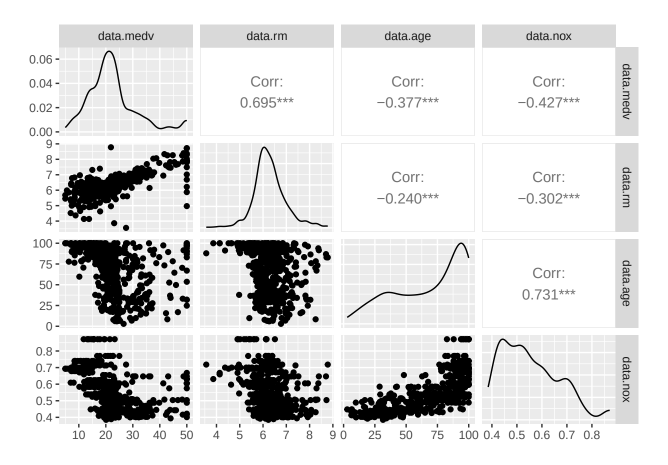
Compulsory exercise 1: Group 12

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```
**Problem 1
Quantative: Time, income earned, horsepower of a car. Qvalitative: Marital status, origin, Gender.
b) KNN, LDA, QDA can be used for multi-class classifications.
d) The nereast neighbour for k=1 is a blue dot, so our classification is blue. For K=3, two of the nearest
neighbors are red and 1 blue. This gives 2/3 red, so it is red. For K=5 we have 3/5 red, so it is red.
library (MASS)
data(Boston)
data = Boston
model = lm(medv ~ rm + age,data=data)
summary(model)
##
## Call:
## lm(formula = medv ~ rm + age, data = data)
## Residuals:
       Min
                1Q Median
                                 3Q
                                        Max
## -20.555 -2.882 -0.274
                              2.293 40.799
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -25.27740
                            2.85676 -8.848 < 2e-16 ***
## rm
                 8.40158
                             0.41208 20.388 < 2e-16 ***
## age
                -0.07278
                             0.01029 -7.075 5.02e-12 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6.316 on 503 degrees of freedom
## Multiple R-squared: 0.5303, Adjusted R-squared: 0.5284
## F-statistic: 283.9 on 2 and 503 DF, p-value: < 2.2e-16
cor_matrix = cor(data.frame(data$medv, data$rm, data$age))
print(cor_matrix)
##
              data.medv
                            data.rm
                                      data.age
## data.medv 1.0000000 0.6953599 -0.3769546
## data.rm
              0.6953599 1.0000000 -0.2402649
## data.age -0.3769546 -0.2402649 1.0000000
```

```
model2 = lm(medv ~ rm + age + nox, data=data)
summary(model2)
##
## Call:
## lm(formula = medv ~ rm + age + nox, data = data)
## Residuals:
      Min
              1Q Median
                            3Q
                                  Max
## -18.343 -3.168 -0.539 2.221 40.260
##
## Coefficients:
             Estimate Std. Error t value Pr(>|t|)
## (Intercept) -19.08308
                        3.33919 -5.715 1.88e-08 ***
             ## rm
             ## age
            -12.47877 3.58434 -3.481 0.000542 ***
## nox
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 6.247 on 502 degrees of freedom
## Multiple R-squared: 0.5413, Adjusted R-squared: 0.5386
## F-statistic: 197.5 on 3 and 502 DF, p-value: < 2.2e-16
library(GGally)
## Loading required package: ggplot2
## Registered S3 method overwritten by 'GGally':
    method from
##
    +.gg ggplot2
ggpairs(data.frame(data$medv, data$rm, data$age, data$nox))
```



e) IV Looking at the correlation between Age and NOX, it is 0.731, which is quite high which suggest it has multicollinearity, which means they give the some of the same information for the model.

*Problem 2 a)

```
model3 = lm(medv ~ poly(rm, 2) + I(age*crim) + age + crim, data=data)
summary(model3)
```

```
##
## Call:
## lm(formula = medv ~ poly(rm, 2) + I(age * crim) + age + crim,
##
       data = data)
##
## Residuals:
##
       Min
                1Q
                                 3Q
                                        Max
                    Median
  -34.314 -2.602
                   -0.369
                                    35.081
##
                              2.136
##
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                  28.096141
                               0.666283
                                         42.169
                                                 < 2e-16 ***
                 123.366564
                                         22.027
                                                 < 2e-16 ***
## poly(rm, 2)1
                               5.600727
## poly(rm, 2)2
                  64.836354
                               5.479728
                                         11.832
                                                  < 2e-16 ***
## I(age * crim)
                   0.005792
                               0.003553
                                          1.630
                                                  0.1037
## age
                  -0.067283
                               0.009342
                                         -7.202 2.19e-12 ***
## crim
                  -0.796544
                                        -2.350
                               0.338946
                                                  0.0192 *
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 5.369 on 500 degrees of freedom
## Multiple R-squared: 0.6626, Adjusted R-squared: 0.6592
## F-statistic: 196.3 on 5 and 500 DF, p-value: < 2.2e-16

?Boston

valuechanged = (-10*(-0.796544)+60*(-0.067283)+0.005792 * (-10 + 60))*1000</pre>
```

If the crime is reduced by 10 and age is 60, and then considering all other factor keeps equal our median value of the property is changed by 4218.06.

**b) First thing we could do is to plot all our data with say ggpairs from GGAlly. We can then look how the data is spread and correlated. We want to carefully identify if we have any outliers. If most our points follow a specific pattern, but then one point stick out, we could carefully consider to remove it. One approach could be to look at the mean and std of a given data columns and if some point lie 2 standard deviations away from the mean we could remove them.

***Problem 3 a)

```
library(titanic)
data("titanic_train")
vars_to_be_removed <- c("PassengerId", "Name", "Ticket", "Cabin", "Embarked")
View(titanic_train)
titanic_train <- titanic_train[, -which(names(titanic_train) %in% vars_to_be_removed)]
titanic_train$Pclass <- as.factor(titanic_train$Pclass)
train_idx <- sample(1:nrow(titanic_train), 0.8 * nrow(titanic_train))
titanic_test <- titanic_train[-train_idx, ]
titanic_train <- titanic_train[train_idx, ]
titanic_train <- na.omit(titanic_train)
titanic_test <- na.omit(titanic_test)
logReg = glm(Survived ~ Pclass + Sex + Age + SibSp + Parch + Fare, data = titanic_train, family = "binot summary(logReg)</pre>
```

```
##
## Call:
## glm(formula = Survived ~ Pclass + Sex + Age + SibSp + Parch +
    Fare, family = "binomial", data = titanic_train)
##
## Coefficients:
##
           Estimate Std. Error z value Pr(>|z|)
## (Intercept) 4.048694 0.553988 7.308 2.71e-13 ***
## Pclass2
          ## Pclass3
          ## Sexmale
          ## Age
                   0.142759 -2.942 0.003256 **
## SibSp
          -0.420064
## Parch
          -0.099062
                   0.131840 -0.751 0.452422
## Fare
          0.002732 0.002796 0.977 0.328516
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
##
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 763.63 on 564 degrees of freedom
##
## Residual deviance: 508.23 on 557 degrees of freedom
## AIC: 524.23
## Number of Fisher Scoring iterations: 5
predicted_probabilities <- predict(logReg, newdata = titanic_test, type = "response")</pre>
predicted_labels <- ifelse(predicted_probabilities > 0.5, 1, 0)
actual_labels <- titanic_test$Survived</pre>
misclassification_error <- mean(predicted_labels != actual_labels)</pre>
accuracy <- 1 - misclassification_error</pre>
print(accuracy)
## [1] 0.7919463
chitest = anova(logReg,test="Chisq")
chitest
## Analysis of Deviance Table
## Model: binomial, link: logit
## Response: Survived
## Terms added sequentially (first to last)
##
##
##
         Df Deviance Resid. Df Resid. Dev Pr(>Chi)
## NULL
                           564
                                    763.63
## Pclass 2 69.483
                           562
                                    694.15 8.163e-16 ***
                                   538.14 < 2.2e-16 ***
## Sex
         1 156.007
                           561
             17.682
                                    520.46 2.611e-05 ***
## Age
           1
                           560
## SibSp 1
             10.866
                            559
                                    509.59 0.0009795 ***
## Parch 1 0.289
                           558
                                    509.30 0.5911460
## Fare
          1 1.074
                           557
                                    508.23 0.3001461
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
?predict
newdata = data.frame(Pclass=as.factor(c(1,3)),Sex=c("female","female"),Age=c(40,40),SibSp=c(1,1),Parch=
newdata
##
              Sex Age SibSp Parch Fare
    Pclass
## 1
     1 female 40
                       1
                                 0 200
         3 female 40
                                 0
                                     20
                           1
casepredict = predict(logReg,newdata = newdata,type="response")
casepredict
```

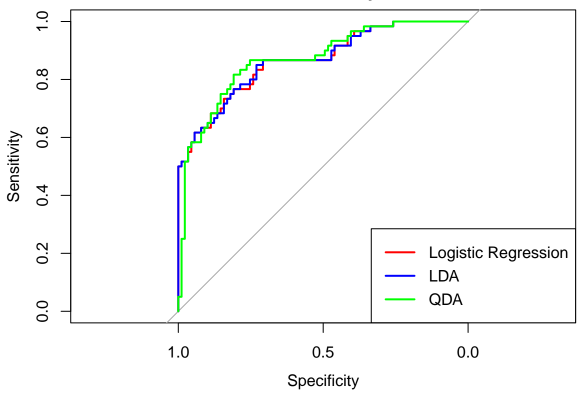
```
## 1 2
## 0.9217204 0.4136693
```

Given a p value of less that $2.2*10^{(-16)}$ suggest it is really relevant predictor. Given that H_0 is predictor could just as well be 0, a low p value would break this hypothesis. Comparing the a woman with the all the same factors, except for class and fare price, the higher class woman has a 0.9217204 chance of survival, while the lower class woman has a 0.4136693 chance of survival.

```
library(MASS)
?lda
ldamodel = lda(Survived ~ ., data = titanic_train)
print(ldamodel)
## Call:
## lda(Survived ~ ., data = titanic_train)
## Prior probabilities of groups:
##
## 0.5929204 0.4070796
##
## Group means:
       Pclass2
                  Pclass3
                            Sexmale
                                          Age
                                                  SibSp
                                                                       Fare
## 0 0.2119403 0.6238806 0.8417910 30.74328 0.5432836 0.3820896 23.76219
## 1 0.2695652 0.2956522 0.3173913 28.71413 0.5000000 0.5217391 53.21353
##
## Coefficients of linear discriminants:
##
                    LD1
## Pclass2 -0.789073856
## Pclass3 -1.504574355
## Sexmale -2.068582677
           -0.026553115
## Age
## SibSp
           -0.251774569
## Parch
           -0.086165420
## Fare
            0.001778469
predicted <- predict(ldamodel, newdata = titanic_test)</pre>
predicted_labels <- predicted$class</pre>
actual_labels <- titanic_test$Survived</pre>
misclassification_error = mean(predicted_labels != actual_labels)
accuracy <- 1 - misclassification_error</pre>
print(accuracy)
## [1] 0.7919463
qdamodel = qda(Survived ~ ., data = titanic_train)
print(qdamodel)
## Call:
## qda(Survived ~ ., data = titanic_train)
## Prior probabilities of groups:
##
           0
                      1
```

```
## 0.5929204 0.4070796
##
## Group means:
                            Sexmale
                                                                        Fare
       Pclass2
                Pclass3
                                          Age
                                                   SibSp
                                                             Parch
## 0 0.2119403 0.6238806 0.8417910 30.74328 0.5432836 0.3820896 23.76219
## 1 0.2695652 0.2956522 0.3173913 28.71413 0.5000000 0.5217391 53.21353
predicted <- predict(qdamodel, newdata = titanic_test)</pre>
predicted_labels <- predicted$class</pre>
actual_labels <- titanic_test$Survived</pre>
misclassification_error = mean(predicted_labels != actual_labels)
accuracy <- 1 - misclassification_error</pre>
print(accuracy)
## [1] 0.7986577
library(pROC)
## Type 'citation("pROC")' for a citation.
## Attaching package: 'pROC'
## The following objects are masked from 'package:stats':
##
##
       cov, smooth, var
logreg probs <- predict(logReg, newdata = titanic test, type = "response")</pre>
lda_probs <- predict(ldamodel, newdata = titanic_test)$posterior[,2]</pre>
qda_probs <- predict(qdamodel, newdata = titanic_test)$posterior[,2]</pre>
actual_outcomes <- titanic_test$Survived</pre>
# ROC for Logistic Regression
roc_logreg <- roc(actual_outcomes, logreg_probs)</pre>
## Setting levels: control = 0, case = 1
## Setting direction: controls < cases
# ROC for LDA
roc_lda <- roc(actual_outcomes, lda_probs)</pre>
## Setting levels: control = 0, case = 1
## Setting direction: controls < cases
# ROC for QDA
roc_qda <- roc(actual_outcomes, qda_probs)</pre>
## Setting levels: control = 0, case = 1
## Setting direction: controls < cases
```

ROC Curves Comparison



```
# AUC for Logistic Regression
auc_logreg <- roc_logreg$auc

# AUC for LDA
auc_lda <- roc_lda$auc

# AUC for QDA
auc_qda <- roc_qda$auc
# Print AUC values
cat("AUC for Logistic Regression:", auc_logreg, "\n")</pre>
```

AUC for Logistic Regression: 0.8677903

```
cat("AUC for LDA:", auc_lda, "\n")
## AUC for LDA: 0.867603
cat("AUC for QDA:", auc_qda, "\n")
```

```
## AUC for QDA: 0.8702247
```

AUC over 0.8 is generally considered good. QDA performs slightly better than LDA and Logistic Regression. However Logistic regression is almost as accurate, but more interpretable, for one it gives clearer relationships between the predictors and they are linear.

```
**b)
```

Diagnostic Paradigm:

The diagnostic paradigm in classification focuses on directly modeling the relationship between the predictors and the outcome (class labels. The primary goal is to diagnose the class of an observation based on its features. Models under this paradigm typically estimate the probability that an observation belongs to a certain class given its features. This approach is more deterministic, where the decision rule is often based on the estimated probabilities and predefined thresholds. Sampling Paradigm:

The sampling paradigm, on the other hand, is based on modeling the distribution of features for each class. It involves estimating how the data is generated or sampled for each class and then uses this information to classify new observations. This approach is more generative, as it tries to understand the underlying data generation process for each class and uses this generative model to make classification decisions.

Key Differences:

Modeling Focus: The diagnostic paradigm focuses on the direct relationship between features and classes, often modeling the conditional probability of a class given the features. The sampling paradigm focuses on modeling the distribution of features for each class, effectively capturing how data for each class is generated. Approach: Diagnostic models are more discriminative, directly seeking a decision boundary between classes. Sampling models are generative, aiming to understand the data generation process for each class. Decision Rule: In the diagnostic paradigm, classification decisions are often made based on estimated probabilities and thresholds. In the sampling paradigm, decisions are made based on likelihoods derived from the modeled distributions of features for each class.

ii) Classification Models and Their Paradigms

Logistic Regression: Belongs to the diagnostic paradigm. It models the probability of a class given the features directly using the logistic function.

KNN (K-Nearest Neighbors): Can be seen as part of the sampling paradigm. It classifies an observation based on the majority class among its K nearest neighbors, effectively using the local distribution of the data around an observation to make a classification decision.

Naive Bayes Classifier: Fits within the sampling paradigm. It is a generative model that assumes independence among features given the class and estimates the distribution of each feature within each class to compute the posterior probability of the class given an observation.

LDA (Linear Discriminant Analysis): Primarily fits within the sampling paradigm. LDA models the distribution of features in each class (assuming a Gaussian distribution with a common covariance matrix across classes) and uses this to classify new observations by finding the class that maximizes the posterior probability.

QDA (Quadratic Discriminant Analysis): Also belongs to the sampling paradigm. Similar to LDA, QDA models the distribution of features for each class, but it allows for class-specific covariance matrices, leading to quadratic decision boundaries. Like LDA, it uses the estimated distributions to compute class probabilities for classification.