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"A comparison of classification models for imbalanced datasets "

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Abstract

Many practical classification problems are imbalanced; i.e., at least one of the classes constitutes only a very small minority of the data. For such problems, the interest usually leans towards correct classification of the minor class. Examples of such problems include fraud detection, rare disease diagnosing, etc. However, the most commonly used classification algorithms do not work well for such problems because they aim to minimize the overall error rate, rather than paying special attention to the minor class. In the master's thesis, a number of models are evaluated with the objective to find those that better address the classification problem of imbalanced datasets. A special focus is given to the investigation of some ways of dealing with imbalanced datasets based on a Logistic Regression.

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Louvain School of Management and Norwegian School of Economics

A comparison of classification models for imbalanced datasets

Research Master's Thesis submitted by **Sergei Kurin**

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with a view of getting the degrees

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Executive Summary

Many practical classification problems are imbalanced; i.e., at least one of the classes constitutes only a very small minority of the data. For such problems, the interest usually leans towards correct classification of the minor class. Examples of such problems include fraud detection, rare disease diagnosing, etc. However, the most commonly used classification algorithms do not work well for such problems because they aim to minimize the overall error rate, rather than paying special attention to the minor class.

In the master's thesis, a number of models are evaluated with the objective to find those that better address the classification problem of imbalanced datasets. Namely, k-Nearest Neighbors (k-NN), Support Vector Machines (SVMs), and Random Forest classification models are discussed. A special focus is given to the investigation of some ways of dealing with imbalanced datasets based on a Logistic Regression. Furthermore, a comparison of different preprocessing techniques such as undersampling, oversampling, synthetic data generation (SMOTE) is made.

The methodology used is based on the analysis of imbalanced datasets and implementation of classification models using statistical software with data mining algorithms (e.g., R). Specifically, yeast datasets from KEEL dataset repository and a real-world dataset 'Credit Card Fraud Detection' available on Kaggle have been studied.

For all the studied *yeast* datasets, both oversampling and the combination of overand undersampling techniques have revealed the highest accuracy in terms of area under Receiver Operating Characteristics (ROC) curve (on average it is equal 0.9610 for both over- and undersampling techniques combined with Random Forest). Undersampling is less competitive when the number of instances is small (as for *yeast*3, *yeast*4, *yeast*5, *yeast*6 datasets) and otherwise for a large number of instances (as for 'Credit Card Fraud Detection' dataset).

Logistic Regression model demonstrates relatively poor accuracy (or very low accuracy of 0.5682 specifically for yeast4 dataset) compared to other classification models if additional adjustment procedure for priors is not applied. However, the same model with adjustment for priors (so called Log-EM model if priors are not known and TP model for known priors) is competitive with more advanced classification models (e.g., Random Forest). Note that Log-EM model procedure has never been studied in the literature for imbalanced datasets. Thus, this work provides the very first results on Log-EM model specifically for imbalanced datasets.

For the future work, it would be important to focus on such related topics as deeper comparison of imbalanced datasets with different number of observations and imbalance ratios as well as more comprehensive study of different sampling techniques including some extensions of SMOTE.

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Introduction

Currently, in the Big Data (BD) era that we are living in, data science algorithms are of great importance to improve the performance of different applications (especially for those areas where data are collected daily and where meaningful knowledge and valuable assistance can be extracted to improve current systems) (Pérez-Ortiz and Jiménez-Fernández, 2016).

Generally, the term data science refers to the extraction of knowledge from data. This involves a wide range of techniques and theories drawn from many research fields within mathematics, statistics and information technology, including statistical learning, data engineering, pattern recognition, uncertainty modeling, probability models, high performance computing, signal processing and machine learning (ML), among others. Precisely, the growth and development of this last research area has made data science more relevant, increasing the necessity of data scientists and the development of novel methods in the scientific community, given the great breadth and diversity of knowledge and applications of this area.

Classification problems and methods have been considered a key part of ML, with a huge amount of applications published in the last few years. The concept of classification in ML has been traditionally treated in a broad sense, very often including supervised, unsupervised and semi-supervised learning problems. Unsupervised learning is focused on the discovery and analysis of the structure of unlabeled data. This paradigm is especially useful to analyze whether there are differentiable groups or clusters present in the data (e.g., for segmentation). In the case of supervised learning, however, each data input object is preassigned a class label. The main task of supervised algorithms is to learn a model that ideally produces the same labeling for the provided data and generalizes well on unseen data (i.e., prediction). In this regard, classification learning applications are widely used to cope with difficult problems arising, for instance, from new renewable energy (RE) sources. A huge amount of RE applications can be found in the literature, such as prediction problems (e.g., solar radiation (Wang and Zhen, 2015) or significant wave height estimation (Fernandez and Salcedo-Sanz, 2015)), optimization algorithms (wind farm or RE devices'

1.1 Motivation and Problem Statement

In many supervised classification learning applications, there is a significant difference between the prior probabilities of different classes, i.e., between the probabilities with which an example belongs to the different classes of the classification problem. This situation is known as the class imbalance problem and it is common in many real problems from telecommunications, web, finance-world, ecology, biology, medicine, and which can be considered one of the top problems in data mining today. Examples of such problems include fraud detection, rare disease diagnosing (Chen and Liaw, 2004), new control techniques or fault diagnosis in RE systems (Panigrahi and Dash, 2009).

The hitch with imbalanced datasets is that standard classification learning algorithms are often biased towards the majority class (known as the "negative" class) and therefore there is a higher misclassification rate for the minority class instances (called the "positive" instances) (Lopéz and Fernández, 2013). Most of the studies on the behavior of several standard classifiers in imbalance domains have shown that significant loss of performance is mainly due to the skewed class distribution, given by imbalance ratio (IR), defined as ratio of the number of instances in the majorty class to the number of examples in the minority class.

The research problem in the master's thesis: classification models reveal low accuracy when dealing with imbalanced datasets. Therefore, a number of models will be evaluated with the objective to find those that better address the classification problem of imbalanced datasets. A special focus will be given to the investigation of some ways of dealing with imbalanced datasets based on a Logistic Regression.

More specifically, the research questions are presented below:

- 1. How do sampling techniques and classification models affect the accuracy of classification?
- 2. How does adjustment procedure affect the outcome of Logistic Regression?
- 3. What evaluation metrics is the most appropriate for imbalanced datasets?

Thus, the main objectives of the master's thesis are as follows:

- 1. The analysis of current problems of imbalanced learning;
- 2. The investigation of different classification models and evaluation metrics for imbalanced datasets;
- 3. The comparison of different classification models for imbalanced datasets;
- 4. The investigation of classification models based on Logistic Regression.

The theoretical results will include a literature review of the problem of imbalanced learning and the analysis of different classification models and their performance measures for imbalanced datasets. The empirical results will include the implementation of classification models using statistical software with data mining algorithms (e.g. R).

1.2 Thesis Structure

The thesis is organized as follows. First, some background knowledge on imbalanced problems and the results of related works will be presented (Chapter 2, Chapter 3, Chapter 4). Then, main results of this study will be discussed (Chapter 5) followed by the conclusion (Chapter 6).

Chapter 2

In this Chapter, different methods to deal with imbalanced datasets will be discussed. A special attention will be paid to some sampling techniques including undersampling, oversampling, and synthetic data generation.

Chapter 3

This Chapter is dedicated to different classification models that will be later evaluated with the objective to find those that better address the classification problem of imbalanced datasets. Namely, k-Nearest Neighbors (k-NN), Support Vector Machines (SVMs), and Random Forest are discussed. It will be shown that some of these classifiers can be oriented towards the class imbalance issues. For example, focused oversampling method is an efficient tool in combination with SVMs or stratified bootstrap can be used for Random Forest model. An important question for imbalanced problems is how to assess the performance of different classifiers. Simple predictive accuracy based on confusion matrix is not appropriate in such situations. It will be shown that the Area Under the ROC Curve (AUC) provides a single measure of a classifier's performance for evaluating which model is better.

Chapter 4

In this Chapter, a special focus will be given to the investigation of some ways of dealing with imbalanced datasets based on a Logistic Regression. An iterative procedure that estimates the new a priori probabilities of a new dataset and adjusts the outputs of a classifier will be proposed for evaluation on imbalanced datasets.

Chapter 5

In this Chapter, the main results of performed study are presented and discussed. First, a comparison of different preprocessing techniques and classification models is made on the basis of several sample imbalanced datasets. Then, Logistic Regression with its adjustment procedure is analyzed. Finally, different classifiers and preprocessing techniques are applied to a real-word dataset 'Credit Card Fraud Detection'.

Chapter 6

This Chapter presents main conclusions and gives proposals for the future work related to the class imbalance problem.

Related Work

Throughout the last years, many solutions have been proposed to deal with the problem of poor performance of imbalanced datasets, both for standard learning algorithms and for ensemble techniques (Lopéz and Fernández, 2013). This can be categorized into three major groups:

- 1. Data sampling in which the training instances are modified in such a way to produce a more or less balanced class distribution that allow classifiers to perform in a similar manner to standard classification.
- 2. Cost-sensitive learning. This type of solutions incorporate approaches at the data level, at the algorithmic level, or at both levels combined, considering higher costs for the misclassification of examples of the positive class with respect to the negative class, and therefore, trying to minimize higher cost errors.
- 3. Algorithmic modification. This procedure is oriented towards the adaptation of base learning methods to be more attuned to class imbalance issues.

In the master's thesis, cost-sensitive learning and algorithmic modification are not analyzed.

2.1 Sampling Methods

Sampling is a popular methodology to counter the problem of class imbalance. In the specialized literature, one can find some papers about resampling techniques studying the effect of changing class distribution in order to deal with imbalanced datasets (Ha and Bunke, 1997). The goal of sampling methods is to create a dataset that has a relatively balanced class distribution, so that traditional classifiers are better able to capture the decision boundary between the majority and the minority classes. Those works have proved empirically that applying a preprocessing step in order to balance the class distribution is usually a useful solution. Furthermore, the main advantage of these techniques is that they are independent of the underlying classifier. Since the sampling methods are used to make the classification of the minority class instances easier, the resulting (sampled) dataset should represent a

"reasonable" approximation of the original dataset. Specifically, the resulting dataset should contain only instances that are, in some sense, similar to those in the original dataset, that is, all instances in the modified dataset should be drawn from the same (or similar) distribution to those originally in the dataset. Note that sampling methods need not create an exactly balanced distribution, merely a distribution that the traditional classifiers are better able to handle.

These sampling methods have acquired higher importance after many researches have proved that balanced data results in improved overall classification performance compared to an imbalanced dataset (Analytics Vidhya Content Team, 2016).

The main sampling methods used to treat imbalanced datasets are presented below:

- 1. Undersampling;
- 2. Oversampling;
- 3. Both undersampling and oversampling;
- 4. Synthetic data generation (e.g., synthetic minority oversampling technique (SMOTE)).

Two of the first sampling methods developed are random undersampling and random oversampling. In the random undersampling, the majority class instances are discarded at random until a more balanced distribution is reached. Consider, for example, a dataset consisting of 10 minority class instances and 100 majority class instances. In random undersampling, one might attempt to create a balanced class distribution by selecting 90 majority class instances at random to be removed. The resulting dataset will then consist of 20 instances: 10 (randomly remaining) majority class instances and (the original) 10 minority class instances.

Alternatively, in random oversampling, minority class instances are copied and repeated in the dataset until a more balanced distribution is reached. Thus, if there are two minority class instances and 100 majority class instances, traditional oversampling would copy the two minority class instances 49 times each. The resulting dataset would then consists of 200 instances: the 100 majority class instances and 100 minority class instances (i.e., 50 copies each of the two minority class instances). While random undersampling and random oversampling create more balanced distributions, they both suffer from serious drawbacks. For example, in random undersampling (potentially), vast quantities of data are discarded. In the random undersampling example mentioned above, for instance, roughly 82% of the data (the 90 majority class instances) was discarded. This can be highly problematic, as

the loss of such data can make the decision boundary between minority and majority instances harder to learn, resulting in a loss in classification performance.

Alternatively, in random oversampling, instances are repeated (sometimes to very high degrees). Consider the random oversampling example mentioned above, where each instance had to be replicated 49 times in order to balance out the class distribution. By copying instances in this way, one can cause drastic ovefitting to occur in the classifier, making the generalization performance of the classifier exceptionally poor. The potential for ovefitting is especially true as the class imbalance ratio becomes worse, and each instance must be replicated more and more often.

Thus, one major drawback of sampling techniques is that one needs to determine how much sampling to apply. An oversampling level must be chosen so as to promote the minority class, while avoiding ovefitting to the given data. Similarly, an undersampling level must be chosen so as to retain as much information about the majority class as possible, while promoting a balanced class distribution.

In order to overcome these limitations, more sophisticated techniques have been developed, e.g., SMOTE.

2.2 Synthetic Minority Oversampling Technique (SMOTE)

Chawla and Bowyer, 2002 developed a method of creating synthetic instances instead of merely copying existing instances in the dataset. This techniques is known as the synthetic minority oversampling technique (SMOTE). This approach is inspired by a technique that proved to be successful in handwritten character recognition (Ha and Bunke, 1997). They created extra training data by performing certain operations on real data. In their case, operations like rotation and skew were natural ways to perturb the training data. In SMOTE, the minority class is oversampled by taking each minority class sample and introducing synthetic examples along the line segments joining any/all of the k minority class nearest neighbors. Depending upon the amount of oversampling required, neighbors from the k nearest neighbors are randomly chosen. There is an implementation that currently uses five nearest neighbors. For instance, if the amount of oversampling needed is 200%, only two neighbors from the five nearest neighbors are chosen and one sample is generated in the direction of each. Synthetic samples are generated in the following way:

- 1. Take the difference between the feature vector (sample) under consideration and its nearest neighbor;
- 2. Multiply this difference by a random number between 0 and 1;

3. Add it to the feature vector under consideration.

Thus, in SMOTE, the training set is altered by adding synthetically generated minority class instances, causing the class distribution to become more balanced. We say that the instances created are synthetic, as they are, in general, new minority instances that have been extrapolated and created out of existing minority class instances.

2.3 Conclusion

There are three main ways to deal with imbalanced datasets: data sampling, costsensitive learning, algorithmic modification. In this work, only data sampling will be studied thoroughly. It includes undersampling, oversampling, both undersampling and oversampling, SMOTE. Classification Models

The general aim of supervised classification algorithms is to separate the classes of the problem (with a margin as wide as possible) using only training data (Pérez-Ortiz and Jiménez-Fernández, 2016). If the output variable has two possible values, the problem is referred to as binary classification. On the other hand, if there are more than two classes, the problem is named multiclass or multinomial classification.

A classification problem can be formally defined as the task of estimating the label y of a K-dimensional input vector \mathbf{x} , where $\mathbf{x} \in X \subseteq \mathbb{R}^K$ and $y \in \Omega = (\omega_1, \omega_2, ..., \omega_n)$. $\omega_1, \omega_2, ..., \omega_n$ denote n classes. This task is accomplished by using a classification rule or function $g: X \to \Omega$ able to predict the label of new patterns. In the supervised setting, we are given a training set of N_t points, represented by D, from which g will be adjusted, $D = (x_i, y_i), i = 1, ..., N_t$.

3.1 k-Nearest Neighbors

In k-NN classification, a point is classified by a majority vote of its neighbors, which means the point will be assigned to the class most common among its k nearest neighbors (Barber, 2012).

In other words, given a query point x_0 , we find the k training points x_r , r=1,...,k closest in distance to x_0 , and then classify using majority vote among the k neighbors. In order to compute the distance between two points a and b, Euclidean distance can be used: $d(a,b) = \sqrt{\sum_i (a_i - b_i)^2}$ (i denotes the i-th coordinate in Euclidean space).

If we assume the Euclidean distance as the dissimilarity measure, the k-NN algorithm considers a hypersphere centred on the test point \mathbf{x} . The radius of the hypersphere is increased until it contains exactly k train inputs. The class label ω_i is then given by the most numerous class within the hypersphere (Fig. 3.1). We centre a hypersphere around the point we wish to classify (here the central dot). The inner circle corresponds to the nearest neighbour, a square. However, using the 3 nearest neighbours, we find that there are two round-class neighbours and one square-class neighbour—and we would therefore classify the central point as round-class.

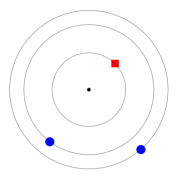


Fig. 3.1: k-Nearest Neighbors model

Whilst there is some sense in making k > 1, there is certainly little sense in making $k = N_t$ (N_t being the number of training points). For k very large, all classifications will become the same – simply assign each novel \mathbf{x} to the most numerous class in the train data. This suggests that there is an optimal intermediate setting of k which gives the best generalisation performance. This can be determined by CV.

3.2 Support Vector Machines

Support vector machine (SVM) is a statistical learning algorithm that aims to identify the optimal decision boundary between classes to minimize misclassification. Owing to its theoretical and practical advantages, such as solid mathematical background, high generalization capability, and ability to find global and nonlinear classification solutions, SVM have been very popular among the machine learning and data-mining researchers (He and Ma, 2013).

The support vector machine (SVM) is an extension of the support vector classifier that results from enlarging the feature space in a specific way, using kernels, in order to accommodate a non-linear boundary between the classes. Thus, prior to classification, a kernel is typically applied to the input feature space to increase separability between classes (Hastie and Tibshirani, 2009).

In this study, SVMs with a radial basis function (RBF) kernel (Johnson and Tateishi, 2012) are used for classification. With the SVMs-RBF classifier, one can adjust the cost parameter (c) and the kernel spread function (γ) prior to classification. A wide range of paired combinations of c values and γ values should be tested. The optimal parameters for each of the SVMs classifications are selected by CV.

Although SVMs often work effectively with balanced datasets, they could produce suboptimal results with imbalanced datasets. More specifically, an SVM classifier trained on an imbalanced dataset often produces models that are biased toward the majority class and have low performance on the minority class. All the data preprocessing methods discussed in Chapter 2 can be used to balance the datasets

before training the SVMs models. These methods include random and focused under/oversampling methods and synthetic data generation methods such as SMOTE. However, more sophisticated preprocessing techniques for SVMs exist. For example, the paper (Batuwita and Palade, 2010) presents an efficient focused oversampling method for SVMs. In this method, first the separating hyperplane found by training an SVM model on the original imbalanced dataset is used to select the most informative examples for a given classification problem, which are the data points lying around the class boundary region. Then, only these selected examples are balanced by oversampling as opposed to blindly oversampling the complete dataset. This method reduces the SVM training time signficantly while obtaining comparable classification results to the original oversampling method.

3.3 Random Forest

Random Forest (RF) is an ensemble of unpruned classification trees, induced from bootstrap samples of the training data, using random feature selection in the tree induction process. The RF approach starts with a standard decision tree technique but it takes this notion to the next level by creating a large number of decision trees (Benyamin, 2012). In this approach illustrated in Fig. 3.2, each tree is induced from a bootstrap sample of the training data, i.e., every observation is fed into every decision tree. Then, the most common outcome for each observation is calculated and used as the final output. Thus, prediction is made by aggregating (majority vote) the predictions of the ensemble.

In this model, there are some sources of randomness in order to make these trees different from one another. Therefore, this approach provides a group of unique trees which all make their classifications differently.

Random Forest generally exhibits a substantial performance improvement over the single tree classifier such as CART and C4.5 (Chen and Liaw, 2004). However, similar to most classiflers, RF can also suffer from the curse of learning from an extremely imbalanced training data set. As it is constructed to minimize the overall error rate, it will tend to focus more on the prediction accuracy of the majority class, which often results in poor accuracy for the minority class. There is a significant probability that a bootstrap sample contains few or even none of the minority class, resulting in a tree with poor performance for predicting the minority class. One of the way of fixing this problem is to use a stratified bootstrap; i.e., sample with replacement from within each class.

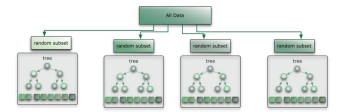


Fig. 3.2: Random Forest model

Let us consider the RF classification model in more detail. RF grows many classification trees (Breiman and Cutler, 2004). To classify a new object from an input vector, put the input vector down each of the trees in the forest. Each tree gives a classification, and one can say the tree "votes" for that class. The forest chooses the classification having the most votes (over all the trees in the forest).

Each tree is grown as follows:

- 1. If the number of cases in the training set is N, sample N cases at random but with replacement, from the original data. This sample will be the training set for growing the tree.
- 2. If there are M input variables, a number m < < M is specified such that at each node, m variables are selected at random out of the M and the best split on these m is used to split the node. The value of m is held constant during the forest growing.
- 3. Each tree is grown to the largest extent possible. There is no pruning.

When the training set for the current tree is drawn by sampling with replacement, about one-third of the cases are left out of the sample. This out-of-sample data is used to get a running unbiased estimate of the classification error as trees are added to the forest. It is also used to get estimates of variable importance.

After each tree is built, all of the data are run down the tree, and proximities are computed for each pair of cases. If two cases occupy the same terminal node, their proximity is increased by one. At the end of the run, the proximities are normalized by dividing by the number of trees. Proximities are used in replacing missing data, locating outliers, and producing illuminating low-dimensional views of the data.

3.4 Accuracy Measure

A classifier is, typically, evaluated by a confusion matrix as illustrated in Table 3.1 (Lopéz and Fernández, 2013). In the confusion matrix, TN is the number of nega-

tive examples correctly classified (True Negatives), FP is the number of negative examples incorrectly classified as positive (False Positives), FN is the number of positive examples incorrectly classified as negative (False Negatives) and TP is the number of positive examples correctly classified (True Positives). Predictive accuracy is defined as Accuracy = (TP+TN)/(TP+FP+TN+FN). However, predictive accuracy might not be appropriate when the data is imbalanced and/or the costs of different errors vary markedly. As an example, consider the classification of pixels in mammogram images as possibly cancerous (Woods and Doss, 1993). A typical mammography dataset might contain 98% normal pixels and 2% abnormal pixels. A simple default strategy of guessing the majority class would give a predictive accuracy of 98%. The nature of the application requires a fairly high rate of correct detection in the minority class and allows for a small error rate in the majority class in order to achieve this (Chawla and Bowyer, 2002). Simple predictive accuracy is clearly not appropriate in such situations.

Tab. 3.1: Confusion matrix for a two-class problem

		Predicted classes				
		Positive prediction	Negative prediction			
True classes	Positive class	True Positive (TP)	False Negative (FN)			
True classes	Negative class	False Positive (FP)	True Negative (TN)			

In imbalanced domains, the evaluation of the classifier's performance must be carried out using specific metrics in order to take into account the class distribution. Concretely, there are four metrics from Table 3.1 to measure the classification performance of both, positive and negative, classes independently:

- True positive rate: %TP = TP/(TP + FN);
 True negative rate: %TN = TN/(FP + TN);
 False positive rate: %FP = FP/(TN + FP);
 False negative rate: %FN = FN/(TP + FN).
- There is a necessity of combining the individual measures of both the positive and negative classes, as none of these measures alone is adequate by itself.

A well-known approach to unify these measures and to produce an evaluation criteria is to use the Receiver Operating Characteristics (ROC) graphic. This graphic allows the visualization of the trade-off between the benefits and costs, as it evidences that any classifier cannot increase the number of true positives without also increasing the false positives. The Area Under the ROC Curve (AUC) provides a single

measure of a classifier's performance for evaluating which model is better on average.

ROC curves can be thought of as representing the family of best decision boundaries for relative costs of TP and FP (Egan, 1975). On an ROC curve the X-axis represents %FP = FP/(TN + FP) and the Y-axis represents %TP = TP/(TP + FN) (see Fig. 3.3).

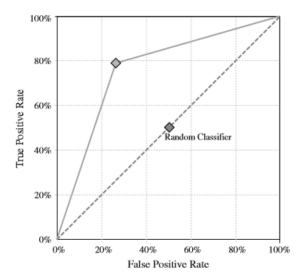


Fig. 3.3: Example of a ROC plot

Points in (0, 0) and (100, 100) are trivial classifiers where the predicted class is always negative and positive one, respectively. The ideal point on the ROC curve would be (0,100), that is all positive examples are classifled correctly and no negative examples are misclassifled as positive. The AUC measure is computed just by obtaining the area of the graphic.

Plotting ROC curves is not the only way to measure the accuracy of imbalanced data. Precision–Recall (PR) Curves are similar to ROC curves in that they explore the trade-off between the well-classifled positive examples and the number of misclassifled negative examples. Precision and recall are defined as follows:

$$Precision = TP/(TP + FP)$$

 $Recall = TP/(TP + FN)$

As the name suggests, PR curves plot the precision of the classifler as a function of its recall, as shown in Fig. 3.4.

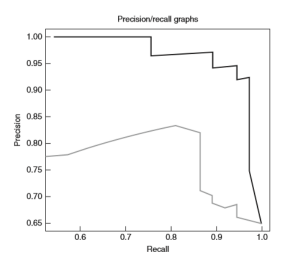


Fig. 3.4: Precision-Recall Curves illustration for two scoring classiflers

In other words, it measures the amount of precision that can be obtained as various degrees of recall are considered. For instance, in the domain of document retrieval systems, PR curves would plot the percentage of relevant documents identified as relevant against the percentage of relevant documents deemed as such with respect to all the documents in the sample. The curves, thus, look different from ROC curves as they have a negative slope. This is because precision decreases as recall increases.

The F-measure is another metric whose purpose is to combine the values of the precision and recall of a classifler on a given domain to a single scalar. More specifically, the F-measure is the weighted harmonic mean of precision and recall, and is formally defined as follows: For any $\alpha \in R$, $\alpha > 0$,

$$F_{\alpha} = \frac{(1+\alpha)[precision \times recall]}{[\alpha \times precision] + recall}$$

where α typically takes the values of 1, 2, or 0.5 to signify that precision and recall are equal in the case of $\alpha=1$, that recall weighs twice as much as precision when $\alpha=2$ and that precision weighs twice as much as recall when $\alpha=0.5$.

In the master's thesis, for the evaluation of different classifiers, ROC curves will be plotted as a single measure of the accuracy. In order to compare different classi-

fication models for imbalanced datasets, cross-validation (CV) is performed. This approach involves randomly m-fold CV dividing the set of observations into m groups, or folds, of approximately equal size (James and Hastie, 2015).

The first fold is treated as a validation set, and the method is fit on the remaining (m-1) folds. The AUC is then computed on the observations in the held-out fold. This procedure is repeated m times; each time, a different group of observations is treated as a validation set. This process results in m estimates of AUC: AUC_1 , AUC_2 ,..., AUC_m . The m-fold CV estimate is computed by averaging these values:

$$CV_{(m)} = \frac{1}{m} \sum_{i=1}^{m} AUC_i$$

In the work, 5-fold CV is performed in most cases.

3.5 Conclusion

Three classification models (k-NN, SVMs-RBF, Random Forest) used for imbalanced datasets in this work are presented. It is shown that some of these classifiers can be oriented towards the class imbalance issues. Many studies have documented the weakness of the most notable threshold metric, accuracy, in comparison to ranking methods and metrics (most notably ROC analysis/AUC) in the case of class imbalances.

Logistic Regression

Many classification methods, including Neural Network classifiers or Logistic Regression models, provide estimates of the a posteriori probabilities of the classes. Following Bayes' rule (Hastie and Tibshirani, 2009), the a posteriori probabilities depend in a nonlinear way on the a priori probabilities. Therefore, the knowledge of the 'true' a priori probabilities in the real-world data is often highly desirable since optimal Bayesian decision making is based on the a posteriori probabilities of the classes given the observation (we have to select the class label that has the maximum estimated a posteriori probability) (Saerens, 2001). Therefore, a change of the a priori probabilities may have an important impact on the a posteriori probabilities of membership, which themselves affect the classification rate. From the preceding point, this means that applying such a classifier as-is on new data having different a priori probabilities from the training set can result in a loss of classification accuracy, in comparison with an equivalent classifier that relies on the 'true' a priori probabilities of the new dataset.

4.1 Estimation of a Posteriori Probabilities

In the master's thesis, in order to directly estimate the a posteriori probabilities of n classes, a Logistic Regression model is used. This model indeed ensures the probabilities sum to one and remain in [0,1] range (Hastie and Tibshirani, 2009). It separates the data by hyperplanes: if the classes are linearly separable, it will separate them and if the classes are not linearly separable, it will find the best linear separation according to the maximum likelihood criterion.

Mathematically it gives:

$$\mathbf{P}(w_n|\mathbf{x}) \approx \hat{y}_n(\mathbf{x}) = \frac{exp(\mathbf{w}_n^T \mathbf{x}')}{\sum\limits_{j=1}^q exp(\mathbf{w}_j^T \mathbf{x}')}$$

where $\mathbf{x}' = [1, x_1, x_2, ..., x_p]^T$ is the vector \mathbf{x} of p features belonging to one of the n possible classes $w_1, w_2, ..., w_n$. Noted also that the membership values (\hat{y}_n) are

distributed according to a multinomial logistic distribution and, therefore,

$$0 \le \hat{y}_n(\mathbf{x}) \le 1$$
 and $\sum_{n=1}^q \hat{y}_n(\mathbf{x}) = 1$

In other words, the model takes the form

$$\log \left(\frac{\mathbf{P}(w_n|\mathbf{x})}{\mathbf{P}(w_{n'}|\mathbf{x})} \right) \approx (\mathbf{w}_n^T - \mathbf{w}_{n'}^T) \mathbf{x}'$$

That is, the log-odds of the posterior probabilities is linear in x.

4.2 Adjustment for the New a Priori Probabilities

In the master's thesis, an iterative procedure that estimates the new a priori probabilities of a new dataset and adjusts the outputs of a classifier (see (Saerens, 2001)) is evaluated on imbalanced datasets. The main goal is to show that this readjustment procedure can be beneficial (i.e. it increases classification accuracy) when a classifier has been trained on a training set that does not reflect the true a priori probabilities of the classes in real-world conditions. The output readjustment procedure can only be applied if the classifier supplies estimates of the a posteriori probabilities (e.g., Logistic Regression).

Let us suppose a classification problem in 2 classes with the class labels taking their value in $\Omega = (\omega_1, \omega_2)$. In order to train a classification model, we rely on a training set, i.e. a collection of observation vectors, \mathbf{x}_k , measured on individuals and allocated to one of the 2 classes $\in \Omega$.

For building this training set, we suppose that, for each class ω_i , observations on N_t^i individuals belonging to the class (with $\sum_{i=1}^n N_t^i = N_t$, the total number of training examples) have been independently recorded according to the within-class probability density $p(\mathbf{x}|\omega_i)$. The a priori probability of belonging to class ω_i in the training set will be denoted as $p_t(\omega_i)$ (subscript t will be used for estimates carried out on the basis of the training set) and is therefore estimated by the class frequency $\hat{p}_t(\omega_i) = \frac{N_t^i}{N_t}$. Let us now assume that the classification model has been trained, i.e. its parameters have been estimated on the basis of the training set (as indicated by subscript t). The classification model could be an artificial neural network, a Logistic

Regression, or any other model that provides as outputs estimates of the a posteriori probabilities of the classes given the observations. We therefore assume that the model has 2 outputs, $g_i(x)$ (i = 1, 2), providing estimated a posteriori probabilities of membership $\hat{p}_t(\omega_i|\mathbf{x}) = g_i(\mathbf{x})$ (i.e., probability of belonging to class ω_i , given that observation vector \mathbf{x} has been observed) in the conditions of the training set.

There will be considered two cases:

- 1. An estimate of the new a priori probabilities is known. In other words, the classifier will be applied on a large number of datasets for which we know the probability distribution of the priors.
- 2. No estimate of the new a priori probabilities is known. In this case, no class can be favored over any other class.

4.3 An Estimate of the New a Priori Probabilities Is Known

Let us now suppose that the trained classification model has to be applied to another dataset from which the class frequencies, estimating the a priori probabilities $p(\omega_i)$ are known. One should make the natural assumption that the generation of the observations within the classes, and thus each within-class density, does not change from the training set to the new dataset $(\hat{p}_t(\mathbf{x}|\omega_i) = \hat{p}(\mathbf{x}|\omega_i))$, only the number of measurements observed from each class has changed. The corrected a posteriori probabilities $\hat{p}(\omega_i|\mathbf{x})$ can be calculated according to the formula (Saerens, 2001):

$$\hat{p}(\omega_i \mid \mathbf{x}) = \frac{\frac{\hat{p}(\omega_i)}{\hat{p}_t(\omega_i)} \hat{p}_t(\omega_i | \mathbf{x})}{\sum_{j=1}^n \frac{\hat{p}(\omega_j)}{\hat{p}_t(\omega_j)} \hat{p}_t(\omega_j | \mathbf{x})}$$

4.4 No Estimate of the New a Priori Probabilities Is Known

In many real-world situations, we ignore what the real-world a priori probabilities $p(\omega_i)$ are since we do not know the class labels for these new data (Saerens, 2001).

However, there is an iterative procedure for a priori and a posteriori probabilities adjustment which is based on the expectation-maximization (EM) algorithm (Dempster and Laird, 1977). Let us define $g_i(\mathbf{x}_k)$ as the model's output value corresponding to class ω_i for observation \mathbf{x}_k from the new data set to be scored (Saerens, 2001). The model's outputs provide an approximation of the a posteriori probabilities of the classes given the observation in the conditions of the training set (subscript t), while the a priori probabilities are estimated by the class frequencies:

$$\hat{p}_t(\omega_i \mid \mathbf{x}_k) = g_i(\mathbf{x}_k)$$
$$\hat{p}_t(\omega_i) = \frac{N_t^i}{N_t}$$

Let us define as $\hat{p}^{(s)}(\omega_i)$ and $\hat{p}^{(s)}(\omega_i|\mathbf{x}_k)$ the estimates of the new a priori and a posteriori probabilities at step s of the iterative procedure. If the $\hat{p}^{(s)}(\omega_i)$ probabilities are initialized by the frequencies of the classes in the training set, the EM algorithm provides the following iterative steps (Saerens, 2001), for each new observation \mathbf{x}_k and each class ω_i :

$$\hat{p}^{(0)}(\omega_i) = \hat{p}_t(\omega_i)$$

$$\hat{p}^{(s)}(\omega_i \mid \mathbf{x}_k) = \frac{\frac{\hat{p}^{(s)}(\omega_i)}{\hat{p}_t(\omega_i)} \hat{p}_t(\omega_i | \mathbf{x}_k)}{\sum_{j=1}^n \frac{\hat{p}^{(s)}(\omega_j)}{\hat{p}_t(\omega_j)} \hat{p}_t(\omega_j | \mathbf{x}_k)}$$

$$\hat{p}^{(s+1)}(\omega_i) = \frac{1}{N} \sum_{k=1}^N \hat{p}^{(s)}(\omega_i \mid \mathbf{x}_k)$$

At each iteration step s, both the a posteriori $(\hat{p}^{(s)}(\omega_i|\mathbf{x}_k))$ and the a priori probabilities $(\hat{p}^{(s)}(\omega_i))$ are re-estimated sequentially for each observation \mathbf{x}_k and each class ω_i . The iterative procedure proceeds until the convergence of the estimated probabilities, ω_i . The condition for convergence is as follows:

$$\sum_{i=1}^{n} | \hat{p}^{(s+1)}(\omega_i) - \hat{p}^{(s)}(\omega_i) | < 0.0001$$

4.5 Conclusion

A special focus is given to the investigation of some ways of dealing with imbalanced datasets based on a Logistic Regression. An iterative procedure that estimates the new a priori probabilities of a new dataset and adjusts the outputs of a classifier is

proposed for evaluation on imbalanced datasets. The procedure is an instance of the EM algorithm and is applied here to the outputs of a Logistic Regression to improve classification accuracy.

Results

In order to achieve well-founded conclusions regarding performance of classifiers on imbalanced datasets, let us make use of four baseline classification models based on the following paradigms: k-NN as an instance-based learning approach, SVMs that enlarges the feature space using kernels, Random Forest that is an extension of a standard decision tree technique, Logistic Regression that gives as an output the estimates of a posteriori probabilities. The analysis is structured in the following way: first, the experimental framework is introduced, that is, the classification algorithms, their parameters and the selected datasets for the study; next, a separate study for preprocessing techniques used is developed.

The parameters of baseline classifiers used in all the experiments are presented below:

- 1. k-NN: in this case, the number of neighbors k for determining the output class is selected by CV (the euclidean distance metric is applied).
- 2. SVMs: RBFs with the cost parameter (c) and the kernel spread function (γ) are chosen by CV.
- 3. Random Forest: the parameter ntree (number of decision trees) is set equal to 100.
- 4. Logistic Regression: it contains no parameters.

Four imbalanced datasets have been gathered, whose features are summarized in Table 5.1, namely the number of observations (#Obs.), number of variables (#Var.) and IR. Estimates of the AUC metric were obtained by means of a 5-fold CV. That is, the dataset was split into 5 folds, each one containing 20% of the patterns of the dataset. For each fold, the alghoritm was trained with the examples contained in the remaining folds and then tested with the current fold. This value is set up with the aim of having enough positive class instances in the different folds, hence avoiding additional problems in the data distribution, especially for highly imbal-

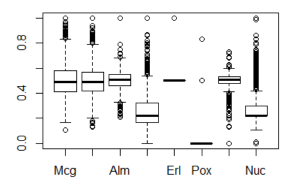


Fig. 5.1: Boxplot of the features of yeast datasets

anced datasets. The datasets are available for download at the KEEL (Knowledge Extraction Evolutionary Learning) dataset repository (KEEL dataset repository, 2017) that include different binary class datasets with imbalance ratio 1.5-9, higher than 9, and multiple class imbalanced datasets.

Table 5.1 presents *yeast* datasets with the same number of observations and variables but different IR.

Tab. 5.1: Summary of imbalanced datasets used (KEEL dataset repository)

Name	#Obs.	#Var.	IR
yeast3	1484	8	8.1
yeast4	1484	8	28.41
yeast5	1484	8	32.78
yeast6	1484	8	39.15

In total, eight features Mcg, Gvh, Alm, Mit, Erl, Pox, Vac, Nuc were used for classification. They are displayed as a boxplot in Fig. 5.1.

Before applying any of resampling techniques, the standardization of the data was performed. It means the data was scaled and centered to have mean zero. The result of this operation is shown in Fig. 5.2.

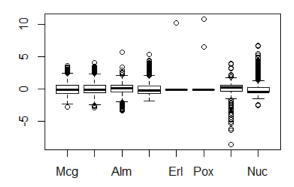


Fig. 5.2: Boxplot of the features of yeast datasets after scaling and centering

5.1 Study on the Preprocessing Methods

In this section, the behavior of the preprocessing methods on imbalanced datasets is analyzed. The representative set of methods is composed by the following techniques:

- 1. Oversampling;
- 2. Undersampling;
- 3. Both oversampling & undersampling;
- 4. SMOTE.

All those methods can be implemented in R using the parameters presented in Table 5.2). The data generated from oversampling have expected amount of repeated observations. Data generated from undersampling is deprived of important information from the original data. To encounter these issues, SMOTE helps us to generate extra positive observations synthetically as well.

Tab. 5.2: Parameters of the preprocessing methods

Name	Undersampling	Oversampling	Both	SMOTE
yeast3	N=326	N = 2642	N=1000, p=0.5	perc.over=100, perc.under=200
yeast4	N=102	N=2866	N=1000, p=0.5	perc.over=100, perc.under=200
yeast5	N=88	N = 2880	N=1000, p=0.5	perc.over=100, perc.under=200
yeast6	N=70	N=2898	N=1000, p=0.5	perc.over=100, perc.under=200

N refers to the number of observations in the resulting balanced set. For instance, for yeast3 dataset, originally there were 163 negative observations. Therefore, with total number N equal to 326, in the resulting balanced dataset (after undersampling) there will be 163 negative and 163 positive instances. The parameter p refers to the probability of positive class in newly generated sample. We set perc.over = 100 to double the quantity of positive cases, and set perc.under = 200 to keep half of what was created as negative cases.

Imbalanced Classification in R

In R, packages such as DMwR or ROSE perform sampling strategies quickly. ROSE (Random Over Sampling Examples) package provides a function named ovun.sample which enables oversampling, undersampling in one go. This package has well defined accuracy functions to do the tasks quickly. One have to set a desirable resampling method in the function ovun.sample (e.g., "under" for undersampling, "over" for oversampling, "both" if both undersampling and oversampling are used). If we do both undersampling and oversampling on the imbalanced data, the minority class is oversampled with replacement and majority class is undersampled without replacement.

Package DMwR allows implementation of SMOTE algorithm in R that generates data synthetically.

First, let us check the accuracy of the prediction using ROC curve without applying any sampling techniques. This will give us a clear picture, if the model is worth. Using the function roc.curve available in this package, for yeast4 dataset after running Random Forest" model we get AUC = 0.5677 which is a terribly low score. In this case, the algorithm gets biased toward the majority class and fails to map minority class. Therefore, it is necessary to balance data before applying a ML algorithm. We will use oversampling and undesampling techniques as well as SMOTE algorithm and balance datasets trying to improve this prediction accuracy. The results of using 4 techniques to balance yeast4 dataset are presented in Fig. 5.3 for Random Forest" model, in Fig. 5.4 for k-NN, in Fig. 5.5 for SVMs-RBF.

From these figures, it is clear that sampling techniques has improved dramatically the performance of all the analyzed classifiers.

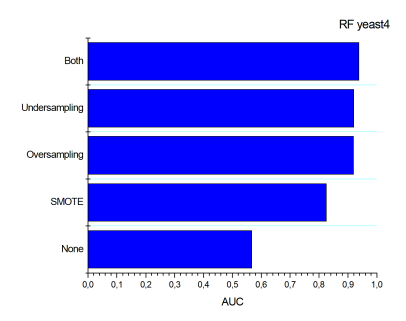


Fig. 5.3: Performance of Random Forest classifer after applying sampling techniques for yeast4 dataset

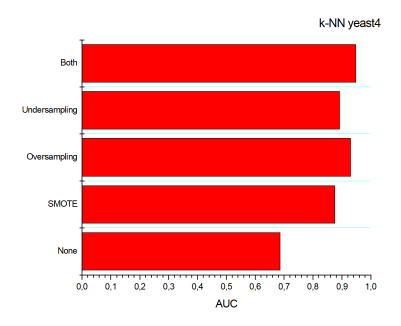


Fig. 5.4: Performance of k-NN classifer after applying sampling techniques for yeast4 dataset

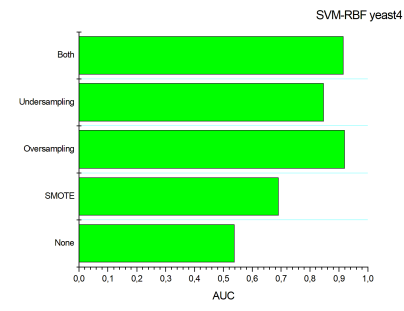


Fig. 5.5: Performance of SVMs-RBF classifer after applying sampling techniques for yeast4 dataset

5.2 Logistic Regression

For Logistic Regression, two cases should be considered:

- 1. An estimate of new priors is known.
- 2. No estimate of new priors is known.

The second case has never been studied for imbalanced datasets in the literature. Thus, this work provides the very first results on Log-EM model specifically for imbalanced datasets.

A full algorithm for classification in both cases consists of the following steps:

- 1. Preprocessing (e.g., oversampling); output: new priors.
- 2. Running Logistic Regression; output: a posteriori probabilities.
- 3. Adjustment of outputs (e.g., using iterative procedure if no estimate of new priors is known).

Let us demonstrate how these steps work on the example of yeast3 dataset.

Initially, the dataset includes 1484 observations with the following number of positive and negative instances:

negative/positive: 1321/163

Thus, initial priors for a chosen dataset are as follows:

negative/positive: 0.89/0.11

All 1484 observations have been split into 5 folds (for 5-fold CV) with the number and distribution of instances in each fold presented in Table 5.3:

Tab. 5.3: Number and distribution of instances in the folds

	Fold1	Fold2	Fold3	Fold4	Fold5
total	297	298	296	296	297
negative	264	265	264	264	264
positive	33	33	32	32	33

After applying a preprocessing technique (oversampling), the number of instances in the dataset has increased up to 2642 with the following numbers of positive and negative instances:

negative/positive: 1057/1585

Thus, the new priors are as follows:

negative/positive: 0.4/0.6

The following procedure will be shown on the example of observation #216. After running Logistic Regression it has a posteriori probability of belonging to a positive class equal to 0.8828.

Then, we can apply the following formula to calculate adjusted a posteriori probability:

$$\hat{p}(\omega_i \mid \mathbf{x}) = \frac{\frac{\hat{p}(\omega_i)}{\hat{p}_t(\omega_i)} \hat{p}_t(\omega_i | \mathbf{x})}{\sum_{j=1}^n \frac{\hat{p}(\omega_j)}{\hat{p}_t(\omega_j)} \hat{p}_t(\omega_j | \mathbf{x})}$$

Here, $\hat{p}(\omega_i \mid \mathbf{x})$ is an adjusted a posteriori probability of belonging to class ω_i , $\hat{p}(\omega_i)$ is initial a priori probability of belonging to class ω_i , $\hat{p}_t(\omega_i)$ is a new a priori probability (in our case, after oversampling), $\hat{p}_t(\omega_i \mid \mathbf{x})$ is a posteriori probability (in our case, the output of Logistic Regression prior to adjustment).

In our example, for the observation #216, $\hat{p}(\omega_1) = 0.89$, $\hat{p}(\omega_2) = 0.11$ ($\omega_1 = negative$, $\omega_2 = positive$), $\hat{p}_t(\omega_1) = 0.4$, $\hat{p}_t(\omega_2) = 0.6$, $\hat{p}_t(\omega_1 \mid \mathbf{x}) = 0.1172$, $\hat{p}_t(\omega_2 \mid \mathbf{x}) = 0.8828$.

The adjusted a posteriori probability $\hat{p}(\omega_i \mid \mathbf{x})$ for the observation #216 is now equal to 0.3857. It means that after adjustment this observation should be labeled as belonging to a negative class (0.3857 < 0.5).

Let us now assume we have no estimates of new priors. Then, the iterative procedure have to be performed to estimate new priors. It is useful to understand how many iterations would be sufficient to achieve convergence. The condition for convergence defined in Chapter 4 is as follows:

$$\sum_{i=1}^{n} | \hat{p}^{(s+1)}(\omega_i) - \hat{p}^{(s)}(\omega_i) | < 0.0001$$

For *yeast*3 dataset, Fig. 5.6 shows that convergence has been achieved after 6 iterations.

At the same time, for *yeast4* dataset, the convergence has been achieved after 16 iterations (Fig. 5.7).

The performance of Logistic Regression after using four techniques to balance yeast3 dataset is presented in Fig. 5.8, for yeast4 - in Fig. 5.9, for yeast5 - in Fig. 5.10, for yeast6 - in Fig. 5.11. LR denote Logistic Regression without adjustment for priors, TP means "True priors" (it corresponds to the case when priors are known), Log-EM corresponds to the case with not known priors. The results are also available in Tables 5.4 - 5.8.

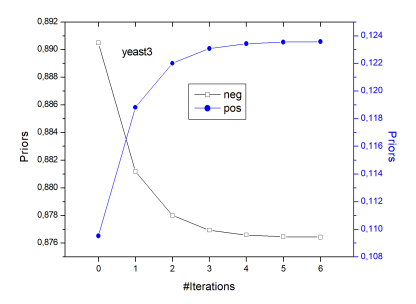


Fig. 5.6: New priors and number of iterations for yeast3 dataset

5.3 Results for All Considered Classification Models and Preprocessing Techniques

The results for yeast3 dataset for all considered classification models and preprocessing techniques are presented in Table 5.4. AUC(Test) means that the alghoritm was trained with the examples contained in the remaining folds and then tested with the current fold to which the accuracy measure AUC is applied.

Tab. 5.4: Results for *yeast*3 dataset (after 5-fold CV)

yeast3	Logistic Regression	True priors	Log-EM (priors are not known)	Random Forest	kNN	SVM-RBF
Preprocessing	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)
None	0.8286	0.9644	0.9644	0.8422	0.8458	0.8642
SMOTE	0.8935	0.967	0.967	0.9189	0.9126	0.9056
Oversampling	0.9112	0.9715	0.9715	0.9829	0.9692	0.9272
Undersampling	0.8974	0.9703	0.9703	0.9695	0.9344	0.9153
Both	0.9113	0.9648	0.9648	0.9724	0.9438	0.9148

SMOTE: perc.over = 100, perc.under = 200; Oversampling: N=2642; Undersampling: N=326; Both: N=1000, p=0.5

For *yeast*3 dataset, a ROC curve after running Logistic Regression with adjustment for not known priors is presented in Fig. 5.12. For *yeast*4, *yeast*5, *yeast*6 datasets ROC curves are similar.

Similar analysis was performed for *yeast*4 dataset (Table 5.5), *yeast*5 dataset (Table 5.6), *yeast*6 dataset (Table 5.7).

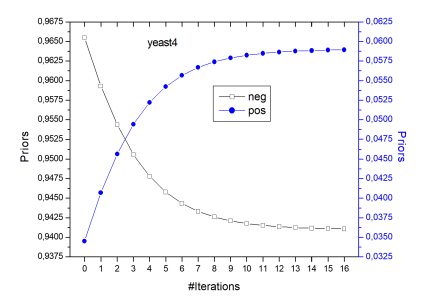


Fig. 5.7: New priors and number of iterations for yeast4 dataset

Tab. 5.5: Results for *yeast4* dataset (after5-fold CV)

yeast4	Logistic Regression	True priors	Log-EM (priors are not known)	Random Forest	kNN	SVM-RBF
Preprocessing	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)
None	0.5682	0.8669	0.8669	0.5677	0.6868	0.538
SMOTE	0.8144	0.867	0.8669	0.8257	0.8752	0.6909
Oversampling	0.8523	0.9013	0.9013	0.9193	0.9304	0.9193
Undersampling	0.8098	0.8941	0.8941	0.9199	0.8917	0.8468
Both	0.8427	0.8995	0.8994	0.9381	0.9479	0.9151

SMOTE: perc.over = 100, perc.under = 200; Oversampling: N=2866; Undersampling: N=102; Both: N=1000, p=0.5

Table 5.8 shows the average AUC results where the best average values per algorithm group are highlighted in bold. From this table, one may conclude the goodness of the use of this type of solution for imbalanced data, as there is a significant difference with respect to the results obtained with the original data.

5.4 Wilcoxon Signed-rank Test

In order to provide a comprehensive comparison of different classification models and preprocessing techniques, the Wilcoxon signed-rank test is performed. It is a

Tab. 5.6: Results for yeast5 dataset (after 5-fold CV)

yeast5	Logistic Regression	True priors	Log-EM (priors are not known)	Random Forest	kNN	SVM-RBF
Preprocessing	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)
None	0.7337	0.9871	0.987	0.7799	0.844	0.7424
SMOTE	0.9299	0.9795	0.9795	0.9434	0.9597	0.9545
Oversampling	0.9757	0.9894	0.9894	0.9868	0.9878	0.9319
Undersampling	0.95	0.9751	0.9751	0.9691	0.9712	0.9708
Both	0.9777	0.9883	0.9883	0.9826	0.9812	0.9316

SMOTE: perc.over = 100, perc.under = 200; Oversampling: N=2880; Undersampling: N=88; Both: N=1000, p=0.5

yeast3 Logistic Regression

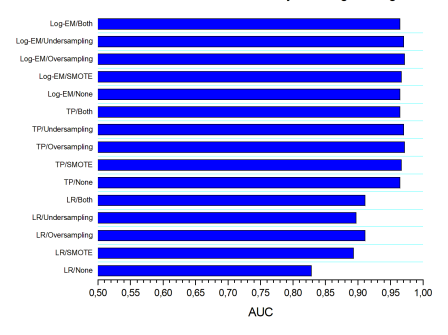


Fig. 5.8: Performance of Logistic Regression after applying sampling techniques and adjustment of priors for *yeast*3 dataset

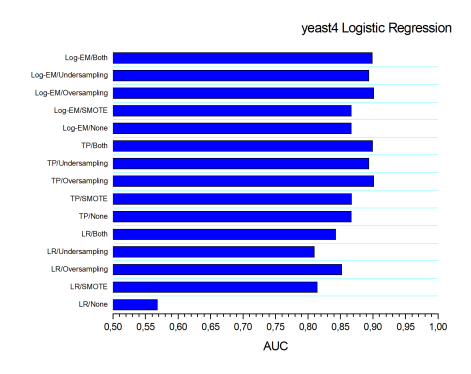


Fig. 5.9: Performance of Logistic Regression after applying sampling techniques and adjustment of priors for *yeast4* dataset

yeast5 Logistic Regression

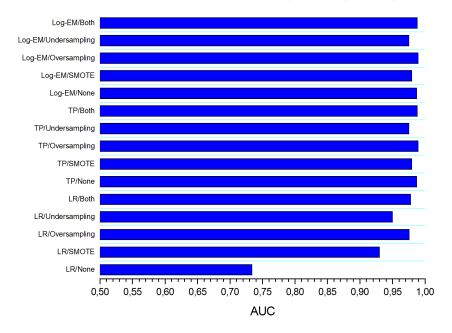


Fig. 5.10: Performance of Logistic Regression after applying sampling techniques and adjustment of priors for *yeast*5 dataset

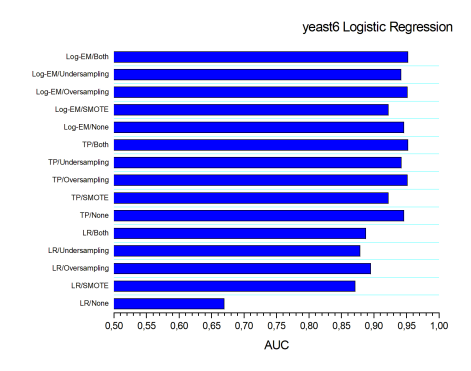


Fig. 5.11: Performance of Logistic Regression after applying sampling techniques and adjustment of priors for *yeast*6 dataset

Fig. 5.12: ROC curve for *yeast*3 dataset after running Logistic Regression with adjustment for not known priors

Tab. 5.7: Results for yeast6 dataset (after 5-fold CV)

yeast6	Logistic Regression	True priors	Log-EM (priors are not known)	Random Forest	kNN	SVM-RBF
Preprocessing	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)
None	0.6694	0.9461	0.9461	0.7126	0.7826	0.6547
SMOTE	0.871	0.9218	0.9218	0.8736	0.8812	0.8297
Oversampling	0.895	0.9513	0.9513	0.9547	0.9537	0.9279
Undersampling	0.8788	0.9419	0.9416	0.9276	0.9	0.876
Both	0.8875	0.9521	0.952	0.9509	0.9593	0.9265

SMOTE: perc.over = 100, perc.under = 200; Oversampling: N=2898; Undersampling: N=70; Both: N=1000, p=0.5

non-parametric statistical hypothesis test used when comparing two related samples, matched samples, or repeated measurements on a single sample to assess whether their population mean ranks differ (i.e. it is a paired difference test). It can be used to determine whether two dependent samples were selected from populations having the same distribution. (Lowry, 1998).

Let N be the sample size, i.e., the number of pairs. Thus, there are a total of 2N data points. For pairs i=1,...,N, let $x_{1,i}$ and $x_{2,i}$ denote the measurements. H_0 and H_1 denote the hypotheses tested.

 H_0 : difference between the pairs follows a symmetric distribution around zero. H_1 : difference between the pairs does not follow a symmetric distribution around zero.

An obtained value of Wilcoxon's test statistic W is calculated as follows:

$$W = \sum_{i=1}^{N} [sgn(x_{2,i} - x_{1,i}) \cdot R_i]$$

Tab. 5.8: Average AUC results for all studied datasets

yeast3-6	Logistic Regression	True priors	Log-EM (priors are not known)	Random Forest	kNN	SVM-RBF
Preprocessing	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)	AUC (Test)
None	0.7000 ± 0.1859	0.9411 ± 0.0887	0.9411 ± 0.0886	0.7256 ± 0.2000	0.7898 ± 0.1268	0.699825 ± 0.234085
SMOTE	0.8772 ± 0.0821	0.9338 ± 0.0865	0.9338 ± 0.0865	0.8904 ± 0.0881	0.9072 ± 0.0656	0.845175 ± 0.195124
Oversampling	0.9086 ± 0.0869	0.9534 ± 0.0646	0.9534 ± 0.0646	0.9609 ± 0.0530	0.9603 ± 0.0413	0.926575 ± 0.008951
Undersampling	0.88840 ± 0.0983	0.9454 ± 0.0631	0.9453 ± 0.0631	0.9465 ± 0.0450	0.9243 ± 0.0616	0.902225 ± 0.091057
Both	0.9048 ± 0.0956	0.9521 ± 0.0638	0.9511 ± 0.0639	0.9610 ± 0.0343	0.9581 ± 0.0285	0.922 ± 0.014264

 R_i denote the rank. W-value should be compared to the critical value in the table (see Appendix I) taking into account the number of subjects N. The obtained value is statistically significant if it is equal to or smaller than the value in the table.

For instance, suppose an obtained value is 22, and there were 15 participants. The critical value in the table is 25: the obtained value is smaller than this, and so one can conclude that the difference between the two conditions in the study was unlikely to occur by chance (p < 0.05 two-tailed test, or p < 0.025, one-tailed test).

For two methods (Log-EM without preprocessing (Method 1) vs. Log-EM with oversampling (Method 2) Table 5.9 shows the results of Wilcoxon signed-rank test. f1, f2, f3, f4, f5 denote CV folds, Abs is an absolute value of difference between the outcomes of Method 1 and Method 2, Sign shows if Method 1 is superior than Method 2 (Sign = 1) or not (Sign = -1).

Tab. 5.9: Results (Wilcoxon)

	Method1	Method2	Sign	Abs	R	SignR
yeast3 - f1	0.9776472	0.9799516	-1	0.0023	9	-9
yeast3 - f2	0.9327365	0.959596	-1	0.0269	18	-18
yeast3 - f3	0.9774372	0.9793529	-1	0.0019	7	-7
yeast3 - f4	0.9657746	0.9701493	-1	0.0044	12	-12
yeast3 - f5	0.9580189	0.960908	-1	0.0029	10	-10
yeast4 - f1	0.8947552	0.9374126	-1	0.0427	19	-19
yeast4 - f2	0.9229965	0.9304878	-1	0.0075	14	-14
yeast4 - f3	0.7857597	0.9203751	-1	0.1346	20	-20
yeast4 - f4	0.8482578	0.8489547	-1	0.0007	3	-3
yeast4 - f5	0.8829268	0.871777	1	0.0111	16	16
yeast5 - f1	0.9870756	0.9878472	-1	0.0008	4	-4
yeast5 - f2	0.9789738	0.9801312	-1	0.0012	5	-5
yeast5 - f3	0.9921875	0.9915365	1	0.0007	2	2
yeast5 - f4	0.9886188	0.9940201	-1	0.0054	13	-13
yeast5 - f5	0.9884259	0.9886188	-1	0.0002	1	-1
yeast6 - f1	0.8984182	0.9087988	-1	0.0104	15	-15
yeast6 - f2	0.9926108	0.9889163	1	0.0037	11	11
yeast6 - f3	0.9568966	0.958867	-1	0.002	8	-8
yeast6 - f4	0.9184729	0.9337438	-1	0.0153	17	-17
yeast6 - f5	0.9642857	0.9655172	-1	0.0012	6	-6

The W-value is 29. The critical value of W for N=20 at $p\leq 0.05$ is 52 (see Appendix I). Therefore, the result is significant at $p\leq 0.05$ and Method 2 therefore revealed a higher accuracy than Method 1.

Similarly, other hypotheses can be tested. For instance, the comparison of Random Forest (Method 1) and Log-EM (Method 2) has shown that Log-EM outperforms Random Forest (the corresponding W-value is 1) (both methods were compared without applying sampling techniques). However, when oversampling is applied, it is impossible to make a conclusion whether this method performs better than Random Forest with oversampling (W-value is 71 that is higher than critical value).

5.5 Dependence of Number of Iterations on Accuracy

In the paper (Gao and Sebastiani, 2016) the authors obtained the results showing that iterations beyond the first, which were carried out in order to obtain convergence, were actually detrimential, instead of benificial, to the accuracy. The authors do not mention which accuracy measure (AUC or classification accuracy based on confusion matrix) should be used. In order to theck this statement, the dependence of number of iterations on accuracy has been studied on yeast3 and yeast5 datasets. Both AUC and classification accuracy based on confusion matrix were calculated.

For yeast3, AUC revealed the same value (0.9644) for both 1 iteration and multiple iterations (i.e., until convergence has been obtained) whereas AUC value was a bit higher for yeast5 with 1 iteration (0.9871) than for the same dataset with miltiple iterations (0.9870).

In contrast, the classification accuracy (based on confusion matrix) was lower for yeast3 with 1 iteration (0.9474) than for yeast3 with multiple iterations (0.9488) when convergence was achieved and higher for yeast5 with 1 iteration (0.9771) than for yeast5 with multiple iterations (0.9744). Sampling techniques were not applied. Thus, the impact of number of iterations on accuracy is ambiguous and additional studies are required to better understand the dependence of given accuracy measure (e.g., AUC) against number of iterations.

5.6 Real-world Dataset

The dataset taken from Kaggle dataset repository (Kaggle dataset repository, 2016) contains transactions made by credit cards in September 2013 by European cardholders. The dataset has been collected and analysed during a research collaboration of Worldline and the Machine Learning Group (http://mlg.ulb.ac.be) of ULB (Université Libre de Bruxelles) on big data mining and fraud detection. This dataset presents transactions that occurred in two days, where 492 frauds out of 284,807 transactions were detected. The dataset is highly imbalanced, the positive class (frauds) account for 0.172% of all transactions.

It contains only numerical input variables which are the result of a principal component analysis (PCA) transformation (James and Hastie, 2015). Unfortunately, due to confidentiality issues, the authors cannot provide the original features and more background information about the data.

PCA is an unsupervised approach that refers to the process by which principal components are computed, and the subsequent use of these components in understanding the data. It finds a low-dimensional representation of a dataset that contains as much as possible of the variation. The idea is that each of the N observations lives in p-dimensional space, but not all of these dimensions are equally interesting.

Features V1, V2, ... V28 (see Fig. 5.13) are the principal components obtained with PCA, the only features which have not been transformed with PCA are 'Time' and 'Amount'. Feature 'Time' contains the seconds elapsed between each transaction and the first transaction in the dataset. The feature 'Amount' is the transaction amount. Feature 'Class' is the response variable and it takes value 1 in case of fraud and 0 otherwise.

The observations have been split into 3 folds (for 3-fold CV) with the number and distribution of instances in each fold presented in Table 5.10:

Tab. 5.10: Number and distribution of instances in the folds for 'Credit Card Fraud Detection' dataset

	Fold1	Fold2	Fold3
total	94936	94935	94936
negative	94772	94771	94772
positive	164	164	164

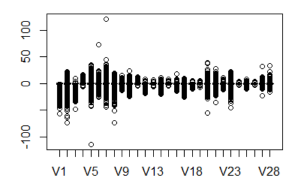


Fig. 5.13: PCA features of 'Credit Card Fraud Detection' dataset

Given the class imbalance ratio, the accuracy has been measured using AUC. Figure 5.14 shows the ROC curve after running Logistic Regression without adjustment for priors. AUC is equal to 0.8007 that demonstrates rather poor prediction accuracy of the algorithm.

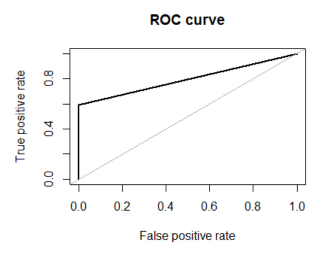


Fig. 5.14: ROC curve for 'Credit Card Fraud Detection' dataset after running Logistic Regression without adjustment for priors

Figure 5.15 shows the ROC curve after undersampling and running Logistic Regression with adjustment for not known priors (Log-EM). AUC is equal to 0.9861.

Obviously, confusion matrix accuracy is not meaningful for imbalanced classification but for reference one can show the confusion matrix for the model Log-EM/Undersampling (Table 5.11). It gives the accuracy of 0.9943.

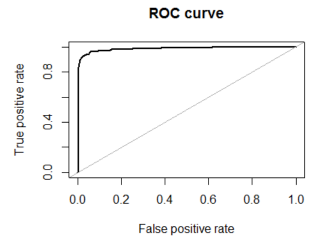


Fig. 5.15: ROC curve for 'Credit Card Fraud Detection' dataset after undersampling and running Logistic Regression with adjustment for not known priors (Log-EM)

Tab. 5.11: Confusion matrix for the model Log-EM/Undersampling

Log EM/Und	orcampling	Predicted classes		
Log-EM/Undersampling		0	1	
True classes	0	282785	1530	
	1	78	414	

The performance of classification models after applying four techniques to balance 'Credit Card Fraud Detection' dataset is presented in Fig. 5.16. The performance was calculated after 3-fold CV. Note that for Random Forest model the parameter ntree was set equal to 10 to reduce computational time. For the same reason, oversampling was not performed for Random Forest and k-NN models. Figure shows that for studied dataset the highest accuracy demonstrated Log-EM and TP models with different sampling techniques. The second best model is the Random Forest with undersampling.

It is important to note that for any dataset with large number of observations, CPU time required for the implementation of algorithm plays crucial role. The comparison of CPU time of processor Intel Core i3-4030U @ 1.90GHz for studied methods is presented in Fig.5.17. The combination of undersampling and Logistic Regression revealed the shortest CPU time that can be considered as an advantage of this approach. On the other hand, Logistic Regression with oversampling revealed the CPU time as long as $512.2\ s.$

In most cases, eight iterations are enough for the convergence of Log-EM algorithm (Fig. 5.18).

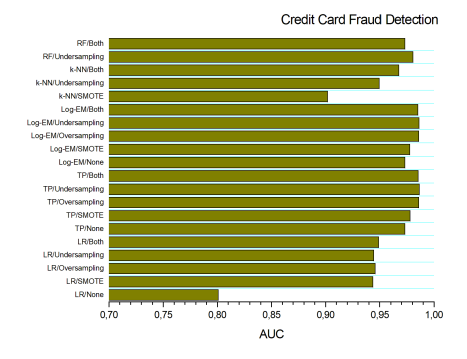


Fig. 5.16: Performance of classification models and sampling techniques

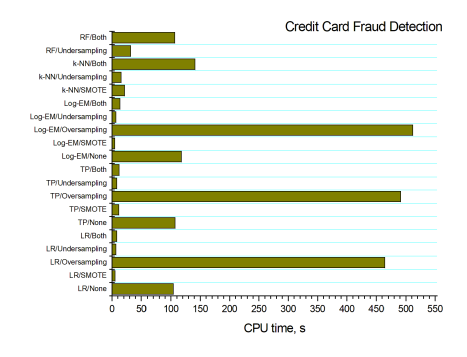


Fig. 5.17: CPU time

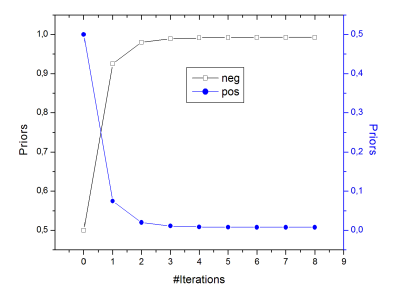


Fig. 5.18: Convergence of Log-EM algorithm

5.7 Conclusion

For all the studied *yeast* datasets, oversampling as well as the combination of overand undersampling techniques have revealed the highest accuracy for all classification models (average AUC reaches 0.9610 for both over- and undersampling techniques combined with Random Forest). Undersampling is less competitive when the number of instances is small (as for *yeast*3, *yeast*4, *yeast*5, *yeast*6 datasets). For a large number of instances (as for 'Credit Card Fraud Detection'), undersampling has revealed the highest accuracy in combination with all studied classification models (AUC reaches 0.98652 for undersampling together with Logistic Regression with known priors).

For all studied datasets, Logistic Regression demonstrates relatively poor accuracy compared to other classification models if adjustment procedure for priors is not applied.

Logistic Regression with adjustment of priors is competitive with such a sophisticated model as Random Forest. Note that for studied datasets there is only a slight difference in performance of Logistic Regression with known and unknown priors.

In terms of CPU time, for 'Credit Card Fraud Detection' dataset, oversampling has revealed the worst result (512.2 s) whereas undersampling and SMOTE performs an order of magnitude faster.

Conclusion

6.1 Main conclusions

In the master's thesis, imbalanced classification problems have been studied. Different classification models have been evaluated on *yeast* datasets and real-world dataset 'Credit Card Fraud Detection'. Namely, Random Forest, k-NN, and SVMs-RBF models have been analyzed as well as preprocessing methods such as oversampling, undersampling, SMOTE. In all the cases, the performance of classification models has been considerably improved after applying preprocessing techniques. More specifically, for all the studied *yeast* datasets and classification models, oversampling and the combination of both over- and undersampling techniques have revealed the highest accuracy. Undersampling seems to be less competitive when the number of instances is small (as for *yeast*3, *yeast*4, *yeast*5, *yeast*6 datasets) and more competitive for a large number of instances (as for 'Credit Card Fraud Detection' dataset).

A special focus has been given to the investigation of some ways of dealing with imbalanced datasets based on Logistic Regression. Different adjustment procedures for priors depending on whether the information on estimates of priors is available (TP model) or not (Log-EM model) have been studied. Note that Log-EM model has never been studied in the literature for imbalanced datasets. Thus, this work provides the very first results on Log-EM model specifically for imbalanced datasets.

One can conclude that Logistic Regression model has demonstrated relatively poor accuracy in comparison with other classification models when additional adjustment procedure for priors is not applied. However, with adjustment for priors it is competitive with more advanced models such as Random Forest.

In terms of CPU time, for 'Credit Card Fraud Detection' dataset, oversampling has revealed the worst result (512.2 s) whereas undersampling and SMOTE performs an order of magnitude faster.

6.2 Future Work

For the future work one may propose the following topics:

- 1. Additional study of adjustment procedure for the priors of Logistic Regression. For instance, it is important to clarify a question regarding the impact of number of iterations on overall performance of Logistic Regression. Furthermore, it is not completely clear the reason behind significant improvement of the performance of Logistic Regression when adjustment procedure is applied. One should study comprehensively the behavior of a bias term which might be responsible for that.
- 2. Study of some extensions of preprocessing methods. For instance, for SMOTE technique one should consider SMOTE+ENN (Batista and Prati, 2004), Bordeline-SMOTE (Border-SMOTE) (H.Han and Wang, 2005), Safe-Level SMOTE (SL-SMOTE) (Bunkhumpornpat and Sinapiromsaran, 2009). In all the cases a level of balance in the training data near to 50:50 distribution should be obtained.
- 3. Study of the cost-sensitive algorithms. It could be beneficial to carry out an analysis regarding cost-sensitive classifiers. Namely, "Weighted-Classifier" (CS-Weighted) (Barandela and Sanchez, 2003; Ting, 2002) and CostSensitive Classifier (CS-Classifier) from the Weka environment (Hall and Frank, 2009). In the first case, the base classifiers are modified usually by wieghting the instances of the dataset to take into account the a priori probabilities, according to the number of samples in each class. In the two latter casses, one can use an input cost-matrix.
- 4. Algorithmic modification can also be important for imbalanced datasets. There were several possible ways of performance improvement when dealing with Random Forest and SVMs-RBF but new techniques or modifications of well-known old techniques could be studied.

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${\bf Appendix}\; {\bf I}$

Table 1: Table of critical values for the Wilcoxon test

	Table 1: Table of critical values for the Wilcoxon test Two tailed significance levels				
N	0.05	0.01			
6	0	-			
7	2	-			
8	4	0			
9	6	2			
10	8	3			
11	11	5			
12	14	7			
13	17	10			
14	21	13			
15	25	16			
16	30	20			
17	35	23			
18	40	28			
19	46	32			
20	52	38			
21	59	43			
22	66	49			
23	73	55			
24	81	61			
25	89	68			

