# Assignment 2

Maggie Wang

2023-10-10

## Setup

```
# Load required libraries
library(ggplot2)
library(ggbiplot)
library(ROCR)
library(corrplot)
library(ISLR)
library(caret)
library(randomForest)
# Read data
ovarian.data <- na.omit(read.delim("ovarian.data", sep=",", header = FALSE))
features <- c("perimeter", "area", "smoothness", "symmetry", "concavity",
              paste("protein", seq(1, 25), sep=""))
names(ovarian.data) <- c("cell_id", "diagnosis", features)</pre>
# pasteO(features, "_mean"), pasteO(features, "_se"), pasteO(features, "_worst"))
dim(ovarian.data)
head(ovarian.data)
```

### Q1. Dimensionality Reduction

#### Q1.1

```
ovarian.pca <- prcomp(ovarian.data[,c(3:32)], center = TRUE,scale. = TRUE)
summary(ovarian.pca)</pre>
```

```
## Importance of components:
##
                             PC1
                                    PC2
                                            PC3
                                                    PC4
                                                            PC5
                                                                   PC6
                                                                           PC7
## Standard deviation
                          3.5820 2.2873 1.62395 1.37410 1.24910 1.0844 0.8306
## Proportion of Variance 0.4277 0.1744 0.08791 0.06294 0.05201 0.0392 0.0230
## Cumulative Proportion 0.4277 0.6021 0.68997 0.75291 0.80492 0.8441 0.8671
##
                              PC8
                                      PC9
                                             PC10
                                                     PC11
                                                            PC12
                                                                   PC13
                          0.74686 0.67762 0.61684 0.60200 0.5771 0.5139 0.5021
## Standard deviation
## Proportion of Variance 0.01859 0.01531 0.01268 0.01208 0.0111 0.0088 0.0084
## Cumulative Proportion 0.88571 0.90101 0.91369 0.92578 0.9369 0.9457 0.9541
                                    PC16
                                          PC17
                                                   PC18
                                                           PC19
                          0.45896 0.3989 0.3834 0.36254 0.32797 0.30949 0.3001
## Standard deviation
```

```
## Proportion of Variance 0.00702 0.0053 0.0049 0.00438 0.00359 0.00319 0.0030
## Cumulative Proportion 0.96110 0.9664 0.9713 0.97569 0.97928 0.98247 0.9855
                                             PC24
                                                            PC26
##
                             PC22
                                     PC23
                                                    PC25
                                                                    PC27
## Standard deviation
                          0.27191 0.26081 0.24722 0.2326 0.22154 0.20068 0.18042
## Proportion of Variance 0.00246 0.00227 0.00204 0.0018 0.00164 0.00134 0.00108
## Cumulative Proportion 0.98794 0.99020 0.99224 0.9940 0.99568 0.99702 0.99811
                             PC29
                                     PC30
## Standard deviation
                          0.17164 0.16532
## Proportion of Variance 0.00098 0.00091
## Cumulative Proportion 0.99909 1.00000
```

#### str(ovarian.pca)

```
## List of 5
## $ sdev
              : num [1:30] 3.58 2.29 1.62 1.37 1.25 ...
   $ rotation: num [1:30, 1:30] -0.22 -0.11 -0.229 -0.222 -0.137 ...
##
    ..- attr(*, "dimnames")=List of 2
     ....$ : chr [1:30] "perimeter" "area" "smoothness" "symmetry" ...
     ....$ : chr [1:30] "PC1" "PC2" "PC3" "PC4" ...
##
   $ center : Named num [1:30] 14.1809 19.3922 92.1982 663.7854 0.0965 ...
##
    ..- attr(*, "names")= chr [1:30] "perimeter" "area" "smoothness" "symmetry" ...
##
            : Named num [1:30] 3.5715 4.2746 24.1993 354.8356 0.0142 ...
   $ scale
    ..- attr(*, "names")= chr [1:30] "perimeter" "area" "smoothness" "symmetry" ...
##
              : num [1:625, 1:30] -4.476 0.448 1.916 1.874 -2.802 ...
##
   $ x
   ..- attr(*, "dimnames")=List of 2
##
     ....$ : chr [1:625] "1" "2" "3" "4" ...
     ....$ : chr [1:30] "PC1" "PC2" "PC3" "PC4" ...
## - attr(*, "class")= chr "prcomp"
```

About 42.77% of the variation in the data is associated with PC1.

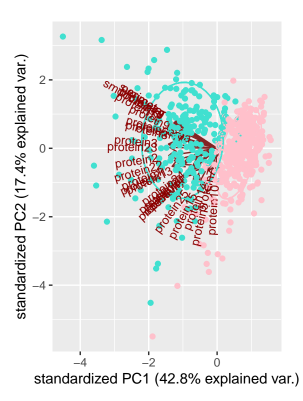
Q1.2 To represent 90% of the variance in the data by dimensionality reduction, you would need about 9 PCs.

#### Q1.3

```
diagnosis <- ovarian.data[,2]

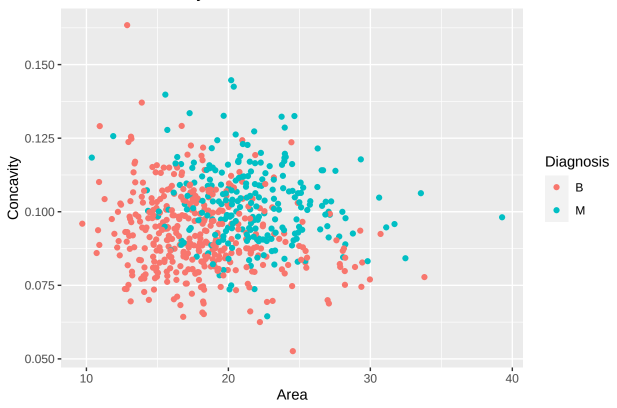
ggbiplot(ovarian.pca, choices=c(1,2), ellipse=TRUE, groups=diagnosis) +
    scale_color_manual(name="Diagnosis", values=c("pink", "turquoise")) +
    scale_shape_manual(name="Variety", values=c(2)) +
    geom_point(aes(colour=diagnosis), size = 0.01) +
    theme(legend.direction ="horizontal",legend.position = "top")</pre>
```





## Q1.4

## Area vs. Concavity of Tumor



Q1.5 The first plot using the first two important PCs has more separation between the classes, while in the second one they are a lot more mixed. This is because the first two PCs have the highest proportion of the variation in the dataset, so they will have the most difference between them.

#### Q1.6

### Q2. Clustering

### Q2.1

```
# Scaling the data
ovarian.scaled <- scale(ovarian.data[,c(3:32)])

# Performing kmeans
km.out <- kmeans(ovarian.scaled, centers = 2, iter.max = 1, nstart = 20)
km.out$cluster <- ifelse(km.out$cluster == 1, "M", "B")
table(ovarian.data$diagnosis, km.out$cluster)

##
## B M
## B 14 371
## M 205 35

mean(ovarian.data$diagnosis == km.out$cluster)</pre>
```

```
## [1] 0.0784
```

There is a good amount of concordance between the identified clusters and the true labels of the cell. 371 benign cells were identified correctly while 14 were mislabelled as malignant. 205 malignant cells were identified correctly while 35 were mislabelled as benign.

#### Q2.2

```
accuracies <- numeric(10)

# Repeat kmeans 10 times
for(i in 1:10){
    km.out <- kmeans(ovarian.scaled, centers = 2, iter.max = 10, nstart = 20)
    km.out$cluster <- ifelse(km.out$cluster == 1, "M", "B")
    accuracies[i] <- mean(ovarian.data$diagnosis == km.out$cluster)
}
mean(accuracies)</pre>
```

#### ## [1] 0.66864

The values change from run to run because the results of the kmeans algorithm is dependent on the initializtion of the centers, which is different each time.

#### Q2.3

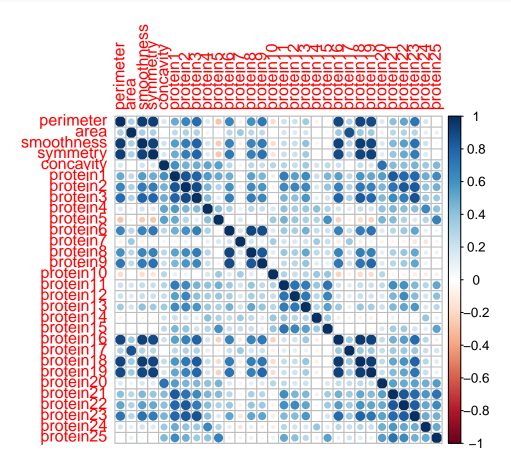
```
# Transform pca results to dataframe
pca.data <- as.data.frame(ovarian.pca$x[,1:5])</pre>
# Perform kmeans analysis
km.out <- kmeans(pca.data, centers = 2, nstart = 20)</pre>
km.out$cluster <- ifelse(km.out$cluster == 1, "M", "B")</pre>
table(ovarian.data$diagnosis, km.out$cluster)
##
##
         В
            М
##
     B 369 16
     M 35 205
##
mean(ovarian.data$diagnosis == km.out$cluster)
## [1] 0.9184
Q2.4
```

#### Q3. Classification

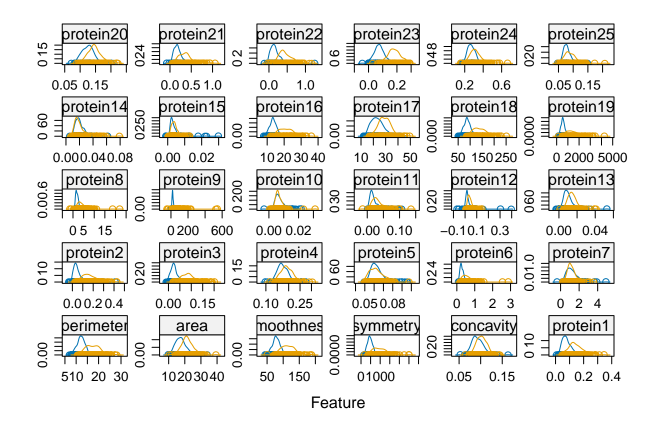
```
# Divide dataset into training and testing sets
ovarian.data.train <- ovarian.data[sample(nrow(ovarian.data))[1:(nrow(ovarian.data)/2)],]
ovarian.data.test <- ovarian.data[sample(nrow(ovarian.data))[(nrow(ovarian.data)/2):(nrow(ovarian.data)</pre>
```

### Q3.1

```
# Plot correlation between pairs of variables
correlations <- cor(ovarian.data[,3:32])
corrplot(correlations, method="circle")</pre>
```



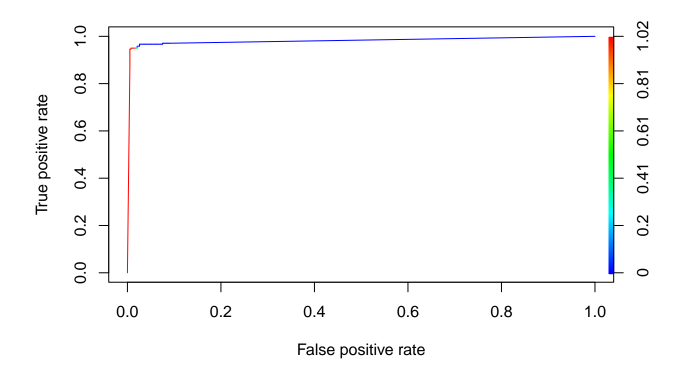
```
# Plot density distribution of each variable, separated by diagnosis
x <- ovarian.data[,3:32]
y <- as.factor(ovarian.data[,2])
scales <- list(x=list(relation="free"), y=list(relation="free"))
featurePlot(x=x, y=y, plot="density", scales=scales)</pre>
```



```
# Change diagnosis column to factors
ovarian.data.train$diagnosis <- as.factor(ovarian.data.train$diagnosis)</pre>
# Logistic regression training model
training.model <- glm(diagnosis ~. -cell_id, data = ovarian.data.train, family = binomial)
# Predicting on testing model
probabilities <- predict(training.model, ovarian.data.test, type = "response")</pre>
predicted.diagnosis <- ifelse(probabilities > 0.5, "M", "B")
prediction <- as.factor(predicted.diagnosis)</pre>
actual <- as.factor(ovarian.data.test$diagnosis)</pre>
# Confusion matrix
table(prediction, actual)
##
             actual
## prediction
##
             B 191
                    10
##
                 4 108
# To calculate accuracy, precision, recall
accuracy <-mean(prediction == actual)</pre>
precision <- posPredValue(prediction, actual, positive='M', negative = 'B')</pre>
recall <- sensitivity(prediction, actual, positive="M")</pre>
accuracy
```

```
## [1] 0.9552716
precision
## [1] 0.9642857
recall
## [1] 0.9152542
Q3.2
# Logistic regression training model using top 5 PCs
pca.training.model <- glm(diagnosis ~ perimeter + area + smoothness + symmetry
                          + concavity, data = ovarian.data.train, family = binomial)
# Predicting on testing set
pca.probabilities <- predict(pca.training.model, ovarian.data.test, type = "response")</pre>
pca.predicted.diagnosis <- ifelse(pca.probabilities > 0.5, "M", "B")
pca.prediction <- as.factor(pca.predicted.diagnosis)</pre>
# Confusion matrix
table(pca.prediction, actual)
##
                 actual
## pca.prediction B M
                B 188 14
##
##
                M 7 104
# To calculate accuracy, precision, recall
pca.accuracy <-mean(pca.prediction == actual)</pre>
pca.precision <- posPredValue(pca.prediction, actual, positive='M', negative = 'B')</pre>
pca.recall <- sensitivity(pca.prediction, actual, positive="M")</pre>
pca.accuracy
## [1] 0.9329073
pca.precision
## [1] 0.9369369
pca.recall
## [1] 0.8813559
Q3.3
Q3.4
Q3.5
```

```
pred.prob <- predict(training.model, ovarian.data, type="response")
predict <- prediction(pred.prob, ovarian.data$diagnosis, label.ordering=c("B","M"))
perform <- performance(predict,"tpr","fpr")
plot(perform,colorize=TRUE)</pre>
```



## $\mathbf{Q3.6}$

```
# Split into training (70%) and testing (30%)
chunk <- sample(nrow(ovarian.data), 0.7 * nrow(ovarian.data))
rf.training <- ovarian.data[chunk, ]
rf.testing <- ovarian.data[-chunk, ]

# Random forest model
rf.training$diagnosis <- as.factor(rf.training$diagnosis)
ovarian.rf <- randomForest(diagnosis ~.-cell_id, rf.training)

# Predicting on train set
pred.train <- predict(ovarian.rf, rf.training, type = "class")

# Checking classification accuracy
table(pred.train, rf.training$diagnosis)</pre>
```

```
## pred.train B M
     B 273 0
##
##
          M 0 164
# Predicting on Validation set
pred.test <- predict(ovarian.rf, rf.testing, type = "class")</pre>
# Checking classification accuracy
mean(pred.test == rf.testing$diagnosis)
## [1] 0.962766
table(pred.test, rf.testing$diagnosis)
##
## pred.test B
##
         B 109
                 4
##
          M 3 72
# Repeat with top 5 PCs
# Random forest model
pca.rf <- randomForest(diagnosis ~ perimeter + area + smoothness + symmetry</pre>
                         + concavity, rf.training)
# Predicting on Validation set
pca.pred.test <- predict(pca.rf, rf.testing, type = "class")</pre>
# Checking classification accuracy
mean(pca.pred.test == rf.testing$diagnosis)
## [1] 0.9308511
table(pca.pred.test, rf.testing$diagnosis)
##
## pca.pred.test B
##
              B 107
                     8
##
              M 5 68
```

#### Contributions