White Wine Quality Evaluation and Prediction

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Columbia University in the City of New York

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**Abstract**

This article deals with the design and creation of models for white wine quality evaluation and prediction. A large dataset is considered, with white wine *vinho verde* sample, which from Portugal. Regression models are built to predict wine quality based on physicochemical tests, and classification models are used to classify wine quality into several categories according to its score. The Random Forest model achieves the most promising result. Apart from these models, other kinds of models may be used to give new relationships between wine quality and the predictors.

Key words: Regression, Random forests, Classification models, Ridge, LASSO

**1.Introduction**

As one kind of luxury goods, white wine is increasingly enjoyed by a wide variety of consumers. To safeguard human health and to develop the technologies of wine making, white wine quality assessments and prediction become more and more important.

Wine certification is generally assessed by physicochemical and sensory test (Appalasamy *et al*., 2012). Based on physicochemical tests, the predictors for white wine include its density, alcohol or pH values. Meanwhile, human experts perform sensory tests such as taste preference (Cortez *et al*., 2009). There are three regression techniques, which are the Support Vector Machine (SVM), multiple regression, and neural networks from machine learning using by the Cortez and his teammates.

We analyze the regression relationships between eleven predictors based on physicochemical tests and wine quality. The physicochemical test use to characterize white wine include *fixed acidity, volatile acidity,* *citric acid*, *residual sugar*, *chlorides*, *free sulfur dioxide*, *total sulfur dioxide*, *density*, *pH*, *sulphates* and *alcohol.* Each sample in this dataset is first evaluated by a minimum of three sensory assessors (using blind tastes) who grade the wine according to a scale that ranges from 0 (very bad) to 10 (excellent) (Appalasamy *et al*., 2012).

For the data we use to assess the quality of white wine, the data is continuous, and it is from Wine Quality Data Set on the UCI Machine Learning Repository website. Moreover, it is collected from May, 2004 to February, 2007 using only protected designation of origin samples that were tested at the official certification entity.

Our main focus in analyzing the white wine data set is to find out which predictors have more influence on determines the quality of white wine and to find out the capable models of prediction quality of white wine.

**2.Materials**

For creation the models, the dataset we use included 11 input variables (based on physicochemical tests) and 1 output variable *Quality* (based on sensory data). The total number of observations is 4898. The *Quality* of white wines is evaluated using a scale from 0 to 10 by integers and all the input variables are numeric, which showed in Table1 in appendix.

**2.1 Input Data Summary**

We summarize the twelve variables in Table 2. Based on Table 2, we conclude that the minimum value of the output variable, *Quality*, is 3 and the maximum value of *Quality* is 9 instead of 0(very bad) and 10(excellent). The mean is greater than the median for all the input variables and Range is much larger compared to the IQR. This phenomenon indicates that there are outliers existing in the data set.

In addition, Figure 1 shows that *alcohol* and *volatile acidity* have a linear relationship with *Quality*, and the values of wine *Quality* is almost a constant with respect to *free sulfur dioxide*, *pH*, and *residual sugar*, which means that the *Quality* has nothing to do with these factors.

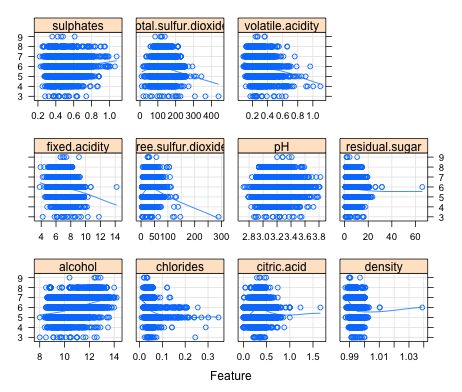


Figure 1 Scatter plots of each predictor versus the wine *Quality*

**2.2 Exploratory Data Analysis and Data Pre-processing**

For missing value checking, based on Table3 in appendix, which describes the twelve variables in summary, there are no missing values for all the variables.

For outlier checking, the box plots, showed in figure 2, indicates that all of the inputs variables except *alcohol* have outliers. *Residual sugar* has a positively skewed distribution and even after eliminating the outliers distribution will still remain skewed. Also we notice that mostly outliers are on the larger side.

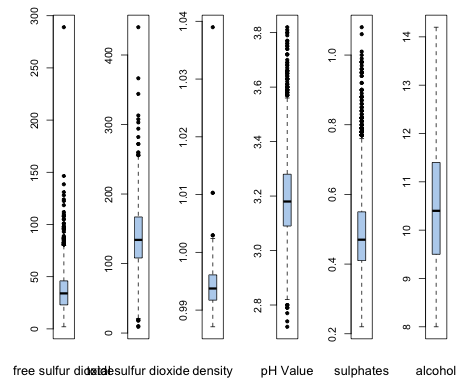
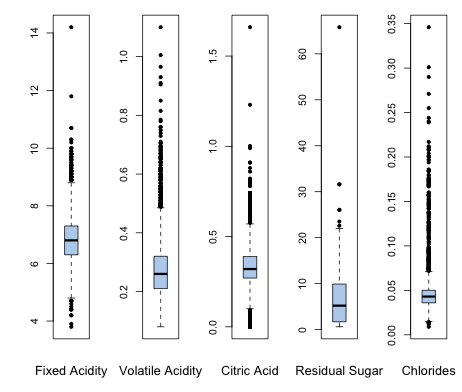


Figure 2. The boxplots of all input variables

For correlation checking, since the variables have non-normal distribution, we consider person correlations, which in Appendix Table 4, spearman rank correlations which in Appendix Table 5 and the correlation plot showed in figure 3 below. There are high correlations, which is greater than or equal to 40% in absolute value, existing. In addition based on figure 3, we observe that *density* and *residual sugar* has a strong positive relationship, *alcohol* and *density* has strong negative relationship.

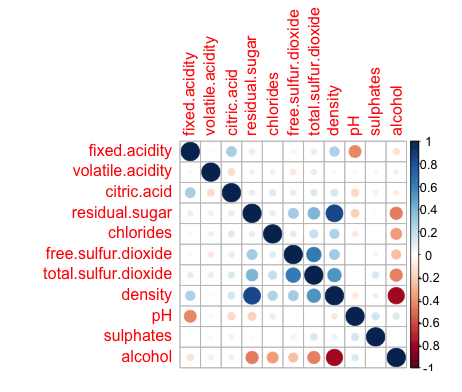


Figure 3. Correlation plot between predictors

**2.3 Data Preparation**

Since most of the input variables have large amounts of outliers and there are correlation existing, data in predictors need to be centered, scaled and transformed to prepare for model building. After transformation, new box plots shows that our data has a better distribution. Then, we split our data to training set and test set with proportion of 3:1. The training set is used to create the model and do the10-fold cross-validation to produce reasonable estimates for model building, and the test set is used to qualify performance.

**3. Methodology and Models**

To evaluate white wine quality based on our data set, a series of models are created and evaluated. The purpose of testing data in a variety of models is to select the best model to predict wine quality. In every model, tuning parameters (if had) are first estimated, and then are used to fit our dataset which after transformation. 10-fold cross-validation method also uses for resampling. Then, predictions are made. Final we obtain our result displayed by R2 for prediction and error rate to test the fitness of the models.

**3.1 Regressions Models**

First, we use linear regression models, including multiple linear regression, Ridge, LASSO and polynomial regression. R2 and error rate are generated for checking whether the model is appropriate for our white wine data set. R2 for prediction and error rate value are summarized in Table 9 below.

Since multiple linear regression is the common model built to understand the association between the input variables and the output variable in the first place and it is easier to interpret so we use multiple linear regression as our first model to study the white wine quality. The result displays in the appendix Table 6. We find that the input variables *citric acid*, *chlorides* and *total sulfur dioxide* are not significant. The R2 for the predictions is nearly 27%, which means the model does not match the condition of the data so well. The error rate for the predictions is nearly 48%, which means there is a lot of error happened for prediction of *Quality*.

Since the result of multiple regression shows that there are some predictors not significant, Ridge and LASSO are taken into consideration to do regularization. The result of Ridge displays in the appendix Table 7 and the result of LASSO marks in the appendix Table 8. From the result of Ridge, none of the predictor is dropped, and there is no improvement from multiple regressions. The *R*2 for the predictions is nearly 27%, and the error rate for the predictions is nearly 48%.

In the result of Lasso, the input predictors *fixed acidity, citric acid, total sulfur dioxide* and *density* are dropped. These parameters dropped are the correlated predictors shown in Table 4, Table 5 and figure 3. The *R*2 for the predictions is nearly 26%, and the error rate for the predictions is nearly 48%, which are still not good.

In order to investigate whether a polynomial relationship fits the model better, a polynomial regression model with squared terms of the significant variables is tried, which improves *R*2 value to nearly 31%, and the error rate for the predictions is nearly 48%. Although polynomial regression model can explain more about our data set, it still cannot be the good model we expect.

Since the result of Multiple Regression is not good in prediction, Ridge and Lasso regression methods do not make much improvement and polynomial regression cannot explain more than 50% of the data then we can conclude that linear regression is not that appropriate for this data set.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | Multiple Linear Regression | Ridge | LASSO | Polynomial Regression |
| Error rate | 0.4818449 | 0.4789009 | 0.484789 | 0.4759568 |
| R2 | 0.2737022 | 0.2784749 | 0.2625427 | 0.3066771 |

Table 9. R2 and error rate for Linear Regression Models

Second, we used non-linear regression models, including Random Forest, neural networks (NN) and Support Vector Machines (SVM). R2 for prediction and error rate are summarized in Table 10 below.

Since the Random Forest is one of the most accurate learning algorithms available, we try this model as our first non-linear regression model. Based on decision trees, the algorithm nature makes Random Forest suitable to regression and classification. We estimate the turning parameter with 10-fold cross-validation method, it turns out that the best turning parameter is 2. The error rate for random forest with regression prediction is almost 31.7% and R2 for prediction we obtain is nearly 52%.

The Random Forest model also provides a method to measure the importance of predictors in prediction. There are two measures to measure the variable importance, mean decrease in accuracy and mean decrease in node impurity. For each tree, the prediction accuracy on the out-of-bag portion of the data is recorded. Then the same is done after permuting each predictor variable. The difference between the two accuracies are then averaged over all trees, and normalized by the standard error. For regression, the MSE is computed on the out-of-bag data for each tree, and then the same compute after permuting a variable. The differences are averaged and normalized by the standard error. The second measure is the total decrease in node impurities from splitting on the variable, averaged over all trees. For classification, the node impurity is measured by the Gini index. For regression, it is measured by residual sum of squares. According to figure 4, in the mean decrease in accuracy measure, the input predictors *volatile acidity*, *alcohol* and *free sulfur dioxide* are the most important while *alcohol*, *density* and *free sulfur dioxide* are the most important predictors in mean decrease in node impurity measure. Therefore, we notice that the input predictors *alcohol*, *free sulfur dioxide* and *volatile acidity* have more influence to the white wine *Quality*.

Since Neural Networks are the powerful nonlinear regression techniques inspired by theories about how the brain works, we also use this model for white wine *Quality* analysis. The outcome is modeled by an intermediary set of unobserved variables called hidden units. These hidden units are linear combinations of the original predictors. However, neural networks have a tendency to over-fit the relationship between the predictors and the response due to the large number of regression coefficients. In this model, the final values used for the model are size = 3 and decay = 0.1. For prediction, the values we used are size=3, decay=0.1, and 5 hidden units. In addition, the error rate for NN we obtain is nearly 50% and the R2 value for prediction is 20%.

For support vector machine (SVM), since it is less sensitive to outliers, it is better than multiple linear regression. In addition, SVM is characterized by usage of kernels, absence of local minima, and capacity control obtained by acting on the margin, or on number of support vectors. The variables are transformed into a high m-dimensional feature space, by using a nonlinear mapping that does not need to be explicitly known but that depends of a kernel function. The result we got includes nearly 47% error rate and approximately 30% R2 value for prediction.

|  |  |  |  |
| --- | --- | --- | --- |
|  | Random Forest | NN | SVM |
| Error rate | 0.3169935 | 0.4996101 | 0.4698320 |
| R2 | 0.5221166 | 0.200876 | 0.2956326 |

Table 10. R2 and error rate for Non-linear Regression Models

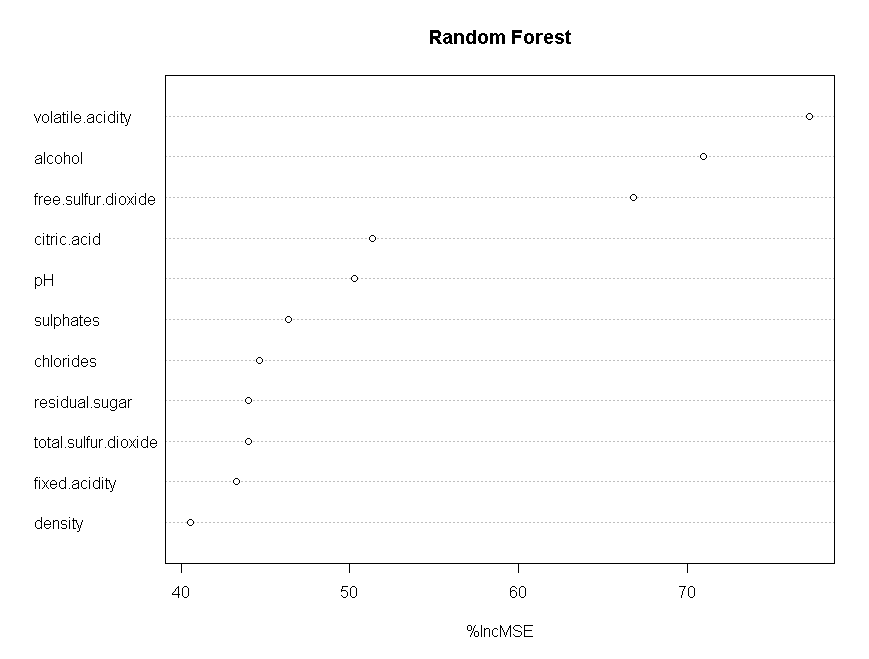
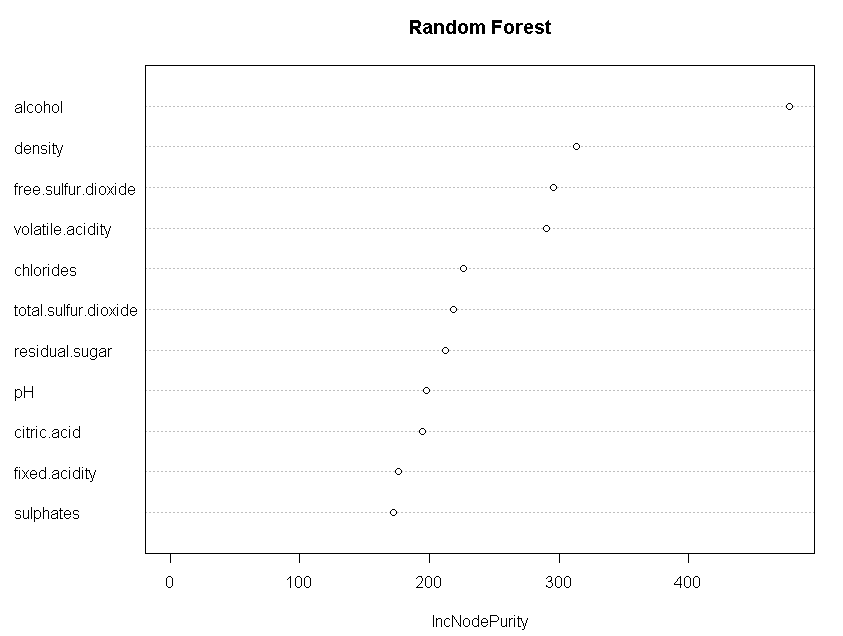


Figure 4. The importance of predictors in prediction based on Random Forest

**3.2 Classification Models**

Moreover, we built classification models to classify white wine *Quality* into 7 categories, from 3 to 9. We use Decision Tree, Random Forest, Support Vector machine (SVM) and Adaboost to classify. In each model, an error rate is generated. The error rate for each model summarize in Table 11 below.

Since the problem can be viewed as a multi-class classification problem, it is very natural to use tree-based method. The very first model using is just the decision tree. Decision tree is a supervised machine learning method. Although it is a greedy algorithm, it can still sometimes show off a very good result. A key advantage of tree-based method is that it is very easy to understand and it is very easy to interpret the result. In every step, the tree will choose both the feature and the cut point that could increase most the purity of notes. And the result is given by the most majority of predicts in this leaf. With this method, the tree will be trained to put the similar wines with similar wine score together and therefore useful in classification. For the white wine evaluation, we get the error rate of 48.65%. It is not so good so we used Random Forest.

The decision tree method only train one tree, but the ensemble method such as Random Forest which combined a lot of trees may get a better result. Random Forest is an algorithm, which is unexcelled in accuracy among current algorithms. Compare to the other classification methods we used for our white wine evaluation dataset, Random Forest performs very well. The error rate is just 31% for this multi-class classification problem, which is the best one among all the algorithms we have tried.

For Adaboost model, it is also an effective machine-learning algorithm that can be used to classification. The algorithm gives every observation a weight to measure how important it is in the classification. The higher the error rate is, the higher the weight of this observation is, so that the classifier will pay more attention to classify the observations that are easy to get wrong result. The result of Adaboost in this white wine quality dataset, which indicated by the error rate, is nearly 48%.

Basically support vector machine (SVM) is typically an algorithm to solve 0-1 classification problems. However, with the method of “one-against-one” or “one-against-all” method, we can also apply SVM into the multi-class classification problem. Here we take the radial bas function (RBF) kernel to solve the problem and the result is showed below.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | Decision Tree | Random Forest | Adaboost | SVM |
| Error rate | 0.4865702 | 0.3109504 | 0.4690873 | 0.3167331 |

Table 11. Error rate for Classification Models

**4. Results**

Compare Table 9 and Table 10, for regression models we can conclude that except Random Forest model, the other regression models are not good fits because the values of R2 for prediction are all below 0.5, which means all the regression models cannot explain half of the white wine data.

Based on Table 11, for classification models, Random Forest have the best performance compare to other classification models. Support Vector Machine also has a good performance.

Compare the result showed in Table 9, Table 10 and Table 11, the regression models and the classification models we observe that Random Forest model do the best job compare to all the other models. It works well in variance reduction and noise robustness. Also Random Forest has several advantages. Random Forest is one of the most accurate learning algorithms available. Random Forest gets its samples from the Bootstrap sample, which the variance between different trees is very large, and it is very good for the ensemble algorithms. Also, Random Forest runs efficiently on large databases and gives estimates of what variables are important in the classification. It can handle thousands of input variables without variable deletion. Moreover, Random Forest is able to generate forests, which can be saved for future use. Therefore, random forest model provides more stable predictions and model accuracy.

**5. Conclusions and Implications**

Based on figure 4, *alcohol*, *free sulfur dioxide* and *volatile acidity* have more influence on white wine *Quality* than other predictors.

Compare all the regression models built, the values of R2 for prediction for the total 7 regression models are not big enough, even Random Forest can only explain 52% of the data set, then regression models cannot fit the data very well. Compare all the classification models built, the SVM model performs much better than the other model except Random Forest. Random Forest is best model we have to cover all the conditions of our data. Finally compare all the models used, the error rate for random forest is the lowest, Random Forest can be a more capable model for predicting *Quality* of the white wine.

In this data set, the values of the response variable are integers mainly from 3 to 9, instead of all the real numbers in this range, so the regression lines between the outcome and predictors are not perfect. In this sense, more adjustments should be applied to get better fits, and other models should be used to fit the data to gain a relatively perfect prediction.

**Appendix**

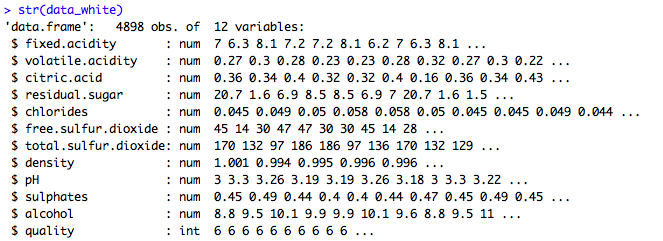


Table1. The structure of white wine data set

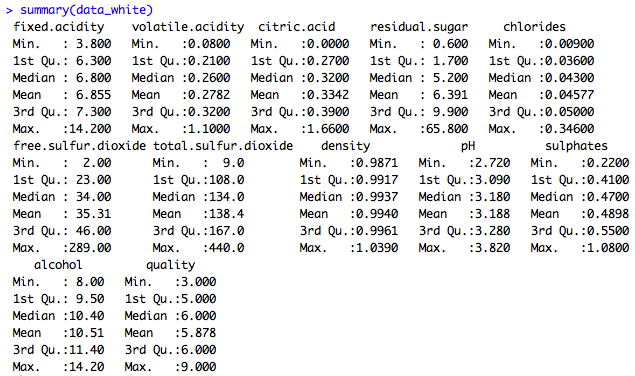
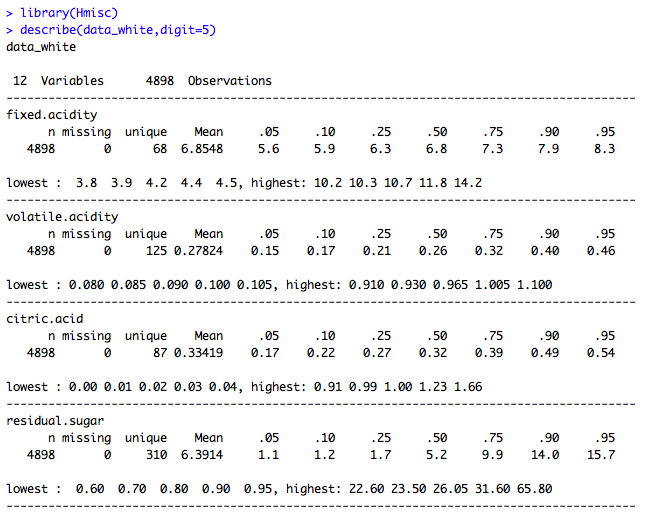
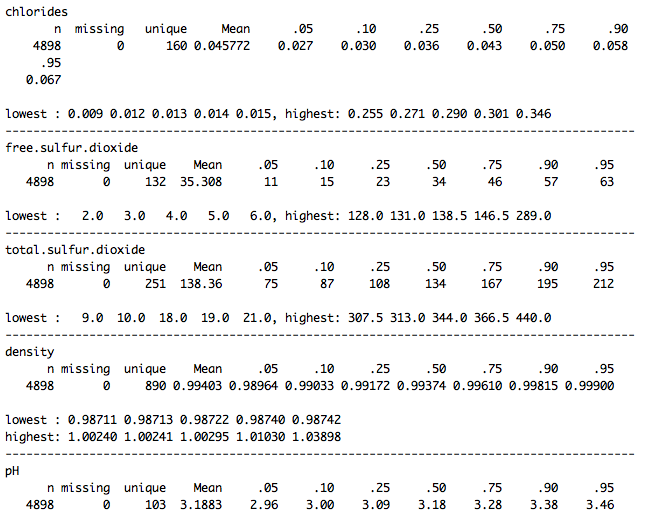


Table 2. The summary of white wine data set

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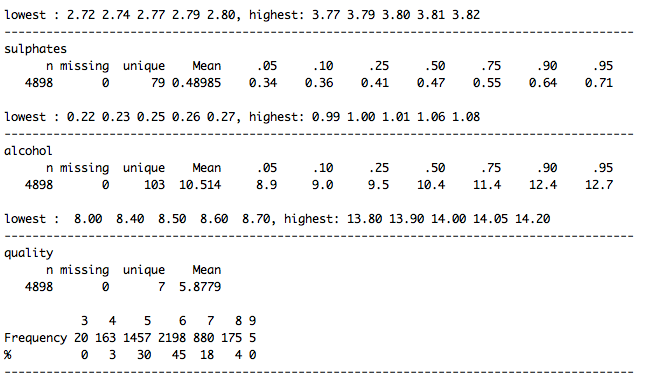
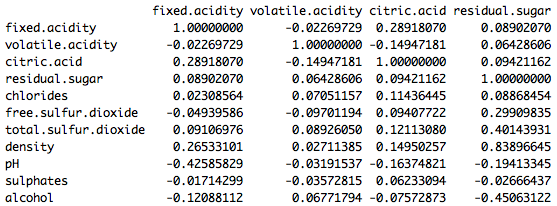
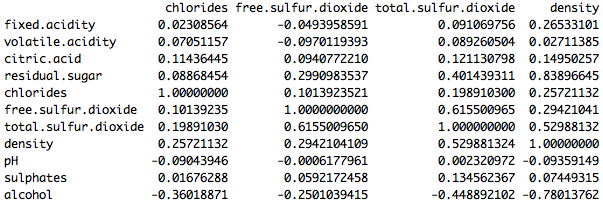
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Table 3. Describe the white wine data set





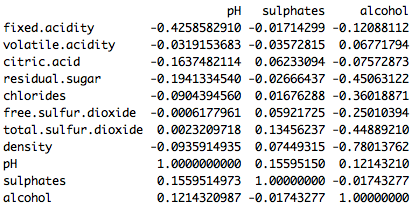
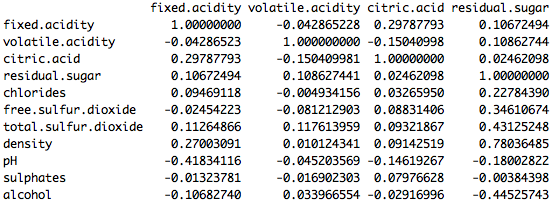
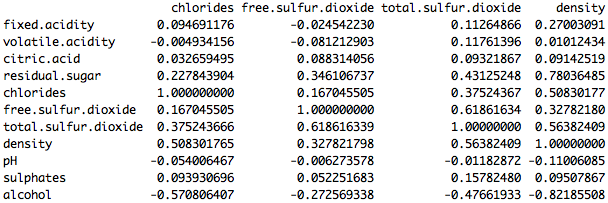


Table 4. Pearson’s Correlation





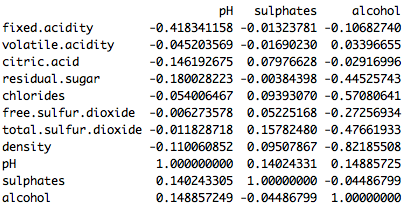
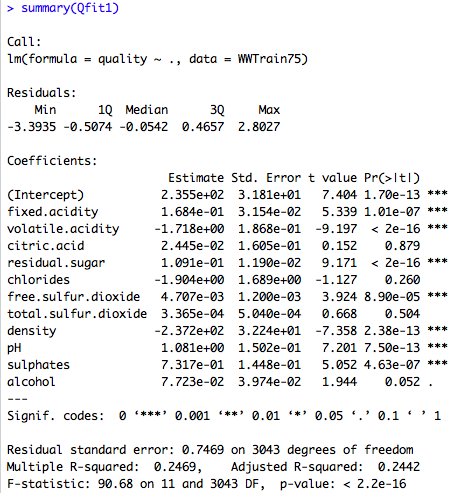


Table 5. Spearman Rank Correlation



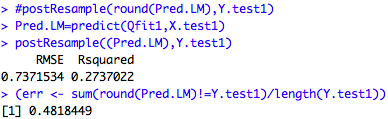
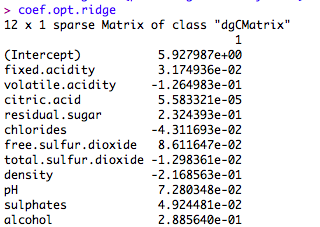


Table 6. The result of multiple linear regression



Screen Shot 2014-11-04 at 9.36.51 PM.png

Screen Shot 2014-11-04 at 9.37.00 PM.png

Table 7. Result of Ridge

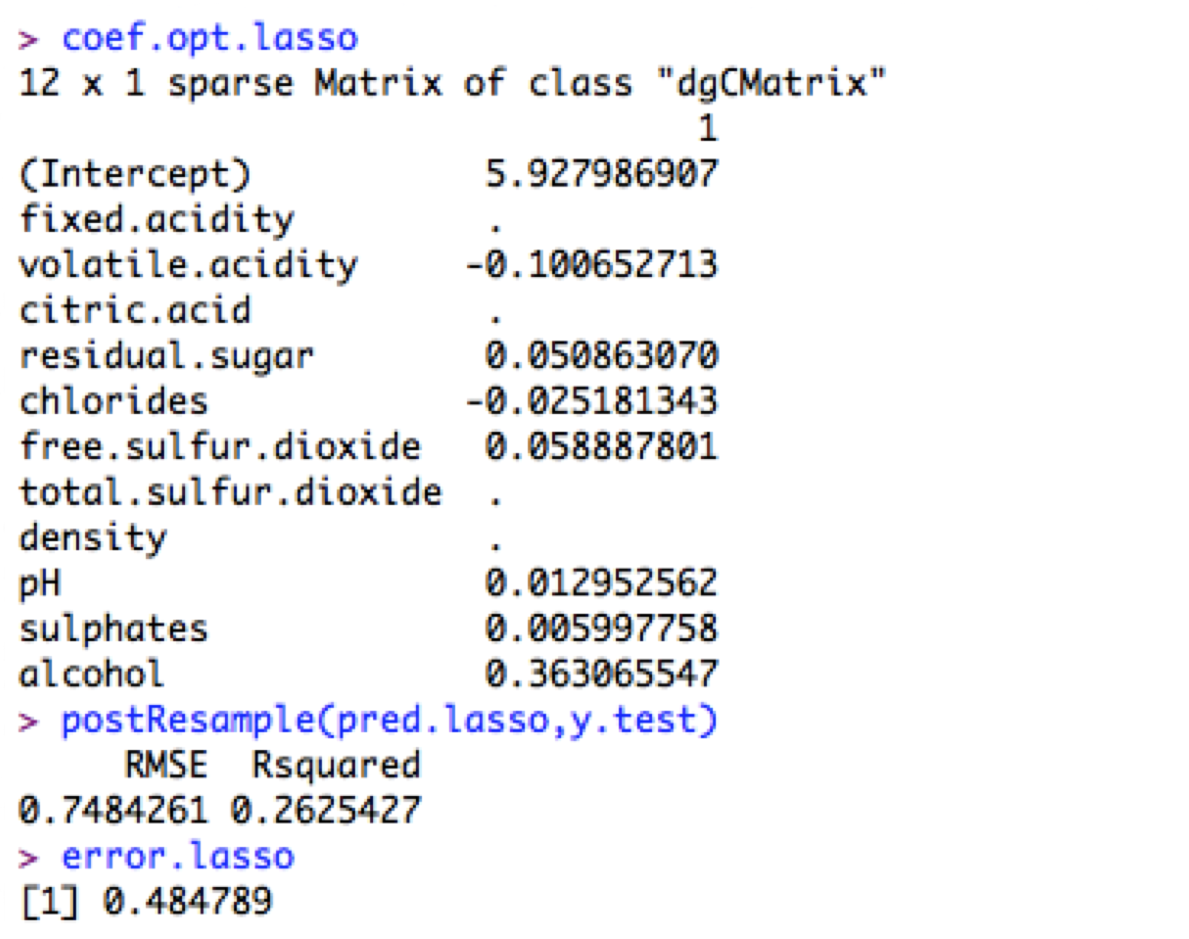


Table 8. Result of LASSO

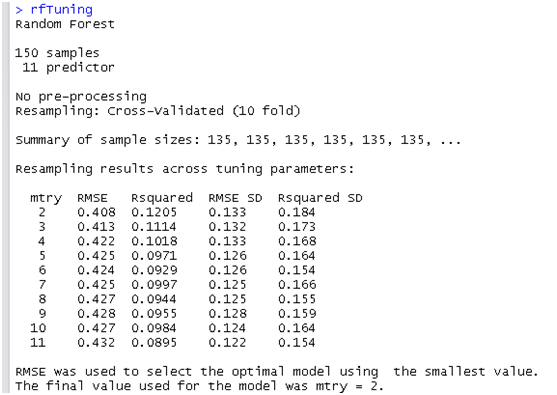


Table 9. Random Forest 10 fold cross validation

R codes for the whole process,

data\_white <- read.table("/Users/Wenxin\_AN/Documents/Master/ada/project/winequality-white.csv",sep=";",header=TRUE)

head(data\_white)

data\_white\_x <- data\_white[,-12]

# Data Source

attach(data\_white)

Class\_variables<- sapply(data\_white,is.numeric)

str(data\_white)

# Sample R code for EDA/Preparation

library(Hmisc)

describe(data\_white,digit=5)

library(caret)

featurePlot(x=data\_white[,-12],y=data\_white[,12],between=list(x=1,y=1),type=c("g","p","smooth"))

# Plot the boxplot for outlier checking

par(mfrow=c(1,5), oma = c(1,1,0,0) + 0.1, mar = c(3,3,1,1) + 0.1)

attach(data\_white)

boxplot(fixed.acidity, col="slategray2", pch=19)

mtext("Fixed Acidity", cex=0.8, side=1, line=2)

boxplot(volatile.acidity, col="slategray2", pch=19)

mtext("Volatile Acidity", cex=0.8, side=1, line=2)

boxplot(citric.acid, col="slategray2", pch=19)

mtext("Citric Acid", cex=0.8, side=1, line=2)

boxplot(residual.sugar, col="slategray2", pch=19)

mtext("Residual Sugar", cex=0.8, side=1, line=2)

boxplot(chlorides, col="slategray2", pch=19)

mtext("Chlorides", cex=0.8, side=1, line=2)

par(mfrow=c(1,6), oma = c(1,1,0,0) + 0.1, mar = c(3,3,1,1) + 0.1)

boxplot(free.sulfur.dioxide, col="slategray2", pch=19)

mtext("free sulfur dioxide", cex=0.8, side=1, line=2)

boxplot(total.sulfur.dioxide, col="slategray2", pch=19)

mtext("total sulfur dioxide", cex=0.8, side=1, line=2)

boxplot(density, col="slategray2", pch=19)

mtext("density", cex=0.8, side=1, line=2)

boxplot(pH, col="slategray2", pch=19)

mtext("pH Value", cex=0.8, side=1, line=2)

boxplot(sulphates, col="slategray2", pch=19)

mtext("sulphates", cex=0.8, side=1, line=2)

boxplot(alcohol, col="slategray2", pch=19)

mtext("alcohol", cex=0.8, side=1, line=2)

# Correlation checking

Correlation<- cor(data\_white[,-12])

Correlation

cor(data\_white[,-12], method="spearman")

par(mfrow=c(1,1))

library(corrplot)

corrplot(Correlation)

# Sample R code for Preparing Data

limout <- rep(0,11)

for (i in 1:11){

t1 <- quantile(data\_white[,i], 0.75)

t2 <- IQR(data\_white[,i], 0.75)

limout[i] <- t1 + 1.5\*t2

}

WhiteWineIndex <- matrix(0, 4898, 11)

for (i in 1:4898)

for (j in 1:11){

if (data\_white[i,j] > limout[j]) WhiteWineIndex[i,j] <- 1

}

WWInd <- apply(WhiteWineIndex, 1, sum)

WhiteWineTemp <- cbind(WWInd, data\_white)

Indexes <- rep(0, 208)

j <- 1

for (i in 1:4898){

if (WWInd[i] > 0) {Indexes[j]<- i

j <- j + 1}

else j <- j

}

WhiteWineLib <-data\_white[-Indexes,]

set.seed(1)

indexes = sample(1:nrow(WhiteWineLib), size=0.75\*nrow(WhiteWineLib))

WWTrain75 <- WhiteWineLib[indexes,]

WWTest25 <- WhiteWineLib[-indexes,]

X.train1 <- WWTrain75[,-12]

X.test1 <- WWTest25[,-12]

Y.train1 <- WWTrain75[,12]

Y.test1 <- WWTest25[,12]

# Sample R code for Multiple Linear Regression

options(digits=7)

library(car)

Qfit1 <- lm(quality ~ ., data=WWTrain75)

summary(Qfit1)

#vif(Qfit1)

#postResample(round(Pred.LM),Y.test1)

Pred.LM=predict(Qfit1,X.test1)

postResample((Pred.LM),Y.test1)

(err <- sum(round(Pred.LM)!=Y.test1)/length(Y.test1))

par(mfrow=c(1,1))

truehist(residuals(Qfit1), h = 0.25, col="slategray3")

#qqPlot(residuals(Qfit1), pch=19, col="darkblue", cex=0.6)

mtext("Distribution of Residuals for Qfit1", outer=T, side=1, line = 2)

# Polynomial Regression

Qfit6 <- lm(quality ~ poly(alcohol,2) + poly(volatile.acidity,2) + residual.sugar + poly(free.sulfur.dioxide,2) + chlorides + sulphates + poly(pH,2), data=WWTrain75)

summary(Qfit6)

Pred.LM\_6=predict(Qfit6,X.test1)

postResample((Pred.LM\_6),Y.test1)

(err <- sum(round(Pred.LM\_6)!=Y.test1)/length(Y.test1))

residualPlots(Qfit6, pch=19, col="blue", cex=0.6)

# Ridge and Lasso

set.seed(1)

indexes = sample(1:nrow(WhiteWineLib), size=0.75\*nrow(WhiteWineLib))

x.train <- WhiteWineLib[indexes,]

x.test <- WhiteWineLib[-indexes,]

y.train=x.train$quality

y.test=x.test$quality

x.train=x.train[,-12]

x.test=x.test[,-12]

x.train=as.data.frame(x.train)

x.test=as.data.frame(x.test)

mean.train <- apply(x.train, 2, mean)

sd.train <- apply(x.train, 2, sd)

x.train <- scale(x.train)

x.train[,sd.train==0] <- 0

x.test <- scale(x.test, center = mean.train, scale=sd.train)

x.test[,sd.train==0] <- 0

library("glmnet")

set.seed(2)

fit.ridge <- cv.glmnet(x.train, y.train, alpha = 0)

#plot(fit.ridge)

coef.opt.ridge <- coef(fit.ridge)

#test

pred.ridge <- predict(fit.ridge,x.test)

pred.ridge.final=round(pred.ridge)

error.ridge=sum(pred.ridge.final!=y.test)/length(y.test)

coef.opt.ridge

postResample(pred.ridge,y.test)

error.ridge

#lasso

set.seed(2)

fit.lasso <- cv.glmnet(x.train, y.train, alpha = 1, standardize=F,family="gaussian")

#plot(fit.lasso)

coef.opt.lasso <- coef(fit.lasso)

#test

pred.lasso <- predict(fit.lasso,x.test)

postResample(pred.lasso,y.test)

pred.lasso.final=round(pred.lasso)

error.lasso=sum(pred.lasso.final!=y.test)/length(y.test)

coef.opt.lasso

postResample(pred.lasso,y.test)

error.lasso

#Nonlinear models

#Random Forest

wine<-read.csv(file="winequality-white.csv",sep=";",header=TRUE)

library(caret)

X<-wine[,-12]

quality<-wine[,12]

pre.x=predict(P,wine[,-12])

Y=wine[,12]

Y=data.frame(Y)

trainset<-createDataPartition(wine[,12],p=3/4,list=F)

train.x<-pre.x[trainset,]

test.x<-pre.x[-trainset,]

train.y<-Y[trainset,]

test.y<-Y[-trainset,]

library(randomForest)

trainx=train.x[1:150,]

trainy=train.y[1:150]

rfTuning=train(trainx,trainy,method="rf",tuneLength=10,trControl=trainControl(method="cv"))

rfModel<-randomForest(train.x,train.y,importance=TRUE,ntrees=1000,mtry=2)

rfcv(train.x,train.y,cv.fold=10,scale="log",step=0.5,mtry=function(p)2,recursive=FALSE)

varImpPlot(rfModel,main="Random Forest")

predict.y<-predict(rfModel,test.x)

postResample(pred=predict.y,obs=test.y)

err<-sum(round(predict.y)!=test.y)/length(test.y)

# NN

library(nnet)

nnetFit=nnet(trainwine,trainqty,size=5,decat=0.01,linout=TRUE,trace=FALSE,maxit=500,MAXNWts=5\*(ncol(trainwine)+1)+5+1)

nnetAvg=avNNet(trainwine,trainqty,size=5,decat=0.01,repeats=5,linout=TRUE,trace=FALSE,maxit=500,MAXNWts=5\*(ncol(trainwine)+1)+5+1)

predfit=predict(nnetFit,testwine)

predavg=predict(nnetAvg,testwine)

nnvaluesfit=data.frame(obs=testqty,pred=predfit)

nnvaluesavg=data.frame(obs=testqty,pred=predavg)

defaultSummary(nnvaluesfit)

defaultSummary(nnvaluesavg)

toohigh<-findCorrelation(cor(trainwine),cutoff=0.7)

trainnnet<-trainwine[,-toohigh]

testnnet<-testwine[,-toohigh]

nnetgrid<-expand.grid(.decay=c(0,0.01,.1),.size=c(1:10))

set.seed(100) nnettune<-train(trainwine,trainqty,method="nnet",tuneGrid=nnetgrid,trControl=trainControl(method="cv"),preProc=c("center","scale"),linout=TRUE,trace=FALSE,MaxNWts=5\*(ncol(trainnnet)+1)+5+1,maxit=500)

nnettune

nnetFit=nnet(trainwine,trainqty,size=5,decat=0.1,linout=TRUE,trace=FALSE,maxit=500,MAXNWts=5\*(ncol(trainwine)+1)+5+1)

> predqtyFIT=predict(nnetFit,trainwine) lmvaluesFIT=data.frame(obs=trainqty,pred=predqtyFIT) defaultSummary(lmvaluesFIT)

predqtyFIT2=predict(nnetFit,testwine)

lmvaluesFIT2=data.frame(obs=testqty,pred=predqtyFIT2)

defaultSummary(lmvaluesFIT2)

error

# Classification Models

data$quality <- as.factor(data$quality)

limout.up <- rep(0,11)

limout.low <- rep(0,11)

for (i in 1:11){

  t1 <- quantile(data[,i], 0.75)

  t2 <- IQR(data[,i], 0.75)

  t3 <- quantile(data[,i],0.25)

  limout.up[i] <- t1 + 1.5\*t2

  limout.low[i]<- t3 - 1.5\*t2

}

dataIndex <- matrix(0, 4898, 11)

for (i in 1:4898)

  for (j in 1:11){

    if (data[i,j] > limout.up[j]||data[i,j]< limout.low[j]) dataIndex[i,j] <- 1

  }

WWInd <- apply(dataIndex, 1, sum)

Indexes <- rep(0, 883)

j <- 1

for (i in 1:4898){

  if (WWInd[i] > 0) {Indexes[j]<- i

                     j <- j + 1}

  else j <- j

}

dataLib <-data[-Indexes,]

indexes = sample(1:nrow(dataLib), size=0.75\*nrow(dataLib))

data <- dataLib

train.data <- data[indexes,]

test.data <- data[-indexes,]

require(rpart)

p<-ncol(data)

fit.tree <- rpart(quality~.,data=train.data,method="class")

predict.tree <- predict(fit.tree,test.data[,-p],type="class")

err.tree<-sum(predict.tree!=test.data$quality)/nrow(test.data)

require(randomForest)

set.seed(123)

fit.rf<-randomForest(quality~.,data=train.data,importance=TRUE,proximity=TRUE,ntree=1000)

round(importance(fit.rf),2)

predict.rf <- predict(fit.rf,test.data[,-p],type="class")

predict.rf$confusion

err.rf <-sum(predict.rf!=test.data$quality)/nrow(test.data)

require(ada)

require(adabag)

fit.ada <- boosting.cv(quality~.,data=train.data)

predict.ada <- predict(fit.ada,newdata=test.data)

err.ada <-sum(predict.ada$class!=test.data$quality)/nrow(test.data)

require(e1071)

fit.svm <- svm(quality~.,data=train.data,gamma=1,type="C",cross=10)

predict.svm <-predict(fit.svm,test.data[,-p])

# table(pred=predict.svm,true=test.data$quality)

err.svm <-sum(predict.svm!=test.data$quality)/nrow(test.data)

err.svm

err.tree

err.rf

err.ada

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