skilled personnel working on the plant and in the table are identified with alphabetic letters in order to understand how many and which outliers are correctly detected by several methods. In the second row the outliers are shown that are present for each considered variable and the other columns refer to outliers detected by each considered method. Finally, table 2 illustrates how many samples have been misclassified as outliers (the so-called *false alarms*) for each method.

Table 2 shows that fuzzy-base approach is able to detect all outliers present in the dataset without fall in false alarms errors. Therefore the obtained results confirm the effectiveness of the fuzzy-based approaches because they outperform traditional techniques and in particular the second fuzzy proposed approach (Cateni et al., 2009) obtains the best results.

	Tapping Temperature	Reheating Temperature	Aluminium addition	
OUTLIERS	A -B - C	A - B - C	A - B - C - D -E	
Mahalanobis distance	В	В	A - D - E	
Fuzzy C-means	A – B	A - B	B – C	
Local Outlier Factor	В	В	A - B - C - E	
Grubbs Test	А	А	B - D - E	
Fuzzy 1 (Yousri et al.,2007)	A - B	A - B	A - B - C - E	
Fuzzy 2 (Cateni et al., 2009)	A - B -C	A - B -C	A - B - C - D -E	

Table 1. Outlier Detection.

	Tapping Temperature	Reheating Temperature	Aluminium addition
OUTLIERS	A -B - C	A - B - C	A - B - C - D -E
Mahalanobis distance	0	0	1
Fuzzy C-means	0	0	1
Local Outlier Factor	1	0	0
Grubbs Test	0	0	2
Fuzzy 1 (Yousri et al.,2007)	0	0	0
Fuzzy 2 (Cateni et al., 2009)	0	0	0

Table 2. Number of false alarms.

3. Imbalanced datasets

In a classification task often there are more instances of a class than others. Class imbalance is mainly connected to the concept of *rarity*. There are two types of rarity, *rare cases* or *outliers* (as seen in the previous paragraph) and *rare class* where a class of interest includes few samples in comparison to the other classes present in the dataset. Outliers and rare classes are not related but there are empirical studies which demonstrate that the minority class contains more outliers than the majority class (Weiss and Provost, 2003). Imbalanced datasets are present in many real-world applications such as detecting cancerous cell (Chan and Stolfo, 1998), fraud detection (Phua et al., 2004), keyword extraction (Turney, 2000), oil-spill detection (Kubat et al., 1998), direct marketing (Ling and Li, 1998), and so on. Many approaches have been proposed in order to solve the imbalance problem (Visa and Ralescu, 2005) and they include resampling the training set, feature selection (Castillo and Serrano, 2004), one class learners (Raskutti and Kowalczyk, 2004) and finally cost-sensitive learners, that take into account the misclassification cost (Zadrozny et al., 2003). In this kind of data one class is significantly larger than other classes and often the targeted class (known as *positive* class) is the smallest one. In this cases classification is difficult because the

conventional computational intelligence methods tend to classify all instances to the majority class (Hong et al., 2007). Moreover this methods (such as Multilayer Perceptron, Radial Basis Functions, Linear Discriminant Analysis...) cannot classify imbalanced dataset because they learn data based on minimization of accuracy without taking into account the error cost of classes (Visa & Ralescu, 2005; Alejo et al., 2006; Xie & Qiu, 2007).

The performance of machine learning is typically calculated by a confusion matrix. An example of the confusion matrix is illustrated in Tab.3 where columns represent the predicted class while the rows the actual class. Most of studies in imbalanced domain are referred to binary classification, as a multi-class problem can be simplified to a two-class problem. Conventionally the class label of the minority class is positive while the class label of the majority class is negative. In the table True Negative value (*TN*) represents the number of negative samples correctly classified, True Positive (*TP*) value is the number of positive samples correctly classified, False Positive (*FP*) is the number of negative samples classified as negative. Other common evaluation measures are *Precision* (Prec) which is a measure of the accuracy providing that a specific class has been predicted and *Recall* (Rec) which is a measure of a prediction model to select instances of a certain class from a data set. In this case, Recall is also referred to as true positive rate and true negative rate is also called *Specificity* (Spec).

	Predicted Negative	Predicted Positive
Negative	TN	FP
Positive	FN	TP

Table 3. Confusion Matrix

Through this matrix the following widely adopted evaluation metrics can be calculated:

$$Accuracy = (TP+TN)/(TP+FN+FP+TN)$$
(10)

$$FP \ rate = Spec = FP/(TN+FP) \tag{11}$$

$$TP \ rate = Rec = TP/(TP+FN)$$
(12)

$$Prec = TP/(TP+FP)$$
(13)

$$F - Measures = [(1+\beta^2)*Rec*Prec]/[(\beta^2*Prec)+Prec]$$
(14)

where β corresponds to the relative importance of *Precision* versus *Recall*. Typically β =1 when false alarms (false positive) and misses (false negative) can be considered equally costly.

3.1 Traditional approaches

In general approaches for imbalanced dataset can be divided in two categories: *external* and *internal* approaches. The external methods do not depend on the learning algorithm to be used: they mainly consist in a pre-processing phase aiming at balancing classes before training classifiers. Different re-sampling methods fall into this category. In

contrast internal method develops variations of the learning algorithm in order to solve the imbalance problem.

3.1.1 External methods

Re-sampling strategies have several advantages. First of all re-sampling methods are competitive (McCharty, 2005) and often similar (Maloof, 2003) to the results obtained choosing the cost-sensitive learning. Moreover re-sampling methods are simple and do not require to modify the internal working of the classifier chosen (Elkan, 2001). Re-sampling methods are pre-processing techniques and they can be divided in two categories: oversampling techniques and undersampling techniques. Oversampling methods balance the classes by adding new points to the minority class while undersampling methods increase the number of samples belonging to the minority class.

The simplest re-sampling techniques are the random oversampling method and the random undersampling methods. The random oversampling method balances the distribution classes by randomly replicating some instances belonging to the minority class but random oversampling can lead to overfitting. The random undersampling technique randomly removes negative examples from the majority class encountering the problem to deleting some important information of the dataset. Both random techniques sample the dataset until the classes are approximately balanced. In order to solve the cited limitations improved resampling techniques have been proposed.

3.1.1.1 Oversampling techniques

Synthetic Minority Oversampling TEcnique (SMOTE) creates minority samples to oversample the minority class avoiding the overfitting problem (Chawla et al., 2002). Instead of replicating existing data points SMOTE generates new samples as follow: for every minority example, its *n* nearest neighbours belonging to the same class are evaluated (in SMOTE *n* is set to 5); then new synthetic data points are randomly generated along the segment joining the original data point and its selected neighbour. Figure 1 shows an example concerning SMOTE algorithm: x represents the selected data point, n_i are the selected nearest neighbours and s_1 , s_2 and s_3 are samples generated by the randomized interpolation.

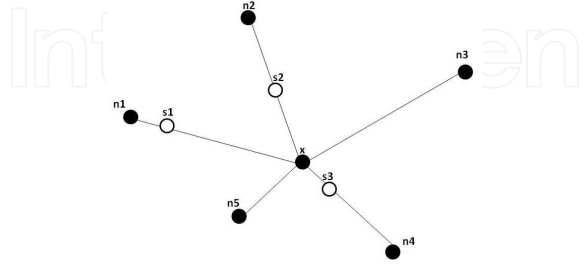


Fig. 1. Example of SMOTE algorithm.

As SMOTE over-generalizes the minority class but does not take into account the distribution of neighbours from the majority class another novel approach, called Borderline-SMOTE (Han et al., 2005), is proposed. This approach is a generalization of SMOTE approach, it focuses the attention on oversampling around samples located in the borderline between classes. This approach is based on the assumption that the positive instances are divided into three regions: *noise, borderline* and *safe* by considering the number of negative examples on k nearest neighbours. If n is the number of negative examples among the k nearest neighbours, the regions are defined as follow:

- Noise: *n*=*k*
- Borderline: $0.5k \le n \le k$
- Safe: 0≤*n*≤0.5*k*

Borderline-SMOTE exploits the same oversampling technique as SMOTE but it oversamples only the instances belonging to the borderline region.

Bunkhumpornpat et al. proposed another approach called Safe-Level-SMOTE (Bunkhumpornpat et al., 2009). This method assigns for each positive data-points a *safe-level* (*sf*) which is defined as the number of positive data-points in the *k* nearest neighbours. If the safe level of an instance is close to *k*, then it considered safe. The safe level ratio is defined as the ratio between the safe level of a positive instance and the safe level of a nearest neighbours. Also each synthetic new point is generated in safe region by considering the safe level ratio of instances. This method is able to outperform both SMOTE and Borderline-SMOTE because they may generate instances in unsuitable positions such as overlapping or noise regions.

Wang et al. (Wang et al., 2006) propose a novel approach that improves the SMOTE algorithm by including the Locally Linear Embedding (LLE) algorithm (Sam & Lawrence, 2000). The SMOTE approach has an important limitation: it assumes that the local space between any positive samples is positive, i.e. belonging to the minority class. This fact could be not true if the training data is not linearly separable. This method maps the training data into a lower dimensional space through the Locally Linear Embedding technique, then SMOTE is applied in order to create the desirable number of synthetic data points and finally the new data points are mapped back to the initial input space. The so-called LLE-based SMOTE algorithm is evaluated on three datasets applying three different classifiers: Naive Bayesian, K-NN classifier and Support Vector Machine (SVM). Experimental results show that the LLE-based SMOTE algorithm outperforms the conventional SMOTE.

Liu and Ghosh (Liu & Ghosh, 2007) propose a novel oversampling method, called generative oversampling, which increases information to the training set by creating artificial minority class points on the basis of the probability distribution to model the minority class. Also, generative oversampling can be used if the data distribution can be approximated with an existing model. Firstly a probability distribution is chosen in order to model the minority class, then parameters for the probability distribution are studied on the basis of the training data and finally synthetic data points are created from the learned probability distribution until the necessary number of points belonging to the minority class has been reached. Authors demonstrate that this approach works well for a range of text classification datasets using as classifier a SVM classifier. This method is simple to develop and it is suitable for several data types by selecting appropriate generative models.

3.1.1.2 Undersampling techniques

A popular undersampling method is the Condensed Nearest Neighbour (CNN) rule (Hart, 1968). CNN is used in order to find a consistent subset of samples. A subset \hat{S} is defined consistent with *S* if, using a one nearest neighbour, \hat{S} correctly classifies the instances in *S*. Fawcett and Provost (Fawcet and provost, 1997) propose an algorithm to extract a subset \hat{S} from *S* and using the approach as an undersampling method. Firstly one example belonging to the majority class is randomly extracted and put with all examples belonging to the minority class in \hat{S} . Then a 1-NN over the examples in \hat{S} is used in order to classify the examples belonging to *S*. If an example in *S* is misclassified, it is moved to \hat{S} . The main aim of this method is to delete the examples belonging to the majority class belonging to the majority class belonging to the majority class belonging to the example in *S* is misclassified, it is moved to \hat{S} . The main aim of this method is to delete the examples belonging to the majority class which are distant from the decision border.

Another undersampling approach is the so-called *Tomek links* (Tomek, 1976). This method can be defined as follow: let us suppose that x_i and x_j are two examples belonging to different classes and $d(x_i, x_j)$ is their distance. A pair (x_i, x_j) is called a Tomek link if there is not an example x_k such that $d(x_i, x_l) < d(x_i, x_j)$ or $d(x_j, x_l) < d(x_i, x_j)$. If two examples are a Tomek link then either one of these is noise or both are borderline. If Tomek Link is used as underline sampling, only samples belonging to the majority class are removed. Kubat and Matwin (Kubat & Matwin, 1997) propose a method, called One-Side Selection (OSS) which uses both Tomek Link and CNN. Tomek Link is used as undersampling technique removing noisy and borderline samples belonging to the majority class. Borderline samples are considered as *unsafe* since noise can make them fall on the wrong side of the decision border. CNN is used to delete samples belonging to the majority class which are distant from the decision border. The remainder samples including *safe* samples of majority class and all samples belonging to the minority class, are used for learning.

3.1.2 Internal methods

Internal methods deal with variations of a learning algorithm in order to make it less sensitive for the class imbalance.

Two common methods Boosting and Cost-Sensitive learning are used in this area.

Boosting is a method used to improve the accuracy of weak classifiers. The most famous boosting algorithm is the so-called AdaBoost (Freund & Schapire, 1997). It is based on the fusion of a set of weak learners, i.e. classifiers which have better performance than random classifiers in a classification task. During the learning phase weak learners are trained and included in the strong learner. The contribution of the added learners is weighted on the basis of their performance. At the end all modified learners contribute to classify unlabelled samples. This approach is suitable to deal with imbalanced dataset because the samples, belonging to the minority class, are most likely to be misclassified and also have higher weights during iterations. In literature several approaches using boosting techniques for imbalanced dataset has been proposed (Guo & Viktor, 2004; Leskovec & Shawe-Taylor, 2003) and results confirm the effectiveness of the method.

Another effective approach is the cost-sensitive learning. In this approach cost is associated with misclassifying samples; the cost matrix is a numerical representation of the penalty of classifying samples from a class to another. A correct classification has not penalty and the

cost of misclassifying minority examples is higher than the cost of misclassifying the majority examples. The aim of this approach is to minimize the overall cost on the training dataset. The cost matrix can balance the dataset by assigning the cost misclassifying a class with inverse proportion to its frequency. Another way is to set the cost matrix by considering the application driven criteria taking into account user requirements. Cost matrix is a general notion which can be exploited within common classifiers such as decision tree (Pazzali et al., 2004; Chawla, 2003) or neural networks (De Rouin et al., 1991).

Soler and Prim (Soler & Prim, 2007) propose a method based on the Rectangular Basis Function network (RecBF) in order to solve the imbalance problem. RecBF networks have been introduced by Berthold and Huber (Berthold & Huber, 1995) and are a particular type of Radial Basis Function (RBF) networks which exploit neurons with hyper-rectangular activation function in the hidden layer.

3.2 Fuzzy based approaches

In classification task with imbalanced dataset SVMs are widely used (Baser et al., 1992). In (Akbani et al., 2004; Japkowicz & Shayu, 2002) the capabilities of SVM and their effect on imbalance have been widely discussed. SVM is a widely used machine learning method which has been applied to many real world problems providing satisfactory results. SVM works effectively with balanced dataset but provides suboptimal classification models considering the imbalanced dataset; several examples demonstrate this conclusion (Veropoulus et al., 1999; Akbani et al., 2004; Wu & Chang, 2003; Wu & Chang, 2005; Raskutti & Kowalczyk, 2004; Imam et al., 2006; Zou et al., 2008; Lin et al., 2009; Kang & Cho, 2006; Liu et al., 2006; Haibo & Garcia, 2009). SVM is biased toward the majority class and provides poor results concerning the minority class.

A limit of the SVM approach is that it is sensitive to outliers and noise by considering all the training samples uniformly. In order to overcome this problem a Fuzzy SVM (FSVM) has been proposed (Lin & Wang, 2002) which is a variant of the traditional SVM algorithm. FSVM associates different fuzzy membership values (called weights) for different training samples in order to assign their importance degree of its class. Subsequently the proposed approach includes these weights inner the SVM learning algorithm in order to reduce the effect of outliers or noise when finding the separating hyperplane.

An extension of this approach is due to Wang et al. (Wang et al.,2005). They introduced two membership values for each training sample defining the membership degree of positive and negative classes. This approach has been proposed again by Hao et al. (Hao et al., 2007) based on the notion of *vague set*.

Spyrou et al. (Spyrou et al., 2005) propose another kind of fuzzy SVM approach which uses a particular kernel function built from fuzzy basis functions. There are also other works which combine fuzzy theory with SVM assigning a membership value to the outputs of the algorithm. For example Xie et al. (Xie et al., 2005) define a membership degree for the output class through the decision value generated by SVM algorithm, while Inoue and Abe (Inoue & Abe, 2001) use the fuzzy output decision for multiclass classification. Finally Mill and Inoue (Mill & Inoue, 2003) propose an approach which generates the fuzzy membership values for the output classes through the strengths support vectors.

Another typology of FSVM regards the extraction of fuzzy rules from the trained SVM model and a lot of works were been proposed (Chiang & Hao, 2004; Chen & Wang, 2003; Chaves et al., 2005; Castro et al., 2007).

The above approaches demonstrate that FSVM outperforms the traditional SVM algorithm avoiding the frequent problem of outliers and noise. However the FSVM technique, such as the SVM method, can be sensitive to the class imbalance problem. Batuwita and Palade (Batuwita & Palade, 2010) propose a novel method which uses the FSVM for class imbalance leraning (CIL). This approach, that is called FSVM-CIL, is able to classify with a satisfactory accuracy solving both the problems of class imbalance and outlier/noise. FSVM-CIL is been improved by Lakshmanan et al. (Lashmanan et al., 2011) extending the approach to multi-class classification problem instead of binary classification.

The general implementation scheme of the proposed approach is represented in figure 2.

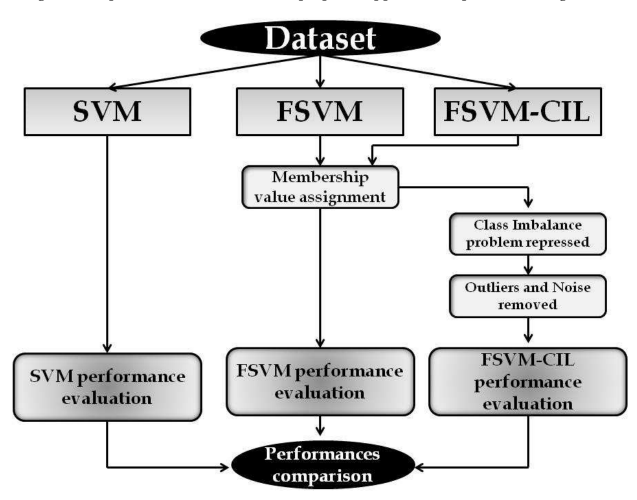


Fig. 2. Implementation diagram

FSVM-CIL method assigns a membership value for training examples in order to suppress the effect of class imbalance and to reflect the within-class importance of different training samples suppressing the effect of outliers and noise. Let us suppose that k_i^+ and k_i^- are the membership values of the positive class sample x_i^+ and negative class sample x_i^- in their own class respectively. They are calculated as follows:

$$k_{i^{+}} = f(x_{i^{+}})^{*}k^{+}$$
 $k_{i^{-}} = f(x_{i^{-}})^{*}k$ (15)

where k^+ and k^- are values which reflect the class imbalance such that $k^+ > k^-$ and $f(x_i)$ is defined considering the distance between x_i and its class centre d_i .

Samples near to the class centre are considered important samples because containing more information and also their $f(x_i)$ value is high. In contrast, samples which lie far away from centres are treated as outliers or noise and their $f(x_i)$ value is low. In FSVM-CIL approach authors use two separate decaying functions of distance to define $f(x_i)$: a linearly decaying function $f_{lin}(x_i)$ and $f_{exp}(x_i)$. The two functions are defined as follows:

$$f_{lin}(x_i) = 1 - [d_i/(max\{d_i\} + a)]$$

where *a* is a small positive value which is introduced in order to avoid that $f_{lin}(x_i)$ could assume null value.

$$f_{exp}(x_i) = 2/(1 + exp\{\beta d_i\}) \tag{17}$$

where β , which can assume values inner the range [0;1], determines the steepness decay and d_i is the Euclidean distance between x_i and its own centre.

The performance measure used in this approach is the geometric mean of sensitivity *GM* and it is represented in equation (18).

$$GM = \sqrt{SE * SP} \tag{18}$$

where *SE* and *SP* are the proportions of positive and negative samples among the correctly classified ones, respectively.

Three datasets (Blanke & Merz, 1988) have been exploited in order to demonstrate the effectiveness of the FSVM-CIL approach. Then a comparison between this method and SVM and FSVM has been evaluated considering both linear and exponential decaying function. Table 4 shows a description of the used datasets while Tab.5 illustrates the obtained results for the several approaches tested.

	# Positive Samples	# Negative Samples	IR
Ecoli	77	259	0.297
Pima Indians	268	500	0.536
Page Blocks	115	5358	0.021

Table 4. Datasets descriptions. IR represents the imbalance ratio, i.e. the ratio between the positive and the negative class.

As already mentioned, SVM classifier favours the majority class over the minority class obtaining an high value of SP and a low value of SE. This result confirms that SVM classifier is sensitive to the imbalance problem. Concerning FSVM results show that the best FSVM setting depends on the considered dataset. For Ecoli and Pima databases the choice of the type of decaying function is irrelevant while treating with page-blocks dataset the exponential decaying function provides better results than the linear one. Finally table shows that concerning FSVM-CIL approach, the use of exponential decaying function provides the best results independently from the considered dataset.

(16)

		SVM (%)	FSVM (lin) (%)	FSVM (exp) (%)	FSVM- CIL (lin) (%)	FSVM- CIL (exp) (%)
	GM	88.59	88.53	85.50	89.08	90.64
ECOLI	SE	78.67	78.18	78.64	90.02	92.45
	SP	93.12	93.60	92.97	86.52	88.19
	GM	70.18	69.78	71.33	71.56	72.74
PIMA	SE	55.04	54.55	55.60	66.94	69.10
	SP	89.50	87.5	82.50	76.14	76.35
	GM	76.47	79.15	81.62	94.51	95.05
PAGE	SE	58.26	61.74	67.06	93.81	93.14
	SP	99.54	91.48	99.01	95.21	95.36

Table 5. Classification results and comparison between several methods

Jesus et al. (Jesus et al., 2006) propose the study of the performance of Fuzzy Rule Based Classification System (FRBCS) in imbalanced datasets (Chi et al., 1996). The authors analyze the synergy of the linguistic FRBCS with some pre-processing techniques using several approaches such as undersampling, oversampling or hybrid models. A FRBCS is composed by a Knowledge Base (KB) and a Fuzzy Reasing Method (FRM). FRM uses the information of KB in order to determine the class for each sample which goes to the system. KB is composed of two elements: the Data Base (DB) which includes the notion of the fuzzy sets associated to the linguistic terms and the Rule Base (RB) which contains a set of classification rules.

The FRM, which is an inference procedure, utilizes the information of KB to predict a class from an unclassified sample. In a classification task the model of FRM includes four operations:

- 1. Compute the compatibility degree of data with the precedent of the rules.
- 2. Compute the association degree of data to the consequent class of each rule. This step consists in the generation of an aggregation function between the compatibility degree and the certainty degree of the rule with the class related.
- 3. Set the association degree of data with the several classes.
- 4. Classify by applying a decision function *F* on the association degree of data with classes.

FRBCS has been proposed in (Chi et al., 1996) and is an extension of the well known Wang & Mendel method (Wang & Mendel, 1992) to classification task. FRBCS finds the relationship between variables of the problem and establishes an association between the space of the features and the space of the classes. The main operations are as follows:

- Firstly a domain of variation of each feature Xi is determined, then fuzzy partitions are computed. This step is important in order to establish the linguistic partitions.
- For each example, a fuzzy rule is generated. Rules are created by considering several main steps.
 - Computing the matching degree of each example to the several fuzzy regions.
 - Assigning the selected example to the fuzzy region with the higher membership degree.

- Creating a rule for each example in this manner: the antecedent is determined by the selected Fuzzy region with the label of class of the sample in the subsequent.
- Compute the certainty degree.

In order to demonstrate the effectiveness of the proposed approach, authors considered several datasets belonging to the UCI repository with different degrees of imbalance.

The proposed method has been divided into three parts: an analysis of the use of preprocessing for imbalanced problems (such as SMOTE, random oversampling, random under-sampling ...), a study of the effect of the FRM and an analysis of the influence of the granularity applied to the linguistic partitions in combination with the inference method.

Results show that in all considered cases the presence of the pre-processing phase improves the behaviour of the learning algorithm.

The main conclusion of the proposed method is that the FRCM algorithm outperforms the other analyzed methods obtaining the best results adding a re-sampling operation before use the FRCM technique; moreover authors have found that FRBCSs perform well again the C4.5 decision tree in the context of very high imbalanced datasets.

An alternative to imbalance problem is the use of Complementary Learning Fuzzy Neural Network (CLFNN) which is proposed in (Tan et al., 2007). The use of fuzzy logic allows to tolerate uncertainty in the data reducing the effect of data imbalance. CLFNN has the main advantage that does not requires data pre-processing and hence, does not make any a prior assumption on data and does not alter the data distribution. This method exploits a neuro-fuzzy system which is based on complementary learning theory (Tan et al., 2005). Complementary learning is a system observed in human brain; with this theory different brain areas, which are segregated and mutually exclusive, are registered in order to recognize different objects (Gauthier, 2000). When an object is seen, registered areas are activated while the irrelevant areas are inhibited; this mechanism is called lateral inhibition. Generally complementary learning has the following characteristics:

- Features extraction of positive and negative examples.
- Separation of positive and negative information.
- Development of lateral inhibition.

According to this concepts, the different approaches that are present in the literature can be divided into three groups:

- 1. Positive/negative systems, where the system builds information on the basis only of the target class.
- 2. Neutral learning systems, where the notion of positive and negative does not exist.
- 3. Complementary learning systems, where the system creates knowledge on the basis of positive and negative classes considering the relation between positive and negative samples.

CLFNN is considered a 9-tuple (*X*, *Y*, *D*, *A*, *R*, *B*, *l*, *s*, *p*). The definition of each element is as follows:

- *X* is the observation, belonging to the input space *U*, which has a set of variables $x=(x_1, x_2, ..., x_l)$.

- Y is the output associated to the input X. If M is the number of possible outputs, $Y \in V$ ($V = V_1 \times V_2 \times ... \times V_M$) where V is the output space. Moreover Y is the union of Y^+ and Y^- which represent respectively the positive and negative output.
- *D* is a set of observation *X* and its relative output *Y* and represents the source knowledge data.
- *A* is a set of knowledge which explains the behaviour amongst samples *X* in *D*.
- *R* represents the rules, i.e. the information learnt from *D* connecting the inputs to the relative outputs.
- *B* is a set of knowledge which explains the behaviour amongst samples *Y* in *D*.
- *l* represents the learning function building the CLFNN based on *D*. This function includes two complementary learning function: *s* and *p*.
- *s* is the structure learning which builds the net. For instance a clustering algorithm can independently obtain the structures from data. Moreover it creates the fuzzy sets and memory elements that describe the underlying data. If the existing knowledge is not sufficient for data description, other nodes are added. The process ends when the structure of data is suitable to reflect the data *D*.
- *p* is the parameter learning which can be any supervised learning algorithm.

In CLFNN the behaviour is characterized by a linguistic term derived by fuzzy set *A*. *A* is a mapping of *X* to the linguistic model of CLFNN; each set *A* identifies a set of locations in the input space characterizing by their membership functions. CLFNN is less subjected to data imbalance problem because it builds its knowledge from positive and negative classes separately and the influence of each class on the other one is minimized. Moreover in CLFNN the inference system uses the lateral inhibition which improves the system performance treating with imbalanced dataset.

In order to demonstrate the efficiency of CLFNN method, four imbalanced datasets are used: Single Photon Emission Computed Tomography (SPECT) (Blake & Merz, 1998), Wisconsin Breast Cancer diagnosis (WBCD) (Blake & Merz, 1998), Fine Needle Aspiration (FNA) (Cross & Harrison, 2006) and Thermogram (THERM).(Ng &Fok, 2003).

The dataset is divided in training set, testing set and validation set maintaining the class distribution. The averaged performance of CLFNN, which is calculated by the F-Measure over three cross-validation sets, is compared with other popular methods: Multilayer Perceptron (MLP)(Rosenblattx, 1958)., Radial Basis Function (RBF) (Powell, 1985), Linear Discriminant Analysis (LDA) (McLachlan, 2004), Decision tree C4.5 (Brodley & Utgoff,P.E.) and Support Vector Machine (SVM).

Table 6 illustrates the description of datasets and the averaged results obtained over the different cross validations with the several approaches. The acronym IR indicates the imbalanced ratio, i.e. the ratio between the number of positive samples and the number of negative samples.

Table 6 shows that CLFNN provides better results than the other approaches. In thermogram dataset none of the system can give satisfactory results because of its very high imbalance ratio but, also in this case, CLFNN outperforms the other approaches. This work confirms that CLFNN provides more consistent results over different data distributions coming a promising tool for handling imbalanced dataset.

	SPECT	WBCD	FNA	THERM
# Positive Samples	212	212	235	5
# Negative Samples	55	357	457	65
IR	3.85	0.59	0.51	0.07
MLP	0.762	0.925	0.917	0.2
RBF	0.8	0.912	0.912	0.2
C4.5	0.763	0.907	0.906	0.2
SVM	0.79	0.93	0.93	0.2
LDA	0.90	0.931	0.918	0.2
CLFNN	0.91	0.96	0.96	0.37

Table 6. Datasets description and averaged results.

4. Conclusions

Fuzzy Systems offer several advantages, among which the possibility to formalise and simulate the expertise of an operator in process control and tuning. Moreover the fuzzy approach provides a simple answer for processes which are not easily modelled. Finally they are flexible and nowadays they can be easily implemented and exploited also for real-time applications. This is the main reason why, in many industrial applications, dealing with processes that are difficult to model, fuzzy theory is widely adopted obtaining satisfactory results.

In particular, in this chapter, applications of FIS to industrial data processing have been presented and discussed, with a particular emphasis on the detection of rare patterns or events. Rare patterns are typically much difficult to identify with respect to common objects and often data mining algorithms have difficulty dealing with them. There are two kind of "rarity": rare case and rare classes. Rare cases, commonly known as outliers, refer to anomalous samples, i.e. observations that deviate significantly from the rest of data. Outliers may be due to sensor noise, process disturbances, human errors and instruments degradation. On the other hand, rare classes or more generally class imbalance, occur when, in a classification problem, there are more samples of some classes than others.

This chapter provides a preliminary review of classical outlier detection methods and then illustrates novel interesting detection methods based on Fuzzy Inference System. Moreover the class imbalance problem is described and traditional techniques are treated. Finally some actual Fuzzy based approaches are considered.

Results demonstrate how, in real world applications, fuzzy theory can effectively provide an optimal tool outperforming other traditional techniques.

5. References

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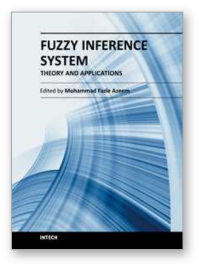
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