STATISTICAL PROGRAMMING LANGUAGES

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Prediction Model Selection for Bank Telemarketing

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1 Introduction

Nowadays, gigantic amounts of data are available. Therefore, companies in almost all kinds of industries engage in the exploitation of data to obtain competitive advantages over rivals (Provost and Fawcett, 2013). The technological progress of computers and digitization of businesses makes corporations able to engage directly with customers, collecting and mining information about them, in order to tailor their products in a more optimal way (Rust et al., 2010). In particular, in the area of marketing the technological developments enable a rethinking of marketing strategies by analyzing available data and customer metrics (Moro et al., 2014). This is exactly what this paper will be aiming at. Optimizing the marketing efforts of the company using data mining.

In this paper, we focus on optimization models that increase company revenue by offering new products to existing customers. Models for data analysis are more suited for increasing revenue through existing customers than for acquiring new customers, because more information is available on the already existing customers (Nobibon et al., 2011). The knowledge of historical data allows the company to implement response models, to predict the probability that a customer will accept the offer of a certain product or service. In the retail banking industry, the banks have access to some of the richest datasets in the world of business (Nobibon et al., 2011). This should enable banks to target their marketing efforts towards customers that are more likely to accept the marketed products, which leads to lower marketing costs and increased profit.

In this paper, we study data related to a telemarketing campaign by a Portuguese Bank in the period of 2008-2010. Specifically we want to show that the bank can successfully classify their customers as accepters or decliners of an offer made through a telemarketing campaign. Several models can do such a classification task, we use the Logistic Regression, Decision Trees, Random Forest and Neural Network. The logistic regression and decision trees have the advantage that they make it easy to interpret which variables affect the classification probability. Random forest and neural network have the advantage that they can model highly complex nonlinear relations which tend to make these two models more accurate compared to the logistic regression and decision trees. However, due to the complexity of these models they are difficult to interpret and understand.

We fit all four models, and by using the AUC measure we conclude on which model does the most accurate predictions.

2 Theory and Design

2.1 Data pre-processing

2.1.1 Data Cleaning and Imputing

Multiple imputation(Rubin, 1987) is the method of choice for complex incomplete data problems. There are two methods for imputing multivariate data: one is joint modeling (JM) and the other one fully conditional specification (FCS)(van Buuren, 2007).

JM imputation(Sharfer, 1997) requires the specification of a parametric joint model $P\left(x^{mis}|x^{obs},\theta\right)$ for the complete data and a prior distribution $P(\theta)$ for parameter θ . Imputations are independent draws from the posterior predictive distribution of the missing data given the observed data $P\left(x|\theta\right)$, which under the ignorability assumption:

$$P(x^{mis}, x^{obs}) = \int P(x^{mis}|x^{obs}, \theta) p(\theta|x^{obs}) dx$$
 (1)

FCS (van Buuren and Groothuis-Oudshoorn, 2015) specifies the multivariate imputation model on a variable-by-variable basis by a set of conditional densities, one for each incomplete variable. Multivariate Imputation by Chained Equations (MICE) is on the basis of FCS:

$$\psi_{1}^{(t)} \sim p(\psi_{1}) p\left(x_{1}^{obs} | x_{2}^{(t-1)}, x_{3}^{(t-1)}, \dots, x_{R}^{(t-1)}, x_{R+1}, \dots, x_{K}, \psi_{1}\right)$$

$$x_{1}^{mis(t)} \sim p\left(x_{1}^{mis} | x_{2}^{(t-1)}, x_{3}^{(t-1)}, \dots, x_{R}^{(t-1)}, x_{R+1}, \dots, x_{K}, \psi_{1}^{(t)}\right)$$

$$\psi_{2}^{(t)} \sim p(\psi_{2}) p\left(x_{2}^{obs} | x_{1}^{(t)}, x_{3}^{(t-1)}, \dots, x_{R}^{(t-1)}, x_{R+1}, \dots, x_{K}, \psi_{2}\right)$$

$$x_{2}^{mis(t)} \sim p\left(x_{2}^{mis} | x_{1}^{(t)}, x_{3}^{(t-1)}, \dots, x_{R}^{(t-1)}, x_{R+1}, \dots, x_{K}, \psi_{2}^{(t)}\right)$$

$$\vdots$$

$$\psi_{R}^{(t)} \sim p(\psi_{R}) p\left(x_{R}^{obs} | x_{1}^{(t)}, x_{2}^{(t)}, \dots, x_{R-1}^{(t)}, x_{R+1}, \dots, x_{K}, \psi_{R}\right)$$

$$x_{R}^{mis(t)} \sim p\left(x_{R}^{mis} | x_{1}^{(t)}, x_{2}^{(t)}, \dots, x_{R-1}^{(t)}, x_{R+1}, \dots, x_{K}, \psi_{R}^{(t)}\right)$$

2.1.2 Imbalanced data resampling

A dataset is imbalanced if the classes are not approximately equally represented. Imbalanced data sets exists in many real-world domains, such as telecommunications management, detection of fraudulent telephone, detection of oil spills in satellite images and so on. In these domains, what we are really interested in is the minority class other than the majority class. Thus, we need a fairly high prediction for the minority class.

There are two main ways addressed the issue of class imbalance. One is cost sensitive learning, the other is to re-sample the original dataset. Here we introduce an approach, SMOTE(Synthetic Minority Class Oversampling), which blends under-sampling of the majority class with a special form of over-sampling the minority class.(Chawla, 2002)

- For each minority class x_i
 - Identify K nearest neighbors of minority class
 - Randomly select one of these neighbors x_{ij}
 - * Calculate feature vector difference
 - * Multiply with random number $\delta \in [0,1]$
 - * Add result to x_i
 - Generate a new point between x_i and x_{ij}

$$x_{new} = x_i + (x_i - x_{ij}) * \delta$$
 (3)

• For the majority class, under-sampled by randomly removing samples from the majority class population until the minority class becomes some specified percentage of the majority class.

2.2 Prediction Models

2.2.1 Logit Regression Model

General linear model (GLM) (McCullagh and Nelder, 1982) usually refers to conventional linear regression models for a continuous response variable given continuous and/or categorical predictors. In logistic regression, probability of the response taking a particular value is modeled based on combination of values taken by the predictors. The form is $y_i \sim N(x_i^T \beta, \alpha^2)$, where x_i contains known covariates and β contains the coefficients to be estimated. These models are fit by least squares and weighted least squares. In these

models, the response variable y_i is assumed to follow an exponential family distribution with mean μ_i , which is assumed to be some (often nonlinear) function of $x_i^T \beta$. Some would call these nonlinear because μ_i is often a nonlinear function of the covariates, but McCullagh and Nelder(McCullagh and Nelder, 1982) consider them to be linear, because the covariates affect the distribution of y_i only through the linear combination $x_i^T \beta$.

Binary Logistic Regression models how binary response variable Y depends on a set of k explanatory variables, $X = (X_1, X_2, \dots, X_k)$.

$$Logit(\pi) = log(\frac{\pi}{1-\pi}) = \beta_0 + \beta x_1 + \dots + \beta x_{k'}$$
 (4)

which models the log odds of probability of "success" as a function of explanatory variables. In (3), the distribution of Y is assumed to be Binomial (n,π) , where π is a probability of "success". X's are explanatory variables (can be continuous, discrete, or both) and are linear in the parameters, e.g., $\beta_0 + \beta_{x_i} \cdots + \beta_{x_k}$. Transformation of the X's themselves are allowed like in linear regression; this holds for any GLM. The Logit Link $(Logit(\pi) = log(\frac{\pi}{1-\pi}))$ models the log odds of the mean, and the mean here is π . Binary logistic regression models are also known as logit models when the predictors are all categorical (McCullagh and Nelder, 1982).

2.2.2 Decision Tree Model

Decision tree induction algorithms have long been popular in machine learning, statistics, and other disciplines for solving classification and related tasks. Decision Tree models where the target variable is a factor are called classification trees. In the classification tree structures, leaves represent class labels and branches represent conjunctions of features that lead to those class labels. Each node represent an splitting rule based on the value of one specific input variable, which is calculated by recursive partitioning. The recursion is completed when splitting no longer adds value to the predictions. This process of top-down induction of decision trees (TDIDT) (Quinlan, 1988) is an example of a greedy algorithm.

2.2.3 Random Forest Model

Random forest is a learning method, which is based on an ensemble of decision trees. It is mainly used for classification and regression problems. There are varied methods to grow random forests but the most popular one is the

method published by Breiman in 2001 (Breiman, 2001). It has a higher performance and better results in compare to other methods (Banfield et al., 2007).

The random forest method generates a lot of classifiers and aggregate the results. In the end, a weighted vote is taken for prediction. The decision trees are created on a bootstrap sample of the training data set. The method is like bootstrap aggregating (Bagging) (Breiman, 1996). The successive decision trees are not related to each other. They are based on a bootstrap sample of the data set. In the end, a simple majority vote is taken for prediction.

The construction of standard decision trees, is seperating each node and using the best split among all variables. Breimans method splits on each node using the best among a subset of predictors randomly chosen at that node. Breiman showed, that the random forest model is robust to overfitting. So it is not affected, if great number of trees are used for creating the forest (Breiman, 2001).

A general random forest can be implemented based on the following algorithm (see alg. 1).

```
Select the number of models to build, m

for i = 1 to m do

Generate a bootstrap sample of the original data

Train a tree model on this sample

for each split do

Randomly select k (< P) of the original predictors

Select the best predictor among the k predictors and partition the data

end

Use typical tree model stopping criteria to determine when a tree is complete (but do not prune)

end
```

Algorithm 1: Random Forest

Each generated decision tree give a prediction for a new sample. The random forest prediction results on the average of the m tree predictions.

2.2.4 Neural Network Model

Neural Network models are mathematical models that are, loosely put, based on how the biological brain works (Baesens et al., 2003). This allows for

highly complex nonlinear relationships between the input and the predicted variables.

A neural network is a system of neurons. Each individual neuron is simple. It receives an input, process it and generates an output. Though each neuron is by itself simple, a network of neurons can produce very complex and intelligent calculations (Shiffman et al., 2012). A neural network typically consists of three layers. An input layer, a hidden layer and an output layer. The need for only three layers is known as the universal approximation theorem and states that a neural network with three layers can approximate any continuous function to arbitrary degrees of accuracy (Hornik et al., 1989).

2.3 Model Performance Evaluation

A class can be assigned from a probabilistic outcome by assigning a threshold D, such that event c is true if $P(c|x_k) > D$. The receiver operating characteristic (ROC) curve shows the performance of a two class classifier across the range of possible threshold (D) values, plotting one minus the specificity versus the sensitivity(Fawcett, 2006). The overall accuracy is given by the area under the curve ($AUC = \int_0^1 ROC dD$), measuring the degree of discrimination that can be obtained from a given model. AUC is a classification index(Martens et al., 2011) that presents advantages of being independent of the class frequency or specific false positive/negative costs. The ideal method should present an AUC of 1.0, while an AUC of 0.5 denotes a random classifier.

3 Implementation

3.1 Data Pre-processing

3.1.1 Data Imputation

Codes for cleaning and imputing data consists of two parts: first one is used to detect the pattern of missing data, in order to choose the right algorithm to impute the incomplete data.

In line 1 above, the report used aggr() function provided by MICE package. The function yielded one histogram and one graph, which were named "Missing Data Ratio" and "Missing Data Pattern" respectively. In the graph, variables without missing value was marked with color navy blue, and incomplete variables were marked with color yellow. Variables were sorted according to the severity of missing.

```
tempData <- mice(data_missing,m=5,maxit=10,meth,seed=500,diagnostics=T)
data_complete <- complete(tempData,1)
```

In line 1 above, based on the pattern of missing value, the report imputes missing value. "tempData" contains complete datasets imputed by mice() function. "m" denotes how many sets of complete dataset imputed and "meth" denotes which method is used. "maxit" means how many iterations are needed, in which normally 10 is good enough. "diagnostics" is used to evaluate how good the imputation is. Codes in line 2 chooses the first dataset from m datasets created by line 1.

3.1.2 Data Balancing

Core part of the code for data balancing

```
library(unbalanced)
library(FNN)

# Calculate the SMOTE resamping scale parameters

sum = length(data_train$y)

ori_min = length(which(data_train$y == 1))

new_min = round(sum*(1/2)) - ori_min

over_s = (round(sum*(1/2)) / ori_min) - 1

un_s = (sum - all_min)/new_min

newdata = ubSMOTE(X,Y, k = 5,

perc.over = 100*over_s,

perc.under = 100*un_s,

verbose = FALSE)
```

Above codes shows how we implement the SMOTE algorithm using the package "unbalanced". From line 4 to 7, function ubSMOTE() is called. First we

specify the input variables X and response variable Y from the imbalanced dataset. The parameter K defines the numbers of nearst neighbors which are randomly choose. The scaling parameter per.over and perc.under control the amount of over-sampling of the minority class and under-sampling of the majority classes, respectively. For instance, if per.over = 200, per.under = 100, it means 200/100 = 2 new minority samples will be generated for each minority sample. And at the same time, 100/100 * 2 = 2 majority samples will be randomly chosen from the majority samples.

3.2 Prediction Models

3.2.1 Logit Regression Model

The core codes in this part divides into two part, one is settings for prediction model, the other one is prediction model:

The report uses caret package to train predict models. The codes in line 1 makes the settings for the train() function:method is cross validation. It means when training the model, the function will randomly choose one part of data as testing data, and test the model trained based on the train data. In this way, the model is improved. number is the number of folds in cross validation. If the chosen number is n, then n-1 folds are used to train the prediction model and the left one is to test whether the model is good enough. classProbs means whether class probabilities be computed for classification models (along with predicted values) in each resample. Since the purpose of the report is to predict whether a customer will or not buy a product, True is chosen in this report. Similarly, twoClassSummary is also chosen to compute performance metrics across resamples. verboseIter is also chosen for printing a training log.

In line 2, specific models are trained in train() function. The target variable is y, "." represents all other variables in the data as predictors. The method is the most important part of the codes. There are several options and the report uses "glm" (generalized linear model) and "nnet" (neural network). trControl is set in Line 1. metric is the method used for evaluate whether the model is good or not. tuneLegth denotes the amount of granularity in the tuning parameter grid.

3.2.2 Decision Tree Model

Core part of the code for building the decision tree

```
1 library(rpart)
  # pre-pruning the tree
  rpart.control= rpart.control(minsplit=5,
          minbucket = round(5/3),
          maxdepth=4,
          cp = 0.001)
  # build the model
  tree <- rpart(y~.,data = train,</pre>
                   method = "class"
10
                   parms=list(split="Gini/Information"),
11
                   control = rpart.control)
12
  # prediction
prediction <- predict(tree ,newdata = test, type = "prob")
```

We use the package rpart, which implement the popular non-parametric CART algorithm.

Parameters defined in line 3 is to pre-prune the tree in case the tree grows to large. We set three stopping criterions. The minsplit parameter is the smallest number of observations in the parent node that could be split further. The parameter minbucket provides the smallest number of observations that are allowed in a terminal node. And maxdepth parameter defined a certain depth of the tree from root to leafs.

From line 6 to 9, we build the decision tree model using function rpart(). As we are building a classification tree, so we choose the parameter method = "class". The parameter parms defines the splitting function. The splitting index can be "Gini" or "Information", which are two functions to calculate the Impurity of each node. Gini will tend to find the largest class, and Information tends to find groups of classes that make up 50% of the data.(detail calculation and equation in (Terry, 2015))

In line 12, the predict function will return the probability of each classes as we define the parameter type = "prob".

3.2.3 Random Forest Model

The "mlr" package is used to build the random forest model in R. The package offers a object oriented structure and is good for parallel computing. This is necessary because the larger the forest the more computational burden.

```
cvDesc <- makeResampleDesc(method = "CV", iters = 5, stratify = T)
rf.lrn = makeLearner("classif.h2o.randomForest",
```

Line 1 of the code defines the cross validation with 5 folds. In line 2 the learner is defined as a random forest classifier which can be built on the h2o instance. In line 6-7 the parameters for the amount of trees and number of splits in each decision tree are set.

Line 1 is creating a h2o instance using all the threads of the server (Intel Xeon E5-2670 @ 2.6 GHz). In this instance the the tuneParams function is running, to find the best parameters, which are defined in rf.PS. The last line is training the model with the optimal parameters.

3.2.4 Neural Network Model

The neural network can be trained in three different ways; supervised learning, unsupervised learning and reinforcement learning. Supervised learning can be used when one already has a dataset containing the answers to the question that we want to predict, and this is the training method that will be used in this paper.

The model attempts to predict outcomes, initiating this procedure with random weights. The errors of the predicted outcomes are then calculated and fed backwards through the model and used to update the weights. Through this process, the weights of the network are updated in such a way that the neurons recognise different patterns of the input space. This process continues until the weights have been optimally set or a maximum number of iterations have been reached. This process is called backward propagation. Backward propagation requires that the activation functions are differentiable. The standard approach is to use a sigmoid function such as a logit regression.

To train the neural network model in R, we make use of the 'caret' package. We first use the 'trainControl' function to specify the options of the neural network algorithm.:

```
model.control<- trainControl(
method = "cv", # 'cv' for cross validation
number = 5, # number of folds in cross validation
classProbs = TRUE, # Calculate class probabilities in each resample
summaryFunction = twoClassSummary, # Compute sensitivity, specificity and
AUC for each resample
returnData = FALSE # The training data will not be included in the ouput
training object
)</pre>
```

In line 2 we specify that we would like to train the model using cross validation, and in line 3 we specify how many folds we would like to use in the cross validation. By choosing five folds, we divide the data into 5 equally large subsets and leave one of them out. This is repeated for all five subsets. Line 4 and 5 specify that we would like to calculate class probabilities and sensitivity/specificity and AUC for all resamples. Line 6 specifies that we are not interested in saving the training data.

Next, we consider the parameters for the neural network.

```
nn.parms <- expand.grid(decay = c(0, 10^seq(-3, 0, 1)), size = seq(3,15,1)
```

To choose the optimal parameters for our neural network, we define a grid of parameters that we would like to test. As previously described, the neural network needs two parameters to be specified, namely weight decay and number of nodes in the hidden layer. To choose the optimal values of the parameters, we use the function 'expand.grid' to specify a grid within which to search for optimal parameters. We specify that we would like to search for a weight decay of (0,001; 0,01; 0,1; 1) and a number of hidden nodes between three and fifteen.

After having specified the options and the parameters we want to test, we initiate the training of the neural network.

```
1  nn <- train(y~., data = train,
2  method = "nnet",
3  maxit = 500,
4  trace = FALSE, # options for nnet function
5  tuneGrid = nn.parms, # parameters to be tested
6  metric = "ROC", trControl = model.control)</pre>
```

In line 1 we specify that the variable we want to predict is 'y' and that we want to use all variables as explanatory variables. In line 5 we give the grid that we want to search for optimal parameters. In line 6 we give the options that we specified previously.

3.3 Model Performance Evaluation

Codes in line 3 predict make the prediction with the model trained in line 2. type, containing "raw" and "prob", defines the output is class predictions or class probabilities. Line 3 outputs class prediction and line 5 class probability. Codes in line 4 compares this prediction with the real behavior of customers. Since the report hopes to predict customer who buys the product, "Yes" is chosen in line 4. In line 5 to 8, ROC curve is calculated based on the confusion matrix.

```
prediction1 <- predict.train(model,newdata)
confusionMatrix(prediction,data$y, positive="yes")
prediction2 <- predict(model, type = "prob")[,2]
roc<-roc(data$y,prediction2)
auc(roc)
plot.roc(roc)</pre>
```

4 Empirical Study/Testing

4.1 Data Input and Description

The report used dataset from http://archive.ics.uci.edu/ml/datasets/Bank+Marketing. The data is related with direct marketing campaigns of a Portuguese banking institution. The time period is May 2008-Nov 2010 and the dataset contains 41188 respondents. The marketing campaigns were based on phone calls, in order to access if the product (bank term deposit) would be ("yes") or not ("no") subscribed(the target variable is "y"). The predictor variables with are listed below in table 1:

Demographic Info	Marketing Info	Macroeconomy Info
age (numeric) job(categrical) marital(categorical) education(categorical) default(categrical) housing(categorical) loan(categorical)	contact(categorical) month(categorical) day.of.week(categorical) duration(numeric) campain(numeric) pdays(numeric) previous(numeric) poutcome(categorical)	emp.yar.rate(numeric) cons.price.idx(numeric) cons.conf.idx(numeric) euribor3m(numeric) nr.employed(numeric)

Table 1: Predictor Variables

Variables "default" and duration" are not included in the final analysis. There are only three respondents had default records, which led to exclusion of "default" due to very low variance (Kuhn and Johnson, 2016). Variable "duration" describes last contact duration. However, this attribute highly affects the output target (e.g., if duration=0 then y="no"). Yet, the duration is not known before a call is performed. Also, after the end of the call "y" is obviously known. Thus, this input is discarded if the intention is to have a realistic predictive model.

4.2 Data Pre-processing

4.2.1 Missing Value Pattern Overview

• Function

• Expected Output

The report expects an overview of missing value, both its ratio and pattern, based on which an appropriate imputation method is chosen. The aggr() function from MICE package will yield two graph, one is a histogram showing the ratio of missing value in every variable, the other one is a graph showing missing values according to observations, i.e., the pattern of missing data.

Actual Output

The Figure 1 indicates missing value pattern of this dataset. On the left side of Figure 1 is the ratio of missing value of each variable. Specifically, variable "education" has more than 4% respondents without educational information. Other variables with missing values are "housing", "loan", "job" and "marital". The missing value ratio of those five variables are all under 10%, which is acceptable in a statistical analysis perspective. On the right side of Figure 1 is missing value pattern from the perspective of observations: the bottom row of graph, with all cubic in color blue, represents 38245 cases with complete data. The row above, with one cubic in yellow and others in blue, represents 1558 observations with educational information missing and other information complete. For the same reason, the third row from the bottom indicates 946 observations with both housing and loan information missing. In a word, the cubic in yellow shows the how the data is missing.

• Comments

According to the pattern showed in Figure 1,the missing value is missing at random(MAR). Therefore, the report decides to impute the missing value with MICE package in the next section.

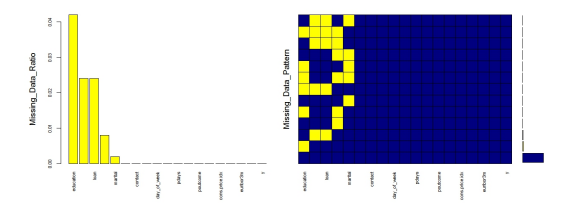


Figure 1: Missing Value Ratio and Pattern

4.2.2 Data Imputation

Function

```
1 # Data Preparation
  n <- nrow(bank)
  sample.size <- ceiling(n*0.8)</pre>
  idx.train <- sample(n, sample.size)</pre>
  bank_train <- bank[idx.train, ]
bank_test <- bank[-idx.train, ]</pre>
  # Data Imputing for Train and Test Dataset
  tempData1 <- mice(bank_train, m=5, maxit=10, meth="polyreg", seed=500,</pre>
       diagnostics=T)
  bankclean_train <- complete(tempData1,1)</pre>
  tempData2 <- mice(bank_test, m=5, maxit=10, meth="polyreg", seed=500,
10
       diagnostics=T)
  pred <- tempData2$predictorMatrix</pre>
  pred[,"y"] <- 0
  tempData3 <- mice(bank_test, pred=pred, pri=F)</pre>
bankclean_test <- complete(tempData3,1)</pre>
```

• Input Description

Because the purpose of the report is to make precise prediction of "y", so predictor variables cannot be affected by the target variable. However, the imputation process is in essence predictions made based on other available variables. Therefore, in test dataset, missing values are predicted only by other predictor variables, while in train dataset, it is allowed to be predicted by all variables including target variable. As a result, before imputing missing value, the report splits the whole dataset into train data set(80%) and test data set(20%). Line 1 to 6 achieved this goal. So far, the dataset "bank", which is originally input, is split into "bank-train" and "bank-test".

Then both incomplete datasets were completed with MICE package function in line 7 to 16. In line 10, there are five sets of predictions are created by mice function, and line 9 indicates the report picks the first set as the basis for further analysis. The same also goes for test dataset, with the exception that target variable "y" is excluded. In line 12 and 13, the report removes "y" as a predictor for missing values in test dataset. So far, both train and test data are free from missing value, and are renamed as "bankclean_train" and "bankclean_test".

• Expected Output

The report expects that all missing values of each variable are correctly and precisely predicted to the best extent.

• Actual Output

Train dataset and test dataset are free of missing value after imputation. Therefore, both of datasets are prepared for further analysis.

Comments

As showed in table 1, even though some variables (for example macro economy index) are in numeric format, they can also be regarded as categorical variables since each of factor contains less than ten levels. Therefore, the imputation process deals with all variables as categorical variables and chooses "polyreg" as imputing method. Unfortunately, "polyreg" makes it more difficult to evaluate the imputation quality, since existing functions provided by MICE is for numeric variables.

4.2.3 Data Balancing

• Function

• Expected Output

We will get the balanced training set with approximately 50% of positive response samples. And the prediction is assumed to be improved with the model trained by balanced data.

• Actual Output

As we can see in table 2, the dataset before balancing is quite skewed. However, after balancing, the ratio of "yes" and "no" is quite close to 50 to 50.

	Imbalanced Data	Balanced Data
No	29270	14223
Yes	3681	14724

Table 2: Data balancing comparison

• Comments

The balancing technique is only used on train dataset rather than test dataset. Balancing helps to improve the accuracy of prediction models, while prediction on test dataset should sticks to the original data structure.

4.3 Building Prediction Models

4.3.1 Logit Regression Model

Function

```
predict.logit.test1<-predict.train(logit,bank.test.dummy)
confusionMatrix(predict.logit.test1, bank.test.class, positive="yes")
predict.logit.test2 <- predict(logit, bank.test.dummy, type="prob")[,2]
logit.test.roc<-roc(bank.test.dummy$y,predict.logit.test2)
auc(logit.test.roc)
plot.roc(logit.test.roc)</pre>
```

• Input Description

The logit regression model requires all factor variables, i.e., categorical variables, need to be changed into dummy variables. As the function above shows, all factor variables are changed by dummyVars function. As a result, there are 55 variables in the new dataset "bank.train.dummy". The test dataset is also transformed with the same code.

To implement logit regression modeling, the report adopts train function provided by caret package in R, then specifies the method as "glm" and family as "binomial" ("logit"). Cross validation is chosen to improve model accuracy. Meanwhile, the dataset is requited to be standardized before logistic analysis, which is also achieved within the train function. After the model is derived, it is evaluated by comparing its prediction with real customer behavior.

• Expected Output

A binary logit analysis is expected to yield the relationship between each predictor variable and the target variable, considering other predictor variables hold still. Additionally, comparing predicted result with real result, the report also expect to evaluate how good the model is. An AUC index and an ROC curve are also expected to evaluate how well the logistic model works with future data.

Actual Output

The equation from binary logistic regression model is showed below in equation (4). Comparing coefficients of each predictor, it is obvious that macro economy situation is significantly important issue when customers considering purchasing bank product. Customer demographic information is relatively less important (most coefficients are less than 0.1). The marketing strategies of bank is of importance. Specifically, the marketing conducted via cellphone is more useful than cellular, and Wednesday is the best time to make marketing calls. Customers who rejected in last marketing campaign

is more likely to reject again.

```
y = 0.21 - 0.06*age - 0.10*job.bluecollar + 0.05*job.retired + 0.07*job.student \\ -0.16*marital.married - 0.04*education.basic4y - 0.03*education.basic9y \\ -0.04*education.highschool - 0.05*education.professionalcours \\ +0.08*housing.no - 0.52*loan.no - 0.10*contact.cellular + 0.10*month.aug \\ +0.18*month.jul + 0.21*month.mar - 0.31*month.may - 0.09*month.nov (5) \\ +0.06*month.oct - 0.05*day\_of\_weekfri - 0.12*day\_of\_weekmon \\ -0.04*day\_of\_weekthu - 0.04*day_of_weektue - 0.10*campaign + 0.34*pdays \\ -0.16*poutcome.nonexistent - 1.36*emp.var.rate + 0.48*cons.price.idx \\ +0.07*cons.conf.idx + 0.70*euribor3m + -0.65*nr.employed
```

How good is the derived model? The matrix in table 2 compares the prediction made by the model with real customer behavior: in train dataset, of 14724 customers who actually deposited, the model predicts 10605 ones correctly, while it predicts 4119 are potential customers while the fact is opposite. Meanwhile, the model predicts 2624 customers will buy the product while they actually did not. In test dataset, the model predict 6506 cases correctly (624 customers who says yes, and 5882 customers says no). Based on the confusion matrix, ROC curve in Figure 2 is created to evaluate the model visually.

	Reference(Traindata/Testdata)	
Prediction	No	Yes
No	11599/5882	4119/335
Yes	2624/1396	10605/624

Table 3: Confusion Matrix of Logit Model

• Comments

An index of AUC can be calculated from ROC curve. The AUC of prediction on train dataset equals to 0.8286 and on test dataset is 0.7769. Figure 2 shows the ROC curve of model performance on test dataset.

4.3.2 Decision Tree Model

• Function

• Expected Output

Here we are building a classification tree as our response variable y is a factor. Since the decision tree can deal with both numerical and categorical variables, we can directly input the balanced training set. We choose to use "information" as splitting criteria, as most of the input variables are categorical variables. After the model is trained, we will test it based on our testing set. And evaluate the prediction performance by calculating the confusion matrix with 0.5 threshold.

As a result of model building, we will get an well-shaped binary tree decision algorithm, which is displayed as a flowchart like tree, where each split represent a "Yes/No" decision. For the prediction test, we will expect an accuracy than better 0,5.

• Actual Output

	Reference(specificity/sensitivity)		
Prediction	No	Yes	
No	6572(90%)	454	
Yes	706	505 (52.6%)	

Table 4: Confusion Matrix of Decision Tree Model

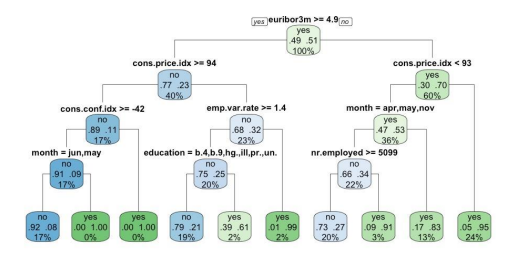


Figure 2: Decision Tree

• Comments

As we can see in Figure 2, the trained decision tree model grows pretty good. We can easily interpret the classification rules from it. For example, the variable at the root is social economic variable, euribor3m, which represent the most import explanatory variables for classification. We can say that, in our training model, if the euribor3m index is smaller than 4.9 and cons.price.idx is bigger than 93, which 24% of the samples fall into, the probability of a successful call is 0.95.

From the confusion matrix showed in Table 4, we can evaluate the prediction performance of the decision tree model. A generally accuracy of 85.9% is arrived, composed with 90% specificity and 52.6% sensitivity, which indicates that its more difficult to predict the positive result from decision tree model.

4.3.3

• Function

```
bank.train <- makeClassifTask(data = train,target="y", positive = "yes")
    #Creating a Classification Task based on the balanced train dataset
bank.test <- makeClassifTask(data = bank_test,target="y", positive = "yes")
    #Creating a Classification Task based on the balanced train dataset
tune.ctrl <- makeTuneControlGrid(tune.threshold = TRUE)
cvDesc <- makeResampleDesc(method = "CV", iters = 5, stratify = T)
ff.lrn = makeLearner("classif.h2o.randomForest",
    predict.type = "prob",</pre>
```

```
fix.factors.prediction = TRUE)

rf.PS = makeParamSet(
    makeDiscreteParam("ntrees", values = seq(500,1000,500)),
    makeDiscreteParam("mtries", values = seq(1,10,1))

h2o.init(nthreads = -1)

rf.para <- tuneParams(rf.lrn, task = bank.train, resampling = cvDesc, par. set = rf.PS, control = tune.ctrl, measures = auc)

h2o.shutdown()

library(kernlab)

lrn = setHyperPars(rf.lrn, par.vals = rf.para$x)

rf.mod <- mlr::train(lrn,task = bank.train)

yhat.rf <- predict(rf.mod, task = bank.train)

predict.rf <- predict(rf.mod, task = bank.test)</pre>
```

• Input Description

The random forest function is using all the independent variables to predict the dependent variable.

• Expected Output

The random forest model is in compare to the decision tree more robust to overfitting. The model is popular because of its precision. The prediction with more created trees should be more precise.

• Actual Output

The confusion matrix shows the results of the random forest predictions on train and test dataset (see table 5). The AUC of the prediction on train dataset is 0.999 and on test dataset 0.7946.

	Reference(Traindata/Testdata)		
Prediction	No	Yes	
No	14194/6731	29/547	
Yes	895/469	13829/490	

Table 5: Confusion matrix of random forest model

• Comment

The Random Forest could be more precise if the amount of created trees are increased. The problem is that the computation time is also accumulating accordingly while the accuracy is only improved to a relatively small extent.

4.3.4 Neural Network

• Function

```
1  nn <- train(y~., data = balancedTrain,
2  method = "nnet",
3  maxit = 500,
4  trace = FALSE, # options for nnet function
5  tuneGrid = nn.parms, # parameters to be tested
6  metric = "ROC", trControl = model.control)</pre>
```

• Input Description

The neural network use all variables as input variables. The categorical variables have to be formatted as factors variables, thus inputting a dummy for each category.

• Expected Output

As previously described, one of the disadvantages of a neural network model is that it is very difficult to interpret which variables influence the predicted probability. On the other hand we would expect that the model does well in terms of predictive accuracy. After having trained the model we can produce a ROC curve that shows the predictive performance in terms of specificity and sensitivity across all thresholds.

• Actual Output

After resampling across the grid of different parameters the best model in terms of achieved AUC is a model with thirteen nodes in the hidden layer and a weight decay of 1. This model achieves an AUC of approx. 0.7866. Setting a threshold of 0.50 gives the below confusion matrix.

	Reference	
Prediction	No	Yes
No	6546	432
Yes	732	527

Table 6: Confusion Matrix of Neural Network

5 Conclusion

In this paper, we have applied four different data analysis techniques on a recent dataset from a Portuguese bank, in order to predict customer behavior. A logit regression, a decision tree, a random forest and a neural network was applied. The goal was to predict whether customers would agree to open a fixed term deposit upon receiving a phonecall from the marketing department. The prediction was attempted based on customer characteristics and a few socioeconomic variables.

The four models were primarily compared by two metrics: the area under the receiver operating characteristic curve (AUC) and confusion tables. For both metrics the neural network had the best performance, which is in line with what we expected. The Random Forest achieved an AUC of 0.7946 which is very high performance. By looking at the logit regression and decision tree it is apparent that the socioeconomic variables are very important in explaining which customers accept and which do not. For example, the decision tree show that the two most important variables are the three month euribor rate (euribor3m) and the consumer price index (cons.price.idx). This suggests that it is the socioeconomic variables rather than individual specific characteristics that determine the behavior of the customers.

Future research should investigate the predictive power of the models across larger time periods, allowing for more variation in the socioeconomic variables.

Summing up, this paper demonstrates that data mining techniques are effectively able to assist companies in targetting the potential customers that are most inclined to accept a given offer.

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Appendix

 $Code: \ \verb|https://github.com/chinne/SPL_bank_marketing|.$