# An algorithmic approach to determining wine quality from chemical data

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### Introduction

"Best Wines in the World" (BWITW) is a wine club that prides itself on being able to provide new quality wines to its members at reasonable prices. It currently relies on testers to judge the quality of its wines which is a slow time consuming and expensive process. Also, by the time that it has tested new wines the market is usually aware of them and the price rises. It has commissioned this report to assess the viability of using chemical analysis to determine whether a wine is "quality" or not quickly and cheaply, so that it can get access to new quality wines before its competition. The management of BWITW consider a wine to be "quality" if it receives a score of 7 out of 10 by a panel of its testers. It is hoped that with the application of chemical analysis and data science, it would be able to quickly screen a large number of new and undiscovered wines, identify those that are "quality" and be able to offer these to its customers.

This report details analysis that has been done to determine whether this combination of chemical testing and data science approach would work and makes a judgement on suitability of this approach and possible next steps.

BWITW have been doing chemical testing on their red wines and recording this information along with a quality score given by its panel of testers. The approach taken in this exercise was to first investigate the dataset to understand its structure and the variables within it, including some visualisations. A very simple regression was performed to see what would be simply possible using very simple techniques. After this a range of more sophisticated machine learning algorithms were applied to see whether this would lead to better results. From these results conclusions have been drawn.

NOTE: The data set used here is from https://archive.ics.uci.edu/ml/datasets/Wine+Quality and P. Cortez, A. Cerdeira, F. Almeida, T. Matos and J. Reis, Modeling wine preferences by data mining from physicochemical properties, in Decision Support Systems, Elsevier, 47(4):547-553, 2009 is cited.

### Methods and Analysis

The first step in the analysis was to ensure that neccesary packages and the dataset were downloaded. The code for this is below.

```
## load packages that are required

if(!require(tidyverse)) install.packages("tidyverse", repos = "http://cran.us.r-project.org")
if(!require(caret)) install.packages("caret", repos = "http://cran.us.r-project.org")
if(!require(data.table)) install.packages("data.table", repos = "http://cran.us.r-project.org")
if(!require(gam)) install.packages("gam", repos = "http://cran.us.r-project.org")
if(!require(nnet)) install.packages("nnet", repos = "http://cran.us.r-project.org")

## set seed at the start to ensure consistency of output

set.seed(8934, sample.kind = "Rounding")

## download the red wine quality dataset
```

```
wine_quality <- read.csv("https://archive.ics.uci.edu/ml/machine-learning-databases/wine-quality/winequ</pre>
                          sep = ";")
```

After this, the size and shape of the dataset were investigated.

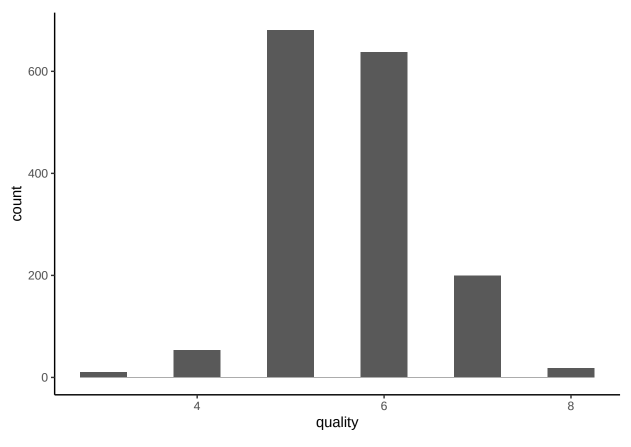
```
## have an initial look at the dataset
head(wine_quality)
##
     fixed.acidity volatile.acidity citric.acid residual.sugar chlorides
## 1
               7.4
                                0.70
                                            0.00
                                                             1.9
                                                                     0.076
## 2
               7.8
                                0.88
                                            0.00
                                                             2.6
                                                                     0.098
               7.8
## 3
                                0.76
                                            0.04
                                                             2.3
                                                                     0.092
## 4
              11.2
                                0.28
                                            0.56
                                                             1.9
                                                                     0.075
               7.4
                                            0.00
## 5
                                                             1.9
                                                                     0.076
                                0.70
## 6
               7.4
                                0.66
                                            0.00
                                                             1.8
                                                                     0.075
##
     free.sulfur.dioxide total.sulfur.dioxide density
                                                        pH sulphates alcohol
## 1
                      11
                                            34 0.9978 3.51
                                                                  0.56
                                                                           9.4
## 2
                      25
                                            67 0.9968 3.20
                                                                  0.68
                                                                           9.8
## 3
                                            54 0.9970 3.26
                      15
                                                                  0.65
                                                                           9.8
## 4
                      17
                                            60 0.9980 3.16
                                                                           9.8
                                                                  0.58
## 5
                      11
                                            34 0.9978 3.51
                                                                  0.56
                                                                           9.4
## 6
                      13
                                            40 0.9978 3.51
                                                                  0.56
                                                                           9.4
##
     quality
## 1
           5
## 2
           5
## 3
           5
## 4
           6
## 5
           5
## 6
           5
dim(wine_quality)
```

```
## [1] 1599
              12
```

It can be seen that there are 1599 records and 12 columns with observations. One of the columns is the wine quality which is a numeric score.

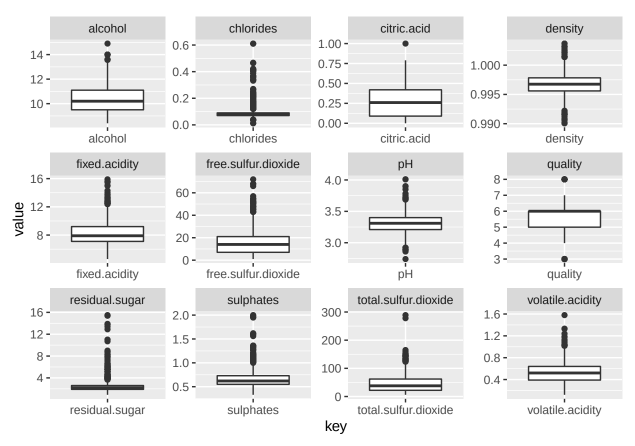
The dataset was investigated with a series of visualisations

```
# first up - take a look at the quality
wine_quality %>% ggplot(aes(x = quality)) +
  geom_histogram(binwidth = 0.5) +
 theme_classic()
```



```
# next - try and box plot to understand the distribution of the different variables
# that we have data for and may determine quality

wine_quality %>%
  gather(key, value) %>%
  ggplot(aes(y = value, x = key)) +
  geom_boxplot() +
  facet_wrap(~ key, scales = "free")
```



It can be seen that there the quality scores approximate a normal distribution and that most scores are near the mean with a 5 or a 6. The other variables in the dataset representing chemical properties/readings are distributed in a range of different ways as shown in the box plots.

BWITW consider a wine to be quality if it receives a score from their panel of testers of 7 or over, so this is defined as success and a new factor with two levels: "quality" where the wine has a score of 7 or over and "sub-quality" where is does not is created.

```
# define a quality wine as having a score of 7 or above

y <- if_else(wine_quality$quality >= 7, "quality", "sub-quality")

y <- factor(y)
data.frame(y) %>%
  group_by(y) %>%
  summarise(n()) %>%
  knitr::kable()
```

у	n()
quality	217
sub-quality	1382

The dataset was split into a training set (80%) and a test set (20%) to start training models.

```
# Validation set will be 20% of the total dataset
```

```
test_index <- createDataPartition(y = wine_quality$quality, times = 1, p = 0.2, list = FALSE)
training_set <- wine_quality[-test_index,]
test_set <- wine_quality[test_index,]
y_training_set <- y[-test_index]
y_test_set <- y[test_index]</pre>
```

As a bit of further data analysis a test was done to check that the percentage of quality wines was roughly the same in both the training set and the test set.

```
# check the proportion of the wines that are quality in both the test set and the training set
mean(y_training_set == "quality")

## [1] 0.1353678

mean(y_test_set == "quality")
```

## [1] 0.1370717

This shows that in both cases, the proportion of "quality" wines is around 14%.

As an initial piece of the analysis the correlation coefficient between the various variables and the quality score for the wine were calculated.

```
# lets now start with the simplest predictive model and see whether a simple correlation would
# give a high degree of accuracy

cor(wine_quality[1:11], wine_quality[12]) %>%
    knitr::kable()
```

	quality
fixed.acidity	0.1240516
volatile.acidity	-0.3905578
citric.acid	0.2263725
residual.sugar	0.0137316
chlorides	-0.1289066
free.sulfur.dioxide	-0.0506561
total.sulfur.dioxide	-0.1851003
density	-0.1749192
pН	-0.0577314
sulphates	0.2513971
alcohol	0.4761663

It can be seen from this that the alcohol content of the wine is the most correlated variable with the wine quality and hence this could be the basis for an intial simple regression model. However, the classification of a a wine into quality or not is categorical rather than continious and hence a relevant model needs to be applied here (and to more complicated models that are developed).

The regression approach can be extended to categorical data. For example, we can try regression to estimate the conditional probability:

```
p(x) = Pr(Y = 1|X = x) = \beta_0 + \beta_1 x
```

Once we have estimates  $\beta_0$  and  $\beta_1$ , we can obtain an actual prediction p(x). Then we can define a specific decision rule to form a prediction. The logical decision rule to apply here would be if  $\hat{p} > 0.5$  and this is done in the code below.

We now need to define success for model. There are several measures that will be used as part of a judgement on the best approach. The accuracy will be looked at, as well as the sensitivity, specificity and the balance accuracy. In the context of this exercise:

- the accuracy represents the total proportion of wine samples that were correctly identified using that approach.
- the sensitivity is the true positive rate and represents the proportion of quality wines that are correctly identified as such.
- the specificity is the true negative rate and measures the proportion of non-quality wines identified as such.
- the balance accuracy makes an adjustment for the fact that there are significantly less "quality" than "sub-quality" wines in the dataset and in effect normalises the results for this and hence shows the proportion of wines that would be correctly identified in a balanced dataset.
- the positive predicted accuracy is the proportion of wines that are predicted to be "quality" that actually are.
- the negative predicted accuracy is the proportion of wines that are predicted of be "sub-quality" that actually are.

The code below also creates a dataframe to store the results from the analysis.

```
# of these variables, it appears that alcohol has the highest correlation with quality, so let's build
# based on this simple correlation
lm_fit <- mutate(training_set, y = as.numeric(quality >= 7)) %>% lm(y ~ alcohol, data = .)
p_hat <- predict(lm_fit, test_set)</pre>
y_hat <- ifelse(p_hat > 0.5, "quality", "sub-quality") %>% factor()
mean(y_hat == y_test_set)
wine_results <- data_frame(method = "regression on alcohol",</pre>
                           accuracy =
                             confusionMatrix(y hat, y test set)$overall["Accuracy"],
                           sensitivity =
                             confusionMatrix(y_hat, y_test_set)$byClass["Sensitivity"],
                           specificity =
                             confusionMatrix(y_hat, y_test_set)$byClass["Specificity"],
                           balanced acc =
                             confusionMatrix(y_hat, y_test_set)$byClass["Balanced Accuracy"],
                           positive_pred_val =
                             confusionMatrix(y_hat, y_test_set)$byClass["Pos Pred Value"],
                           negative_pred_val =
                             confusionMatrix(y_hat, y_test_set)$byClass["Neg Pred Value"])
```

The simple regression model on alcohol gives a base against which to compare the other approaches. The following machine learning algorithms/approaches were subsequently run using all the variables in the dataset:

- A Generalised Linear Model
- A Linear Discriminant Analysis Model
- A Quadratic Discriminant Analysis Model
- A Generalized Additive Model using LOESS

- A Random Forest Model
- A k-Nearest Neighbours Model
- An AdaBoost Classification Trees Model
- A Neural Network Model
- An Ensemble Model which takes the average results of all the models above

The code to generate these is shown below and note that it does take some time to run. The first section in the code below removes the quality score from the dataset, so that this is not included in dataset as it is obviously highly correlated with the "quality" clasification that has been established.

```
# Remove the quality value from the data table to ease the application of machine learning approaches
training_set <- training_set[,-12]
test_set <- test_set[,-12]</pre>
# start with a simple linear regression model to see what results that will give
train_glm <- train(training_set, y_training_set,</pre>
                   method = "glm")
glm_preds <- predict(train_glm, test_set)</pre>
mean(glm_preds == y_test_set)
wine_results <- bind_rows(wine_results,
                           data_frame(method="general linear model",
                                       accuracy =
                                         confusionMatrix(glm_preds,
                                                          y_test_set)$overall["Accuracy"],
                                       sensitivity =
                                         confusionMatrix(glm_preds,
                                                         y_test_set)$byClass["Sensitivity"],
                                       specificity =
                                         confusionMatrix(glm_preds,
                                                          y_test_set)$byClass["Specificity"],
                                       balanced acc =
                                         confusionMatrix(glm_preds,
                                                          y_test_set)$byClass["Balanced Accuracy"],
                                       positive pred val =
                                         confusionMatrix(glm_preds,
                                                          y_test_set)$byClass["Pos Pred Value"],
                                       negative_pred_val =
                                         confusionMatrix(glm_preds,
                                                         y_test_set)$byClass["Neg Pred Value"]))
# next try the LDA model
train_lda <- train(training_set, y_training_set,</pre>
                    method = "lda")
lda_preds <- predict(train_lda, test_set)</pre>
mean(lda_preds == y_test_set)
wine_results <- bind_rows(wine_results,</pre>
                           data_frame(method="LDA model",
                                       accuracy =
                                         confusionMatrix(lda_preds,
                                                         y_test_set)$overall["Accuracy"],
                                       sensitivity =
```

```
confusionMatrix(lda_preds,
                                                         y_test_set)$byClass["Sensitivity"],
                                      specificity =
                                         confusionMatrix(lda_preds,
                                                         y_test_set)$byClass["Specificity"],
                                      balanced acc =
                                         confusionMatrix(lda_preds,
                                                          y_test_set)$byClass["Balanced Accuracy"],
                                      positive_pred_val =
                                         confusionMatrix(lda_preds,
                                                         y_test_set)$byClass["Pos Pred Value"],
                                      negative_pred_val =
                                         confusionMatrix(lda_preds,
                                                         y_test_set)$byClass["Neg Pred Value"]))
# and the QDA model
train_qda <- train(training_set, y_training_set,</pre>
                    method = "qda")
qda_preds <- predict(train_qda, test_set)</pre>
mean(qda_preds == y_test_set)
wine_results <- bind_rows(wine_results,</pre>
                           data_frame(method="QDA model",
                                      accuracy =
                                         confusionMatrix(qda_preds,
                                                         y_test_set)$overall["Accuracy"],
                                      sensitivity =
                                         confusionMatrix(qda_preds,
                                                         y_test_set)$byClass["Sensitivity"],
                                      specificity =
                                         confusionMatrix(qda_preds,
                                                          y_test_set)$byClass["Specificity"],
                                      balanced_acc =
                                         confusionMatrix(qda_preds,
                                                         y_test_set)$byClass["Balanced Accuracy"],
                                      positive_pred_val =
                                         confusionMatrix(qda_preds,
                                                         y_test_set)$byClass["Pos Pred Value"],
                                      negative_pred_val =
                                         confusionMatrix(qda_preds,
                                                         y_test_set)$byClass["Neg Pred Value"]))
# and a Loess model
train_loess <- train(training_set, y_training_set,</pre>
                   method = "gamLoess")
loess_preds <- predict(train_loess, test_set)</pre>
mean(loess_preds == y_test_set)
wine_results <- bind_rows(wine_results,</pre>
                           data_frame(method="Loess",
                                      accuracy =
                                         confusionMatrix(loess_preds,
```

```
y_test_set)$overall["Accuracy"],
                                      sensitivity =
                                         confusionMatrix(loess_preds,
                                                         y_test_set)$byClass["Sensitivity"],
                                      specificity =
                                         confusionMatrix(loess_preds,
                                                         y_test_set)$byClass["Specificity"],
                                      balanced acc =
                                         confusionMatrix(loess_preds,
                                                         y_test_set)$byClass["Balanced Accuracy"],
                                      positive_pred_val =
                                         confusionMatrix(loess_preds,
                                                         y_test_set)$byClass["Pos Pred Value"],
                                      negative_pred_val =
                                         confusionMatrix(loess_preds,
                                                         y_test_set)$byClass["Neg Pred Value"]))
# and a random forest model
train_rf <- train(training_set, y_training_set,</pre>
                      method = "rf")
rf_preds <- predict(train_rf, test_set)</pre>
mean(rf_preds == y_test_set)
wine_results <- bind_rows(wine_results,</pre>
                           data frame (method="Random Forest",
                                      accuracy =
                                         confusionMatrix(rf_preds,
                                                         y_test_set)$overall["Accuracy"],
                                      sensitivity =
                                         confusionMatrix(rf_preds,
                                                         y_test_set)$byClass["Sensitivity"],
                                      specificity =
                                         confusionMatrix(rf_preds,
                                                         y_test_set)$byClass["Specificity"],
                                      balanced_acc =
                                         confusionMatrix(rf_preds,
                                                         y_test_set)$byClass["Balanced Accuracy"],
                                      positive_pred_val =
                                         confusionMatrix(rf_preds,
                                                         y_test_set)$byClass["Pos Pred Value"],
                                      negative_pred_val =
                                         confusionMatrix(rf_preds,
                                                         y_test_set)$byClass["Neg Pred Value"]))
# and a knn model
train_knn <- train(training_set, y_training_set,</pre>
                  method = "knn")
knn_preds <- predict(train_knn, test_set)</pre>
mean(knn_preds == y_test_set)
wine_results <- bind_rows(wine_results,</pre>
                           data_frame(method="k nearest neighbour",
```

```
accuracy =
                                        confusionMatrix(knn preds,
                                                         y_test_set)$overall["Accuracy"],
                                      sensitivity =
                                        confusionMatrix(knn_preds,
                                                         y_test_set)$byClass["Sensitivity"],
                                      specificity =
                                        confusionMatrix(knn_preds,
                                                         y_test_set)$byClass["Specificity"],
                                      balanced_acc =
                                        confusionMatrix(knn_preds,
                                                         y_test_set)$byClass["Balanced Accuracy"],
                                      positive_pred_val =
                                         confusionMatrix(knn_preds,
                                                         y_test_set)$byClass["Pos Pred Value"],
                                      negative_pred_val =
                                        confusionMatrix(knn preds,
                                                         y_test_set)$byClass["Neg Pred Value"]))
# and an adaboost model
train_adaboost <- train(training_set, y_training_set,</pre>
                   method = "adaboost")
adaboost_preds <- predict(train_adaboost, test_set)</pre>
mean(adaboost_preds == y_test_set)
wine_results <- bind_rows(wine_results,</pre>
                           data_frame(method="adaboost",
                                      accuracy =
                                        confusionMatrix(adaboost_preds,
                                                         y_test_set)$overall["Accuracy"],
                                      sensitivity =
                                        confusionMatrix(adaboost_preds,
                                                         y_test_set)$byClass["Sensitivity"],
                                      specificity =
                                        confusionMatrix(adaboost_preds,
                                                         y_test_set)$byClass["Specificity"],
                                      balanced_acc =
                                        confusionMatrix(adaboost_preds,
                                                         y_test_set)$byClass["Balanced Accuracy"],
                                      positive_pred_val =
                                        confusionMatrix(adaboost_preds,
                                                         y_test_set)$byClass["Pos Pred Value"],
                                      negative_pred_val =
                                        confusionMatrix(adaboost_preds,
                                                         y_test_set)$byClass["Neg Pred Value"]))
# and a neural network model
train_nnet <- train(training_set, y_training_set,</pre>
                        method = "nnet")
nnet_preds <- predict(train_nnet, test_set)</pre>
mean(nnet_preds == y_test_set)
wine_results <- bind_rows(wine_results,</pre>
                           data_frame(method="neural network",
```

```
accuracy =
                                       confusionMatrix(nnet preds,
                                                        y_test_set)$overall["Accuracy"],
                                     sensitivity =
                                       confusionMatrix(nnet_preds,
                                                        y_test_set)$byClass["Sensitivity"],
                                     specificity =
                                       confusionMatrix(nnet_preds,
                                                        y_test_set)$byClass["Specificity"],
                                     balanced_acc =
                                       confusionMatrix(nnet_preds,
                                                        y_test_set)$byClass["Balanced Accuracy"],
                                     positive_pred_val =
                                        confusionMatrix(nnet_preds,
                                                        y_test_set)$byClass["Pos Pred Value"],
                                     negative_pred_val :
                                       confusionMatrix(nnet preds,
                                                        y_test_set)$byClass["Neg Pred Value"]))
# finally include an ensemble model, which combines takes an average across all the models
ensemble <- cbind(glm = glm preds == "quality", lda = lda preds == "quality",</pre>
                  qda = qda_preds == "quality",
                  loess = loess_preds == "quality", rf = rf_preds == "quality",
                  knn = knn_preds == "quality", adaboost = adaboost_preds == "quality",
                  nnet = nnet_preds == "quality")
ensemble_preds <- ifelse(rowMeans(ensemble) > 0.5, "quality", "sub-quality") %>% factor()
mean(ensemble_preds == y_test_set)
wine_results <- bind_rows(wine_results,
                          data_frame(method="ensemble",
                                     accuracy =
                                       confusionMatrix(ensemble_preds,
                                                        y_test_set)$overall["Accuracy"],
                                     sensitivity =
                                       confusionMatrix(ensemble preds,
                                                                    y_test_set)$byClass["Sensitivity"],
                                     specificity =
                                       confusionMatrix(ensemble_preds,
                                                        y_test_set)$byClass["Specificity"],
                                     balanced acc =
                                        confusionMatrix(ensemble_preds,
                                                        y_test_set)$byClass["Balanced Accuracy"],
                                     positive_pred_val =
                                       confusionMatrix(ensemble_preds,
                                                        y_test_set)$byClass["Pos Pred Value"],
                                     negative_pred_val =
                                       confusionMatrix(ensemble_preds,
                                                        y_test_set)$byClass["Neg Pred Value"]))
```

### Results

The final table of results can be see below. This shows that the Random Forest model delivers the best results looking across all of the models. It delivers the best results across all the measures defined above.

```
# print the results
wine_results %>%
knitr::kable()
```

method	accuracy	sensitivity	specificity	$balanced\_acc$	$positive\_pred\_val$	negative_pred_val
regression on alcohol	0.8598131	0.0227273	0.9927798	0.5077535	0.3333333	0.8647799
general linear model	0.8753894	0.4090909	0.9494585	0.6792747	0.5625000	0.9100346
LDA model	0.8691589	0.4545455	0.9350181	0.6947818	0.5263158	0.9151943
QDA model	0.8411215	0.4772727	0.8989170	0.6880948	0.4285714	0.9154412
Loess	0.8816199	0.4318182	0.9530686	0.6924434	0.5937500	0.9134948
Random Forest	0.9127726	0.5000000	0.9783394	0.7391697	0.7857143	0.9249147
k nearest neighbour	0.8722741	0.1818182	0.9819495	0.5818838	0.6153846	0.8831169
adaboost	0.9034268	0.4318182	0.9783394	0.7050788	0.7600000	0.9155405
neural network	0.8660436	0.3863636	0.9422383	0.6643010	0.5151515	0.9062500
ensemble	0.8753894	0.3181818	0.9638989	0.6410404	0.5833333	0.8989899

The Random Forest model gets the correct results in 91% of cases in the test set. It can correctly identify quality wines in 50% of cases and correctly identify sub-quality wines in 98% of them. However, the most interesting figure here is probably the positive predicted value wich is 78%. This represents that percentage of wines that are identified as quality that actually are. Whilst the approach discussed here would help to reduce the number of wines that need to be screened, the fact that around 1 in 5 wines identified as quality are not, would mean that human testers would probably still be required before wines could be sent to customers.

#### Conclusion

The Random Forest model combined with the chemical analysis should allow BWITW to quickly screen a wide number of wines to identify which of these are highly likely to be quality and meet their requirements. However, they do need to be careful as some of those that are expected to be quality via this approach will not be and they need to assess the risk of sending these to their customers.

## Possibilities for further analysis

The following are possibilities for future analysis:

- The parameters of the model could be changed, so that positive predicted value is ranked more highly as it is tuned. This would mean that less quality wines overall would be identified, but if the sensitivity increases then it would mean that it was less likely that a non-quality wine was identified as quality. If the positive predicted value were to be increased enough then it may be possible to identify wines purely via the chemical analysis with the Random Forest model and dispatch these directly to customers. This could enable significant additional cost savings and possibly remove the need for having any testers.
- The use of clustering approaches (either hierarchical clustering or k-means clustering) and then using some functionality from the "recommenderlab" package to create a model for predicting.