5110_Final_Project

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```
library(tidyverse)
## -- Attaching packages ------ 1.3.1 --
## v ggplot2 3.3.5
                             0.3.4
                   v purrr
## v tibble 3.1.6
                   v dplyr 1.0.7
## v tidyr 1.1.4
                  v stringr 1.4.0
## v readr
          2.1.1
                    v forcats 0.5.1
## -- Conflicts ------ tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                  masks stats::lag()
library(modelr)
library(gridExtra)
##
## Attaching package: 'gridExtra'
## The following object is masked from 'package:dplyr':
##
##
      combine
library(randomForest)
## randomForest 4.7-1
## Type rfNews() to see new features/changes/bug fixes.
## Attaching package: 'randomForest'
## The following object is masked from 'package:gridExtra':
##
##
      combine
## The following object is masked from 'package:dplyr':
##
##
      combine
```

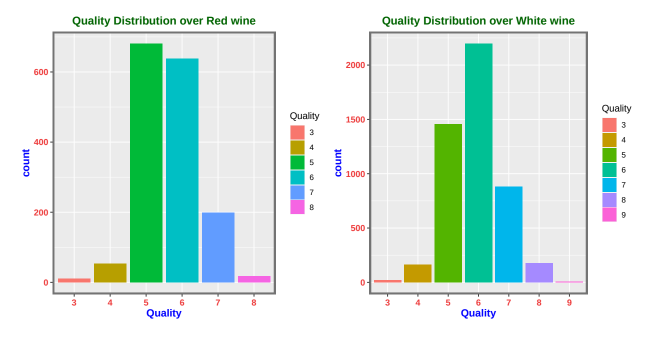
```
## The following object is masked from 'package:ggplot2':
##
##
       margin
library(glmnet)
## Loading required package: Matrix
## Attaching package: 'Matrix'
## The following objects are masked from 'package:tidyr':
##
##
       expand, pack, unpack
## Loaded glmnet 4.1-4
library(caret)
## Loading required package: lattice
##
## Attaching package: 'caret'
## The following object is masked from 'package:purrr':
##
       lift
library(broom)
##
## Attaching package: 'broom'
## The following object is masked from 'package:modelr':
##
##
       bootstrap
dir <- "C:/Users/harsh/Desktop/NEU/IDMP/Final Project/winequality/winequality"</pre>
dir1 <- file.path(dir, "winequality-red.csv")</pre>
dir2 <- file.path(dir, "winequality-white.csv")</pre>
red_wine_df <- read.csv(dir1, header = TRUE, sep = ";")</pre>
white_wine_df <- read.csv(dir2, header = TRUE, sep = ";")</pre>
which(is.na(red_wine_df))
```

integer(0)

Quality Distribution over Red wine

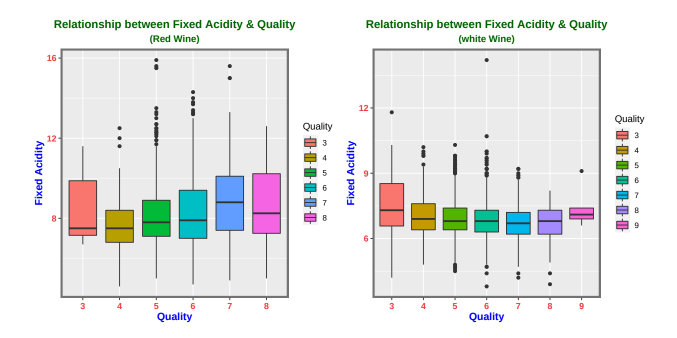
```
qua_red <- ggplot(red_wine_df) +</pre>
          geom_bar(aes(x = factor(quality), fill = factor(quality))) +
          labs(title = "Quality Distribution over Red wine",
               x = "Quality",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
qua_white <- ggplot(white_wine_df) +</pre>
          geom_bar(aes(x = factor(quality), fill = factor(quality))) +
          labs(title = "Quality Distribution over White wine",
               x = "Quality",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
```

```
color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(qua_red, qua_white, nrow = 1)
```



Fixed Acidity Vs Quality

```
plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p1w <- ggplot(white_wine_df) +</pre>
          geom_boxplot(aes(x = factor(quality)), y = fixed.acidity, fill = factor(quality))) +
          labs(title = "Relationship between Fixed Acidity & Quality",
               subtitle = "(white Wine)",
               x = "Quality",
               y = "Fixed Acidity",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p1, p1w, nrow = 1)
```



Volatile Acidity Vs Quality

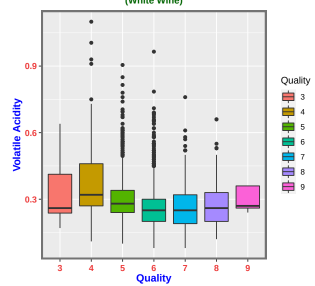
```
p2 <- ggplot(red_wine_df) +</pre>
          geom_boxplot(aes(x = factor(quality), y = volatile.acidity, fill = factor(quality))) +
          labs(title = "Relationship between Volatile Acidity & Quality",
               subtitle = "(Red Wine)",
               x = "Quality",
               y = "Volatile Acidity",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p2w <- ggplot(white_wine_df) +</pre>
          geom_boxplot(aes(x = factor(quality), y = volatile.acidity, fill = factor(quality))) +
```

```
labs(title = "Relationship between Volatile Acidity & Quality",
               subtitle = "(White Wine)",
               x = "Quality",
               v = "Volatile Acidity",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p2, p2w, nrow = 1)
```

Relationship between Volatile Acidity & Quality

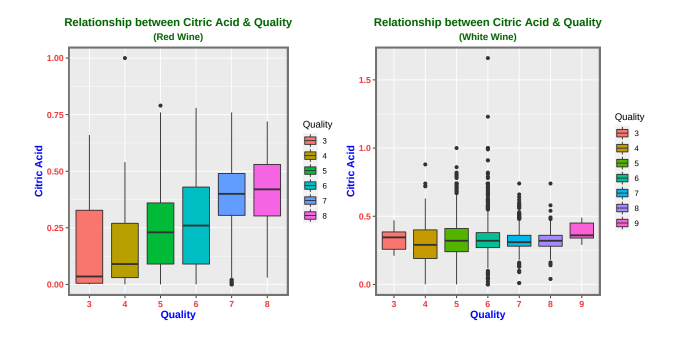
Quality Quality Quality Quality

Relationship between Volatile Acidity & Quality (White Wine)



Citric Acid Vs Quality

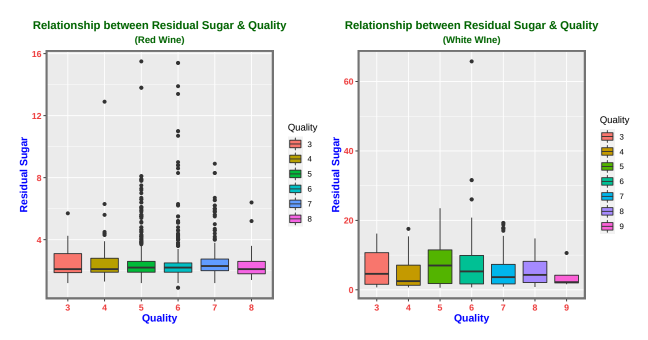
```
fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p3w <- ggplot(white_wine_df) +
          geom_boxplot(aes(x = factor(quality)), y = citric.acid, fill = factor(quality))) +
          labs(title = "Relationship between Citric Acid & Quality",
               subtitle = "(White Wine)",
               x = "Quality",
               y = "Citric Acid",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p3, p3w, nrow = 1)
```



Residual Sugar Vs Quality

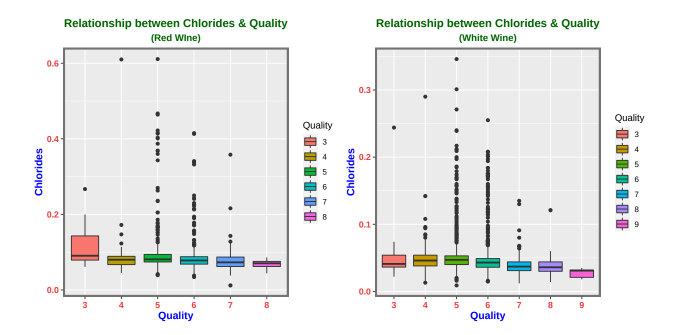
```
p4 <- ggplot(red_wine_df) +
          geom_boxplot(aes(x = factor(quality)), y = residual.sugar, fill = factor(quality))) +
          labs(title = "Relationship between Residual Sugar & Quality",
               subtitle = "(Red Wine)",
               x = "Quality",
               y = "Residual Sugar",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p4w <- ggplot(white_wine_df) +
          geom_boxplot(aes(x = factor(quality)), y = residual.sugar, fill = factor(quality))) +
          labs(title = "Relationship between Residual Sugar & Quality",
```

```
subtitle = "(White WIne)",
               x = "Quality",
               y = "Residual Sugar",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p4, p4w, nrow = 1)
```



Chlorides Vs Quality

```
subtitle = "(Red WIne)",
               x = "Quality",
               y = "Chlorides",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p5w <- ggplot(white_wine_df) +
          geom_boxplot(aes(x = factor(quality), y = chlorides, fill = factor(quality))) +
          labs(title = "Relationship between Chlorides & Quality",
               subtitle = "(White Wine)",
               x = "Quality",
               y = "Chlorides",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p5, p5w, nrow = 1)
```



Fixed Acidity Vs Quality

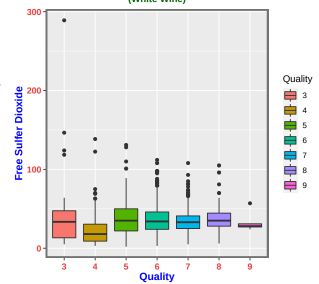
```
p6 <- ggplot(red_wine_df) +
          geom_boxplot(aes(x = factor(quality), y = free.sulfur.dioxide, fill = factor(quality))) +
          labs(title = "Relationship between Free Sulfur Dioxide & Quality",
               subtitle = "(Red Wine)",
               x = "Quality",
               y = "Free Sulfer Dioxide",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p6w <- ggplot(white_wine_df) +</pre>
          geom_boxplot(aes(x = factor(quality), y = free.sulfur.dioxide, fill = factor(quality))) +
```

```
labs(title = "Relationship between Free Sulfur Dioxide & Quality",
               subtitle = "(White Wine)",
               x = "Quality",
               y = "Free Sulfer Dioxide",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p6,p6w, nrow = 1)
```



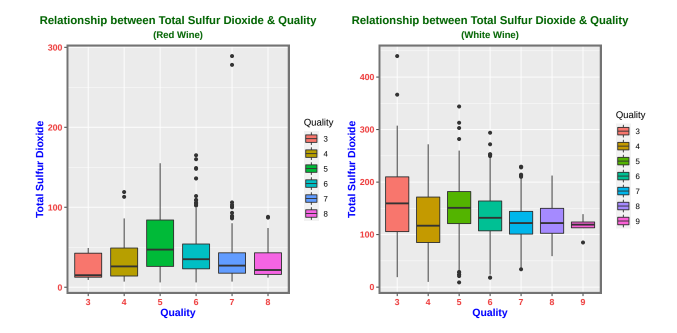
enter Dixord 40 approximation and the second secon

Relationship between Free Sulfur Dioxide & Quality (White Wine)



Total Sulfur Dioxide Vs Quality

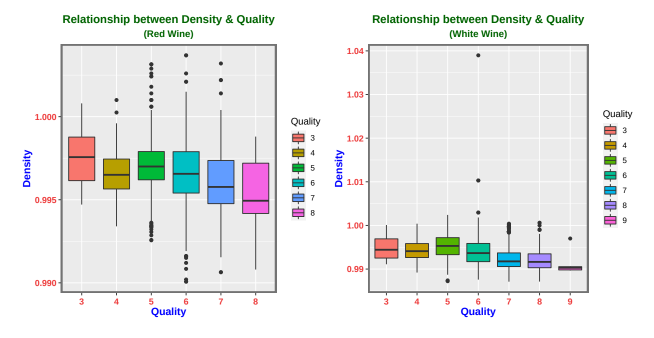
```
labs(title = "Relationship between Total Sulfur Dioxide & Quality",
               subtitle = "(Red Wine)",
               x = "Quality",
               y = "Total Sulfur Dioxide",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p7w <- ggplot(white wine df) +
          geom_boxplot(aes(x = factor(quality), y = total.sulfur.dioxide, fill = factor(quality))) +
          labs(title = "Relationship between Total Sulfur Dioxide & Quality",
               subtitle = "(White Wine)",
               x = "Quality",
               y = "Total Sulfur Dioxide",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p7,p7w, nrow = 1)
```



Density Vs Quality

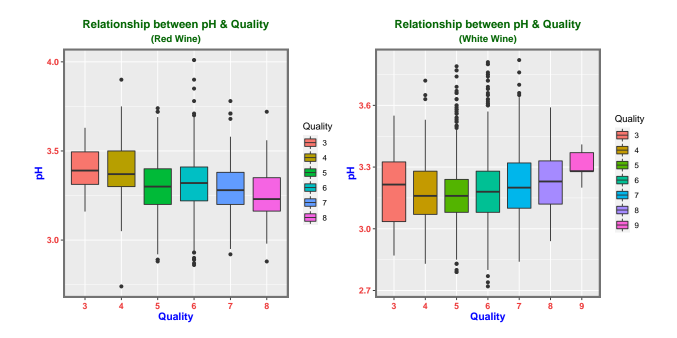
```
p8 <- ggplot(red_wine_df) +
          geom_boxplot(aes(x = factor(quality), y = density, fill = factor(quality))) +
          labs(title = "Relationship between Density & Quality",
               subtitle = "(Red Wine)",
               x = "Quality",
               y = "Density",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p8w <- ggplot(white_wine_df) +</pre>
          geom_boxplot(aes(x = factor(quality), y = density, fill = factor(quality))) +
```

```
labs(title = "Relationship between Density & Quality",
               subtitle = "(White Wine)",
               x = "Quality",
               y = "Density",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p8,p8w, nrow = 1)
```



pH Vs Quality

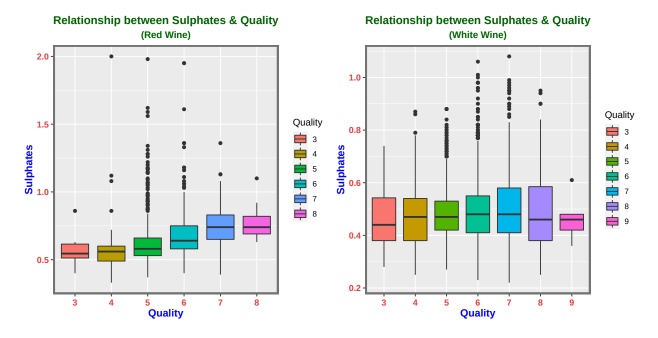
```
labs(title = "Relationship between pH & Quality",
               subtitle = "(Red Wine)",
               x = "Quality",
               v = "Hq" = v
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p9w <- ggplot(white wine df) +
          geom_boxplot(aes(x = factor(quality), y = pH, fill = factor(quality))) +
          labs(title = "Relationship between pH & Quality",
               subtitle = "(White Wine)",
               x = "Quality",
               y = "pH",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p9, p9w, nrow = 1)
```



Sulphates Vs Quality

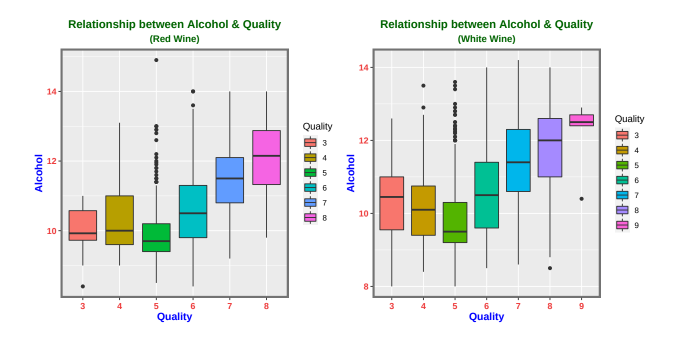
```
p10 <- ggplot(red_wine_df) +
          geom_boxplot(aes(x = factor(quality), y = sulphates, fill = factor(quality))) +
          labs(title = "Relationship between Sulphates & Quality",
               subtitle = "(Red Wine)",
               x = "Quality",
               y = "Sulphates",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p10w <- ggplot(white_wine_df) +
          geom_boxplot(aes(x = factor(quality), y = sulphates, fill = factor(quality))) +
          labs(title = "Relationship between Sulphates & Quality",
```

```
subtitle = "(White Wine)",
               x = "Quality",
               y = "Sulphates",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p10, p10w, nrow = 1)
```



Alcohol Vs Quality

```
subtitle = "(Red Wine)",
               x = "Quality",
               y = "Alcohol",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
p11w <- ggplot(white_wine_df) +
          geom_boxplot(aes(x = factor(quality), y = alcohol, fill = factor(quality))) +
          labs(title = "Relationship between Alcohol & Quality",
               subtitle = "(White Wine)",
               x = "Quality",
               y = "Alcohol",
               fill = "Quality")+
          theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=2))
grid.arrange(p11, p11w, nrow = 1)
```



Stepwise Model Selection for red wine

```
s1 <- step("quality", preds, cands, partition_rw)</pre>
model_rw <- c(model_rw, attr(s1, "best"))</pre>
##
          fixed.acidity
                             volatile.acidity
                                                         citric.acid
                                     0.7855048
##
              0.8515544
                                                           0.8398467
                                     chlorides free.sulfur.dioxide
##
         residual.sugar
##
              0.8578792
                                     0.8472211
                                                           0.8532622
## total.sulfur.dioxide
                                      density
                                                                  pН
##
              0.8401893
                                     0.8376948
                                                           0.8553145
##
              sulphates
                                       alcohol
##
              0.8296731
                                   0.7346630
## attr(,"best")
## alcohol
## 0.734663
preds <- "alcohol"</pre>
cands <- c("fixed.acidity","volatile.acidity","citric.acid","residual.sugar",</pre>
           "chlorides", "free.sulfur.dioxide", "total.sulfur.dioxide", "density",
           "pH", "sulphates")
s1 <- step("quality", preds, cands, partition_rw)</pre>
model_rw <- c(model_rw, attr(s1, "best"))</pre>
##
          fixed.acidity
                             volatile.acidity
                                                        citric.acid
##
              0.7226665
                                    0.6902853
                                                           0.7215572
         residual.sugar
##
                                     chlorides free.sulfur.dioxide
##
              0.7347169
                                    0.7342695
                                                         0.7340596
## total.sulfur.dioxide
                                      density
                                                                  pН
##
              0.7308255
                                    0.7322296
                                                         0.7188688
##
              sulphates
              0.7190395
##
## attr(,"best")
## volatile.acidity
          0.6902853
preds <- c("alcohol", "volatile.acidity")</pre>
cands <- c("fixed.acidity","citric.acid","residual.sugar",</pre>
           "chlorides", "free.sulfur.dioxide", "total.sulfur.dioxide", "density",
           "pH", "sulphates")
s1 <- step("quality", preds, cands, partition_rw)</pre>
model_rw <- c(model_rw, attr(s1, "best"))</pre>
s1
                                   citric.acid
##
          fixed.acidity
                                                      residual.sugar
##
              0.6878592
                                     0.6903075
                                                           0.6911506
##
              chlorides free.sulfur.dioxide total.sulfur.dioxide
##
              0.6902750
                                     0.6890341
                                                           0.6862892
##
                                                           sulphates
                 density
                                            рΗ
```

```
0.6892822
##
                                  0.6877442
                                                          0.6848121
## attr(,"best")
## sulphates
## 0.6848121
preds <- c("alcohol","volatile.acidity","sulphates")</pre>
cands <- c("fixed.acidity","citric.acid","residual.sugar",</pre>
           "chlorides", "free.sulfur.dioxide", "total.sulfur.dioxide", "density",
           "pH")
s1 <- step("quality", preds, cands, partition_rw)</pre>
model_rw <- c(model_rw, attr(s1, "best"))</pre>
##
          fixed.acidity
                                  citric.acid
                                                     residual.sugar
##
              0.6831365
                                    0.6846930
                                                          0.6857594
              chlorides free.sulfur.dioxide total.sulfur.dioxide
##
##
              0.6831365
                                    0.6830200
                                                          0.6794726
##
               density
                                           рΗ
##
              0.6849305
                                  0.6831672
## attr(,"best")
## total.sulfur.dioxide
              0.6794726
preds <- c("alcohol", "volatile.acidity", "sulphates", "total.sulfur.dioxide")</pre>
cands <- c("fixed.acidity","citric.acid","residual.sugar",</pre>
           "chlorides", "free.sulfur.dioxide", "density",
           ("Ha"
s1 <- step("quality", preds, cands, partition_rw)</pre>
model rw <- c(model rw, attr(s1, "best"))</pre>
         fixed.acidity
                              citric.acid
                                                  residual.sugar
                                                                            chlorides
##
             0.6784623
                                                  0.6805401
                                                                            0.6774864
##
                                0.6795727
## free.sulfur.dioxide
                                  densitv
                                                              Нq
                                0.6796242
                                                     0.6776286
            0.6806207
## attr(,"best")
## chlorides
## 0.6774864
preds <- c("alcohol","volatile.acidity","sulphates","total.sulfur.dioxide","chlorides")</pre>
cands <- c("fixed.acidity","citric.acid","residual.sugar",</pre>
           "free.sulfur.dioxide", "density",
           "pH")
s1 <- step("quality", preds, cands, partition_rw)</pre>
model_rw <- c(model_rw, attr(s1, "best"))</pre>
##
         fixed.acidity
                              citric.acid
                                                  residual.sugar free.sulfur.dioxide
```

0.6785556

0.6780041

0.6775808

##

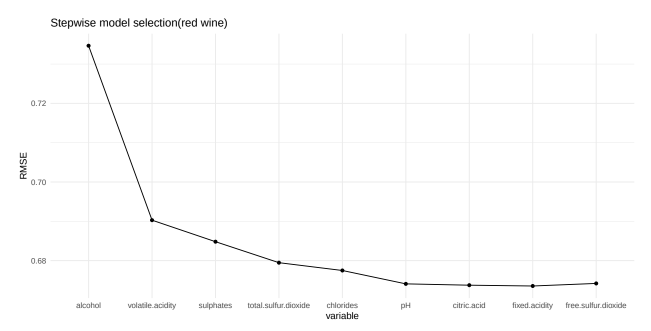
0.6760903

```
##
               density
                                           Нq
                                  0.6740840
##
             0.6776890
## attr(,"best")
##
         рΗ
## 0.674084
preds <- c("alcohol","volatile.acidity","sulphates","total.sulfur.dioxide","chlorides","pH")</pre>
cands <- c("fixed.acidity","citric.acid","residual.sugar",</pre>
            "free.sulfur.dioxide", "density")
s1 <- step("quality", preds, cands, partition_rw)</pre>
model_rw <- c(model_rw, attr(s1, "best"))</pre>
##
         fixed.acidity
                                 citric.acid
                                                   residual.sugar free.sulfur.dioxide
##
             0.6741904
                                  0.6737470
                                                         0.6752222
                                                                              0.6748061
##
                density
##
             0.6747926
## attr(,"best")
## citric.acid
##
      0.673747
preds <- c("alcohol","volatile.acidity","sulphates","total.sulfur.dioxide",</pre>
           "chlorides", "pH", "citric.acid")
cands <- c("fixed.acidity", "residual.sugar",</pre>
           "free.sulfur.dioxide", "density")
s1 <- step("quality", preds, cands, partition_rw)</pre>
model_rw <- c(model_rw, attr(s1, "best"))</pre>
##
         fixed.acidity
                              residual.sugar free.sulfur.dioxide
                                                                                 density
             0.6735421
                                   0.6750824
                                                        0.6743678
                                                                              0.6738133
## attr(,"best")
## fixed.acidity
       0.6735421
##
preds <- c("alcohol","volatile.acidity","sulphates","total.sulfur.dioxide",</pre>
           "chlorides", "pH", "citric.acid", "fixed.acidity")
cands <- c("residual.sugar", "free.sulfur.dioxide", "density")</pre>
s1 <- step("quality", preds, cands, partition_rw)</pre>
model_rw <- c(model_rw, attr(s1, "best"))</pre>
##
        residual.sugar free.sulfur.dioxide
                                                           density
##
             0.6749624
                                   0.6741934
                                                         0.6742341
## attr(,"best")
## free.sulfur.dioxide
             0.6741934
```

Model stopped improving at:

• fit = quality \sim alcohol + volatile acidity + sulphates + total sulfur dioxide + chlorides + pH + citric acid + fixed acidity

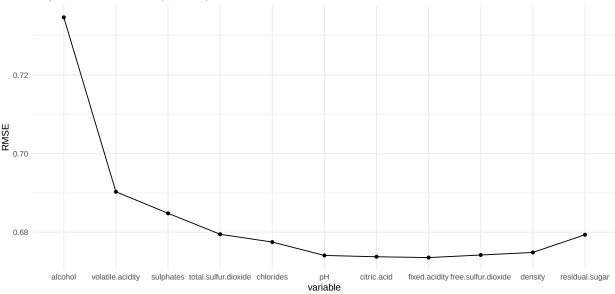
Visualizing how adding each variable affects the RMSE.



visualizing how each variable(including the predictors) affects the RMSE for Red wine

```
preds <- c("alcohol","volatile.acidity","sulphates","total.sulfur.dioxide",</pre>
            "chlorides", "pH", "citric.acid", "fixed.acidity", "free.sulfur.dioxide",
           "density")
cands <- c("residual.sugar")</pre>
s1 <- step("quality", preds, cands, partition_rw)</pre>
temp_model_rw <- c(temp_model_rw, attr(s1, "best"))</pre>
## residual.sugar
        0.6793522
##
## attr(,"best")
## residual.sugar
        0.6793522
##
#Representation of RMSE for all the variables
step_model <- tibble(index=seq_along(temp_model_rw),</pre>
                      variable=factor(names(temp_model_rw),
                                       levels=names(temp_model_rw)),
                      RMSE=temp_model_rw)
ggplot(step_model, aes(y=RMSE)) +
  geom_point(aes(x=variable)) +
  geom_line(aes(x=index)) +
  labs(title="Stepwise model selection(red wine)") +
  theme_minimal()
```

Stepwise model selection(red wine)



Consider the following fits and extract the best fit model:

- fit1 <- quality ~ alcohol + volatile acidity + sulphates + total sulfur dioxide + chlorides + pH
- fit2 <- quality \sim alcohol + volatile acidity + sulphates + total sulfur dioxide + chlorides + pH + citric acid
- fit 3 <- quality \sim alcohol + volatile acidity + sulphates + total sulfur dioxide + chlorides + pH + citric acid + fixed acidity

Cross Validation

```
set.seed(2020)
#partition_rw_train <- red_wine_std[-partition_rw$test$idx,]</pre>
#redwine_cv <- crossv_kfold(partition_rw_train, 5)</pre>
#redwine cv
redwine_cv <- crossv_kfold(red_wine_std, 5)</pre>
redwine cv
## # A tibble: 5 x 3
     train
                                                      .id
                              test
##
                                                     <chr>>
     <named list>
                              <named list>
## 1 <resample [1,279 x 12] > <resample [320 x 12] > 1
## 2 <resample [1,279 x 12] > <resample [320 x 12] > 2
## 3 <resample [1,279 x 12] > <resample [320 x 12] > 3
## 4 <resample [1,279 x 12] > <resample [320 x 12] > 4
## 5 <resample [1,280 x 12] > <resample [319 x 12] > 5
#Calculating RMSE for each fold of data
cv_rw <- redwine_cv %>%
 mutate(fit = purrr::map(train,
                    ~ lm(quality ~ alcohol + volatile.acidity + sulphates +
                           total.sulfur.dioxide + chlorides + pH, data = .)),
         rmse = purrr::map2_dbl(fit, test, ~ rmse(.x, .y)))
cv_rw
## # A tibble: 5 x 5
##
     train
                              test
                                                     .id
                                                            fit.
                                                                           rmse
     <named list>
                              <named list>
                                                     <chr> <named list> <dbl>
## 1 <resample [1,279 x 12] > <resample [320 x 12] > 1
                                                            <1m>
                                                                          0.673
## 2 <resample [1,279 x 12] > <resample [320 x 12] > 2
                                                            <1m>
                                                                         0.659
## 3 <resample [1,279 x 12] > <resample [320 x 12] > 3
                                                            <1m>
                                                                         0.587
## 4 <resample [1,279 x 12] > <resample [320 x 12] > 4
                                                            <1m>
                                                                         0.670
## 5 <resample [1,280 x 12] > <resample [319 x 12] > 5
                                                            <1m>
                                                                         0.666
#Average of RMSEs
mean(cv_rw$rmse)
## [1] 0.650961
```

Comparing models using CV

Calling the function

```
fit1_rmse <- do_redwine_cv(quality ~ alcohol + volatile.acidity + sulphates +
               total.sulfur.dioxide + chlorides + pH )
fit2_rmse <- do_redwine_cv(quality ~ alcohol + volatile.acidity + sulphates +
               total.sulfur.dioxide + chlorides + pH + citric.acid)
fit3_rmse <- do_redwine_cv(quality ~ alcohol + volatile.acidity + sulphates +
               total.sulfur.dioxide + chlorides + pH + citric.acid + fixed.acidity)
fit1_rmse
## # A tibble: 1 x 3
   cv_rmse cv_rsq cv_mae
      <dbl> <dbl> <dbl>
## 1 0.651 0.349 0.506
fit2_rmse
## # A tibble: 1 x 3
   cv_rmse cv_rsq cv_mae
      <dbl> <dbl> <dbl>
## 1 0.651 0.349 0.506
fit3_rmse
## # A tibble: 1 x 3
    cv_rmse cv_rsq cv_mae
      <dbl> <dbl> <dbl>
## 1 0.651 0.348 0.507
```

Best fit model:

• quality \sim alcohol + volatile.acidity + sulphates + total.sulfur.dioxide + chlorides + pH

Stepwise Model Selection for white wine

```
#Partition of data
white_wine_std <- data.frame(scale(white_wine_df[1:11]))
white_wine_std$quality <- white_wine_df$quality</pre>
```

```
set.seed(10)
partition_ww <- resample_partition(white_wine_std,</pre>
                                  p=c(train=0.5,
                                      valid=0.25,
                                      test=0.25))
model_ww <- NULL
preds <- "1"
cands <- c("fixed.acidity","volatile.acidity","citric.acid","residual.sugar",</pre>
            "chlorides", "free.sulfur.dioxide", "total.sulfur.dioxide", "density",
            "pH", "sulphates", "alcohol")
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
##
          fixed.acidity
                              volatile.acidity
                                                         citric.acid
##
               0.9046956
                                    0.9040842
                                                           0.9115613
##
         residual.sugar
                                     chlorides free.sulfur.dioxide
               0.9083983
                                     0.8910796
                                                           0.9115526
##
## total.sulfur.dioxide
                                                                   рΗ
                                       density
##
               0.8967295
                                     0.8741188
                                                           0.9066175
##
               sulphates
                                       alcohol
               0.9097914
                                   0.8153880
##
## attr(,"best")
## alcohol
## 0.815388
preds <- "alcohol"</pre>
cands <- c("fixed.acidity","volatile.acidity","citric.acid","residual.sugar",</pre>
            "chlorides", "free.sulfur.dioxide", "total.sulfur.dioxide", "density",
            "pH", "sulphates")
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
##
          fixed.acidity
                              volatile.acidity
                                                         citric.acid
                                     0.8013749
##
               0.8129728
                                                           0.8146745
                                     chlorides free.sulfur.dioxide
         residual.sugar
               0.8095793
                                     0.8142981
                                                         0.8090157
## total.sulfur.dioxide
                                       density
                                                                   Нq
                                                          0.8140691
##
              0.8153080
                                    0.8141830
##
              sulphates
               0.8129943
##
## attr(,"best")
## volatile.acidity
##
          0.8013749
preds <- c("alcohol", "volatile.acidity")</pre>
cands <- c("fixed.acidity","citric.acid","residual.sugar",</pre>
```

```
"chlorides", "free.sulfur.dioxide", "total.sulfur.dioxide", "density",
           "pH", "sulphates")
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
##
          fixed.acidity
                                   citric.acid
                                                      residual.sugar
##
              0.7986474
                                     0.8022197
                                                           0.7901090
              chlorides free.sulfur.dioxide total.sulfur.dioxide
##
##
              0.8008642
                                     0.7956051
                                                           0.8001968
##
                                                           sulphates
                 density
                                            рΗ
##
              0.7969973
                                   0.8006140
                                                           0.7994550
## attr(,"best")
## residual.sugar
##
         0.790109
preds <- c("alcohol", "volatile.acidity", "residual.sugar")</pre>
cands <- c("fixed.acidity","citric.acid",</pre>
           "chlorides", "free.sulfur.dioxide", "total.sulfur.dioxide", "density",
           "pH", "sulphates")
s2 <- step("quality", preds, cands, partition_ww)</pre>
model ww <- c(model ww, attr(s2, "best"))</pre>
s2
##
          fixed.acidity
                                   citric.acid
                                                           chlorides
              0.7866903
                                     0.7912216
                                                           0.7899837
##
## free.sulfur.dioxide total.sulfur.dioxide
                                                              density
##
              0.7870588
                                    0.7900478
                                                           0.7885526
##
                                    sulphates
                      рΗ
              0.7880903
                                   0.7879418
##
## attr(,"best")
## fixed.acidity
       0.7866903
preds <- c("alcohol","volatile.acidity","residual.sugar","fixed.acidity")</pre>
cands <- c("citric.acid", "chlorides", "free.sulfur.dioxide",</pre>
           "total.sulfur.dioxide", "density", "pH", "sulphates")
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
s2
##
            citric.acid
                                     chlorides free.sulfur.dioxide
              0.7875729
                                     0.7865390
                                                         0.7842173
## total.sulfur.dioxide
                                       density
                                                                   pН
##
              0.7865822
                                     0.7873280
                                                          0.7861562
##
              sulphates
##
              0.7845328
## attr(,"best")
## free.sulfur.dioxide
            0.7842173
##
```

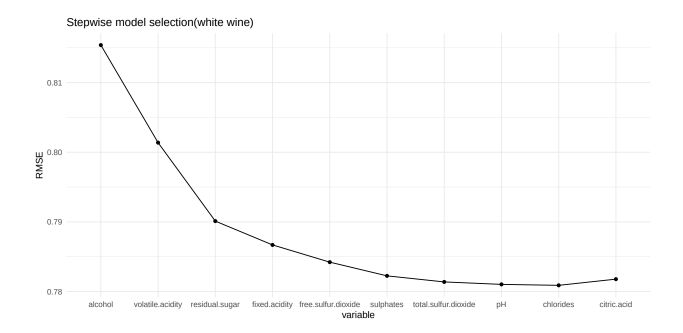
```
preds <- c("alcohol","volatile.acidity","residual.sugar","fixed.acidity",</pre>
           "free.sulfur.dioxide")
cands <- c("citric.acid", "chlorides",</pre>
           "total.sulfur.dioxide", "density", "pH", "sulphates")
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
                                     chlorides total.sulfur.dioxide
##
            citric.acid
                                     0.7840036
##
              0.7850955
                                                           0.7837253
##
                                                           sulphates
                 density
                                   0.7837374
              0.7851699
                                                           0.7822481
## attr(,"best")
## sulphates
## 0.7822481
preds <- c("alcohol","volatile.acidity","residual.sugar","fixed.acidity",</pre>
           "free.sulfur.dioxide", "sulphates")
cands <- c("citric.acid", "chlorides",</pre>
           "total.sulfur.dioxide", "density", "pH")
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
            citric.acid
                                     chlorides total.sulfur.dioxide
##
##
              0.7831977
                                     0.7820405
                                                           0.7813755
##
                 density
                                   0.7820393
              0.7827787
## attr(,"best")
## total.sulfur.dioxide
              0.7813755
preds <- c("alcohol", "volatile.acidity", "residual.sugar", "fixed.acidity",</pre>
           "free.sulfur.dioxide", "sulphates", "total.sulfur.dioxide")
cands <- c("citric.acid","chlorides",</pre>
            "density", "pH")
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
## citric.acid chlorides
                                 density
                                                   рΗ
   0.7822786
                  0.7811962
                               0.7824629 0.7810240
## attr(,"best")
##
         pН
## 0.781024
preds <- c("alcohol","volatile.acidity","residual.sugar","fixed.acidity",</pre>
            "free.sulfur.dioxide", "sulphates", "total.sulfur.dioxide", "pH")
cands <- c("citric.acid","chlorides","density")</pre>
```

```
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
## citric.acid
                  chlorides
                                  density
                  0.7808872
                               0.7861522
     0.7819109
## attr(,"best")
## chlorides
## 0.7808872
preds <- c("alcohol","volatile.acidity","residual.sugar","fixed.acidity",</pre>
            "free.sulfur.dioxide", "sulphates", "total.sulfur.dioxide",
            "pH", "chlorides")
cands <- c("citric.acid", "density")</pre>
s2 <- step("quality", preds, cands, partition_ww)</pre>
model_ww <- c(model_ww, attr(s2, "best"))</pre>
## citric.acid
                    density
##
     0.7817750
                  0.7866456
## attr(,"best")
## citric.acid
##
      0.781775
```

Model stopped improving at:

• fit = quality \sim alcohol + volatile acidity + residual sugar + fixed acidity + free sulfur dioxide + sulphates + total sulfur dioxide + pH + chlorides

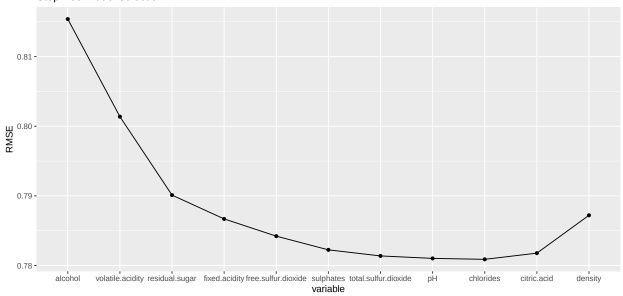
Visualizing how adding each variable affects the RMSE.



visualizing how each variable(including the predictors) affects the RMSE for Red wine

```
temp_model_ww <- model_ww</pre>
preds <- c("alcohol","volatile.acidity","residual.sugar","fixed.acidity",</pre>
            "free.sulfur.dioxide", "sulphates", "total.sulfur.dioxide",
            "pH", "chlorides", "citric.acid")
cands <- c("density")</pre>
s2 <- step("quality", preds, cands, partition_ww)</pre>
temp_model_ww <- c(temp_model_ww, attr(s2, "best"))</pre>
s2
     density
##
## 0.7872114
## attr(,"best")
     density
## 0.7872114
step_model_ww <- tibble(index=seq_along(temp_model_ww),</pre>
                      variable=factor(names(temp_model_ww), levels=names(temp_model_ww)),
                      RMSE=temp_model_ww)
ggplot(step_model_ww, aes(y=RMSE)) +
  geom_point(aes(x=variable)) +
  geom_line(aes(x=index)) +
  labs(title="Stepwise model selection")
```

Stepwise model selection



Consider the following fits and extract the best fit model:

- fit1 <- quality \sim alcohol + volatile.acidity + residual.sugar + fixed.acidity + free.sulfur.dioxide + sulphates + total.sulfur.dioxide + pH
- fit2 <- quality \sim alcohol + volatile.acidity + residual.sugar + fixed.acidity + free.sulfur.dioxide + sulphates + total.sulfur.dioxide + pH + chlorides

Cross Validation

```
set.seed(2020)
#partition_ww_train <- white_wine_std[-partition_ww$test$idx,]</pre>
#whitewine_cv <- crossv_kfold(partition_ww_train, 5)</pre>
#whitewine_cv
whitewine_cv <- crossv_kfold(white_wine_std, 5)</pre>
whitewine_cv
## # A tibble: 5 x 3
##
                                                      .id
     train
                               test
     <named list>
                               <named list>
## 1 <resample [3,918 x 12] > <resample [980 x 12] > 1
## 2 <resample [3,918 x 12] > <resample [980 x 12] > 2
## 3 <resample [3,918 x 12] > <resample [980 x 12] > 3
## 4 <resample [3,919 x 12] > <resample [979 x 12] > 4
## 5 <resample [3,919 x 12] > <resample [979 x 12] > 5
#Calculating RMSE for each fold of data
cv_rw <- whitewine_cv %>%
 mutate(fit = purrr::map(train,
```

~ lm(quality ~ alcohol + volatile.acidity + residual.sugar +

```
fixed.acidity + free.sulfur.dioxide +
                              sulphates + total.sulfur.dioxide + pH, data = .)),
         rmse = purrr::map2_dbl(fit, test, ~ rmse(.x, .y)))
cv_rw
## # A tibble: 5 x 5
##
    train
                                                    .id
                                                          fit
                             test
                                                                         rmse
                             <named list>
##
     <named list>
                                                    <chr> <named list> <dbl>
## 1 <resample [3,918 x 12] > <resample [980 x 12] > 1
                                                          <1m>
                                                                        0.747
## 2 <resample [3,918 x 12] > <resample [980 x 12] > 2
                                                          <1m>
                                                                        0.806
## 3 <resample [3,918 x 12] > <resample [980 x 12] > 3
                                                          <1m>
                                                                        0.743
## 4 <resample [3,919 x 12] > <resample [979 x 12] > 4
                                                          <1m>
                                                                        0.727
## 5 <resample [3,919 x 12] > <resample [979 x 12] > 5
                                                          <1m>
                                                                        0.762
#Average of RMSEs
mean(cv_rw$rmse)
```

Comparing models using CV

[1] 0.7568375

Calling the function

```
## # A tibble: 1 x 3
## cv_rmse cv_rsq cv_mae
## <dbl> <dbl> <dbl> <dbl> ## 1 0.757 0.269 0.588
```

```
fit2_rmse_ww
## # A tibble: 1 x 3
     cv_rmse cv_rsq cv_mae
##
      <dbl> <dbl> <dbl>
## 1
      0.757 0.269 0.588
Best fit Model
  • quality ~ alcohol + volatile.acidity + residual.sugar + fixed.acidity + free.sulfur.dioxide + sulphates
    + total.sulfur.dioxide + pH
fit_rw <- lm(quality ~ alcohol + volatile.acidity + sulphates + total.sulfur.dioxide + chlorides + pH,
summary(fit_rw)
##
## Call:
## lm(formula = quality ~ alcohol + volatile.acidity + sulphates +
       total.sulfur.dioxide + chlorides + pH, data = red_wine_std)
##
## Residuals:
##
       Min
                  1Q
                       Median
## -2.60575 -0.35883 -0.04806 0.46079 1.95643
##
## Coefficients:
##
                        Estimate Std. Error t value Pr(>|t|)
                                   0.01622 347.419 < 2e-16 ***
## (Intercept)
                        5.63602
## alcohol
                         0.30976
                                    0.01791 17.291 < 2e-16 ***
## volatile.acidity
                        -0.18590 0.01798 -10.338 < 2e-16 ***
## sulphates
                                    0.01865
                                            8.076 1.31e-15 ***
                         0.15064
## total.sulfur.dioxide -0.07803
                                    0.01666 -4.684 3.05e-06 ***
## chlorides
                        -0.09424
                                    0.01874 -5.030 5.46e-07 ***
## pH
                        -0.06719
                                    0.01791 -3.750 0.000183 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' '1
## Residual standard error: 0.6487 on 1592 degrees of freedom
## Multiple R-squared: 0.3572, Adjusted R-squared: 0.3548
## F-statistic: 147.4 on 6 and 1592 DF, p-value: < 2.2e-16
fit_ww <- lm(quality ~ alcohol + volatile.acidity + residual.sugar + fixed.acidity + free.sulfur.dioxid
summary(fit_ww)
##
## Call:
## lm(formula = quality ~ alcohol + volatile.acidity + residual.sugar +
##
       fixed.acidity + free.sulfur.dioxide + sulphates + total.sulfur.dioxide +
      pH, data = white_wine_std)
##
##
```

```
## Residuals:
              1Q Median
      Min
                             30
                                   Max
## -3.8931 -0.4982 -0.0358 0.4644 3.1821
## Coefficients:
##
                      Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                      5.87791 0.01080 544.038 < 2e-16 ***
                                 0.01304 34.834 < 2e-16 ***
                      0.45433
## alcohol
## volatile.acidity
                     -0.19887
                                0.01126 -17.665 < 2e-16 ***
                                0.01283 10.349 < 2e-16 ***
## residual.sugar
                      0.13274
## fixed.acidity
                     -0.04222
                                 0.01220 -3.461 0.000542 ***
## free.sulfur.dioxide 0.08078
                                 0.01427 5.662 1.58e-08 ***
                      0.04736
                                0.01108 4.273 1.97e-05 ***
## sulphates
## total.sulfur.dioxide -0.03866
                                0.01581 -2.445 0.014529 *
## pH
                      ## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 0.7561 on 4889 degrees of freedom
## Multiple R-squared: 0.2722, Adjusted R-squared: 0.2711
## F-statistic: 228.6 on 8 and 4889 DF, p-value: < 2.2e-16
```

Random Forest Regression

```
#Red wine
install.packages("randomForest")
library("randomForest")
fit_rw_rf <- randomForest(quality ~ alcohol + volatile.acidity +</pre>
                            sulphates + total.sulfur.dioxide +
                            chlorides + pH, red_wine_std,
                          mtry = 3,
                           importance = TRUE,
                          na.action = na.omit)
fit_rw_rf
##
## Call:
   randomForest(formula = quality ~ alcohol + volatile.acidity +
                                                                         sulphates + total.sulfur.dioxide
                  Type of random forest: regression
##
##
                        Number of trees: 500
## No. of variables tried at each split: 3
##
             Mean of squared residuals: 0.3207565
##
                       % Var explained: 50.79
#White Wine - Random forest regression
fit_ww_rf <- randomForest(quality ~ alcohol + volatile.acidity +</pre>
                            residual.sugar + fixed.acidity +
```

free.sulfur.dioxide + sulphates +

mtry = 3,

total.sulfur.dioxide + pH, white_wine_std,

```
importance = TRUE,
                          na.action = na.omit)
fit_ww_rf
##
## Call:
  randomForest(formula = quality ~ alcohol + volatile.acidity +
                                                                      residual.sugar + fixed.acidity +
                 Type of random forest: regression
##
##
                       Number of trees: 500
## No. of variables tried at each split: 3
##
            Mean of squared residuals: 0.3529432
##
##
                      % Var explained: 54.99
Ridge Regression
#Ridge regression for red wine
set.seed(1)
train <- createDataPartition(red_wine_std$quality, p=0.6, list=FALSE)
table(red_wine_std$quality[train])
        4 5 6 7
     5 35 407 383 119 12
```

glmnet

961 samples

6 predictor

Pre-processing: centered (6), scaled (6)

Resampling: Cross-Validated (10 fold, repeated 10 times)

##

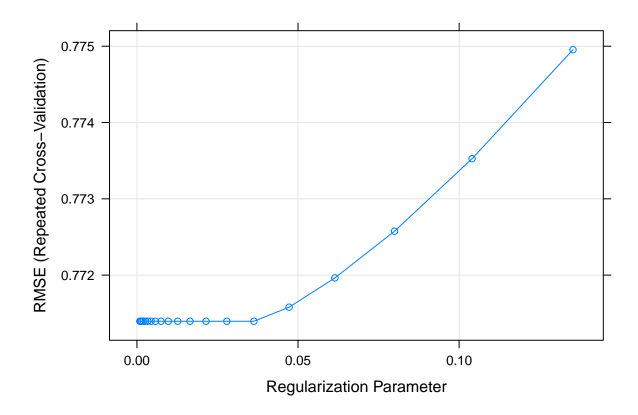
##

##

```
## Summary of sample sizes: 866, 865, 865, 865, 865, 865, ...
## Resampling results across tuning parameters:
##
##
   lambda
                   Rsquared
            RMSE
                           MAE
##
   ##
   0.001543529  0.6608113  0.3426984  0.5170648
##
   0.002008180 0.6608113 0.3426984 0.5170648
##
   ##
##
   ##
   ##
   0.005753800 0.6608113 0.3426984 0.5170648
##
   ##
##
   ##
   0.016485680 0.6608113 0.3426984 0.5170648
##
   ##
   ##
   ##
   ##
   ##
   0.079953042  0.6612124  0.3428183  0.5188810
   ##
   ##
##
## Tuning parameter 'alpha' was held constant at a value of 0
## RMSE was used to select the optimal model using the smallest value.
## The final values used for the model were alpha = 0 and lambda = 0.02790506.
#Ridge regression for white wine
train_w <- createDataPartition(white_wine_std$quality, p=0.6, list=FALSE)
table(white_wine_std$quality[train_w])
##
##
                        9
    3
                 7
                     8
          5
              6
     101 869 1319 518 113
##
whitedf_train <- white_wine_std[as.integer(train_w),]</pre>
whitedf_test <- white_wine_std[-as.integer(train_w),]</pre>
ctrl <- trainControl(method="repeatedcv", number=10, repeats=10)</pre>
grd <- expand.grid(lambda=exp(seq(from=-7, to=-2, length.out=20)),</pre>
            alpha=0)
set.seed(1)
fit_ww_ridge <- train(quality ~ alcohol + volatile.acidity +</pre>
                   residual.sugar + fixed.acidity +
                   free.sulfur.dioxide + sulphates +
                   total.sulfur.dioxide + pH, data=whitedf_train,
         method="glmnet",
```

```
preProcess=c("center", "scale"),
             trControl=ctrl, tuneGrid=grd)
fit_ww_ridge
## glmnet
##
## 2939 samples
##
     8 predictor
## Pre-processing: centered (8), scaled (8)
## Resampling: Cross-Validated (10 fold, repeated 10 times)
## Summary of sample sizes: 2646, 2645, 2644, 2644, 2645, 2645, ...
## Resampling results across tuning parameters:
##
##
    lambda
                 RMSE
                           Rsquared
                                      MAE
##
    0.000911882 0.7713956 0.2644750 0.5963063
##
    0.001186388 0.7713956 0.2644750 0.5963063
##
    0.001543529 0.7713956
                           0.2644750
                                      0.5963063
##
    0.002008180 0.7713956 0.2644750 0.5963063
##
    0.002612707 0.7713956 0.2644750
                                     0.5963063
    0.003399216 0.7713956 0.2644750
                                      0.5963063
##
##
    0.004422489
                 0.7713956
                           0.2644750
                                      0.5963063
##
    0.005753800 0.7713956 0.2644750 0.5963063
##
    0.007485879 0.7713956 0.2644750
                                      0.5963063
##
    0.009739369 0.7713956 0.2644750
                                     0.5963063
##
    0.012671232 0.7713956 0.2644750
                                      0.5963063
##
    0.016485680 0.7713956 0.2644750 0.5963063
##
    0.021448399 0.7713956 0.2644750 0.5963063
##
    0.027905057 0.7713956 0.2644750 0.5963063
##
    ##
    0.047234459 0.7715800 0.2644133 0.5964922
##
    0.061453549 0.7719637 0.2642660
                                     0.5968524
##
    0.079953042 0.7725760
                           0.2640283
                                      0.5973675
##
    0.104021477 0.7735260 0.2636500
                                      0.5981261
##
    0.135335283 0.7749544 0.2630627
                                      0.5992021
##
## Tuning parameter 'alpha' was held constant at a value of \mathbf{0}
## RMSE was used to select the optimal model using the smallest value.
## The final values used for the model were alpha = 0 and lambda = 0.03630538.
```

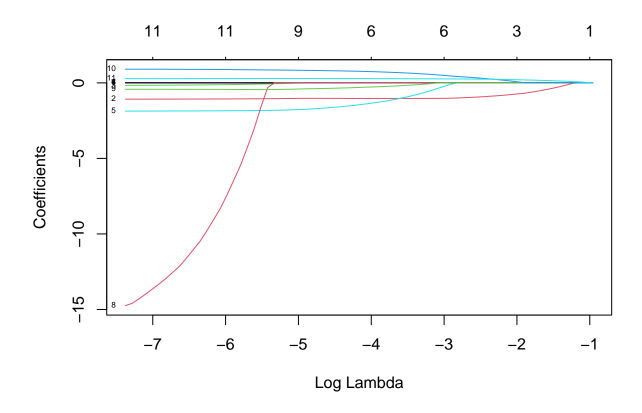
plot(fit_ww_ridge,xvar="lambda",label=TRUE)



Lasso Regression

```
#Lasso regression for red wine
ctrl <- trainControl(method="repeatedcv", number=10, repeats=10)</pre>
grd <- expand.grid(lambda=exp(seq(from=-7, to=-2, length.out=20)),</pre>
                    alpha=1)
set.seed(1)
fit_rw_lasso <- train(quality ~ alcohol + volatile.acidity +</pre>
                             sulphates + total.sulfur.dioxide +
                             chlorides + pH, data=reddf_train,
              method="glmnet",
              preProcess=c("center", "scale"),
               trControl=ctrl, tuneGrid=grd)
fit_rw_lasso
## glmnet
##
## 961 samples
##
     6 predictor
## Pre-processing: centered (6), scaled (6)
```

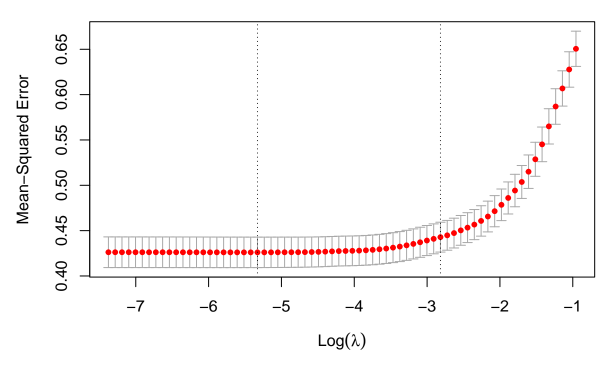
```
## Resampling: Cross-Validated (10 fold, repeated 10 times)
## Summary of sample sizes: 866, 865, 865, 865, 865, 865, ...
## Resampling results across tuning parameters:
##
##
   lambda
             RMSE
                     Rsquared
##
   0.000911882    0.6609175    0.3424848    0.5159115
   0.001186388    0.6609175    0.3424847    0.5159115
##
   ##
##
   ##
   ##
   0.003399216  0.6609532  0.3424168  0.5162430
##
   0.004422489 0.6609853 0.3423684 0.5164219
##
   0.005753800 0.6610418 0.3422918 0.5166667
   ##
##
   ##
   ##
   0.016485680 \quad 0.6621144 \quad 0.3410409 \quad 0.5191288
##
   ##
   ##
##
   0.047234459 \quad 0.6706015 \quad 0.3302082 \quad 0.5312242
##
   0.079953042 \quad 0.6790352 \quad 0.3242652 \quad 0.5433772
##
##
   0.5528960
##
   ## Tuning parameter 'alpha' was held constant at a value of 1
## RMSE was used to select the optimal model using the smallest value.
## The final values used for the model were alpha = 1 and lambda = 0.000911882.
actual_quality_rw<-red_wine_df$quality</pre>
response_rw<-red_wine_df$quality
predictors_rw<-data.matrix(red_wine_df[,c("fixed.acidity","volatile.acidity","citric.acid",</pre>
                   "residual.sugar", "chlorides", "free.sulfur.dioxide",
                   "total.sulfur.dioxide", "density", "pH", "sulphates",
                   "alcohol")])
red_ls_model<-glmnet(predictors_rw,response_rw,alpha=1)</pre>
plot(red_ls_model,xvar="lambda",label=TRUE)
```



```
# Using k-fold cv to get best optimal lambda value
red_cv_ls_model<-cv.glmnet(predictors_rw,response_rw,alpha=1)

#produce plot of MSE values for each lambda
plot(red_cv_ls_model)</pre>
```

11 11 11 11 9 8 7 7 7 7 4 3 3 2 2 0



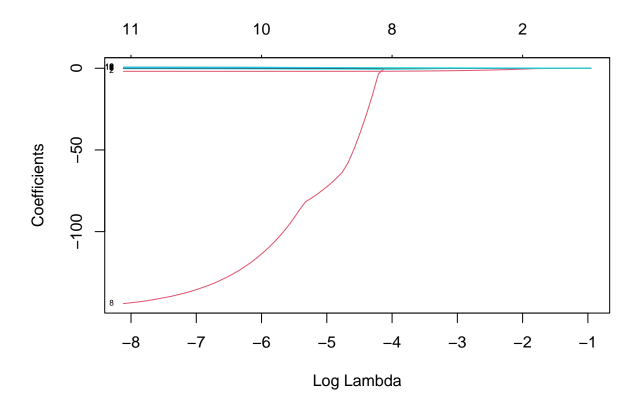
```
#find optimal lambda value that minimizes the MSE
best_lambda_rw<-red_cv_ls_model$lambda.min
best_lambda_rw</pre>
```

[1] 0.004850794

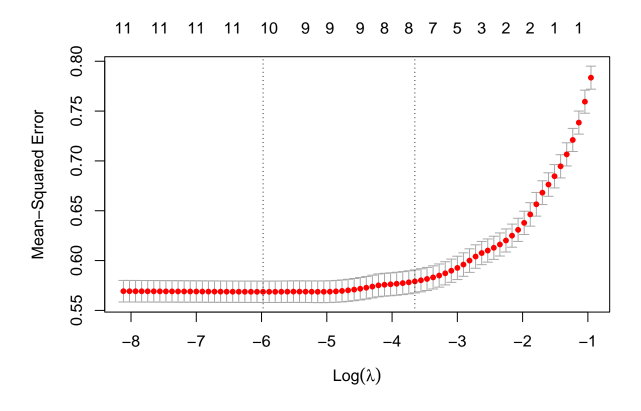
best_model_ls_rw<-glmnet(predictors_rw,response_rw,alpha=1,lambda=best_lambda_rw)
coef(best_model_ls_rw)</pre>

```
## 12 x 1 sparse Matrix of class "dgCMatrix"
##
                                   s0
## (Intercept)
                         4.311058572
## fixed.acidity
                         0.001590786
## volatile.acidity
                        -1.045885442
## citric.acid
                        -0.056043048
## residual.sugar
                         0.004545097
## chlorides
                        -1.812127522
## free.sulfur.dioxide
                         0.003379133
## total.sulfur.dioxide -0.002981089
## density
                        -0.437661346
## pH
## sulphates
                         0.850821345
## alcohol
                         0.287823262
```

```
# To find the rsquare of our best model
#To obtain predicted values
predicted_quality_rw<-predict(best_model_ls_rw, s=best_lambda_rw, newx=predictors_rw)</pre>
\#Finding\ the\ SST\ and\ SSE\ and\ Rsquare
sst_rw<-sum((actual_quality_rw-mean(actual_quality_rw))^2)</pre>
sse_rw<-sum((actual_quality_rw-predicted_quality_rw)^2)</pre>
rsq_ls_rw<-1-(sse_rw/sst_rw)
rsq_ls_rw
## [1] 0.3595207
#Finding the RMSE and MAE
n_rw<-nrow(red_wine_df)</pre>
rmse_ls_rw<-sqrt(sum((actual_quality_rw-predicted_quality_rw)^2)/n_rw)</pre>
rmse_ls_rw
## [1] 0.6460953
mae_ls_rw<-sum(abs(actual_quality_rw-predicted_quality_rw))/n_rw</pre>
mae_ls_rw
## [1] 0.5020461
view(data.frame(RMSE=rmse_ls_rw,MAE=mae_ls_rw,R_Square=rsq_ls_rw))
# Lasso Regression on White wine
response_ww<-white_wine_df$quality
actual_quality_ww<-white_wine_df$quality</pre>
predictors_ww<-data.matrix(white_wine_df[,c("fixed.acidity","volatile.acidity","citric.acid",</pre>
                              "residual.sugar", "chlorides", "free.sulfur.dioxide",
                              "total.sulfur.dioxide", "density", "pH", "sulphates",
                              "alcohol")])
white_ls_model<-glmnet(predictors_ww,response_ww,alpha=1)</pre>
plot(white_ls_model,xvar="lambda",label=TRUE)
```



```
# Using k-fold cv to get best optimal lambda value
library("glmnet")
white_cv_ls_model<-cv.glmnet(predictors_ww,response_ww,alpha=1)
plot(white_cv_ls_model)</pre>
```



```
#find optimal lambda value that minimizes the MSE
best_lambda_ww<-white_cv_ls_model$lambda.min
best_lambda_ww</pre>
```

[1] 0.002537796

best_model_ls_ww<-glmnet(predictors_ww,response_ww,alpha=1,lambda=best_lambda_ww)
coef(best_model_ls_ww)</pre>

```
## 12 x 1 sparse Matrix of class "dgCMatrix"
##
                                    s0
## (Intercept)
                         1.135530e+02
## fixed.acidity
                         3.015296e-02
## volatile.acidity
                        -1.875380e+00
## citric.acid
## residual.sugar
                         6.637850e-02
## chlorides
                        -3.934651e-01
## free.sulfur.dioxide
                         3.599613e-03
## total.sulfur.dioxide -2.344057e-04
## density
                        -1.129171e+02
## pH
                         5.208469e-01
## sulphates
                         5.556340e-01
## alcohol
                         2.329256e-01
```

```
# To find the rsquare of our best model
#To obtain predicted values
predicted_quality_ww<-predict(best_model_ls_ww,s=best_lambda_ww,newx=predictors_ww)</pre>
#find the SST and SSE
sst_ww<-sum((actual_quality_ww-mean(actual_quality_ww))^2)</pre>
sse_ww<-sum((actual_quality_ww-predicted_quality_ww)^2)</pre>
rsq_ls_ww<-1-(sse_ww/sst_ww)
rsq_ls_ww
## [1] 0.2811733
n_ww<-nrow(white_wine_df)</pre>
rmse_ls_ww<-sqrt(sum((actual_quality_ww-predicted_quality_ww)^2)/n_ww)</pre>
rmse_ls_ww
## [1] 0.7508001
mae ls ww<-sum(abs(actual quality ww-predicted quality ww))/n ww
mae_ls_ww
## [1] 0.5843081
ls_results_df <- tibble(wine_type = c("Red Wine","White wine"),</pre>
                             rmse = c(rmse ls rw,rmse ls ww),
                             mae = c(mae_ls_rw,rmse_ls_rw),
                             rsquare = c(rsq_ls_rw,rsq_ls_ww))
view(ls_results_df)
#Lasso regression on white wine
response_w<-white_wine_df$quality</pre>
predictors_w<-white_wine_df[,c("alcohol","volatile.acidity","sulphates",</pre>
                                 "total.sulfur.dioxide", "chlorides", "pH")]
#Fitting the model using lasso regression
#Lasso regression for white wine
ctrl <- trainControl(method="repeatedcv", number=10, repeats=10)</pre>
grd <- expand.grid(lambda=exp(seq(from=-7, to=-2, length.out=20)),</pre>
                    alpha=1)
set.seed(1)
fit_ww_lasso <- train(quality ~ alcohol + volatile.acidity +</pre>
                             sulphates + total.sulfur.dioxide +
                             chlorides + pH, data=whitedf_train,
              method="glmnet",
              preProcess=c("center", "scale"),
              trControl=ctrl, tuneGrid=grd)
fit ww lasso
```

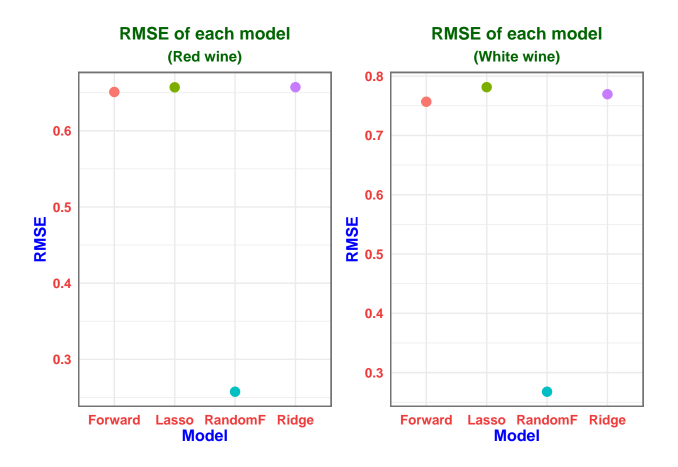
```
## glmnet
##
## 2939 samples
##
      6 predictor
##
## Pre-processing: centered (6), scaled (6)
## Resampling: Cross-Validated (10 fold, repeated 10 times)
## Summary of sample sizes: 2646, 2645, 2644, 2644, 2645, 2645, ...
## Resampling results across tuning parameters:
##
##
     lambda
                  RMSE
                             Rsquared
                                        MAE
##
     0.000911882
                 0.7822624
                             0.2430067
                                        0.6066094
##
     0.001186388 0.7822613
                            0.2430103
                                        0.6066094
##
     0.001543529 0.7822576 0.2430178 0.6066177
##
     0.002008180 0.7822549
                            0.2430244
                                       0.6066303
##
     0.002612707
                 0.7822548
                             0.2430289
                                        0.6066483
##
     0.003399216  0.7822624  0.2430237
                                        0.6066773
##
     0.004422489 0.7822841 0.2430002 0.6067220
##
     0.005753800 0.7823249 0.2429555 0.6067869
##
     0.007485879 0.7823976 0.2428766
                                       0.6068780
##
     0.009739369  0.7825276  0.2427323  0.6070186
##
     ##
     0.016485680 0.7831567 0.2419908 0.6075839
##
     0.021448399 0.7838665 0.2410816 0.6081751
##
     0.027905057  0.7849774  0.2396470  0.6091487
##
     0.036305375 0.7861201 0.2386237 0.6104414
##
     0.047234459 0.7876512 0.2376062 0.6121281
##
     0.061453549 0.7900493 0.2361036
                                       0.6143003
##
     0.079953042 0.7937151 0.2340606
                                       0.6169961
##
     0.104021477
                 0.7998325 0.2297565
                                        0.6209385
##
     ##
## Tuning parameter 'alpha' was held constant at a value of 1
## RMSE was used to select the optimal model using the smallest value.
## The final values used for the model were alpha = 1 and lambda = 0.002612707.
fit fs rmse rw<-as.numeric(fit1 rmse["cv rmse"])</pre>
fit_fs_rsq_rw<-as.numeric(fit1_rmse["cv_rsq"])</pre>
fit_fs_rmse_ww<-as.numeric(fit1_rmse_ww["cv_rmse"])</pre>
fit_fs_rsq_ww<-as.numeric(fit1_rmse_ww["cv_rsq"])</pre>
fit_fs_mae_rw<-as.numeric(fit1_rmse["cv_mae"])</pre>
fit_fs_mae_ww<-as.numeric(fit1_rmse_ww["cv_mae"])</pre>
fit_rd_rmse_rw<-rmse(fit_rw_ridge,reddf_train)</pre>
fit_rd_rsq_rw<-rsquare(fit_rw_ridge,reddf_train)</pre>
fit_rd_rmse_ww<-rmse(fit_ww_ridge,whitedf_train)</pre>
fit_rd_rsq_ww<-rsquare(fit_ww_ridge,whitedf_train)</pre>
fit_rd_mae_rw<-mae(fit_rw_ridge,reddf_train)</pre>
fit_rd_mae_ww<-mae(fit_ww_ridge,whitedf_train)</pre>
fit_ls_rmse_rw<-rmse(fit_rw_lasso,reddf_train)</pre>
fit_ls_rsq_rw<-rsquare(fit_rw_lasso,reddf_train)</pre>
fit ls rmse ww<-rmse(fit ww lasso, whitedf train)
fit_ls_rsq_ww<-rsquare(fit_ww_lasso,whitedf_train)</pre>
```

```
fit_ls_mae_rw<-mae(fit_rw_lasso,reddf_train)</pre>
fit_ls_mae_ww<-mae(fit_ww_lasso,whitedf_train)</pre>
fit_rf_rmse_rw<-rmse(fit_rw_rf,red_wine_std)</pre>
fit_rf_rsq_rw<- 0.5061
fit_rf_rmse_ww<-rmse(fit_ww_rf,white_wine_std)</pre>
fit_rf_rsq_ww<- 0.551
fit_rf_mae_rw<-mae(fit_rw_rf,red_wine_std)</pre>
fit_rf_mae_ww<-mae(fit_ww_rf,white_wine_std)</pre>
rmse_of_plots_rw<-data.frame(rmse_value=c(fit_fs_rmse_rw,fit_rd_rmse_rw,fit_ls_rmse_rw,fit_rf_rmse_rw),
rmse_of_plots_ww<-data.frame(rmse_value=c(fit_fs_rmse_ww,fit_rd_rmse_ww,fit_ls_rmse_ww,fit_rf_rmse_ww),
rsq_of_plots_rw<-data.frame(rsq_value=c(fit_fs_rsq_rw,fit_rd_rsq_rw,fit_ls_rsq_rw,fit_rf_rsq_rw),model=
rsq_of_plots_ww<-data.frame(rsq_value=c(fit_fs_rsq_ww,fit_rd_rsq_ww,fit_ls_rsq_ww,fit_rf_rsq_ww),model=
mae_of_plots_rw<-data.frame(mae_value=c(fit_fs_mae_rw,fit_rd_mae_rw,fit_ls_mae_rw,fit_rf_mae_rw),model=
mae_of_plots_ww<-data.frame(mae_value=c(fit_fs_mae_ww,fit_rd_mae_ww,fit_ls_mae_ww,fit_rf_mae_ww),model=
rmse_plot_rw<-ggplot(data=rmse_of_plots_rw)+</pre>
  geom_point(aes(x=model,y=rmse_value,color=model),size=3)+
  theme minimal()+
  labs(x="Model",
       y="RMSE",
       title="RMSE of each model",
       subtitle="(Red wine)")+
   theme(plot.title = element_text(hjust = 0.5,
                                            color = "darkgreen",
                                            face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                            color = "darkgreen",
                                            face = "bold"),
                axis.title = element_text(color = "blue",
                                            face = "bold",
                                            size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                            vjust = 0.5,
                                            color = "brown2",
                                           face = "bold",
                                            size = 10),
                axis.text.y = element_text(color = "brown2",
                                            face = "bold",
                                            size = 10),
                panel.border = element_rect(colour = "grey45",
                                            fill=NA, size=1),
         legend.position = "none")
rmse_plot_ww<-ggplot(data=rmse_of_plots_ww)+</pre>
  geom_point(aes(x=model,y=rmse_value,color=model),size=3)+
  theme_minimal()+
  labs(x="Model",
       y="RMSE",
```

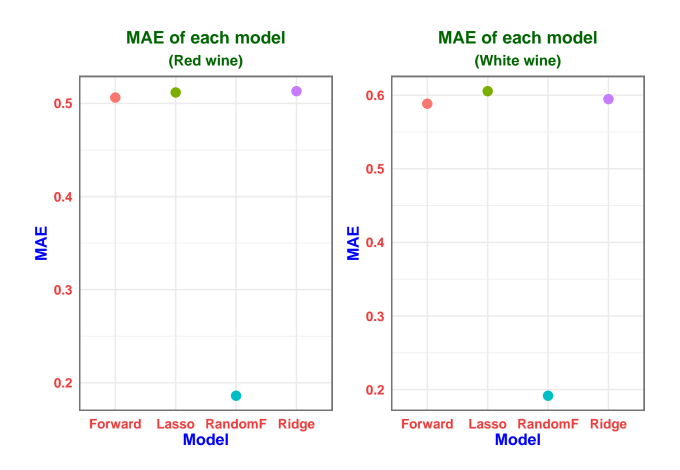
```
title="RMSE of each model",
       subtitle="(White wine)")+
   theme(plot.title = element text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=1),
         legend.position = "none")
rsq_plot_rw<-ggplot(data=rsq_of_plots_rw)+</pre>
  geom_point(aes(x=model,y=rsq_value,color=model),size=3)+
  theme minimal()+
  labs(x="Model",
       y="RSquared Value",
       title="RSquared of each model",
       subtitle="(Red wine)")+
   theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=1),
         legend.position = "none")
rsq_plot_ww<-ggplot(data=rsq_of_plots_ww)+</pre>
  geom_point(aes(x=model,y=rsq_value,color=model),size=3)+
```

```
theme_minimal()+
  labs(x="Model",
       y="RSquared Value",
       title="RSquare of each model",
       subtitle="(White wine)")+
   theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=1),
         legend.position = "none")
mae_plot_rw<-ggplot(data=mae_of_plots_rw)+</pre>
  geom_point(aes(x=model,y=mae_value,color=model),size=3)+
  theme_minimal()+
  labs(x="Model",
       y="MAE",
       title="MAE of each model",
       subtitle="(Red wine)")+
  theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=1),
         legend.position = "none")
```

```
mae_plot_ww<-ggplot(data=mae_of_plots_ww)+</pre>
  geom_point(aes(x=model,y=mae_value,color=model),size=3)+
  theme_minimal()+
  labs(x="Model",
       y="MAE",
       title="MAE of each model",
       subtitle="(White wine)")+
   theme(plot.title = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                plot.subtitle = element_text(hjust = 0.5,
                                           color = "darkgreen",
                                           face = "bold"),
                axis.title = element_text(color = "blue",
                                           face = "bold",
                                           size = 12),
                axis.text.x = element_text(hjust = 0.5,
                                           vjust = 0.5,
                                           color = "brown2",
                                           face = "bold",
                                           size = 10),
                axis.text.y = element_text(color = "brown2",
                                           face = "bold",
                                           size = 10),
                panel.border = element_rect(colour = "grey45",
                                           fill=NA, size=1),
         legend.position = "none")
grid.arrange(rmse_plot_rw,rmse_plot_ww,nrow = 1)
```



grid.arrange(mae_plot_rw,mae_plot_ww,nrow = 1)



grid.arrange(rsq_plot_rw,rsq_plot_ww,nrow = 1)

