# Class08-mini-project

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Today we will complete analysis of some breast cancer biopsy data but first let's revisit the main PCA function in R prcomp() and see what scale=TRUE/FALSE does.

### head(mtcars)

	mpg	cyl	disp	hp	${\tt drat}$	wt	qsec	٧s	$\mathtt{am}$	gear	carb
Mazda RX4	21.0	6	160	110	3.90	2.620	16.46	0	1	4	4
Mazda RX4 Wag	21.0	6	160	110	3.90	2.875	17.02	0	1	4	4
Datsun 710	22.8	4	108	93	3.85	2.320	18.61	1	1	4	1
Hornet 4 Drive	21.4	6	258	110	3.08	3.215	19.44	1	0	3	1
Hornet Sportabout	18.7	8	360	175	3.15	3.440	17.02	0	0	3	2
Valiant	18.1	6	225	105	2.76	3.460	20.22	1	0	3	1

Find the mean value per column of this dataset?

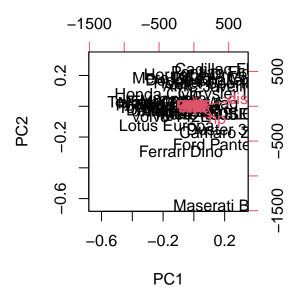
```
apply(mtcars, 2, sd)
```

```
mpg
                   cyl
                               disp
                                              hp
                                                        drat
                                                                       wt
6.0269481
            1.7859216 123.9386938
                                     68.5628685
                                                   0.5346787
                                                                0.9784574
     qsec
                    ٧s
                                           gear
                                                        carb
1.7869432
            0.5040161
                         0.4989909
                                      0.7378041
                                                   1.6152000
```

It is clear that "disp" and "hp" have the highest mean values and the highest standard deviation. They will likely dominate any analysis I do on this dataset. Let's see

```
pc.noscale <- prcomp(mtcars, scale = F)
pc.scale <- prcomp(mtcars, scale = T)</pre>
```

### biplot(pc.noscale)



### pc.noscale\$rotation[,1]

mpg cyl disp hp drat wt
-0.038118199 0.012035150 0.899568146 0.434784387 -0.002660077 0.006239405
qsec vs am gear carb
-0.006671270 -0.002729474 -0.001962644 -0.002604768 0.005766010

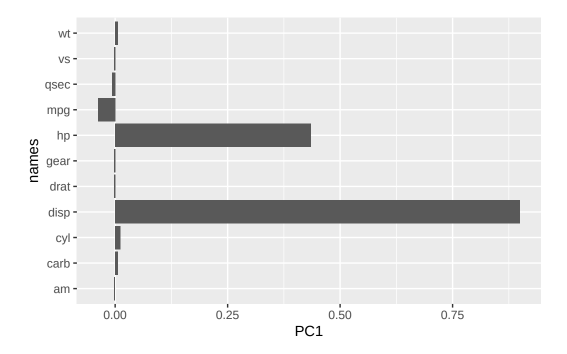
Plot the loadings

```
library(ggplot2)

r1 <-as.data.frame(pc.noscale$rotation)
r1$names <-rownames(pc.noscale$rotation)
r1$names</pre>
```

```
[1] "mpg" "cyl" "disp" "hp" "drat" "wt" "qsec" "vs" "am" "gear" [11] "carb"
```

```
ggplot(r1) +
aes(PC1, names) +
geom_col()
```

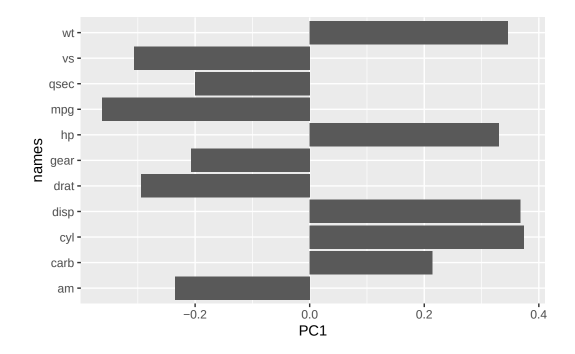


```
library(ggplot2)

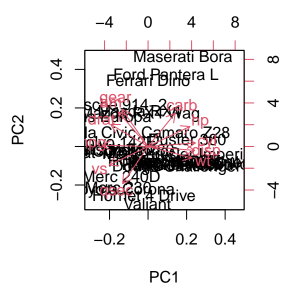
r2 <-as.data.frame(pc.scale$rotation)
r2$names <-rownames(pc.scale$rotation)
r2$names

[1] "mpg" "cyl" "disp" "hp" "drat" "wt" "qsec" "vs" "am" "gear"
[11] "carb"</pre>
```

```
ggplot(r2) +
aes(PC1, names) +
geom_col()
```



biplot(pc.scale)



Take-home: Generally we alawys want to set scale = TRUE when we do this type

of analysis to oavoid our anlyses being dominated by individual varioables with the largest variance just due to their unit of measurement.

### FNA breast cancer data

Load the data into R.

```
# Save your input data file into your Project directory
fna.data <- "WisconsinCancer.csv"

# Complete the following code to input the data and store as wisc.df
wisc.df <- read.csv(fna.data, row.names = 1)
head(wisc.df)</pre>
```

	diagnosis ra	adius_mean	texture_mean	perimeter_mean	area_mean		
842302	M	17.99	10.38	122.80	1001.0		
842517	M	20.57	17.77	132.90	1326.0		
84300903	M	19.69	21.25	130.00	1203.0		
84348301	M	11.42	20.38	77.58	386.1		
84358402	M	20.29	14.34	135.10	1297.0		
843786	M	12.45	15.70	82.57	477.1		
	smoothness_m	mean compac	tness_mean co	ncavity_mean co	oncave.poir	nts_mean	
842302	0.11	1840	0.27760	0.3001		0.14710	
842517	0.08	3474	0.07864	0.0869		0.07017	
84300903	0.10960		0.15990	0.1974		0.12790	
84348301	0.14250		0.28390	0.2414		0.10520	
84358402	0.10030		0.13280	0.1980		0.10430	
843786	0.12	2780	0.17000	0.1578		0.08089	
	symmetry_mea	an fractal_	dimension_mea	n radius_se tex	kture_se pe	erimeter_se	
842302	0.241	19	0.0787	1 1.0950	0.9053	8.589	
842517	0.1812		0.0566	7 0.5435	0.7339	3.398	
84300903	0.2069		0.0599	9 0.7456	0.7869	4.585	
84348301	0.2597		0.0974	4 0.4956	1.1560	3.445	
84358402	0.1809		0.0588	3 0.7572	0.7813	5.438	
843786	0.208	37	0.0761	3 0.3345	0.8902	2.217	
	area_se smoo	othness_se	compactness_s	e concavity_se	concave.po	oints_se	
842302	153.40	0.006399	0.0490	4 0.05373		0.01587	
842517	74.08	0.005225	0.0130	0.01860		0.01340	
84300903	94.03	0.006150	0.0400	6 0.03832		0.02058	
84348301	27.23	0.009110	0.0745	0.05661		0.01867	

84358402	94.44	0.011490	0.02461	0.05688	0.01885			
843786	27.19	0.007510	0.03345	0.03672	0.01137			
	symmetry_se f:	ractal_dimens	sion_se rad:	ius_worst textu	re_worst			
842302	0.03003	0.	006193	25.38	17.33			
842517	0.01389	0.	003532	24.99	23.41			
84300903	0.02250	0.	004571	23.57	25.53			
84348301	0.05963	0.	009208	14.91	26.50			
84358402	0.01756	0.	005115	22.54	16.67			
843786	0.02165	0.	005082	15.47	23.75			
	perimeter_wor	st area_worst	smoothnes	s_worst compact:	ness_worst			
842302	184.	60 2019.0	)	0.1622	0.6656			
842517	158.	80 1956.0	)	0.1238	0.1866			
84300903	152.	50 1709.0	)	0.1444	0.4245			
84348301	98.	87 567.7	•	0.2098	0.8663			
84358402	152.	20 1575.0	)	0.1374	0.2050			
843786	103.	40 741.6	3	0.1791	0.5249			
	concavity_wor	st concave.po	ints_worst	symmetry_worst				
842302	0.71	19	0.2654	0.4601				
842517	0.24	16	0.1860	0.2750				
84300903	0.45	04	0.2430	0.3613				
84348301	0.68	69	0.2575	0.6638				
84358402	0.40	00	0.1625	0.2364				
843786	0.53		0.1741	0.3985				
fractal_dimension_worst								
842302		0.11890						
842517		0.08902						
84300903		0.08758						
84348301		0.17300						
84358402		0.07678						
843786		0.12440						

## dim(wisc.df)

[1] 569 31

Q1. How many observations are in this dataset?

### nrow(wisc.df)

[1] 569

#### dim(wisc.df)

#### [1] 569 31

There are 569 observations in this dataset. > Q2. How many of the observations have a malignant diagnosis?

#### table(wisc.df\$diagnosis)

```
B M
357 212
```

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

#### ncol(wisc.df)

#### [1] 31

#### colnames(wisc.df)

```
[1] "diagnosis"
                                "radius_mean"
 [3] "texture_mean"
                                "perimeter_mean"
 [5] "area_mean"
                                "smoothness_mean"
                                "concavity_mean"
 [7] "compactness_mean"
 [9] "concave.points_mean"
                                "symmetry_mean"
[11] "fractal_dimension_mean"
                                "radius_se"
[13] "texture_se"
                                "perimeter_se"
[15] "area_se"
                                "smoothness_se"
[17] "compactness_se"
                                "concavity_se"
[19] "concave.points_se"
                                "symmetry_se"
[21] "fractal dimension se"
                                "radius worst"
[23] "texture_worst"
                                "perimeter_worst"
                                "smoothness worst"
[25] "area_worst"
[27] "compactness_worst"
                                "concavity_worst"
[29] "concave.points_worst"
                                "symmetry_worst"
[31] "fractal_dimension_worst"
```

A useful function for this is grep()

```
length(grep("_mean", colnames(wisc.df)))
```

[1] 10

Before we go any further we need to exclude the diagnosis column from any future analysis this tells us whether a sample to cancer or non-cancer.

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

[1] M M M M M M M Levels: B M

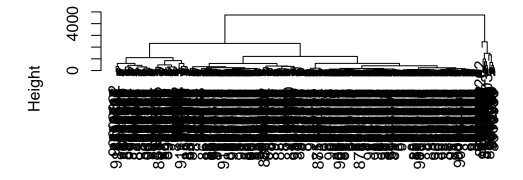
Removing diagnosis column

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

Let's see if we can cluster the wisc.data to find some structure in the dataset.

```
hc <- hclust(dist(wisc.data))
plot(hc)</pre>
```

## **Cluster Dendrogram**



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

## **Principal Component Analysis (PCA)**

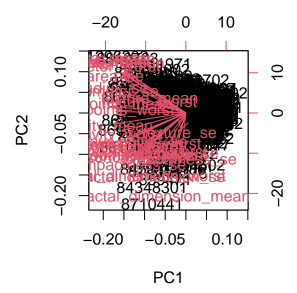
Jump right into pca

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

#### Importance of components:

```
PC1
                                 PC2
                                         PC3
                                                 PC4
                                                         PC5
                                                                 PC6
                                                                          PC7
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
                                                                PC13
                                                                        PC14
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
                                          PC17
                                                  PC18
                                                          PC19
                                                                   PC20
                                  PC16
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
                                         PC24
                                                         PC26
                          PC22
                                  PC23
                                                 PC25
                                                                 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
                       0.99749 0.99830 0.9989 0.99942 0.99969 0.99992 0.99997
Cumulative Proportion
                          PC29
                                  PC30
Standard deviation
                       0.02736 0.01153
Proportion of Variance 0.00002 0.00000
Cumulative Proportion 1.00000 1.00000
```

```
biplot(wisc.pr)
```



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

44.27% of proportion of the originial variance is captured by the first principa components (PC1).

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

The first three principal components 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?

The first seven principal components are required to describe at least 90% of the original variance in the data.

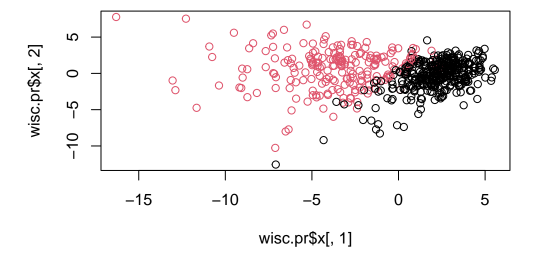
Q7. What stands out to you about this plot? Is it easy or difficult to understand? Why?

This biplot sucks! Everything is so jumbled up it is difficult to parse it. We need to build our own PCA store plot of PC1 vs PC2

head(wisc.pr\$x)

```
PC1
                         PC2
                                    PC3
                                             PC4
                                                        PC5
                                                                    PC6
842302
       -9.184755 -1.946870 -1.1221788 3.6305364 1.1940595 1.41018364
842517
        -2.385703
                   3.764859 -0.5288274 1.1172808 -0.6212284 0.02863116
84300903 -5.728855
                   1.074229 -0.5512625 0.9112808 0.1769302 0.54097615
84348301 -7.116691 -10.266556 -3.2299475 0.1524129 2.9582754 3.05073750
84358402 -3.931842
                    1.946359 1.3885450 2.9380542 -0.5462667 -1.22541641
843786
        -2.378155 -3.946456 -2.9322967 0.9402096 1.0551135 -0.45064213
                PC7
                            PC8
                                        PC9
                                                 PC10
                                                            PC11
                                                                       PC12
         2.15747152 0.39805698 -0.15698023 -0.8766305 -0.2627243 -0.8582593
842302
842517
         0.01334635 -0.24077660 -0.71127897 1.1060218 -0.8124048 0.1577838
84300903 -0.66757908 -0.09728813 0.02404449 0.4538760 0.6050715 0.1242777
84348301 1.42865363 -1.05863376 -1.40420412 -1.1159933 1.1505012 1.0104267
84358402 -0.93538950 -0.63581661 -0.26357355 0.3773724 -0.6507870 -0.1104183
843786
         0.49001396  0.16529843  -0.13335576  -0.5299649  -0.1096698  0.0813699
               PC13
                            PC14
                                        PC15
                                                    PC16
                                                                PC17
842302
         0.10329677 -0.690196797 0.601264078 0.74446075 -0.26523740
842517
        -0.94269981 -0.652900844 -0.008966977 -0.64823831 -0.01719707
84300903 -0.41026561 0.016665095 -0.482994760 0.32482472 0.19075064
84348301 -0.93245070 -0.486988399 0.168699395 0.05132509 0.48220960
84358402 0.38760691 -0.538706543 -0.310046684 -0.15247165 0.13302526
843786
        -0.02625135 0.003133944 -0.178447576 -0.01270566 0.19671335
               PC18
                          PC19
                                     PC20
                                                  PC21
                                                              PC22
842302
        -0.54907956 0.1336499 0.34526111 0.096430045 -0.06878939
         0.31801756 -0.2473470 -0.11403274 -0.077259494 0.09449530
842517
84300903 -0.08789759 -0.3922812 -0.20435242 0.310793246 0.06025601
84348301 -0.03584323 -0.0267241 -0.46432511 0.433811661 0.20308706
84358402 -0.01869779 0.4610302 0.06543782 -0.116442469
                                                        0.01763433
843786
        -0.29727706 -0.1297265 -0.07117453 -0.002400178 0.10108043
                                        PC25
                                                     PC26
               PC23
                            PC24
                                                                 PC27
842302
         0.08444429 0.175102213 0.150887294 -0.201326305 -0.25236294
        -0.21752666 -0.011280193 0.170360355 -0.041092627 0.18111081
842517
84300903 -0.07422581 -0.102671419 -0.171007656 0.004731249 0.04952586
84348301 -0.12399554 -0.153294780 -0.077427574 -0.274982822 0.18330078
84358402 0.13933105 0.005327110 -0.003059371 0.039219780 0.03213957
         0.03344819 - 0.002837749 - 0.122282765 - 0.030272333 - 0.08438081
843786
                 PC28
                              PC29
                                           PC30
842302
        842517
         0.0325955021 -0.005682424 0.0018662342
84300903 0.0469844833 0.003143131 -0.0007498749
84348301 0.0424469831 -0.069233868 0.0199198881
84358402 -0.0347556386 0.005033481 -0.0211951203
         0.0007296587 -0.019703996 -0.0034564331
843786
```

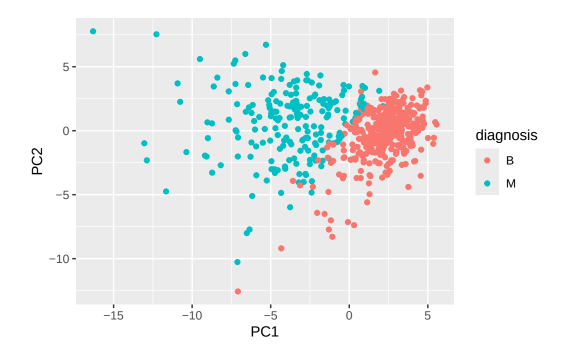
```
plot(wisc.pr$x[,1],wisc.pr$x[,2], col = diagnosis)
```



Make a ggplot version of this score plot

```
pc <- as.data.frame(wisc.pr$x)

ggplot(pc) +
  aes(PC1, PC2, col = diagnosis) +
  geom_point()</pre>
```

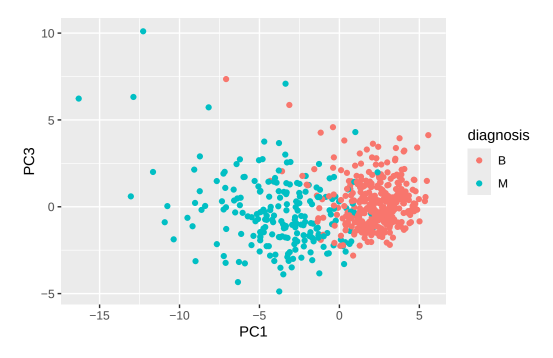


Q8. Generate a similar plot for principal components 1 and 3. What do you notice about these plots?

Since PC2 explains more variance than PC3, the plot with PC2 has better separation between the benign and malignant.

```
pc <- as.data.frame(wisc.pr$x)

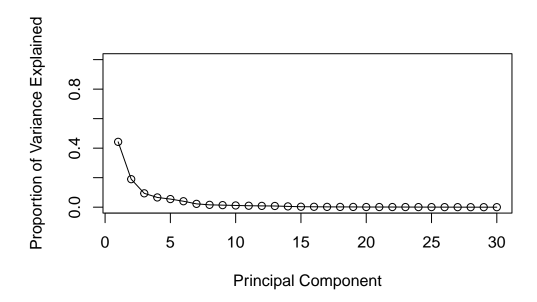
ggplot(pc) +
  aes(PC1, PC3, col = diagnosis) +
  geom_point()</pre>
```

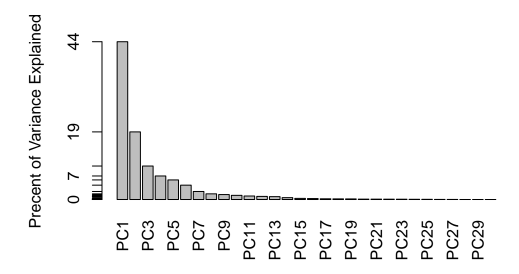


### $\#\# Variance\ Explained$

```
# 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

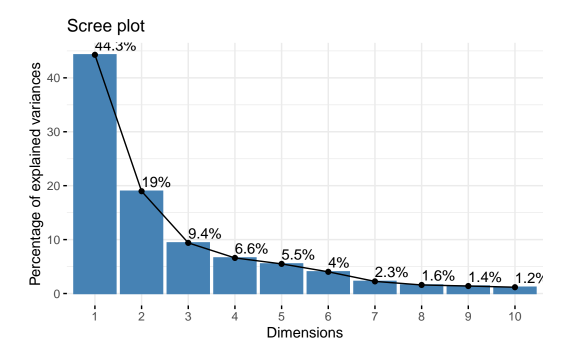




```
## ggplot based graph
#install.packages("factoextra")
library(factoextra)
```

 ${\tt Welcome!\ Want\ to\ learn\ more?\ See\ two\ factoextra-related\ books\ at\ https://goo.gl/ve3WBa}$ 

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



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

The component of the loading vector for the feature concave.points\_mean is -0.2608638.

### wisc.pr\$rotation["concave.points\_mean",1]

#### [1] -0.2608538

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

The minimum number of components is five principal components required to explain 80% of the variance of the data.

```
summary(wisc.pr)
```

### Importance of components:

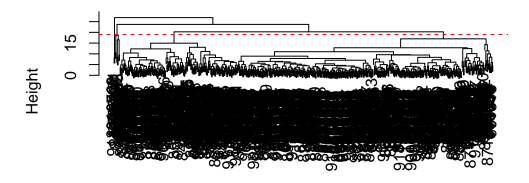
PC1 PC2 PC3 PC4 PC5 PC6 PC7 Standard deviation 3.6444 2.3857 1.67867 1.40735 1.28403 1.09880 0.82172 Proportion of Variance 0.4427 0.1897 0.09393 0.06602 0.05496 0.04025 0.02251 Cumulative Proportion 0.4427 0.6324 0.72636 0.79239 0.84734 0.88759 0.91010

```
PC8
                                  PC9
                                         PC10
                                                PC11
                                                        PC12
                                                                PC13
                                                                        PC14
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
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
```

### #Hiearchical 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, method = "complete")</pre>
```

```
plot(wisc.hclust)
abline(h = 19, col="red", lty=2)
```



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

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

The height at which the cluster model has 4 clusters is height 19.

```
wisc.hclust.clusters <- cutree(wisc.hclust, k = 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
```

```
cluster <- cutree(wisc.hclust, k = 10)
table(cluster, diagnosis)</pre>
```

```
diagnosis cluster B M
```

```
1
     12
         86
2
      0
         59
3
           3
      0
4
   331
         39
5
         20
      0
6
      2
7
     12
           0
8
      0
           2
9
      0
           2
10
      0
           1
```

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

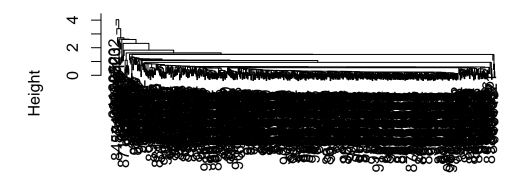
The other numbers of clusters don't do that well.

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

The other methods of "single", "complete", and "average" are more difficult in identify a good cluster while "ward.D2" works with this data.dist dataset.

### Clustering in PC Space

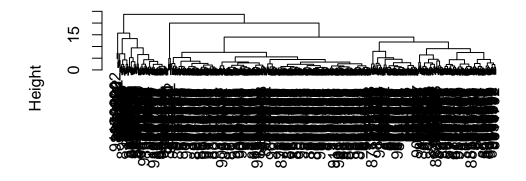
```
hb <- hclust(dist(wisc.pr$x[,1:2]), method = "single")
plot(hb)</pre>
```



dist(wisc.pr\$x[, 1:2]) hclust (\*, "single")

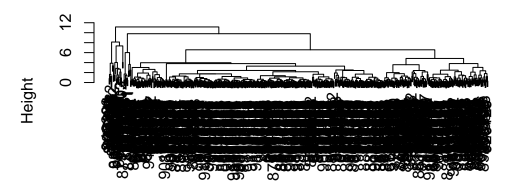
hd <- hclust(dist(wisc.pr\$x[,1:2]), method = "complete")
plot(hd)</pre>

# **Cluster Dendrogram**



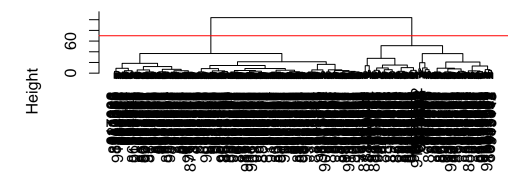
dist(wisc.pr\$x[, 1:2])
hclust (\*, "complete")

```
he <- hclust(dist(wisc.pr$x[,1:2]), method = "average")
plot(he)</pre>
```



dist(wisc.pr\$x[, 1:2])
hclust (\*, "average")

```
hc <- hclust(dist(wisc.pr$x[,1:2]), method = "ward.D2")
plot(hc)
abline(h = 70, col = "red")</pre>
```



dist(wisc.pr\$x[, 1:2])
hclust (\*, "ward.D2")

## K-means clustering

```
wisc.km <- kmeans(scale(wisc.data), centers= 2, nstart= 20)
table(wisc.km$cluster, diagnosis)</pre>
```

diagnosis

B M
1 343 37
2 14 175

# Cluster membership vector

```
grps <- cutree(hc, h = 70)
table(grps)</pre>
```

```
grps
1 2
195 374
```

### table(diagnosis)

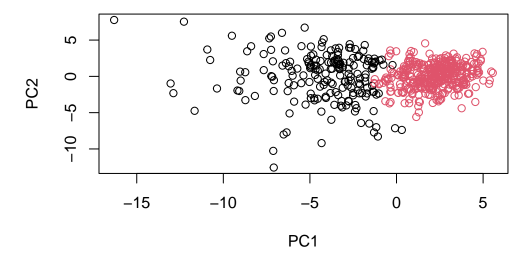
```
diagnosis
B M
357 212
```

Cross-table to see how my clustering groups correspond to expert diagnosis vector of M and B values

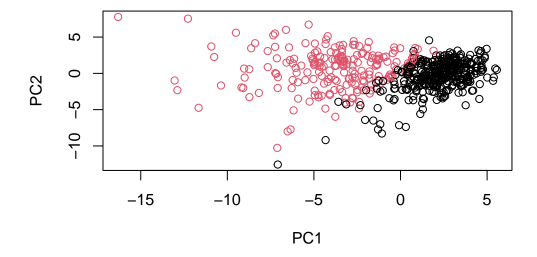
### table(grps, diagnosis)

```
diagnosis
grps B M
1 18 177
2 339 35
```

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



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



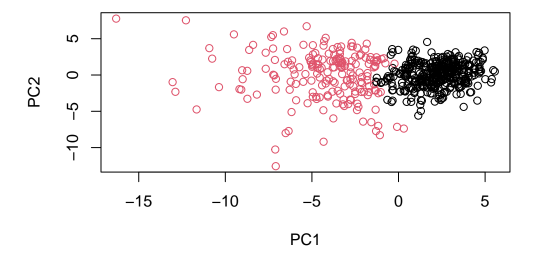
```
g <- as.factor(grps)
levels(g)</pre>
```

[1] "1" "2"

```
g <- relevel(g,2)
levels(g)</pre>
```

[1] "2" "1"

```
# Plot using our re-ordered factor
plot(wisc.pr$x[,1:2], col=g)
```



Positive => cancer M Negative => non-cancer B

True = cluster/group 1 False = group 2

True Positive 188 False Positive 28 True Negative 329 False Negative 24

We can use our PCA results (wisc.pr) to make predictions on our new unseen data.

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

There is a good decent amount of True positives and True negatives but there is still a non-significant amount of false negatives and false positives.

```
## 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" )
wisc.pr.hclust.clusters <- cutree(wisc.pr.hclust, k=2)
table(wisc.pr.hclust.clusters, diagnosis)</pre>
```

```
diagnosis
wisc.pr.hclust.clusters B M
1 28 188
2 329 24
```

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.

The k-means and hierarchical clustering models do mostly the same in terms of separating the diagnoses as their values for true positives and negatives, and false positives and negatives are pretty close.

```
table(wisc.km$cluster, diagnosis)
```

```
diagnosis
B M
1 343 37
2 14 175
```

```
table(wisc.hclust.clusters, diagnosis)
```

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

```
summary(diagnosis)
```

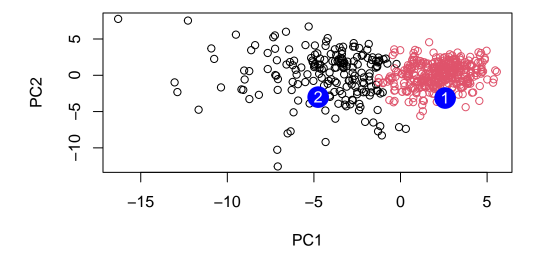
B M 357 212

```
summary(wisc.hclust.clusters)
```

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 1.000 1.000 3.000 2.369 3.000 4.000
```

```
#url <- "new_samples.csv"
url <- "https://tinyurl.com/new-samples-CSV"
new <- read.csv(url)
npc <- predict(wisc.pr, newdata=new)</pre>
```

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



### 343/(343+37)

[1] 0.9026316

### 343/(343+40)

[1] 0.8955614

### 329/(329+24)

[1] 0.9320113

### 175/(175+37)

[1] 0.8254717

```
172/(172+40)
```

[1] 0.8113208

```
188/(188+24)
```

#### [1] 0.8867925

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

Principal clustering analysis has the best specificity at 88.68% and the best sensitivity at 93.20%.

### Sensitivity (TP/(TP+FN))

K-means - 343/(343+37) = 0.9026316 H clustering - 343/(343+40) = 0.8955614 PCA -  $329/(329+24) = \mathbf{0.9320113}$ 

### Specificity (TN/(TN+FN))

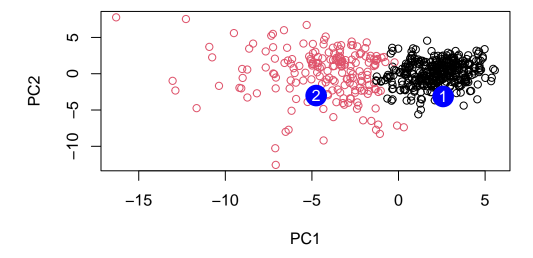
K-means - 175/(175+37) = 0.8254717 H clustering - 172/(172+40) = 0.8113208 PCA -  $188/(188+24) = \mathbf{0.8867925}$ 

```
#url <- "new_samples.csv"
url <- "https://tinyurl.com/new-samples-CSV"
new <- read.csv(url)
npc <- predict(wisc.pr, newdata=new)
npc</pre>
```

```
PC1
                  PC2
                           PC3
                                     PC4
                                              PC5
                                                        PC6
                                                                  PC7
[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
          PC8
                   PC9
                           PC10
                                    PC11
                                             PC12
                                                              PC14
                                                      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
        PC15
                  PC16
                             PC17
                                       PC18
                                                  PC19
                                                           PC20
[1,] 0.3216974 -0.1743616 -0.07875393 -0.11207028 -0.08802955 -0.2495216
```

```
PC21
                      PC22
                                 PC23
                                            PC24
                                                        PC25
                                                                     PC26
     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
            PC27
                         PC28
                                      PC29
                                                   PC30
[1,] 0.220199544 -0.02946023 -0.015620933 0.005269029
[2,] -0.001134152
                  0.09638361 0.002795349 -0.019015820
```

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



Q18. Which of these new patients should we prioritize for follow up based on your results?

We should follow up with group 2 as they are the ones that have the predicted malignant samples.