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A Hybrid Evolutionary Preprocessing Method for Imbalanced Datasets

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Abstract

Imbalanced datasets are commonly encountered in real-world classification problems. As many machine learning algorithms are originally designed for well-balanced datasets, re-sampling has become an important step to pre-process imbalanced data. It aims at balancing the datasets by increasing the samples of the smaller class or decreasing the samples of the larger class, which are known as over-sampling and under-sampling respectively. In this paper, a sampling strategy based on both over-sampling and under-sampling is proposed, in which the new samples of the smaller class are created based on fuzzy logic. Improvement of the datasets are done by the evolutionary computational method of Cross-generational elitist selection, Heterogeneous recombination and Cataclysmic mutation (CHC)

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that under-samples both the minority and majority samples. As a result, a hybrid preprocessing method is proposed to re-sample imbalanced datasets. The evaluation is done by applying the Support Vector Machine (SVM), C4.5 decision tree and nearest neighbor rule to train a classification model from the re-sampled training sets. From the experimental results, it can be seen that our proposed method improves both the F-measure and AUC. The over-sampling rate and complexity of the classification model are also compared. Our proposed method is found to be superior to all other methods under comparison, and is more robust in different classifiers.

1. Introduction

The classification of imbalanced datasets is a popular topic in recent years [22] and [27]. Most of the machine learning tools, such as neural networks and support vector machines, are originally designed for well-balanced datasets. If the dataset is imbalanced, the performance of the classifier can be poor. The reason for this is apparent. For example, considering a dataset with 99% of data from class A and only 1% of data from class B, the accuracy is 99% if the classifier ignores the data from class B and labels the whole dataset as class A. It is already very hard to achieve an accuracy above 99% by using most of the learning algorithms. However, the minority class of datasets is usually more important and meaningful. For example, there are much less samples of people with a particular disease than

those of healthy people in a medical problem. If a classifier is needed to label whether some people are infected or not, it is obvious that the minority class (people with a particular disease) is the class of interest.

Problems with imbalanced datasets can be easily found in the real world, such as intrusion detection [9], speech recognition [26], identification of power distribution fault causes [41], and bioinformatics problems [16]. There are two main approaches to solve the problems caused by imbalanced datasets. One is the data level approach and the other is the algorithm level approach. The data level approaches [3], [8], [18], and [28] include balancing the class distribution by over-sampling the minority class or under-sampling the majority class. The algorithm level approaches improve the existing machine learning methods by adjusting the probabilistic estimate [38], modifying the cost per class [32], adding some penalty constants [25], or learning from one class instead of two classes [35] and [30].

Many experiments [12], [15], and [42] show that re-sampling is a good data level approach to handle imbalanced data. Moreover, it is more flexible as it does not depend on the chosen classifier. Therefore, we focus on re-sampling in this paper. There are three main types of strategies for re-sampling data. The first one is over-sampling, which can be done randomly or by the method of Synthetic Minority Over-sampling Technique (SMOTE) [8]. The second one is under-sampling, which includes Tomek links [37] and Neighborhood Cleaning Rule (NCL) [24]. The last one is the hybrid method, which combines the two previous methods

(over-sampling and under-sampling methods).

The importance of designing sampling strategies has been discussed in [31], which may affect the successful learning of different classes. Hybrid re-sampling methods, reportedly, have advantage on treating datasets with a high imbalanced ratio [3] and [6]. Although some hybrid methods [3], [34], and [40] have been proposed to reduce the over-generalization problem from over-sampling methods, most of these methods are based on SMOTE and the results may be limited by the synthetic samples of SMOTE. Therefore, a hybrid re-sampling method is proposed in this paper. Fuzzy logic, which is a useful tool to treat imbalanced datasets [12], is used to over-sample the minority class samples instead of SMOTE. A fuzzy rule base is formed based on the samples of the minority class. Then, a rule is selected randomly with reference to the effectiveness of each rule. The selected rule is used as the criteria to generate a new sample of the minority class. The above steps will repeat until the sizes of the majority class and minority class are the same.

However, the large over-sampled training dataset will increase the complexity of the classification model and decrease the efficiency of the learning algorithm. It will also cause over-generalization easily, especially for some noisy dataset. This is because the decision boundary could become narrow or the overlapping area between the majority class and minority class could become large after the over-sampling. Therefore, an evolutionary algorithm (EA) is applied to both the

synthetic samples and majority samples to under-sample the dataset. The chosen EA is the CHC (Cross-generational elitist selection, Heterogeneous recombination and Cataclysmic mutation) algorithm [11] since it shows the ability of selecting the most representative instances among many algorithms studied in [5].

Experiments are carried out to compare our proposed method with three SMOTE-extended over-sampling methods, four hybrid re-sampling methods and one under-sampling method. They are SMOTE, Safe-Level-SMOTE [4], Adaptive Synthetic Sampling [21], SMOTE+Tomek Links [3], SMOTE+Rough Set [34], SMOTE+CHC (sCHC) [40], agglomerative hierarchical clustering [10], and EUSCHC [14]. 44 imbalanced datasets from UCI Repository [2] are used in the experiments. The Support Vector Machine (SVM) [7], C4.5 decision tree [33], and nearest neighbor rule (1NN) are used as the tools for reaching a classification model for each re-sampled dataset so as to evaluate each re-sampling method. The evaluation measures are based on F-measure and area under the receiver operating characteristic curve (AUC). Although there exists many hybrid pre-processing methods, only some of them are like our method that consider and focus on the data size. In this paper, CHC is used to reduce the data size and achieve a good performance. Additionally, the proposed method enhances the performance in the over-sampling stage by taking advantage of the fuzzy rule base.

This paper is organized as follows: In Section 2, some preprocessing methods and CHC are reviewed. Section 3 presents the details of the proposed re-sampling

strategy and the evaluation method. To show the effectiveness of our proposed approach, the comparisons with other methods and the results are discussed in Section 4. A conclusion is drawn in Section 5.

2. Previous Work

This section describes some previous works about re-sampling methods, which will be used to compare with our proposed method in the experiments later. The ideas about CHC will also be discussed.

2.1. Re-sampling Methods

As discussed in the previous section, there are three main strategies for re-sampling data.

2.1.1. Over-sampling Methods

Some instances are produced for the minority class to balance the class distribution. The simplest one is a non-heuristic method (random over-sampling) that replicates samples of the original minority class to generate the new instances. This method causes over-fitting easily since the new instances copy exactly from the original minority class. Synthetic Minority Over-sampling Technique (SMOTE) [8] is a well-known method which creates the new instances by interpolating several minority samples that join together. This method makes use

of each minority class sample and inserts synthetic samples along the line segments joining any/all of the k minority class nearest neighbors to over-sample the minority class. An example is shown in Fig. 1. Five nearest neighbors are used in it, where x_i is a selected sample of minority class, x_{i1} to x_{i5} are the 5 nearest neighbors of x_i and s_1 to s_5 are the synthetic samples created by interpolation. If the degree of over-sampling required is 300%, three synthetic examples are selected randomly from s_1 to s_5 .

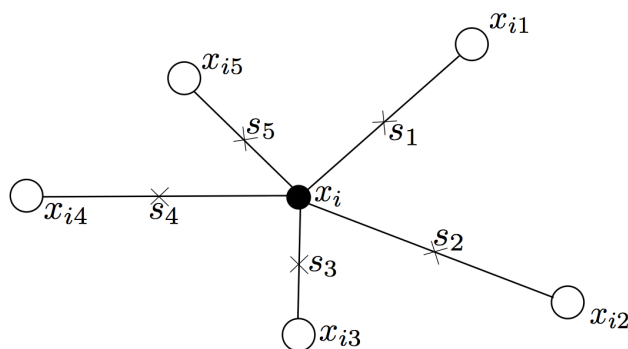


Figure 1: Example of SMOTE with 5 nearest neighbors.

Since the synthetic samples provide a less specific and larger decision region, the over-fitting problem can be reduced. However, this method may introduce more minority synthetic samples in the area of majority class where the minority class is very sparse with respect to the majority class. This causes the problem of over-generalization, which means the decision boundary is very narrow or there is a large overlapping area between the majority class and minority class. Therefore, some methods are developed based on SMOTE to overcome this limitation, such

as Borderline-SMOTE (sBorder) [19], Adaptive Synthetic Sampling (ADASYN) [21], Safe-Level-SMOTE (sSafe) [4], and SPIDERS [29].

2.1.2. Under-sampling Methods

Some instances of majority class are eliminated in order to balance the class distribution. The simplest method is random under-sampling (RUS), which aims to balance the datasets by randomly removing samples of the majority class. However, this method may easily remove some useful data. The other representative methods include (i) condensed nearest neighbor rule (CNN) [20], which eliminates the majority class samples that are distant from the decision border, (ii) Tomek links (TL) [37], which edits out noisy and borderline majority class samples, (iii) one-sided selection (OSS) [23], which is an integrated method of TL and CNN, and (iv) neighborhood cleaning rule (NCL) [24], which is based on the Wilson's Edited Nearest Neighbor Rule (ENN) [39] to remove the majority class samples that lead to misclassification.

2.1.3. Hybrid Methods

Although both over-sampling and under-sampling can balance the class distribution, different drawbacks like over-generalization and removal of useful data are also introduced. Therefore, some hybrid methods are developed to combine SMOTE and under-sampling as a data cleaning method to reduce the problem. Example hybrid methods include SMOTE+Tomek links (sTL), which uses TL

to remove samples of both classes to increase the area of decision border, and SMOTE+ENN (sENN) [3], which uses ENN to remove the samples that are misclassified by their nearest neighbors. Rough set theory (sRST) [34] and evolutionary algorithm (sCHC) [40] have also been applied on SMOTE to select the samples to increase the accuracy of classification.

Most of the above hybrid methods make use of SMOTE to perform over-sampling. Clustering techniques are also developed to perform under-sampling and over-sampling, such as agglomerative hierarchical clustering (AHC) [10].

2.2. CHC [11]

CHC is a kind of EAs that combines a selection strategy with a highly disruptive recombination operator. To avoid premature convergence and maintain diversity, incest prevention and cataclysmic mutation are introduced. The process of CHC can be described as follows. Firstly, a population set of chromosomes P is created. Each chromosome $p_i = (p_{i1}, p_{i2}, \dots, p_{in})$ is an n -dimensional vector, which is a set of genes, where p_{ij} is the j th gene value ($j = 1, 2, \dots, n$) of the i th chromosome in the population ($i = 1, 2, \dots, m$); m is the population size and n is the number of genes.

Secondly, the chromosomes are evaluated by a defined fitness function. The form of fitness function depends on the application. Thirdly, an intermediate population set of chromosomes C , which is of the same size as P , is generated by

copying all members of P in a random order.

Then, a uniform crossover (HUX) operator is applied on C to form C' . HUX exchanges half of the genes randomly between the chromosomes one by one to form C' . CHC also uses an additional method for incest prevention. Before applying HUX to the chromosomes, the Hamming distance between them is calculated. If half of that distance is larger than a difference threshold d , HUX is applied; otherwise these two chromosomes are deleted from C . Therefore, the size of C' may be smaller than that of P or C . The initial threshold d is set at $n/4$. After C' has formed, it is evaluated by the fitness function and an elitist selection is taken. Only the best chromosomes from both P and C' are selected to form the offspring population in the next generation. If the offspring population is the same as P , the difference threshold d is decreased by one.

CHC is different from the traditional genetic algorithm. Mutation is not performed at the recombination stage. CHC performs partial reinitialization (divergence) when the search becomes trapped (i.e., the difference threshold d becomes zero and no new offspring population is formed for several generations). The population is reinitialized, based on the best chromosome, by changing the elements' values randomly with a user-defined divergence rate D_{rate} . For example, if D_{rate} equals to 0.35, the values of 35% elements will be changed randomly. The search is then resumed with a new difference threshold $d = D_{rate} * (1 - D_{rate}) * n$. This process is called cataclysmic mutation.

CHC has shown the ability of selecting the most representative instances among the other algorithms studied in [5]. Therefore, it is chosen as the algorithm to improve the outcome of over-sampling in this paper.

3. Methodology

In this section, the proposed hybrid preprocessing method and the evaluation methods used in this paper are discussed. The proposed method involves two stages. The minority samples of the training sets are firstly over-sampled based on fuzzy logic to form a fuzzy rule base (FRB). To improve the performance, CHC is then implemented to reduce both the synthetic samples and majority samples.

3.1. Fuzzy Rule Base (FRB)

In this paper, let the *positive* class be the minority class and only λ training samples (X_α) of positive class are considered, where $X_\alpha = (x_{\alpha 1}, \dots, x_{\alpha \gamma})$ is an γ -dimensional vector, $\alpha = 1, 2, \dots, \lambda$ and $x_{\alpha \beta}$ is the β th attribute value ($\beta = 1, 2, \dots, \gamma$) of the α th training sample. The θ th fuzzy if-then rule is written as follows:

$$\begin{aligned} \text{Rule } \theta : \text{ IF } z_1 \text{ is } A_1^\theta \text{ AND } \dots \text{ AND } z_\gamma \text{ is } A_\gamma^\theta \\ \text{ THEN class} = \text{positive with } w_\theta \end{aligned} \quad (1)$$

where A_β^θ is a fuzzy term of the θ th rule corresponding to the attribute z_β , $\beta = (1, 2, \dots, \gamma)$ and $z = (z_1, z_2, \dots, z_\gamma)$ is a γ -dimensional attribute vector, and w_θ

is the rule weight. The regular triangular membership functions are used for the fuzzy terms. In this paper, the fuzzy terms A_β^θ are derived based on the samples of positive class. The minimum and maximum values of each attribute are first found. The fuzzy terms are the triangular membership functions within the range of each attribute. The fuzzy terms also depend on the number of labels. Since regular triangular membership functions are used, the fuzzy terms are distributed evenly within the range of each attribute.

The fuzzy rules are generated based on the samples of positive class. For each sample, the label with the highest membership value is selected to form the corresponding rule for each attribute. The maximum number of rules depends on the number of labels and attributes.

The rule weight w_θ is used to reflect the degree of matching of each fuzzy rule over all the positive samples, so that the importance of each rule can be evaluated. First, the fuzzy value of each sample is calculated. The fuzzy value of X_α for the θ th fuzzy rule is defined as follows:

$$\mu_{A^\theta}(X_\alpha) = T(\mu_{A_1^\theta}(x_{\alpha 1}), \dots, \mu_{A_\gamma^\theta}(x_{\alpha \gamma})), \quad (2)$$

where the product T-norm is used. The rule weight (w_θ) is calculated by adding all the fuzzy values of samples.

$$w_\theta = \sum_{\alpha=1}^{\lambda} (\mu_{A^\theta}(X_\alpha)). \quad (3)$$

After the rule base of the positive class is generated, the rules are randomly drawn based on the rule weight. The rule with a higher rule weight will have a higher probability to be chosen. Then, a new sample is generated within the area of the selected rule. These processes are repeated until the number of positive samples is the same as that of the negative samples.

To illustrate the idea more clearly, Fig. 2 shows the distribution of two classes with two attributes as an example of the formulation of fuzzy rules. The x-axis and y-axis govern the values of the two different attributes and regular triangular membership functions with five labels are used. The circle dots correspond to the negative class and the square dots correspond to the positive class. The dashed lines show the minimum or maximum value of the corresponding attribute of the positive samples. As only the attribute vectors of the positive class are considered to generate fuzzy rules, totally ten rules can be formed in this example:

- Rule 1: IF z_1 is $A_1^1 = L1_1$ AND z_2 is $A_2^1 = L2_4$. THEN class = positive with 0.897
- Rule 2: IF z_1 is $A_1^2 = L1_2$ AND z_2 is $A_2^2 = L2_3$. THEN class = positive with 1.147
- Rule 3: IF z_1 is $A_1^3 = L1_2$ AND z_2 is $A_2^3 = L2_4$. THEN class = positive with 1.508
- Rule 4: IF z_1 is $A_1^4 = L1_3$ AND z_2 is $A_2^4 = L2_3$. THEN class = positive with 1.230
- Rule 5: IF z_1 is $A_1^5 = L1_3$ AND z_2 is $A_2^5 = L2_4$. THEN class = positive with 2.344
- Rule 6: IF z_1 is $A_1^6 = L1_3$ AND z_2 is $A_2^6 = L2_5$. THEN class = positive with 1.607
- Rule 7: IF z_1 is $A_1^7 = L1_4$ AND z_2 is $A_2^7 = L2_1$. THEN class = positive with 0.727
- Rule 8: IF z_1 is $A_1^8 = L1_4$ AND z_2 is $A_2^8 = L2_4$. THEN class = positive with 1.319
- Rule 9: IF z_1 is $A_1^9 = L1_4$ AND z_2 is $A_2^9 = L2_5$. THEN class = positive with 1.731
- Rule 10: IF z_1 is $A_1^{10} = L1_5$ AND z_2 is $A_2^{10} = L2_4$. THEN class = positive with 1.399

where z_1 and z_2 represent Attribute 1 and Attribute 2 for the x-axis and y-axis respectively in Fig. 2, $L1_i$ is the i -th label of z_1 attribute, $L2_i$ is the i -th label of z_2 attribute. Rule 5 has the highest rule weight and rule 7 has the lowest rule weight in this example.

For generating the synthetic samples, a rule out of these ten rules is chosen with the probability of selection depending on the rule weight. Then, this rule sets the criteria of the highest and lowest value of each attribute. The new sample is generated randomly within these criteria. This process is repeated until the num-

ber of the positive class is the same as that of the negative class. Fig. 3 shows the samples distribution after over-sampling. The triangle dots represent the synthetic samples. It is found that the spread of the synthetic samples is similar to that of the original positive samples (shown as the square dots). The synthetic samples in Fig. 3 are dense in the area of rule 5.

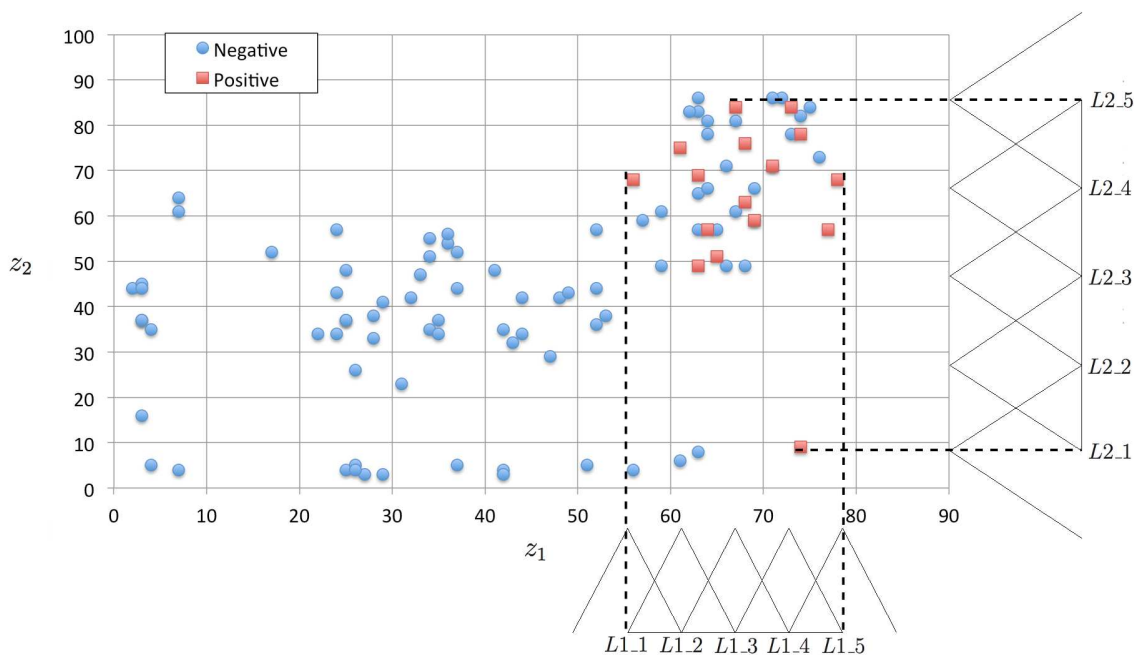


Figure 2: Example of the distribution of imbalanced dataset. The y-axis represents the values of z_2 and x-axis represents the value of z_1 .

3.2. Setting of CHC

After the over-sampling, the number of minority samples is the same as that of majority samples and CHC is then applied. There are two important issues that need to be addressed before the algorithm is employed: the representation of each

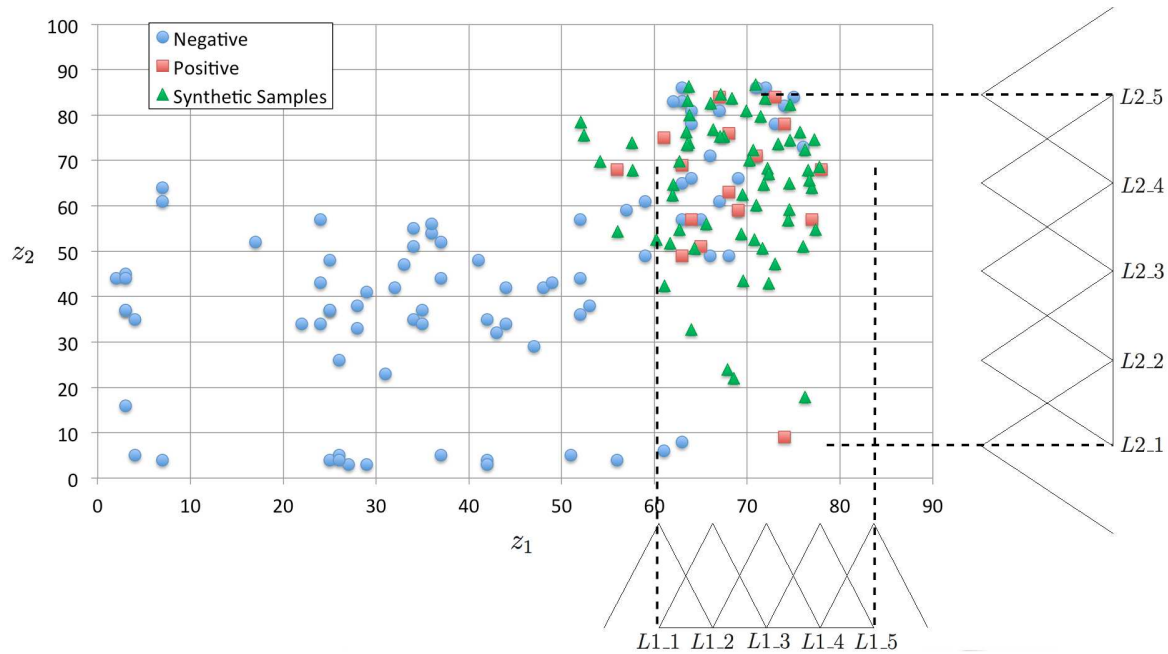


Figure 3: Distribution of the samples after over-sampling. The y-axis represents the values of z_2 and x-axis represents the value of z_1 .

chromosome and the definition of fitness function. Fig. 4 shows the block diagram of the process of FRB+CHC.

3.2.1. Chromosome Representation

CHC is used to reduce the synthetic samples as well as the majority class samples. Therefore, the chromosomes are to represent subsets of these samples. It can be carried out by a binary representation. Each chromosome is an n -dimensional vector. In this section, n is the number of synthetic samples plus majority class samples. Each vector element shows whether the corresponding sample exists in the subset of the training set or not. Therefore, there are two possible values for

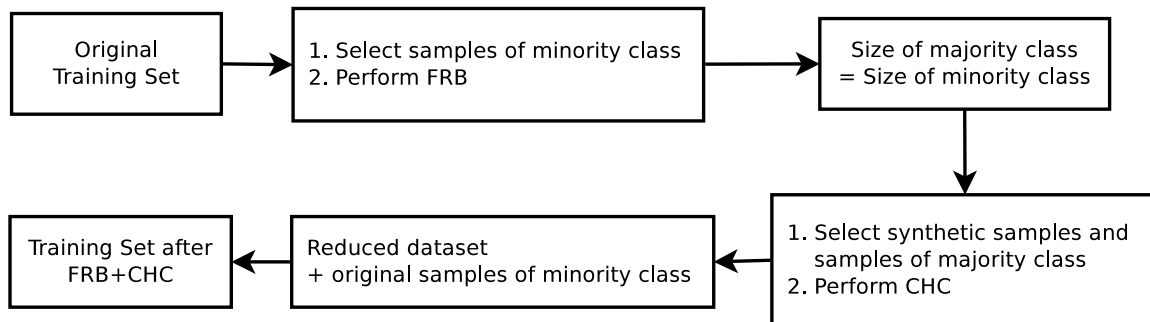


Figure 4: Block Diagram of FRB+CHC.

each element: 0 and 1. If the value is 1, the corresponding sample is included in the subset of the training set. If the value is 0, the sample does not exist in the subset.

3.2.2. Fitness function

In this study, the k-NN classifier is used as the evaluation method of CHC to obtain the subset with the highest classification rate. Normally, accuracy (ratio of correctly classified samples to total number of samples) would be used as the measure of classification rate. However, it may cause difficulty for the imbalanced datasets when doing testing later since the correct classification rate of the majority samples may affect the accuracy more significantly than that of the minority samples. Therefore, some other measures are used in this paper. These measures are commonly employed to analyze problems with imbalanced datasets.

Firstly, precision and recall are introduced [17]. Their definitions are given as

follows:

$$Precision = \frac{TP}{TP + FP} \quad (4)$$

$$Recall = \frac{TP}{TP + FN} \quad (5)$$

where TP is the number of true positives, FP is the number of false positives and FN is the number of false negatives. A high value of precision indicates that the predicted positive samples are most likely relevant. A high value of recall indicates that most of the positive samples can be predicted correctly.

A popular evaluation metric for imbalanced problems is $F - measure$ [17], which is a function of precision and recall. In principle, $F - measure$ represents a harmonic mean between precision and recall. A high value of $F - measure$ means both the precision and recall values are high and do not differ very much. It is an important measure for imbalanced datasets since a high value of it can imply that the method classifies the positive samples correctly at a high rate with little misclassified negative samples. It is defined as follows:

$$F - measure = \frac{2 * Precision * Recall}{Precision + Recall} \quad (6)$$

The area under the receiver operating characteristic curve (AUC) is also commonly used to measure the performance of classification. The AUC measure [13] is the probability of correctly identifying a random sample, and it can be defined

as follows:

$$AUC = \frac{1 + Recall - FP_{rate}}{2} \quad (7)$$

where *Recall* is defined in (5) and $FP_{rate} = \frac{FP}{FP+TN}$, *TN* is the number of true negatives. FP_{rate} defines the percentage of true negatives cases misclassified as positives. A high value of *AUC* implies small values of *FN* and *FP*, meaning that the corresponding classifier is effective.

Since both *F - measure* and *AUC* are important measures on imbalanced datasets, a multi-objective fitness function is used here. The chromosome with both higher values of *F - measure* and *AUC* obviously has a higher rank. If a chromosome *X* has a higher value of *F - measure* ($F_X > F_Y$) and a lower value of *AUC* ($A_X < A_Y$) than that of chromosome *Y*, the difference between the chromosomes' *F - measure* ($|F_X - F_Y|$) and the difference between the chromosomes' *AUC* ($|A_X - A_Y|$) will be compared. If $|F_X - F_Y| > |A_X - A_Y|$, chromosome *X* will be regarded as a better one; otherwise chromosome *Y* will be regarded as a better one. The above setting is also applied in sCHC for the comparison in Section 4.

3.3. Evaluation

3.3.1. *F - measure and AUC measures*

To show the performance of our proposed method, *F - measure* in (6) and *AUC* in (7) are used. The main drawback of over-sampling or hybrid sampling

methods is that the number of training samples are increased greatly. This may cause the increase of complexity of the learning model. Therefore, the over-sampling rates of different methods are also compared. Define:

$$Rate_{over} = \frac{(N_{sampled} - N_{original})}{N_{original}} * 100\% \quad (8)$$

where $N_{sampled}$ is the number of samples in the re-sampled training set and $N_{original}$ is the number of samples in the original training set. The over-sampling rate in (8) shows the increase rate of the number of the training samples. When a support vector machine is used to form the classification mode, the increase rate of the support vectors can be used to evaluate the complexity of the learning model. This rate is calculated based on the support vectors generated.

$$Rate_{SV} = \frac{(SV_{sampled} - SV_{original})}{SV_{original}} \quad (9)$$

where $SV_{sampled}$ is the number of support vectors trained by the re-sampled training set and $SV_{original}$ is the number of support vectors trained by the original training set. It should be noted that the CHC fitness evaluation for data size reduction (by k-NN) and the training of the classification model based on the resampled data (by SVM) are two separated processes. K-NN is used in the fitness evaluation because it is simple with minimal computation effort. SVM is a commonly used method to obtain the classification model.

4. Experimental Study

In this section, we present the experiments that are carried out to compare our proposed method with other hybrid sampling methods and the CHC under-sampling method. The datasets used can be found in UCI Repository [2].

The experiments involve different kinds of hybrid methods, including SMOTE, ADASYN, sTL, sSafe, sRST, sCHC, AHC and our proposed method, which is named as Fuzzy Rule Base+CHC (FRB+CHC). CHC, which is used as an under-sampling method in [14] (EUSCHC), is also compared in the experiment. To measure the performance of the preprocessing methods, the same learning tool should be applied among all the experiments. In this study, three different tools are used. They are Support Vector Machine (SVM), 1 Nearest Neighbor (1NN), and C4.5 decision tree. The programs of all testing methods and the learning tools are based on KEEL, which is an open source software available in the Web [1]. $F - measure$ and AUC are used as measures to analyze the results. The average values of these measures for each method will be calculated. As the expansion of re-sampled training datasets may increase the computational time and complexity of the classification model, the over-sampling rate and the number of support vectors formed from SVM will also be compared.

4.1. Datasets

To study the methods on different datasets, 44 datasets with different imbalance ratio (IR) are chosen. IR is the ratio of the number of majority class to the number of minority class. Table 1 shows the details of the selected datasets, where the number of samples ($N_{samp.}$), the number of attributes ($N_{attr.}$), the distribution of the minority and majority classes, and IR for each dataset can be found.

Table 1: Details of the Selected Imbalanced Datasets.

Dataset	$N_{samp.}$	$N_{attr.}$	Min., Maj.(%)	IR
ecoli034vs5	200	7	(10, 90)	9
yeast2vs4	514	8	(9.92, 90.08)	9.08
ecoli067vs35	222	7	(9.91, 90.09)	9.09
ecoli0234vs5	202	7	(9.9, 90.1)	9.1
glass015vs2	172	9	(9.88, 90.12)	9.12
yeast0359vs78	506	8	(9.88, 90.12)	9.12
yeast0256vs3789	1004	8	(9.86, 90.14)	9.14
yeast02579vs368	1004	8	(9.86, 90.14)	9.14
ecoli046vs5	203	6	(9.85, 90.15)	9.15
ecoli01vs235	244	7	(9.83, 90.17)	9.17
ecoli0267vs35	224	7	(9.82, 90.18)	9.18
glass04vs5	92	9	(9.78, 90.22)	9.22
ecoli0346vs5	205	7	(9.76, 90.24)	9.25
ecoli0347vs56	257	7	(9.73, 90.27)	9.28
yeast05679vs4	528	8	(9.66, 90.34)	9.35
vowel0	988	13	(9.01, 90.99)	9.98
ecoli067vs5	220	6	(9.09, 90.91)	10

Dataset	$N_{samp.}$	$N_{attr.}$	Min., Maj.(%)	IR
glass016vs2	192	9	(8.85, 91.15)	10.29
ecoli0147vs2356	336	7	(8.63, 91.37)	10.59
led7digit02456789vs1	443	7	(8.35, 91.65)	10.97
ecoli01vs5	240	6	(8.33, 91.67)	11
glass06vs5	108	9	(8.33, 91.67)	11
glass0146vs2	205	9	(8.29, 91.71)	11.06
glass2	214	9	(7.94, 92.06)	11.59
ecoli0147vs56	332	6	(7.53, 92.47)	12.28
cleveland0vs4	177	13	(7.34, 92.66)	12.62
ecoli0146vs5	280	6	(7.14, 92.86)	13
shuttlec0vsc4	1829	9	(6.72, 93.28)	13.87
yeast1vs7	459	7	(6.53, 93.47)	14.3
glass4	214	9	(6.07, 93.93)	15.47
ecoli4	336	7	(5.95, 94.05)	15.8
page_blocks13vs4	472	10	(5.93, 94.07)	15.86
abalone918	731	8	(5.65, 94.25)	16.4
glass016vs5	184	9	(4.89, 95.11)	19.44
shuttlec2vsc4	129	9	(4.65, 95.35)	20.5
yeast1458vs7	693	8	(4.33, 95.67)	22.1
glass5	214	9	(4.2, 95.8)	22.78
yeast2vs8	482	8	(4.15, 95.85)	23.1
yeast4	1484	8	(3.43, 96.57)	28.1
yeast1289vs7	947	8	(3.16, 96.84)	30.57
yeast5	1484	8	(2.96, 97.04)	32.73
ecoli0137vs26	281	7	(2.49, 97.51)	39.14
yeast6	1484	8	(2.36, 97.64)	41.4

Dataset	$N_{samp.}$	$N_{attr.}$	Min., Maj.(%)	IR
abalone19	4174	8	(0.77, 99.23)	129.44

4.2. Setup of Experiment

For over-sampling, the rules of the minority samples are associated with regular triangular membership functions with five fuzzy terms. For CHC, the values of the parameters are:

- Population size: 50.
- Divergence rate: 0.35.
- Threshold decreasing rate: 0.001.
- k of k-NN classifier used as evaluation: 1.
- Number of evaluations: 5,000.

In this paper, SVM, 1NN, and C4.5 are used to weigh the influence of each preprocessing method. For SVM, a radial basis function (RBF) is used as the kernel since a non-linear classification model is needed and RBF is a common kernel to handle this problem. The RBF is defined as follows:

$$RBF = \exp\left(-\frac{1}{\sigma}\|\mathbf{x}_i - \mathbf{x}\|^2\right) \quad (10)$$

where $\sigma > 0$ is the parameter to determine the width of the radial basis function.

It controls the flexibility of the classifier. When σ decreases, the flexibility of

the resulting classifier in fitting the training data increases, and this might lead to over-fitting easily. The value of σ is set as 0.01. The tradeoff between training error and margin of SVM is set as 100. The above values are chosen through experiments. For C4.5, the confidence level is set as 0.25, the minimum number of item-sets per leaf is set to 2 and pruning is used as well to obtain the final tree. For 1NN, the Euclidean distance metric is used.

A 5-fold cross validation model is used to compare the classification results from different preprocessing methods. Each dataset are first divided into five parts randomly. Four of them are combined to form a training set and the remaining subset forms a testing set. The process is then repeated five times, so that each subset is used once as a testing set. All the methods involve some random parameters, so five experiments are carried out for each 5-fold cross validation model and the average value are calculated as the results, i.e. totally 25 experiments were done.

4.3. Results

4.3.1. *F* – measure and AUC measures

Tables 2 and 3 show the SVM results on *F* – measure and AUC for each re-sampling method on the 44 datasets respectively. The results of the original datasets are shown in the second column and the best value for each dataset are highlighted in bold. The last row shows the average value of each sampling method for the datasets. The performance of the FRB over-sampling method

are also included (in the rightmost column) for comparison with FRB+CHC. It can be seen that the average values of $F - measure$ and AUC in both FRB and FRB+CHC are higher than other methods. The performance of sCHC and FRB+CHC are similar. This shows that CHC has good performance as a data cleaning method after over-sampling, especially for the results in $F - measure$. The AUC values of SMOTE, sTL, sSafe, sRST and sCHC are similar since they all use SMOTE to perform over-sampling. ADASYN gets the lowest average values of $F - measure$, which means the precision is low and the difference between precision and recall is large.

In this experiment, the performance of FRB and FRB+CHC is very similar, which shows the advantages of FRB over the other hybrid or over-sampling methods. However, the data size will be very large if only FRB is used as the pre-processing method. FRB+CHC can reduce the data size without a large effect to the performance. Therefore, only FRB+CHC will be considered in the following section.

Table 4 shows the average rankings by means of $F - measure$ and AUC using Friedman's method [36]. The highest value of each dataset is ranked as 1. If a certain method obtains the ranking 3, 6, 2, and 1 on four datasets, the average ranking is $(3 + 6 + 2 + 1)/4 = 3$. Therefore, a lower average ranking indicates that the corresponding method is better among the other methods. FRB+CHC obtains the best ranking by AUC and sCHC obtains the best ranking by $F -$

measure. Note that the highest average values of *AUC* or *F – measure* do not imply the best ranking results since the ranking shows the comparison results among all the methods of each dataset. For example, EUSCHC has the lowest *AUC* average values but its ranking is better than ADASYN. Since EUSCHC is an under-sampling method, it easily ignores some useful samples of majority class.

Table 2: SVM: Average F-measure of Testing Datasets among Different Sampling Methods.

Dataset	Original	SMOTE	ADASYN	sTL	sSafe	sRST	sCHC	EUSCHC	AHC	FRB+CHC	FRB
ecoli034vs5	0	0.5629	0.2667	0.5901	0.5578	0.5007	0.5054	0.6591	0.3111	0.5829	0.6337
yeast2vs4	0.6384	0.6824	0.5446	0.6683	0.6824	0.6787	0.6996	0.7418	0.6937	0.7015	0.6971
ecoli067vs35	0.0000	0.4540	0.3975	0.5122	0.4609	0.4447	0.5108	0.4758	0.3692	0.4308	0.6171
ecoli0234vs5	0.0000	0.5176	0.2917	0.5240	0.5012	0.4734	0.5577	0.6682	0.2667	0.6142	0.6462
glass015vs2	0.0000	0.3094	0.3181	0.3103	0.3301	0.3419	0.2137	0.1015	0.2850	0.2049	0.2275
yeast0359vs78	0.3481	0.3541	0.2965	0.3379	0.3580	0.3529	0.4117	0.3481	0.3666	0.3470	0.3481
yeast0256vs3789	0.1782	0.5282	0.4391	0.5206	0.5286	0.5325	0.5624	0.6033	0.5263	0.5899	0.2589
yeast02579vs368	0.8152	0.7199	0.5213	0.7179	0.7189	0.7201	0.7437	0.7487	0.7264	0.7747	0.85
ecoli046vs5	0.0000	0.3901	0.2000	0.3958	0.4084	0.4214	0.3827	0.6786	0.0667	0.5225	0.6584
ecoli01vs235	0.0000	0.4325	0.1648	0.4396	0.4352	0.4264	0.4844	0.5691	0.1385	0.4224	0.5536
ecoli0267vs35	0.0000	0.3158	0.2269	0.3257	0.2902	0.3253	0.3856	0.4035	0.1469	0.4592	0.5337
glass04vs5	1.0000	0.8793	0.8679	0.8747	0.9228	0.9209	0.9933	0.7854	1.0000	0.9631	0.9655
ecoli0346vs5	0.0000	0.5446	0.3636	0.6397	0.5741	0.5642	0.5985	0.7382	0.3404	0.6766	0.6768
ecoli0347vs56	0.0000	0.5743	0.4743	0.5628	0.5576	0.5104	0.5913	0.6669	0.1846	0.5176	0.5913
yeast05679vs4	0.0000	0.4327	0.4265	0.4282	0.4333	0.4250	0.5066	0.4996	0.4189	0.4786	0.5355
vowel0	1.0000	0.9936	0.9796	0.9905	0.9890	0.9816	0.9833	0.9396	1.0000	0.9060	0.9387
ecoli067vs5	0.0000	0.3260	0.2973	0.3463	0.3444	0.3225	0.3787	0.6848	0.2308	0.6173	0.6873
glass016vs2	0.0000	0.3196	0.3203	0.2686	0.3048	0.2963	0.2102	0.1395	0.3404	0.2001	0.2857
ecoli0147vs2356	0.0000	0.4230	0.3014	0.4960	0.4354	0.4435	0.5021	0.2230	0.0500	0.4043	0.5074
led7digit02456789vs1	0.7748	0.5707	0.6197	0.5226	0.5766	0.5156	0.7308	0.5691	0.5961	0.6746	0.7224
ecoli01vs5	0.0000	0.4138	0.2588	0.4482	0.4103	0.4946	0.4392	0.4140	0.2069	0.6843	0.7811
glass06vs5	1.0000	0.9057	0.9655	0.8953	0.8857	0.9083	0.9866	0.8654	1.0000	0.9783	0.9474
glass0146vs2	0.0000	0.2463	0.2512	0.2247	0.2473	0.2814	0.2823	0.1747	0.2931	0.2597	0.2768
glass2	0.0000	0.2477	0.2362	0.2329	0.2478	0.2988	0.2484	0.1131	0.3233	0.2019	0.26

Dataset	Original	SMOTE	ADASYN	sTL	sSafe	sRST	sCHC	EUSCHC	AHC	FRB+CHC	FRB
ecoli0147vs56	0.0000	0.5757	0.4148	0.6288	0.6022	0.5103	0.5164	0.6148	0.0571	0.6762	0.7609
cleveland0vs4	0.0000	0.1539	0.1556	0.1560	0.1263	0.1600	0.0923	0.2621	0.0000	0.1687	0.2030
ecoli0146vs5	0.0000	0.4280	0.1920	0.4112	0.4356	0.4422	0.3762	0.6993	0.3000	0.7456	0.7758
shuttlec0vsc4	0.9490	0.9740	0.8937	0.9749	0.9740	0.9817	0.9724	0.9707	0.9675	0.7964	0.8763
yeast1vs7	0.0000	0.2926	0.2870	0.2865	0.2939	0.2738	0.3120	0.0000	0.2861	0.3161	0.3381
glass4	0.8560	0.6633	0.6565	0.6590	0.6613	0.6463	0.8190	0.7164	0.8471	0.7273	0.7197
ecoli4	0.7500	0.6352	0.5082	0.6354	0.6389	0.6491	0.7931	0.7372	0.7109	0.7356	0.6617
page_blocks13vs4	0.2270	0.2033	0.1907	0.2010	0.2034	0.1894	0.3563	0.0832	0.2270	0.1816	0.1907
abalone918	0.0444	0.4522	0.4172	0.4206	0.4474	0.4570	0.5221	0.2643	0.5303	0.5732	0.5561
glass016vs5	0.6650	0.5674	0.6592	0.5601	0.5668	0.6551	0.7548	0.4688	0.7273	0.7694	0.6857
shuttlec2vsc4	0.4	0.7152	0.7152	0.7152	0.7152	0.7288	0.6103	0.1593	0.4	0.6126	0.7395
yeast1458vs7	0	0.1318	0.1261	0.1260	0.1323	0.1344	0.1585	0	0.1398	0.1557	0.1187
glass5	0.7	0.5937	0.4551	0.5495	0.5932	0.4838	0.6583	0.3542	0.7	0.7533	0.8
yeast2vs8	0.6967	0.5972	0.2079	0.5905	0.5989	0.5984	0.7068	0.6967	0.6570	0.6967	0.6967
yeast4	0	0.2703	0.2464	0.2648	0.2715	0.2711	0.3076	0.0308	0.2714	0.3533	0.3398
yeast1289vs7	0	0.1395	0.1363	0.1357	0.1397	0.1308	0.1851	0	0.1488	0.1967	0.1776
yeast5	0	0.4843	0.4611	0.4742	0.4818	0.4751	0.5146	0.5802	0.5012	0.4476	0.4415
ecoli0137vs26	0	0.3976	0.2400	0.4681	0.4292	0.3636	0.4306	0.3158	0.1	0.3465	0.3826
yeast6	0	0.2698	0.2014	0.2606	0.2705	0.2670	0.3577	0	0.2756	0.3288	0.2759
abalone19	0	0.0408	0.0406	0.0403	0.0409	0.0486	0.0437	0	0.0411	0.0482	0.0445
Mean	0.2510	0.4711	0.3917	0.4734	0.4733	0.4693	0.5090	0.4492	0.4039	0.5179	0.5451

Table 3: SVM: Average AUC of Testing Datasets among Different Sampling Methods.

Dataset	Original	SMOTE	ADASYN	sTL	sSafe	sRST	sCHC	EUSCHC	AHC	FRB+CHC	FRB
ecoli034vs5	0.4972	0.7069	0.5889	0.7236	0.7047	0.6799	0.6747	0.8111	0.5972	0.8217	0.8472
yeast2vs4	0.7362	0.8924	0.8788	0.8900	0.8931	0.8892	0.8656	0.8804	0.8885	0.8757	0.8424
ecoli067vs35	0.5000	0.6860	0.6625	0.7063	0.6790	0.6700	0.6943	0.6675	0.6200	0.7860	0.8325
ecoli0234vs5	0.4972	0.6978	0.6140	0.7081	0.6943	0.6820	0.7181	0.8014	0.5917	0.8289	0.8112
glass015vs2	0.5000	0.7152	0.7352	0.7284	0.7376	0.7496	0.5905	0.4911	0.6484	0.5530	0.575
yeast0359vs78	0.6067	0.7344	0.6936	0.7281	0.7391	0.7334	0.7289	0.6067	0.7371	0.6062	0.6067
yeast0256vs3789	0.5486	0.7960	0.7734	0.7972	0.7965	0.7993	0.8038	0.8064	0.7918	0.7691	0.5761
yeast02579vs368	0.8695	0.9057	0.8610	0.9085	0.9035	0.9071	0.9041	0.9135	0.9052	0.9125	0.9078
ecoli046vs5	0.4973	0.6496	0.5614	0.6488	0.6574	0.6696	0.6395	0.7461	0.5195	0.7880	0.8427
ecoli01vs235	0.4955	0.6606	0.5377	0.6628	0.6598	0.6616	0.6758	0.7423	0.5405	0.7866	0.8659
ecoli0267vs35	0.5000	0.6073	0.5826	0.6093	0.6020	0.6113	0.6405	0.7035	0.5450	0.8176	0.8483
glass04vs5	1.0000	0.9754	0.9754	0.9728	0.9842	0.9830	0.9988	0.9570	1.0000	0.9732	0.9938
ecoli0346vs5	0.4973	0.6974	0.6115	0.7421	0.7124	0.7127	0.7170	0.7878	0.6169	0.8459	0.8656
ecoli0347vs56	0.5000	0.7569	0.7028	0.7594	0.7444	0.7294	0.7511	0.8071	0.5579	0.7888	0.8310
yeast05679vs4	0.5000	0.7869	0.7902	0.7862	0.7861	0.7797	0.7934	0.7860	0.7754	0.7899	0.7786
vowel0	1.0000	0.9993	0.9978	0.9990	0.9988	0.9981	0.9982	0.9933	1.0000	0.9892	0.9933
ecoli067vs5	0.5000	0.6103	0.6100	0.6155	0.6175	0.6106	0.6245	0.8000	0.5725	0.8125	0.845
glass016vs2	0.5000	0.7529	0.7529	0.7106	0.7464	0.7322	0.6239	0.5733	0.7517	0.6114	0.6552
ecoli0147vs2356	0.4984	0.6509	0.6154	0.6920	0.6580	0.6629	0.6891	0.6504	0.5102	0.8054	0.8441
led7digit02456789vs1	0.8788	0.8819	0.8867	0.8799	0.8856	0.8650	0.8946	0.9055	0.8600	0.8844	0.8921
ecoli01vs5	0.4977	0.6602	0.5864	0.6786	0.6566	0.6875	0.6659	0.7091	0.5727	0.8159	0.8432
glass06vs5	1.0000	0.9774	0.9950	0.9574	0.9629	0.9436	0.9895	0.9397	1.0000	0.9840	0.95
glass0146vs2	0.5000	0.6823	0.6849	0.6594	0.6821	0.7142	0.6717	0.5519	0.7153	0.6336	0.6808
glass2	0.5000	0.7132	0.6981	0.6938	0.7127	0.7607	0.6648	0.5248	0.7868	0.6078	0.6875

Dataset	Original	SMOTE	ADASYN	sTL	sSafe	sRST	sCHC	EUSCHC	AHC	FRB+CHC	FRB
ecoli0147vs56	0.5000	0.7160	0.6352	0.7460	0.7335	0.7053	0.6905	0.7722	0.5167	0.8578	0.9171
cleveland0vs4	0.4969	0.5622	0.5575	0.5526	0.5321	0.5421	0.5210	0.5991	0.4811	0.5857	0.6034
ecoli0146vs5	0.4981	0.6440	0.5654	0.6394	0.6467	0.6558	0.6260	0.7731	0.5962	0.8371	0.8442
shuttlec0vsc4	0.9515	0.9747	0.9872	0.9755	0.9747	0.9845	0.9731	0.9715	0.9749	0.9812	0.9897
yeast1vs7	0.5000	0.7583	0.7744	0.7632	0.7602	0.7500	0.6777	0.5000	0.7261	0.6932	0.7579
glass4	0.9092	0.9148	0.9176	0.9113	0.9143	0.9163	0.9333	0.9251	0.9350	0.9230	0.8942
ecoli4	0.8000	0.9101	0.9149	0.9143	0.9171	0.9426	0.9244	0.9528	0.9279	0.9368	0.9231
page_blocks13vs4	0.5700	0.7528	0.7320	0.7493	0.7531	0.7298	0.6847	0.5609	0.5689	0.7141	0.732
abalone918	0.5125	0.8961	0.8860	0.8863	0.8939	0.8916	0.8745	0.5792	0.9144	0.8597	0.83
glass016vs5	0.8443	0.8856	0.9186	0.8791	0.8853	0.9221	0.8979	0.8071	0.8943	0.9186	0.8886
shuttlec2vsc4	0.7	0.9548	0.9548	0.9548	0.9548	0.9590	0.9440	0.6957	0.7	0.9493	0.9632
yeast1458vs7	0.5	0.6427	0.6373	0.6396	0.6444	0.6539	0.6638	0.5	0.6546	0.5958	0.5954
glass5	0.8451	0.8760	0.8256	0.8807	0.8845	0.8515	0.8515	0.8768	0.8451	0.8967	0.8927
yeast2vs8	0.7739	0.7628	0.7242	0.7614	0.7633	0.7770	0.7852	0.7739	0.8381	0.7739	0.7739
yeast4	0.5	0.8156	0.8102	0.8227	0.8160	0.8124	0.8177	0.5093	0.8127	0.7991	0.7663
yeast1289vs7	0.5	0.7141	0.7145	0.7133	0.7109	0.6968	0.7201	0.5	0.7202	0.6990	0.7453
yeast5	0.5	0.9668	0.9635	0.9655	0.9665	0.9655	0.9683	0.7976	0.9691	0.9621	0.9611
ecoli0137vs26	0.5	0.7118	0.5927	0.7390	0.7413	0.6909	0.7294	0.6427	0.5463	0.6655	0.6945
yeast6	0.5	0.8742	0.8597	0.8716	0.8744	0.8736	0.8735	0.5	0.8761	0.8880	0.8880
abalone19	0.5	0.7177	0.7170	0.7163	0.7180	0.7715	0.7166	0.5	0.6881	0.7016	0.7063
Mean	0.6141	0.7784	0.7519	0.7805	0.7795	0.7801	0.7703	0.7248	0.7339	0.8020	0.8133

Table 4: Friedman Rankings of AUC and F-measure.

Preprocessing Method	AUC	F-measure
Original	8.841	7.568
SMOTE	4.636	5.341
ADASYN	6.159	7.341
sTL	4.864	5.636
sSafe	4.624	4.932
sRST	4.659	5.318
sCHC	4.886	3.477
EUSCHC	6	5.659
AHC	5.773	5.5
FRB+CHC	4.455	3.886

Although the hybrid sampling methods can get better results, the main drawback of them is that the size of training set is expanded greatly. If IR of the dataset is large, the size of the re-sampled training set can be nearly double of the original one. This drawback may increase the computational time and complexity of the learning model. Table 5 shows the over-sampling rates of different methods on each dataset and the mean rate of each method. A negative value means the size of re-sampled training set is smaller than that of original one. A value greater than 100% means the size of re-sampled training set is more than 2 times of the original set. Both sCHC and FRB+CHC shrink most of the dataset while the over-sampling rates of the other methods are similar. This shows that both sCHC and FRB+CHC can use less training samples to achieve high performance. Table 6

shows the details of the re-sampled training sets after applying FRB+CHC. It reveals the decrease rate of the majority class, the increase rate of the minority class, and the updated IR. The IR values of re-sampled training sets are not always equal to one because CHC makes use of a fitness function to select a subset of samples. The range of IR is between 0.9 and 1.5.

Table 7 shows the increase rate of the number of support vectors used to form the classification model. The number of support vectors can reflect the complexity of the classification model formed by SVM. When the number of support vectors is smaller, the classification model is more easily applied. Some negative values can be found since the number of support vectors for the re-sampled dataset is less than that of the original dataset. Both sCHC and FRB+CHC have the smallest increase rate of the number of support vectors on average. The average number of support vectors are only increased by around 0.776 times and 0.948 times of the original datasets; while most of the other methods have the number increased by over 2 times.

The results of sCHC and FRB+CHC are similar from the above tables. To show the difference of these two methods, Fig. 5 reveal the average *AUC* results obtained from the training and testing sets (sorted by the nonlinearity of the 1NN classifier.) The x-axis shows the selected 44 datasets. The solid lines in the figures represent the average *AUC* results for the testing set; the dashed lines represent the average *AUC* results for the training set. FRB+CHC shows the advantage

Table 5: Over-sampling Rate (%) of Training Sets among Different Sampling Methods.

Dataset	SMOTE	ADASYN	sTL	sSafe	sRST	sCHC	AHC	FRB+CHC
ecoli0347vs56	80.53	80	77.63	80.53	100.77	-5.60	80	-3.06
yeast2vs4	80.16	80.16	76.85	80.16	80.16	-3.13	80.16	-3.90
ecoli067vs35	80.18	80.18	77.14	80.18	94.13	-5.19	80.18	-4.16
ecoli0234vs5	80.20	80.20	77.10	80.20	89.36	-4.46	80.20	-5.50
glass015vs2	80.23	80.23	70.79	80.23	80.23	-2.06	80.23	-2.28
yeast0359vs78	80.24	80.24	71.49	80.24	80.34	-4.90	80.24	-2.14
yeast0256vs3789	80.28	80.28	74.25	80.28	80.73	-5.70	80.28	-0.92
yeast02579vs368	80.28	80.28	76.97	80.28	80.28	-3.46	80.28	-2.52
ecoli046vs5	80.30	80.30	77.09	80.30	112.06	-3.99	80.30	-2.72
ecoli0147vs2356	84.01	80.33	80.02	84.01	119.57	-4.05	80.33	-1.50
ecoli0267vs35	80.36	80.36	76.56	80.36	94.65	-4.03	80.36	-2.62
glass04vs5	80.44	80.44	77.18	80.44	96.76	-4.40	80.44	-2.91
ecoli034vs5	80.20	80.49	77.72	80.20	86.63	-4.43	80.49	-2.73
ecoli0346vs5	80.39	80.54	78.18	80.39	87.98	-3.54	80.54	-3.12
yeast05679vs4	80.68	80.68	74.48	80.68	80.68	-5.71	80.68	-2.48
vowel0	81.78	81.78	81.78	81.78	84.56	-4.82	81.78	-4.29
ecoli067vs5	81.82	81.82	75.68	81.82	87.27	-4.78	81.82	-1.68
glass016vs2	82.29	82.29	73.18	82.29	82.29	-0.78	82.29	-2.11
ecoli0137vs26	90.48	82.74	87.65	90.48	169.85	-1.33	82.74	0.28
led7digit02456789vs1	83.30	83.30	78.39	83.30	94.02	-3.93	83.30	-7.90
ecoli0147vs56	83.97	83.33	79.91	83.97	137.03	-3.39	83.33	-1.73
glass06vs5	83.34	83.34	80.79	83.34	91.22	-2.71	83.34	-1.67
glass0146vs2	83.41	83.41	74.63	83.41	83.41	-0.94	83.41	-1.40
glass2	84.11	84.11	75.93	84.11	84.11	-2.07	84.11	-0.42
ecoli0146vs5	89.30	84.94	86.00	89.30	139.65	-3.34	84.94	-2.73
cleveland0vs4	84.97	84.97	80.35	84.97	205.49	-3.27	84.97	-1.13
ecoli01vs5	92.75	85.71	90.61	92.75	186.47	-3.58	85.71	-1.22
shuttlec0vsc4	86.55	86.55	86.50	86.55	136.58	-3.32	86.55	-3.17
yeast1458vs7	91.81	86.93	87.16	91.81	91.81	-1.13	86.93	-2.51
glass4	87.85	87.85	83.65	87.85	112.84	-2.68	87.85	-0.95
ecoli4	88.10	88.10	86.46	88.10	88.39	-1.59	88.10	-1.40
page_blocks13vs4	88.14	88.14	86.60	88.14	157.10	-2.88	88.14	-0.86
abalone918	88.58	88.58	83.38	88.58	88.58	-1.72	88.58	-1.44
glass016vs5	90.22	90.22	88.45	90.22	94.57	-2.84	90.22	-1.03
shuttlec2vsc4	90.70	90.70	89.92	90.70	113.19	-3.86	90.70	-3.43
yeast1289vs7	92.32	91.34	88.44	92.32	92.32	-2.50	91.34	-2.35
glass5	91.59	91.59	89.37	91.59	92.76	-1.58	91.59	0.12
yeast2vs8	91.70	91.70	89.83	91.70	98.44	-1.86	91.70	-0.46
yeast4	93.13	93.13	90.09	93.13	93.13	-1.69	93.13	-1.16
yeast1vs7	89.62	93.66	85.04	89.62	89.62	-3.39	93.66	-3.31
yeast5	94.07	94.07	92.62	94.07	94.07	-1.66	94.07	-2.09
ecoli01vs235	82.59	95.02	78.95	82.59	106.04	-4.04	95.02	-0.59
yeast6	95.28	95.28	93.36	95.28	95.28	-2.54	95.28	-1.88
abalone19	98.47	98.47	97.32	98.47	98.47	1.04	98.47	-0.31
Mean	85.70	85.40	81.94	85.70	103.47	-3.13	85.40	-2.17

Table 6: The Details of the Re-sampled Datasets After FRB+CHC.

Dataset	Decrease Rate of	Increase Rate of	Updated IR
	Majority Class	Minority Class	
ecoli034vs5	0.474	3.700	1.010
yeast2vs4	0.469	3.865	0.995
ecoli067vs35	0.500	4.224	0.874
ecoli0234vs5	0.488	3.463	1.048
glass015vs2	0.405	2.989	1.369
yeast0359vs78	0.418	3.995	1.064
yeast0256vs3789	0.452	4.005	1.002
yeast02579vs368	0.456	3.841	1.028
ecoli046vs5	0.469	3.900	1.001
ecoli01vs235	0.445	3.939	1.030
ecoli0267vs35	0.500	3.756	0.969
glass04vs5	0.464	4.325	0.934
ecoli0346vs5	0.476	4.188	0.935
ecoli0347vs56	0.452	3.940	1.032
yeast05679vs4	0.441	3.938	1.064
vowel0	0.514	4.619	0.863
ecoli067vs5	0.461	4.025	1.076
glass016vs2	0.393	3.635	1.357
ecoli0147vs2356	0.424	4.651	1.084
led7digit02456789vs1	0.499	4.603	0.983
ecoli01vs5	0.470	4.975	0.978
glass06vs5	0.482	5.339	0.910
glass0146vs2	0.392	4.568	1.212
glass2	0.449	4.826	1.101
ecoli0147vs56	0.475	5.510	0.992
cleveland0vs4	0.442	6.049	0.984
ecoli0146vs5	0.470	5.550	1.053
shuttlec0vsc4	0.507	6.530	0.908
yeast1vs7	0.420	5.525	1.277
glass4	0.489	6.651	1.038
ecoli4	0.463	7.300	1.024
page_blocks13vs4	0.480	7.236	1.003
abalone918	0.478	7.658	0.994
glass016vs5	0.486	9.296	0.974
shuttlec2vsc4	0.514	9.900	0.936
yeast1458vs7	0.391	7.908	1.513
glass5	0.478	10.321	1.052
yeast2vs8	0.452	10.500	1.105
yeast4	0.468	12.397	1.117
yeast1289vs7	0.418	11.967	1.373
yeast5	0.480	14.768	1.081
ecoli0137vs26	0.467	19.633	1.023
yeast6	0.451	17.693	1.217
abalone19	0.487	61.247	1.067

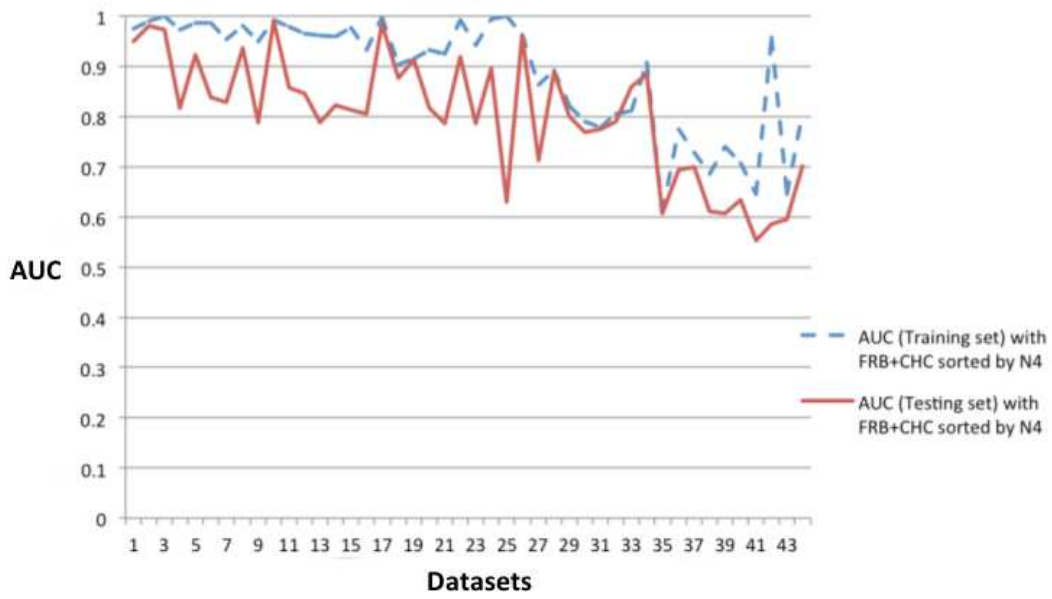
Table 7: The Increase Rate of Number of Support Vectors of the Classification Model formed by SVM.

Dataset	SMOTE	ADASYN	sTL	sSafe	sRST	sCHC	AHC	FRB+CHC
ecoli0347vs56	0.245	0.529	0.418	0.143	0.476	-0.117	0.178	0.026
yeast2vs4	2.662	4.354	2.507	2.981	2.698	1.180	2.377	1.000
ecoli067vs35	0.252	0.506	0.468	0.249	0.465	-0.103	0.215	0.021
ecoli0234vs5	0.272	0.538	0.472	0.122	0.473	-0.097	0.197	0.011
glass015vs2	3.731	3.687	3.193	3.996	3.583	1.705	3.310	1.279
yeast0359vs78	1.384	1.676	1.149	1.434	1.418	0.251	1.361	0.093
yeast0256vs3789	3.555	5.151	3.240	3.669	3.577	1.414	3.416	0.990
yeast02579vs368	2.142	5.240	1.919	2.251	2.188	0.783	1.990	0.521
ecoli046vs5	0.263	0.477	0.460	0.125	0.490	-0.100	0.177	0.031
ecoli0147vs2356	0.229	0.566	0.435	0.170	0.494	-0.103	0.186	0.053
ecoli0267vs35	0.279	0.521	0.490	0.234	0.467	-0.092	0.209	0.040
glass04vs5	1.839	1.477	1.467	4.104	1.242	-0.043	0.122	0.306
ecoli034vs5	0.269	0.526	0.456	0.135	0.477	-0.108	0.163	0.029
ecoli0346vs5	0.286	0.424	0.488	0.140	0.497	-0.094	0.150	0.031
yeast05679vs4	3.500	4.147	3.113	3.561	3.575	1.401	3.474	1.469
vowel0	1.116	1.333	0.752	2.311	0.972	-0.027	0.448	1.631
ecoli067vs5	0.227	0.440	0.391	0.212	0.438	-0.109	0.144	0.055
glass016vs2	3.868	3.927	3.296	4.209	3.660	1.840	3.291	1.530
ecoli0137vs26	0.175	0.404	0.338	0.089	0.439	-0.191	0.160	-0.146
led7digit02456789vs1	3.332	3.322	2.864	4.139	2.838	0.360	2.608	0.563
ecoli0147vs56	0.218	0.487	0.391	0.121	0.522	-0.145	0.147	0.031
glass06vs5	1.494	1.272	1.124	2.924	1.204	0.021	0.077	0.250
glass0146vs2	3.967	4.008	3.396	4.271	3.725	1.915	3.391	1.643
glass2	3.820	3.893	3.221	4.036	3.671	1.881	3.444	1.712
ecoli0146vs5	0.189	0.379	0.361	0.105	0.394	-0.155	0.137	0.014
cleveland0vs4	0.540	0.741	0.712	0.247	0.505	-0.077	0.326	0.008
ecoli01vs5	0.148	0.494	0.309	0.082	0.417	-0.128	0.149	0.032
shuttlec0vsc4	0.221	2.557	0.264	0.001	0.450	-0.109	0.103	1.326
yeast1458vs7	3.121	6.093	2.947	3.233	3.100	1.064	5.311	0.966
glass4	1.968	1.873	0.836	4.223	1.852	0.084	0.675	0.344
ecoli4	2.173	2.998	2.020	2.668	2.394	0.772	1.796	1.012
page_blocks13vs4	0.836	1.042	0.863	0.057	1.198	0.017	0.523	0.110
abalone918	8.586	9.046	8.058	10.025	8.077	3.752	6.750	4.309
glass016vs5	2.084	1.760	1.440	3.498	1.846	0.226	0.492	0.512
shuttlec2vsc4	0.616	1.428	1.388	0.153	1.310	0.282	0.229	0.422
yeast1289vs7	4.846	2.993	4.611	5.170	5.087	1.802	2.825	1.491
glass5	2.307	2.169	1.884	4.623	2.151	0.249	0.524	0.574
yeast2vs8	4.905	8.609	4.920	5.282	5.169	2.101	4.436	1.184
yeast4	3.133	3.547	2.902	3.379	3.173	0.883	2.966	0.919
yeast1vs7	5.196	4.672	4.870	5.523	5.382	2.673	4.088	2.422
yeast5	3.178	3.458	2.617	3.676	3.265	1.578	2.930	2.007
ecoli01vs235	0.286	0.268	0.463	0.153	0.484	-0.096	0.085	0.041
yeast6	6.825	9.553	6.376	7.216	6.900	2.278	6.479	2.632
abalone19	13.156	13.209	12.829	14.221	12.973	5.535	10.986	8.226
Mean	2.351	2.859	2.198	2.708	2.403	0.776	1.887	0.948

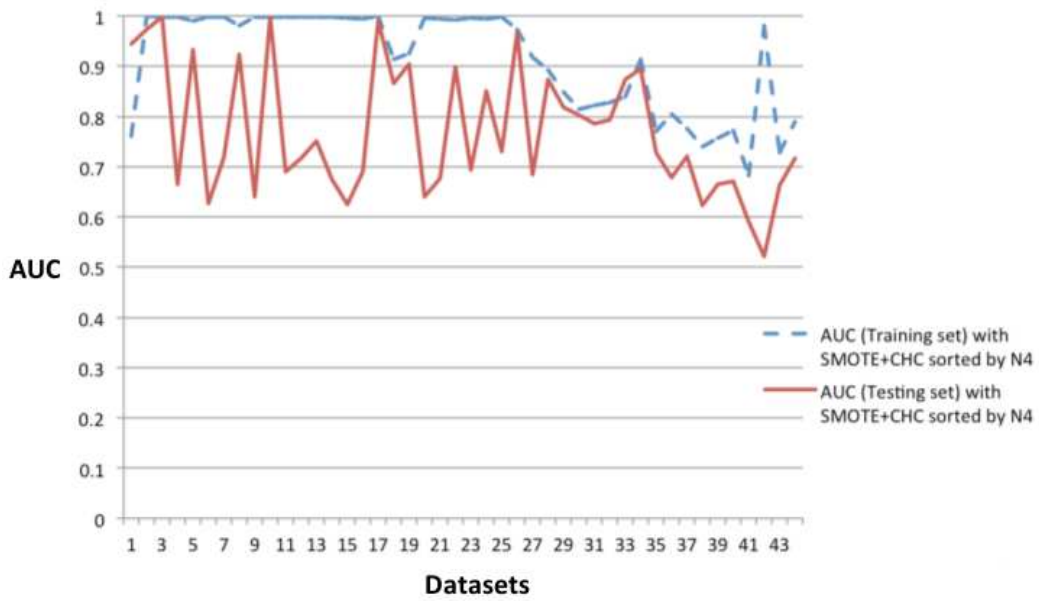
on relaxing the over-fitting problem since the performances on training set and testing set are similar.

Fig. 6 and 7 show an example of the distribution of the positive samples and negative samples after the re-sampling of FRB+CHC and sCHC respectively. The circle dots correspond to the samples of the majority class. The square dots correspond to the samples of the original minority class. The triangle dots correspond to the synthetic samples. Fig. 7 show that the synthetic samples are generated densely around some of the original minority samples. On the contrary, the synthetic samples in Fig. 6 are distributed more evenly in the area of the original minority samples. Therefore, sCHC runs into the over-fitting problem more easily.

Figs. 8 and 9 show the overall results in terms of $F - measure$ and AUC for different classifiers respectively. Only a small difference of the results for 1NN among all the preprocessing methods is revealed. FRB+CHC obtains the highest value of AUC for both C4.5 and 1NN. An improvement by FRB+CHC in terms of $F - measure$ is shown. In addition, a robust behavior of FRB+CHC is shown when the results of the three classifiers only have a small difference. Most of the preprocessing methods can perform better than the original datasets in terms of the average values of $F - measure$ and AUC . This confirms that preprocessing is an important step to deal with imbalanced datasets.



(a) AUC results with FRB+CHC



(b) AUC results with sCHC

Figure 5: Average AUC results obtained from training and testing sets.

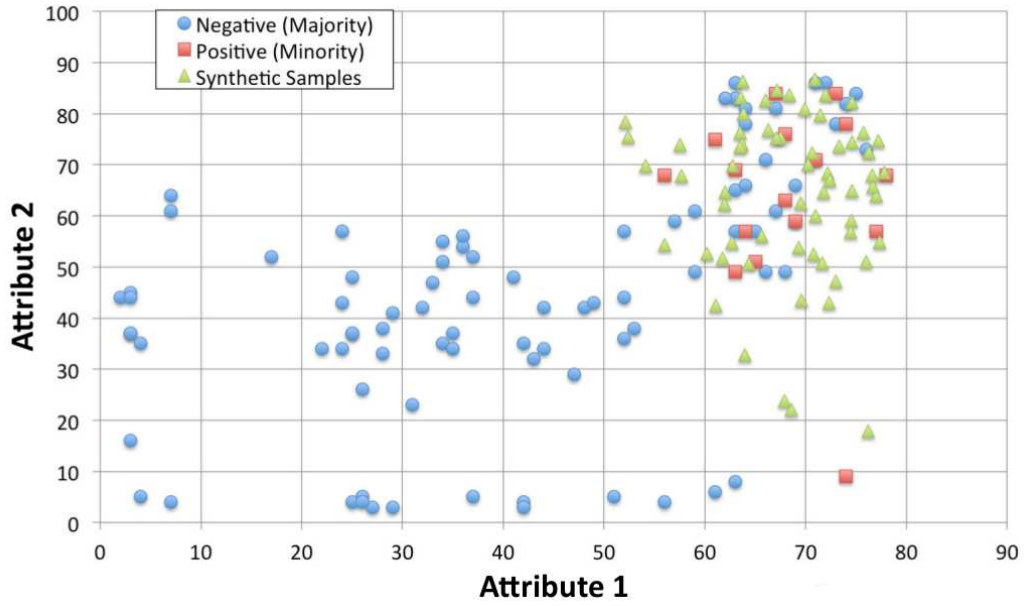


Figure 6: Distribution of the samples after the implementation of FRB+CHC.

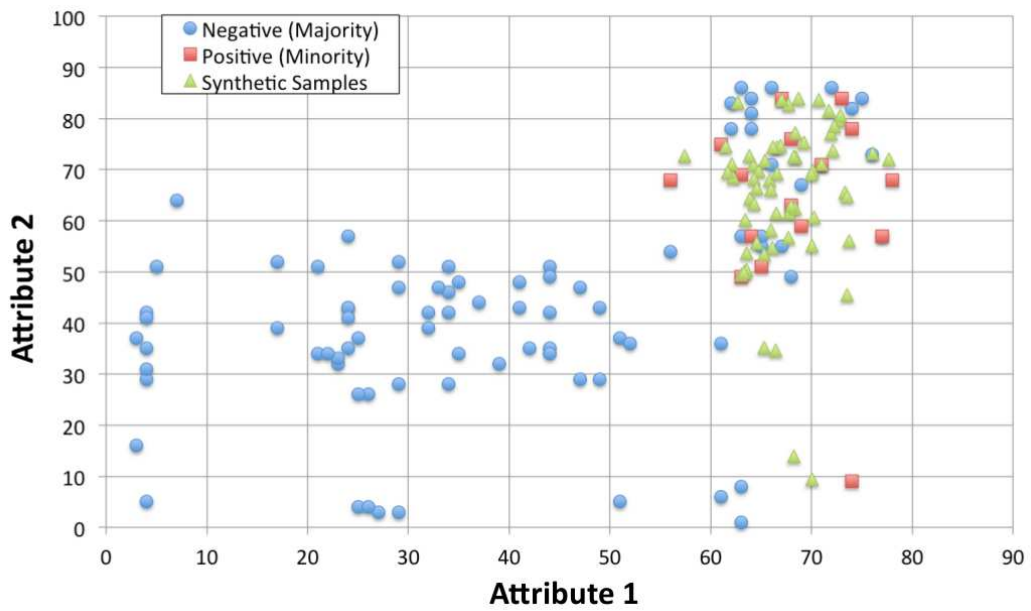


Figure 7: Distribution of the samples after the implementation of sCHC.

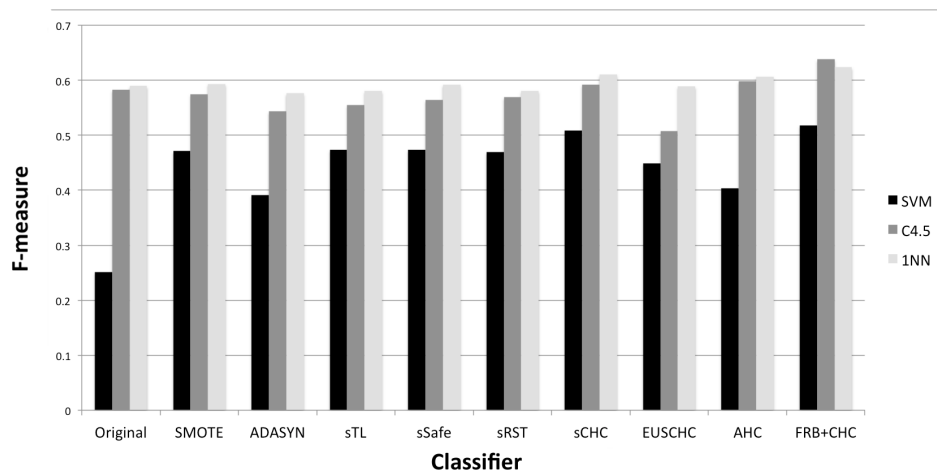


Figure 8: Average F-measure for different classifiers.

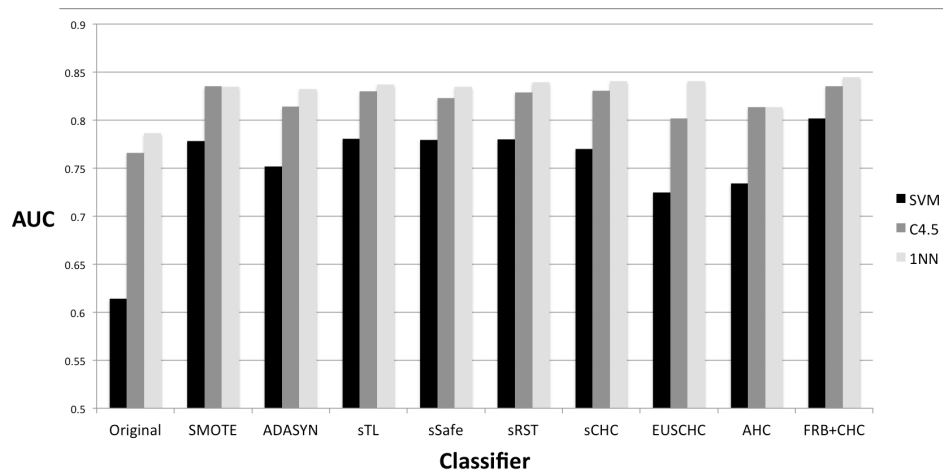


Figure 9: Average AUC for different classifiers.

5. Conclusion

A hybrid re-sampling method developed based on both over-sampling and under-sampling has been proposed. The new synthetic samples of the minority class are generated based on fuzzy logic. To minimize the size of datasets, CHC has been employed over the new samples and the majority samples as a cleaning method to the over-sampled training set.

The proposed sampling method (FRB+CHC) is compared to SMOTE, ADASYN, sTL, sSafe, sRST, sCHC, EUSCHC, and AHC on 44 datasets. To evaluate the performance of these nine sampling methods, the same SVM classifier has been used to obtain the experimental results. It is shown that FRB and FRB+CHC outperforms the other sampling methods on both $F - measure$ and AUC . FRB shows its advantage to act as an over-sampling method. If data size is not a consideration, FRB is a better choice of pre-processing method.

FRB+CHC obtains the best ranking by means of AUC . FRB+CHC and sCHC have similar performance in $F - measure$, which indicates that CHC is a good choice of data cleaning method. The AUC results of SMOTE, sTL, sSafe, sRST, and sCHC are similar since all of them use SMOTE to perform over-sampling. To show the advantages of the proposed method, the over-sampling rate and the number of support vectors formed from SVM for different methods are also compared. In addition, the C4.5 and 1NN classifiers are used and FRB+CHC shows a robust

behavior among different classifiers. FRB+CHC achieves good results under the above criteria, which reflects that FRB+CHC achieves a good balance between accuracy and over-sampling rate. It also has a low impact to the complexity of the learning model. The major reason is that CHC only selects the samples to increase the performance of the datasets, but not considering the locations of the samples. Therefore, the most representative samples are selected to form the training sets.

Acknowledgment

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