## class09

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## Exploratory data analysis

Use the read.csv() function to read the CSV (comma-separated values) file containing the data (available from our class website: WisconsinCancer.csv )

```
fna.data <-read.csv("https://bioboot.github.io/bimm143_S20/class-material/WisconsinCancer.csv")</pre>
```

Complete the following code to input the data and store as wisc.df

```
wisc.df <- data.frame (fna.data, row.names=1)
```

We can use -1 here to remove the first column

```
wisc.data <- wisc.df[,-1]
```

Create diagnosis vector for later

```
diagnosis <- factor(wisc.df$diagnosis)
diagnosis</pre>
```

Q1. How many observations are in this dataset?

```
dim (wisc.data)
```

```
## [1] 569 30
```

Q2. How many of the observations have a malignant diagnosis?

#### table(diagnosis)

```
## diagnosis
## B M
## 357 212
```

Q3. How many variables/features in the data are suffixed with \_mean?

```
grep("_mean", colnames(wisc.data))
```

```
## [1] 1 2 3 4 5 6 7 8 9 10
```

## Principal Component Analysis

### Performing PCA

Check column means and standard deviations

#### colMeans(wisc.data)

```
##
               radius_mean
                                         {\tt texture\_mean}
                                                                perimeter_mean
##
               1.412729e+01
                                         1.928965e+01
                                                                   9.196903e+01
##
                  area_mean
                                      smoothness_mean
                                                              compactness_mean
##
               6.548891e+02
                                         9.636028e-02
                                                                   1.043410e-01
##
             concavity_mean
                                 concave.points_mean
                                                                  symmetry_mean
##
               8.879932e-02
                                         4.891915e-02
                                                                   1.811619e-01
    {\tt fractal\_dimension\_mean}
##
                                            radius_se
                                                                     texture_se
##
               6.279761e-02
                                         4.051721e-01
                                                                   1.216853e+00
##
               perimeter_se
                                                                  smoothness_se
                                              area_se
##
               2.866059e+00
                                         4.033708e+01
                                                                   7.040979e-03
##
             compactness_se
                                         concavity_se
                                                             concave.points_se
##
               2.547814e-02
                                         3.189372e-02
                                                                   1.179614e-02
##
                symmetry_se
                                fractal_dimension_se
                                                                  radius_worst
##
               2.054230e-02
                                         3.794904e-03
                                                                  1.626919e+01
##
             texture_worst
                                     perimeter_worst
                                                                     area_worst
```

```
##
              2.567722e+01
                                        1.072612e+02
                                                                 8.805831e+02
##
          smoothness_worst
                                   compactness_worst
                                                              concavity_worst
##
              1.323686e-01
                                        2.542650e-01
                                                                 2.721885e-01
##
      concave.points_worst
                                      symmetry_worst fractal_dimension_worst
##
              1.146062e-01
                                        2.900756e-01
                                                                 8.394582e-02
apply(wisc.data,2,sd)
##
               radius_mean
                                                               perimeter_mean
                                        texture_mean
##
              3.524049e+00
                                        4.301036e+00
                                                                 2.429898e+01
##
                                     smoothness_mean
                                                             compactness_mean
                  area_mean
##
              3.519141e+02
                                        1.406413e-02
                                                                 5.281276e-02
                                concave.points_mean
##
            concavity mean
                                                                symmetry_mean
##
              7.971981e-02
                                        3.880284e-02
                                                                 2.741428e-02
##
    fractal_dimension_mean
                                           radius_se
                                                                   texture_se
##
              7.060363e-03
                                        2.773127e-01
                                                                 5.516484e-01
##
              perimeter_se
                                                                smoothness_se
                                             area_se
##
              2.021855e+00
                                        4.549101e+01
                                                                 3.002518e-03
##
            compactness se
                                        concavity se
                                                            concave.points_se
##
              1.790818e-02
                                        3.018606e-02
                                                                 6.170285e-03
##
               symmetry_se
                               fractal_dimension_se
                                                                 radius_worst
```

#### Perform PCA on wisc.data by completing the following code

8.266372e-03

6.146258e+00

2.283243e-02

6.573234e-02

texture\_worst

smoothness\_worst

concave.points\_worst

##

##

##

##

##

##

##

```
wisc.pr <- prcomp(wisc.data, scale=TRUE)
summary(wisc.pr)</pre>
```

2.646071e-03

3.360254e+01

1.573365e-01

6.186747e-02

symmetry\_worst fractal\_dimension\_worst

perimeter\_worst

compactness\_worst

4.833242e+00

5.693570e+02

2.086243e-01

1.806127e-02

concavity\_worst

area\_worst

```
## Importance of components:
                                    PC2
                                             PC3
                                                     PC4
                                                             PC5
                                                                     PC6
                                                                             PC7
                             PC1
## Standard deviation
                          3.6444 2.3857 1.67867 1.40735 1.28403 1.09880 0.82172
## Proportion of Variance 0.4427 0.1897 0.09393 0.06602 0.05496 0.04025 0.02251
  Cumulative Proportion
                          0.4427 0.6324 0.72636 0.79239 0.84734 0.88759 0.91010
                              PC8
                                     PC9
                                             PC10
                                                   PC11
                                                            PC12
## Standard deviation
                          0.69037 0.6457 0.59219 0.5421 0.51104 0.49128 0.39624
  Proportion of Variance 0.01589 0.0139 0.01169 0.0098 0.00871 0.00805 0.00523
  Cumulative Proportion 0.92598 0.9399 0.95157 0.9614 0.97007 0.97812 0.98335
##
                             PC15
                                     PC16
                                             PC17
                                                      PC18
                                                              PC19
                                                                      PC20
                                                                             PC21
## Standard deviation
                          0.30681 0.28260 0.24372 0.22939 0.22244 0.17652 0.1731
## Proportion of Variance 0.00314 0.00266 0.00198 0.00175 0.00165 0.00104 0.0010
## Cumulative Proportion 0.98649 0.98915 0.99113 0.99288 0.99453 0.99557 0.9966
##
                             PC22
                                     PC23
                                            PC24
                                                     PC25
                                                             PC26
                                                                     PC27
                                                                             PC28
## Standard deviation
                          0.16565 0.15602 0.1344 0.12442 0.09043 0.08307 0.03987
## Proportion of Variance 0.00091 0.00081 0.0006 0.00052 0.00027 0.00023 0.00005
## Cumulative Proportion 0.99749 0.99830 0.9989 0.99942 0.99969 0.99992 0.99997
```

```
## PC29 PC30
## Standard deviation 0.02736 0.01153
## Proportion of Variance 0.00002 0.00000
## Cumulative Proportion 1.00000 1.00000
```

Q4. From your results, what proportion of the original variance is captured by the first principal components (PC1)?

#### $\mathbf{44.27}\%$

Q5. How many principal components (PCs) are required to describe at least 70% of the original variance in the data?

3 PCs are required to describe at least 70% of the original variance in the data.

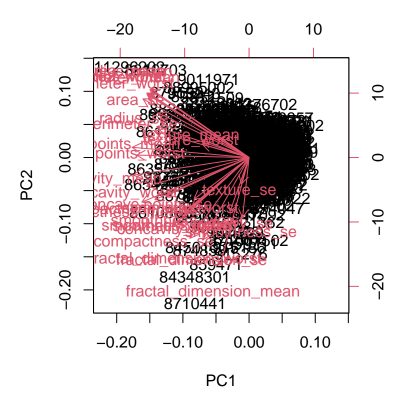
Q6. How many principal components (PCs) are required to describe at least 90% of the original variance in the data?

7 PCs are to describe at least 90% of the original variance in the data.

### Interpreting PCA results

Create a biplot of the wisc.pr using the biplot() function.

biplot(wisc.pr)

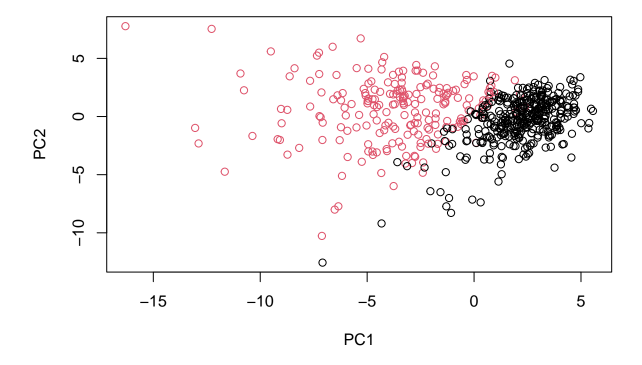


Q7. What stands out to you about this plot? Is it easy or difficult to understand? Why?

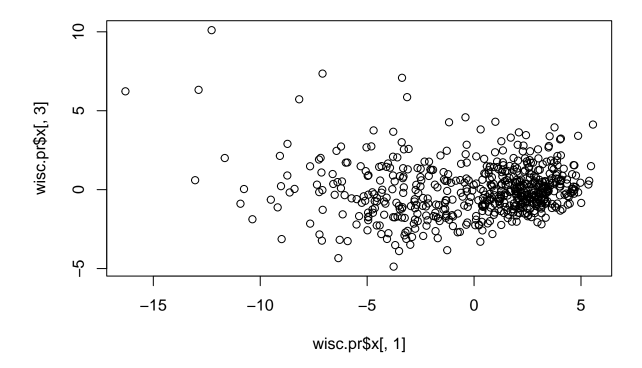
This is a hot mess of a plot and it's diffcult to interpret.

Scatter plot observations by components 1 and 2

```
plot(wisc.pr$x [,1:2], xlab = "PC1", ylab = "PC2", col=diagnosis)
```

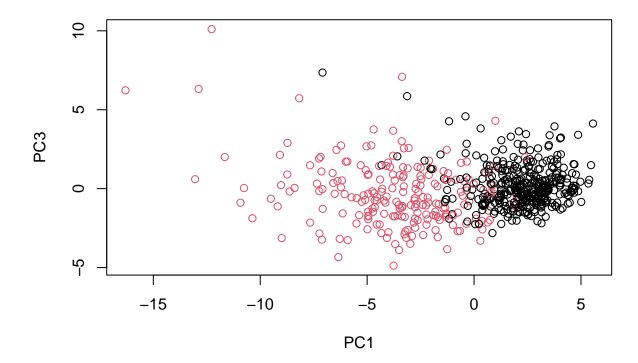


plot(wisc.pr\$x [,1], wisc.pr\$x [,3])



Q8. Generate a similar plot for principal components 1 and 3. What do you notice about these plots?

## Repeat for components 1 and 3 $\,$



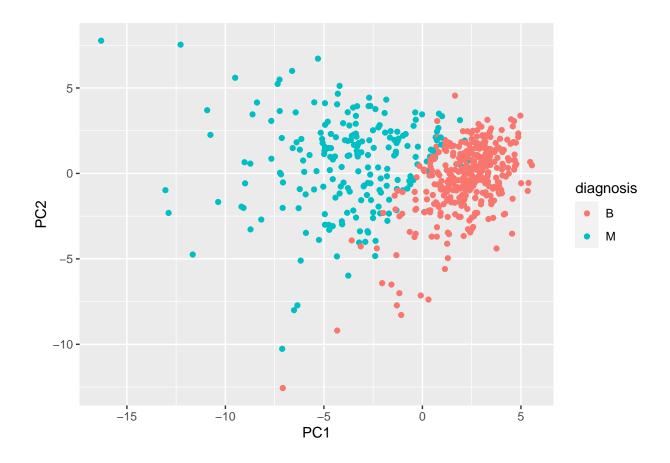
The first plot (PC2 vs PC1) has a cleaner cut separating the two subgroups than this polt, since PC 2 explains more variance in the original data than PC 3.

Create a data.frame for  $\operatorname{ggplot}$ 

```
df <- as.data.frame(wisc.pr$x)
df$diagnosis <- diagnosis</pre>
```

Make a scatter plot colored by diagnosis

```
library("ggplot2")
ggplot(df) +
  aes(PC1, PC2, col=diagnosis) +
  geom_point()
```



## Variance explained

### Calculate variance of each component

```
pr.var <- wisc.pr$sdev^2
head(pr.var)</pre>
```

**##** [1] 13.281608 5.691355 2.817949 1.980640 1.648731 1.207357

### Variance explained by each principal component: pve

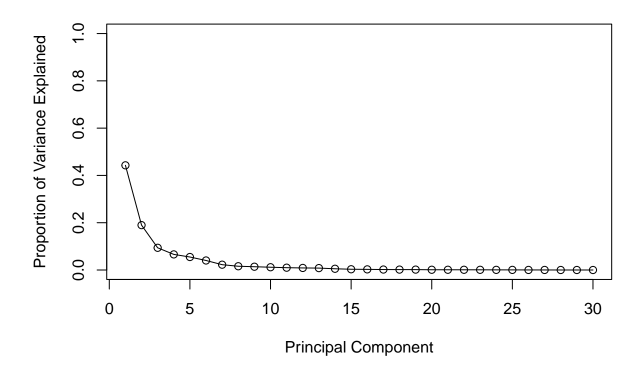
```
pve <- pr.var / sum(pr.var)
pve

## [1] 4 4272030-01 1 8971180-01 9 3931630-02 6 6021350-02 5 4957680-02
```

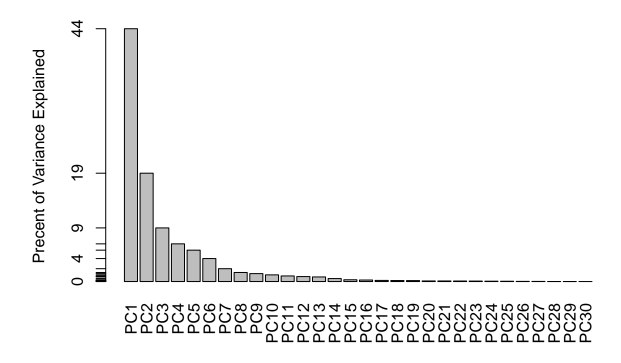
```
## [1] 4.427203e-01 1.897118e-01 9.393163e-02 6.602135e-02 5.495768e-02 ## [6] 4.024522e-02 2.250734e-02 1.588724e-02 1.389649e-02 1.168978e-02 ## [11] 9.797190e-03 8.705379e-03 8.045250e-03 5.233657e-03 3.137832e-03 ## [16] 2.662093e-03 1.979968e-03 1.753959e-03 1.649253e-03 1.038647e-03 ## [21] 9.990965e-04 9.146468e-04 8.113613e-04 6.018336e-04 5.160424e-04 ## [26] 2.725880e-04 2.300155e-04 5.297793e-05 2.496010e-05 4.434827e-06
```

Plot variance explained for each principal component

```
plot(pve, xlab = "Principal Component",
    ylab = "Proportion of Variance Explained",
    ylim = c(0, 1), type = "o")
```



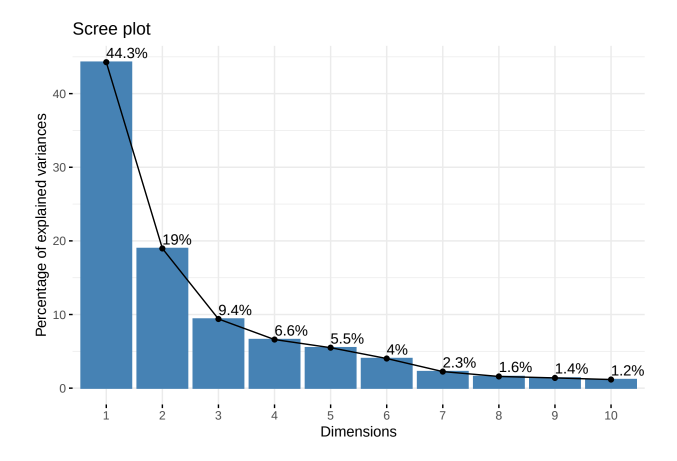
Alternative scree plot of the same data, note data driven y-axis



## ggplot based graph

#install.packages("factoextra")

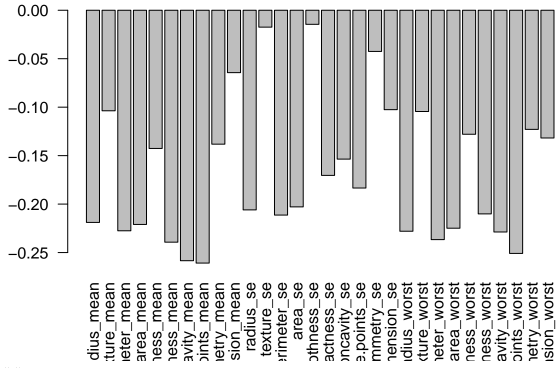
```
library(factoextra)
fviz_eig(wisc.pr, addlabels = TRUE)
```



# Communicating PCA results

Q9. For the first principal component, what is the component of the loading vector (i.e. wisc.prran(1)) for the feature concave.points\_mean?

barplot(wisc.pr\$rotation[,1], las=2, mar=c(10, 3, 3, 20))



### concave.points\_mean

Q10. What is the minimum number of principal components required to explain 80% of the variance of the data

```
var <- summary(wisc.pr)
var$importance</pre>
```

```
##
                               PC1
                                         PC2
                                                  PC3
                                                           PC4
                                                                     PC5
                                                                              PC6
                          3.644394 2.385656 1.678675 1.407352 1.284029 1.098798
## Standard deviation
  Proportion of Variance 0.442720 0.189710 0.093930 0.066020 0.054960 0.040250
##
  Cumulative Proportion
                          0.442720 0.632430 0.726360 0.792390 0.847340 0.887590
##
                                PC7
                                           PC8
                                                     PC9
                                                              PC10
                                                                         PC11
## Standard deviation
                          0.8217178 0.6903746 0.6456739 0.5921938 0.5421399
## Proportion of Variance 0.0225100 0.0158900 0.0139000 0.0116900 0.0098000
  Cumulative Proportion
                          0.9101000 0.9259800 0.9398800 0.9515700 0.9613700
                                                    PC14
##
                                PC12
                                          PC13
                                                              PC15
                                                                         PC16
## Standard deviation
                          0.5110395 0.4912815 0.3962445 0.3068142 0.2826001
## Proportion of Variance 0.0087100 0.0080500 0.0052300 0.0031400 0.0026600
                          0.9700700 0.9781200 0.9833500 0.9864900 0.9891500
  Cumulative Proportion
##
##
                               PC17
                                          PC18
                                                    PC19
                                                              PC20
                                                                         PC21
                          0.2437192 0.2293878 0.2224356 0.1765203 0.1731268
## Standard deviation
## Proportion of Variance 0.0019800 0.0017500 0.0016500 0.0010400 0.0010000
                          0.9911300 0.9928800 0.9945300 0.9955700 0.9965700
##
  Cumulative Proportion
##
                                PC22
                                          PC23
                                                    PC24
                                                              PC25
                                                                         PC26
                          0.1656484 0.1560155 0.1343689 0.1244238 0.0904303
## Standard deviation
```

```
## Proportion of Variance 0.0009100 0.0008100 0.0006000 0.0005200 0.0002700
## Cumulative Proportion 0.9974900 0.9983000 0.9989000 0.9994200 0.9996900
## PC27 PC28 PC29 PC30
## Standard deviation 0.08306903 0.0398665 0.02736427 0.01153451
## Proportion of Variance 0.00023000 0.0000500 0.00002000 0.00000000
## Cumulative Proportion 0.99992000 0.9999700 1.000000000 1.000000000

sum(var$importance[3,] <0.8)
```

## [1] 4

So, four at least.

### Hierarchical clustering

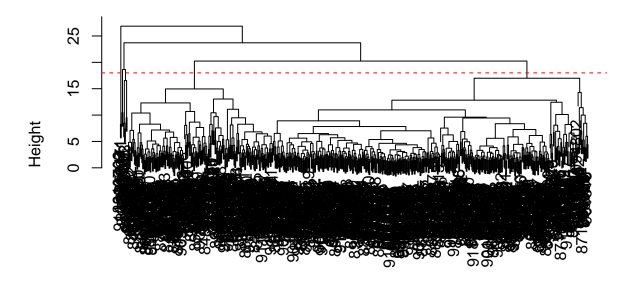
Scale the wisc.data data using the "scale()" function???

```
data.scaled <-scale(wisc.data)
data.dist <- dist(data.scaled)
wisc.hclust <- hclust(data.dist)</pre>
```

Results of hierarchical clustering

Q11. Using the plot() and abline() functions, what is the height at which the clustering model has 4 clusters?

```
plot(wisc.hclust)
abline(wisc.hclust, h=18, col="red", lty=2)
```



data.dist hclust (\*, "complete")

### Selecting number of clusters

Use cutree() to cut the tree so that it has 4 clusters. Assign the output to the variable wisc.hclust.clusters.

```
wisc.hclust.clusters <- cutree(wisc.hclust, 4)
table(wisc.hclust.clusters, diagnosis)</pre>
```

```
## diagnosis
## wisc.hclust.clusters B M
## 1 12 165
## 2 2 5
## 3 343 40
## 4 0 2
```

Q12. Can you find a better cluster vs diagnoses match by cutting into a different number of clusters between 2 and 10?

#### Perhaps no.

```
wisc.hclust.clusters <- cutree(wisc.hclust, 5)
table(wisc.hclust.clusters, diagnosis)</pre>
```

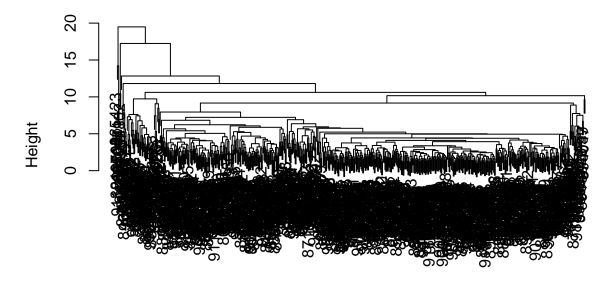
```
## diagnosis
## wisc.hclust.clusters B M
## 1 12 165
## 2 0 5
## 3 343 40
## 4 2 0
## 5 0 2
```

Q13. Which method gives your favorite results for the same data.dist dataset? Explain your reasoning.

The method "ward.D2" looked much better, as he two major clusters are more clearly separated in this method

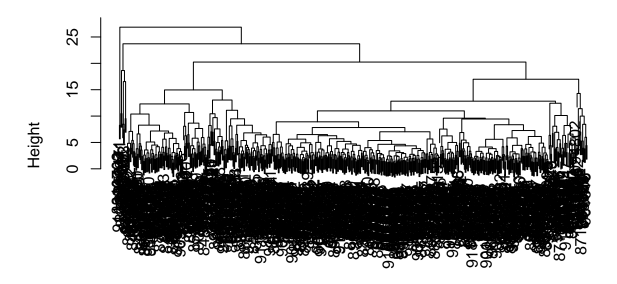
```
plot(hclust(data.dist, method = "average"))
```

# **Cluster Dendrogram**



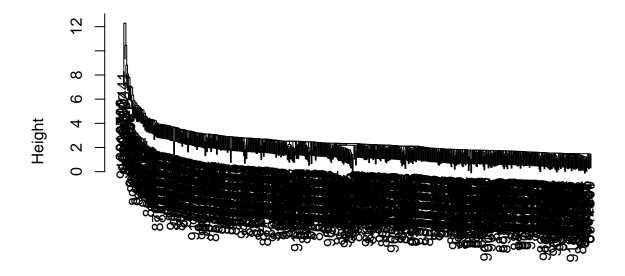
data.dist hclust (\*, "average")

```
plot(hclust(data.dist, method = "complete"))
```



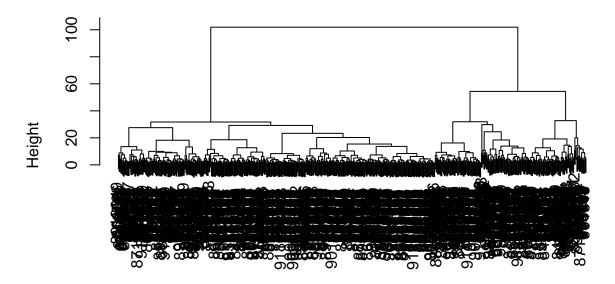
data.dist hclust (\*, "complete")

plot(hclust(data.dist, method = "single"))



data.dist hclust (\*, "single")

plot(hclust(data.dist, method = "ward.D2"))



data.dist hclust (\*, "ward.D2")

## **OPTIONAL:** K-means clustering

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## K-means clustering and comparing results

```
wisc.km <- kmeans(wisc.data, centers= 2, nstart= 20)</pre>
table(wisc.km$cluster, diagnosis )
      diagnosis
##
##
         В
##
     1 356 82
         1 130
table(wisc.km$cluster, wisc.hclust.clusters)
##
      wisc.hclust.clusters
##
       68
                      2
                          0
##
             3 365
```

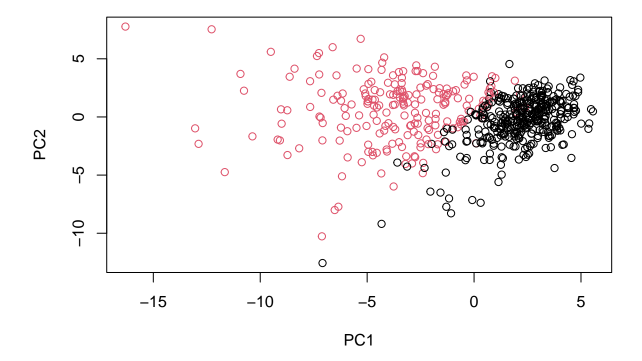
Q14. How well does k-means separate the two diagnoses? How does it compare to your hclust results?

It works well, but less accurate that hclust.

## Combining methods

Here we aim to combine our PCA results with clustering. Essentially, we are going to cluster in "PC", that is cluster on the results wisc.pr.

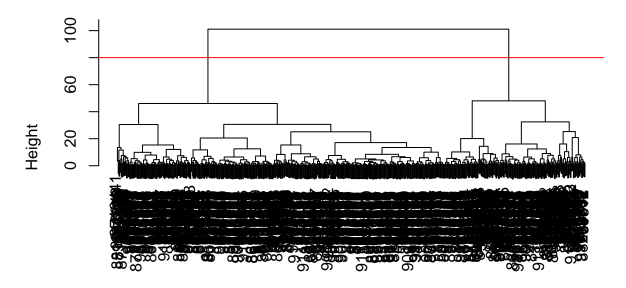
```
plot(wisc.pr$x[,1:2], col= diagnosis)
```



## Clustering on PCA results

I will use 4 PCs and hclust() and dist() as an input.

```
wisc.pr.hclust <- hclust(dist(wisc.pr$x[,1:4]), method = "ward.D2")
plot(wisc.pr.hclust)
abline(h=80, col="red")</pre>
```



dist(wisc.pr\$x[, 1:4]) hclust (\*, "ward.D2")

Let's find our cluster membership vector by cutting this tree into k=2 groups.

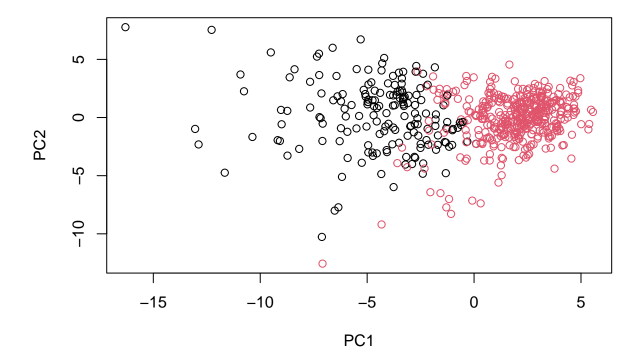
```
grps <- cutree(wisc.pr.hclust, k=2)
table(grps)

## grps
## 1 2
## 171 398

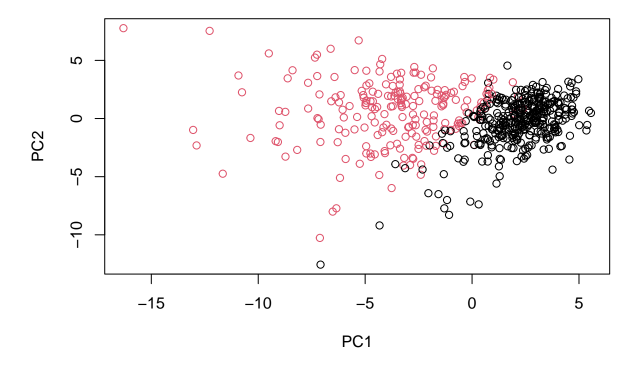
table(grps, diagnosis)

## diagnosis
## grps B M
## 1 6 165
## 2 351 47

plot(wisc.pr$x[,1:2], col=grps)</pre>
```



plot(wisc.pr\$x[,1:2], col=diagnosis)



To match things up we can turn our groups into a factor and reorder the levels so cluster 2 comes first and thus gets the first color (black) and cluster 1 gets the second color (red).

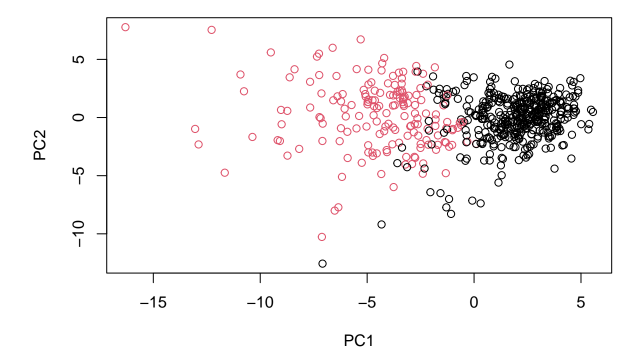
```
g <- as.factor(grps)
levels(g)

## [1] "1" "2"

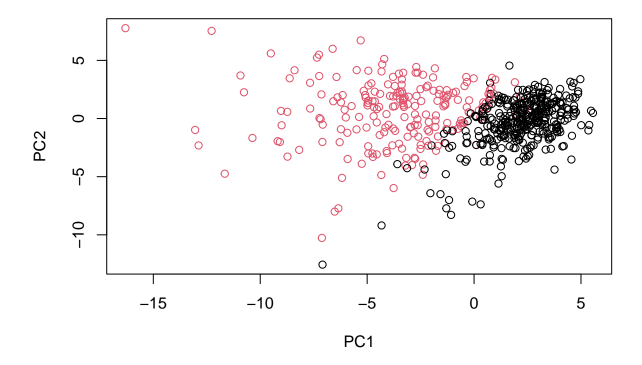
g <- relevel(g,2)
levels(g)

## [1] "2" "1"

plot(wisc.pr$x[,1:2], col=g)</pre>
```



plot(wisc.pr\$x[,1:2], col=diagnosis)



### diagnosis

grps B M 1 6 165 2 351 47

TP: 165, FP:6 TN: 351, FN: 47

Accuracy, essentially how many did we get correct?

Sensitivity: TP/(TP+FN) Specificity: TN/(TN+FN)

```
# **Accuracy**: (TP+TN)/total cases
# **Sensitivity**: TP/(TP+FN)
# **Specificity**: TN/(TN+FN)
Accuracy <- (165+351)/nrow(wisc.data)*100
Sensitivity <- (165)/(165+47)*100
Specificity <- (351)/(351+47)*100
evulation <- rbind(Accuracy, Sensitivity, Specificity)
evulation</pre>
```

```
## [,1]
## Accuracy 90.68541
## Sensitivity 77.83019
## Specificity 88.19095
```

## Use the distance along the first 7 PCs for clustering i.e. wisc.pr\$x[, 1:7]

```
wisc.pr.hclust <- hclust(dist(wisc.pr$x[,1:7]), method="ward.D2")</pre>
wisc.pr.hclust.clusters <- cutree(wisc.pr.hclust, k=2)</pre>
table(wisc.pr.hclust.clusters, diagnosis)
##
                           diagnosis
                              В
## wisc.pr.hclust.clusters
                                  М
##
                          1 28 188
                          2 329 24
##
# **Accuracy**: (TP+TN)/total cases
# **Sensitivity**: TP/(TP+FN)
# **Specificity**: TN/(TN+FN)
Accuracy <- (188+329)/nrow(wisc.data)*100
Sensitivity <- (188)/(188+24)*100
Specificity <- (329)/(329+24)*100
evulation <- rbind(Accuracy, Sensitivity, Specificity)</pre>
evulation
##
                    [,1]
               90.86116
## Accuracy
## Sensitivity 88.67925
## Specificity 93.20113
```

Q15. How well does the newly created model with four clusters separate out the two diagnoses?

It looked good, with higher sensitivity and specificity.

Q16. How well do the k-means and hierarchical clustering models you created in previous sections (i.e. before PCA) do in terms of separating the diagnoses? Again, use the table() function to compare the output of each model (wisc.km\$cluster and wisc.hclust.clusters) with the vector containing the actual diagnoses.

#### table(wisc.km\$cluster, diagnosis)

```
##
      diagnosis
##
         В
             М
     1 356 82
##
         1 130
##
# **Accuracy**: (TP+TN)/total cases
# **Sensitivity**: TP/(TP+FN)
# **Specificity**: TN/(TN+FN)
Accuracy <- (130+356)/nrow(wisc.data)*100</pre>
Sensitivity <- (130)/(130+82)*100
Specificity <- (356)/(356+82)*100
evulation <- rbind(Accuracy, Sensitivity, Specificity)</pre>
evulation
```

```
##
                   [,1]
               85.41301
## Accuracy
## Sensitivity 61.32075
## Specificity 81.27854
table(wisc.hclust.clusters, diagnosis)
##
                        diagnosis
## wisc.hclust.clusters
                          В
                               Μ
##
                          12 165
                       1
                       2
                          0
##
                               5
##
                      3 343 40
##
                       4
                           2
                               0
                               2
                           0
##
# **Accuracy**: (TP+TN)/total cases
# **Sensitivity**: TP/(TP+FN)
# **Specificity**: TN/(TN+FN)
Accuracy <- (165+343)/nrow(wisc.data)*100
Sensitivity <- (165)/(165+40)*100
Specificity <- (343)/(343+40)*100
evulation <- rbind(Accuracy, Sensitivity, Specificity)</pre>
evulation
##
                   [,1]
## Accuracy
               89.27944
## Sensitivity 80.48780
## Specificity 89.55614
```

Q17. Which of your analysis procedures resulted in a clustering model with the best specificity? How about sensitivity?

Using the distance along the first 7 PCs for clustering give the best specificity and sensitivity.

### Prediction

We will use the predict() function that will take our PCA model from before and new cancer cell data and project

```
#url <- "new_samples.csv"</pre>
url <- "https://tinyurl.com/new-samples-CSV"</pre>
new <- read.csv(url)</pre>
npc <- predict(wisc.pr, newdata=new)</pre>
npc
                         PC2
                                                 PC4
                                                            PC5
                                                                                   PC7
##
              PC1
                                     PC3
                                                                       PC6
## [1,] 2.576616 -3.135913 1.3990492 -0.7631950 2.781648 -0.8150185 -0.3959098
## [2,] -4.754928 -3.009033 -0.1660946 -0.6052952 -1.140698 -1.2189945 0.8193031
                          PC9
                                     PC10
                                               PC11
                                                          PC12
                                                                     PC13
```

## [1,] -0.2307350 0.1029569 -0.9272861 0.3411457 0.375921 0.1610764 1.187882

```
## [2,] -0.3307423 0.5281896 -0.4855301 0.7173233 -1.185917 0.5893856 0.303029
                        PC16
                                    PC17
                                                PC18
##
             PC15
                                                            PC19
                                                                       PC20
## [1,] 0.3216974 -0.1743616 -0.07875393 -0.11207028 -0.08802955 -0.2495216
## [2,] 0.1299153 0.1448061 -0.40509706 0.06565549 0.25591230 -0.4289500
              PC21
                         PC22
                                    PC23
                                               PC24
                                                           PC25
## [1,] 0.1228233 0.09358453 0.08347651 0.1223396 0.02124121 0.078884581
## [2,] -0.1224776 0.01732146 0.06316631 -0.2338618 -0.20755948 -0.009833238
                                         PC29
                PC27
                            PC28
                                                      PC30
## [1,] 0.220199544 -0.02946023 -0.015620933 0.005269029
## [2,] -0.001134152  0.09638361  0.002795349 -0.019015820
```

Now add these new samples to our PCA plot

```
plot(wisc.pr$x[,1:2], col=g)
points(npc[,1], npc[,2], col="blue", pch=16, cex=3)
text(npc[,1], npc[,2], labels=c(1,2), col="white")
```

