Class 08 Breast Cancer Mini Project

Irene Hsieh (PID: A16197563)

Before dive into breast cancer project, we will finish class 7 (where we left off) first.

```
url2 <- "https://tinyurl.com/expression-CSV"
rna.data <- read.csv(url2, row.names=1)
head(rna.data)

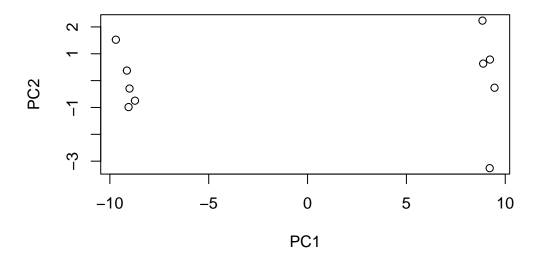
wt1 wt2 wt3 wt4 wt5 ko1 ko2 ko3 ko4 ko5
gene1 439 458 408 429 420 90 88 86 90 93
gene2 219 200 204 210 187 427 423 434 433 426
gene3 1006 989 1030 1017 973 252 237 238 226 210
gene4 783 792 829 856 760 849 856 835 885 894
gene5 181 249 204 244 225 277 305 272 270 279
gene6 460 502 491 491 493 612 594 577 618 638
```

Q10: How many genes and samples are in this data set? 6 genes, and 10 samples.

##Run PCA

```
## Again we have to take the transpose of our data
pca <- prcomp(t(rna.data), scale=TRUE)

## Simple un polished plot of pc1 and pc2
plot(pca$x[,1], pca$x[,2], xlab="PC1", ylab="PC2")</pre>
```



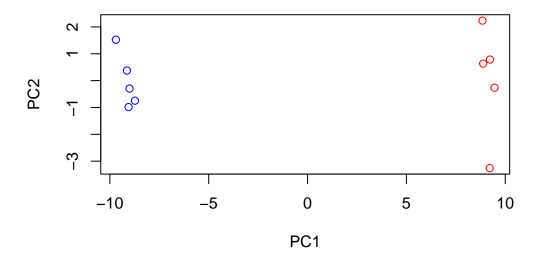
summary(pca)

```
Importance of components:
```

```
PC2
                                         PC3
                                                 PC4
                                                         PC5
                          PC1
                                                                 PC6
                                                                          PC7
Standard deviation
                       9.6237 1.5198 1.05787 1.05203 0.88062 0.82545 0.80111
Proportion of Variance 0.9262 0.0231 0.01119 0.01107 0.00775 0.00681 0.00642
Cumulative Proportion 0.9262 0.9493 0.96045 0.97152 0.97928 0.98609 0.99251
                           PC8
                                   PC9
                                            PC10
Standard deviation
                       0.62065 0.60342 3.345e-15
Proportion of Variance 0.00385 0.00364 0.000e+00
Cumulative Proportion 0.99636 1.00000 1.000e+00
```

```
# We have 5wt and 5ko samples
mycols <- c(rep("blue",5), rep("red",5))
mycols</pre>
```

[1] "blue" "blue" "blue" "blue" "red" "red" "red" "red" plot(pca\$x[,1], pca\$x[,2], xlab="PC1", ylab="PC2", col = mycols)



I could examine which genes contribute to this first PC

```
head(sort(abs(pca$rotation[,1]), decreasing = T))
```

gene100 gene66 gene45 gene68 gene98 gene60 0.1038708 0.1038455 0.1038402 0.1038395 0.1038372 0.1038055

#Analysis of Breast Cancer FNA data.

```
wisc.df <- read.csv("WisconsinCancer.csv", row.names = 1)
head(wisc.df)</pre>
```

	diagnosis radi	us_mean ·	texture_mean	<pre>perimeter_mean</pre>	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_mea	n compac	tness_mean co	oncavity_mean c	oncave.poin	ts_mean
842302	0.1184	-0	0.27760	0.3001		0.14710

842517	0.08	171	0.0786	1	0.0869	1	0.07017
84300903	0.10		0.1599		0.1974		0.12790
84348301	0.10		0.1399		0.1974		0.10520
84358402	0.14		0.2839		0.1980		0.10320
843786	0.12		0.1700		0.1578		0.08089
0.40000	symmetry_mea						-
842302	0.241			07871	1.0950	0.9053	8.589
842517	0.181			05667	0.5435	0.7339	3.398
84300903	0.206			05999	0.7456	0.7869	4.585
84348301	0.259			09744			3.445
84358402	0.180			05883	0.7572		5.438
843786	0.208			07613	0.3345	0.8902	2.217
	area_se smoo		_		•		-
842302	153.40	0.006399		04904	0.053		0.01587
842517	74.08	0.005225		01308	0.018		0.01340
84300903		0.006150		04006	0.038		0.02058
84348301		0.009110		07458	0.056		0.01867
84358402	94.44	0.011490	0.	02461	0.056	888	0.01885
843786	27.19	0.007510	0.	03345	0.036	372	0.01137
	symmetry_se	fractal_d	${\tt imension_s}$	e radi	us_worst t	exture_wors	t
842302	0.03003		0.00619	3	25.38	17.3	3
842517	0.01389		0.00353	2	24.99	23.4	1
84300903	0.02250		0.00457	1	23.57	25.5	3
84348301	0.05963		0.00920	8	14.91	26.5	0
84358402	0.01756		0.00511	5	22.54	16.6	7
843786	0.02165		0.00508	2	15.47	23.7	5
	perimeter_wo	rst area_	worst smoo	thness	s_worst com	pactness_wo	rst
842302	184	.60 20	019.0		0.1622	0.6	656
842517	158	.80 19	956.0		0.1238	0.1	866
84300903	152	.50 1	709.0		0.1444	0.4	245
84348301	98	.87	567.7		0.2098	0.8	663
84358402	152	.20 1	575.0		0.1374	0.2	050
843786	103	.40	741.6		0.1791	0.5	249
	concavity_wo	rst conca	ve.points_	worst	symmetry_w	orst	
842302	0.7		-	.2654	•	4601	
842517	0.2	416	0	.1860	0.	2750	
84300903	0.4	504	0	.2430	0.	3613	
84348301	0.6	869	0	.2575	0.	6638	
84358402	0.4			.1625		2364	
843786	0.5			.1741		3985	
	fractal_dime						
842302	_	0.118					
842517		0.089					
·		3.030	- •				

```
84300903
                           0.08758
84348301
                           0.17300
84358402
                           0.07678
843786
                           0.12440
  wisc.data <- wisc.df[,-1]
  diagnosis <- as.factor(wisc.df$diagnosis)</pre>
     Q1. How many observations are in this dataset? (columns)
  nrow(wisc.df)
[1] 569
  ncol(wisc.data)
[1] 30
30 observations
     Q2. How many of the observations have a malignant diagnosis?
212 malignant diagnosis
  table(wisc.df$diagnosis)
  В
      Μ
357 212
     Q3. How many variables/features in the data are suffixed with _mean?
10 features
  length(grep("_mean$", colnames(wisc.data), value = TRUE))
[1] 10
```

##Principal Component Analysis

Here we will use precomp() on the wisc.data object - the one without the diagnosis column.

First, we have decide whether to use the scale = TRUE argument when we run precomp()

We can look at the means and sd of each column. If they are similar than we are all good to go. If not we should not use scale = TRUE

Check column means and standard deviations
colMeans(wisc.data)

radius_mean	texture_mean	perimeter_mean
1.412729e+01	1.928965e+01	9.196903e+01
area_mean	${\tt smoothness_mean}$	compactness_mean
6.548891e+02	9.636028e-02	1.043410e-01
${\tt concavity_mean}$	concave.points_mean	symmetry_mean
8.879932e-02	4.891915e-02	1.811619e-01
fractal_dimension_mean	radius_se	texture_se
6.279761e-02	4.051721e-01	1.216853e+00
perimeter_se	area_se	smoothness_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	${\tt compactness_worst}$	concavity_worst
1.323686e-01	2.542650e-01	2.721885e-01
concave.points_worst	symmetry_worst	${\tt fractal_dimension_worst}$
1.146062e-01	2.900756e-01	8.394582e-02

apply(wisc.data,2,sd)

radius_mean 3.524049e+00	texture_mean 4.301036e+00	perimeter_mean 2.429898e+01
area_mean	smoothness_mean	compactness_mean
3.519141e+02	1.406413e-02	5.281276e-02
concavity_mean	concave.points_mean	symmetry_mean
7.971981e-02	3.880284e-02	2.741428e-02
${\tt fractal_dimension_mean}$	radius_se	texture_se

```
7.060363e-03
                                 2.773127e-01
                                                         5.516484e-01
        perimeter_se
                                      area_se
                                                         smoothness_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
        8.266372e-03
                                 2.646071e-03
                                                          4.833242e+00
       texture_worst
                             perimeter_worst
                                                            area_worst
        6.146258e+00
                                 3.360254e+01
                                                          5.693570e+02
    smoothness_worst
                            compactness_worst
                                                      concavity_worst
        2.283243e-02
                                 1.573365e-01
                                                          2.086243e-01
concave.points_worst
                               symmetry_worst fractal_dimension_worst
                                 6.186747e-02
        6.573234e-02
                                                          1.806127e-02
```

```
# Perform PCA on wisc.data
wisc.pr <- prcomp( wisc.data, scale = TRUE)
# Look at summary of results
summary(wisc.pr)</pre>
```

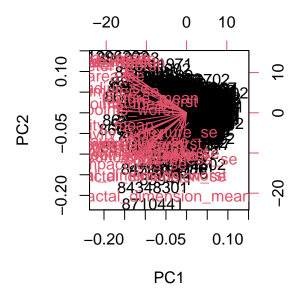
Importance of components:

```
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
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
```

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

- 44.27% of the original variance is captured by the first PC.
 - 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 required to describe at least 90% of the original variance in the data.

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



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

It is messy, we need to build our own plot. Since it is labeled by patients, it is hard to read in this plot.

```
attributes(wisc.pr)
```

\$names

[1] "sdev" "rotation" "center" "scale" "x"

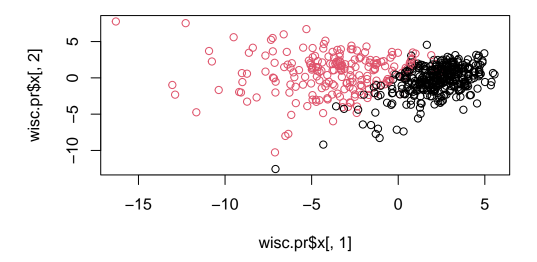
\$class

[1] "prcomp"

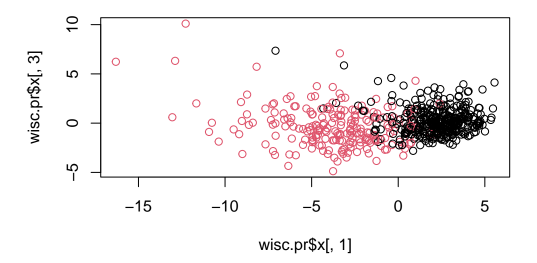
Q8. Generate a similar plot for principal components 1 and 3. What do you notice about these plots?

The plots are more condensed on the lower side of the x-axis whiling comparing with PC1 and 2.

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



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



```
library(ggplot2)

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

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



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?

-0.2608538

```
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?

```
tbl <- summary(wisc.pr)
which (tbl$importance[3,] > 0.8)[1]
```

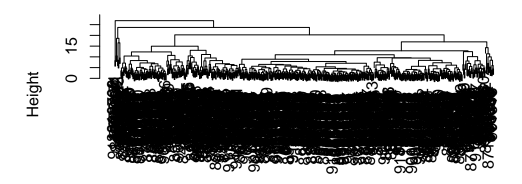
PC5 5

Hierarcal clustering

The main function for Hierarchical clustering is called hclust() it takes a distance matrix as outcome

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

Cluster Dendrogram



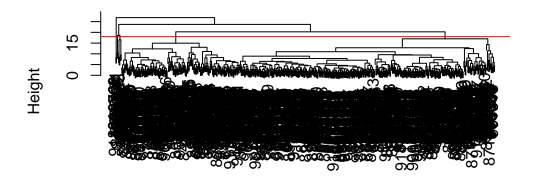
d hclust (*, "complete")

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

18

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

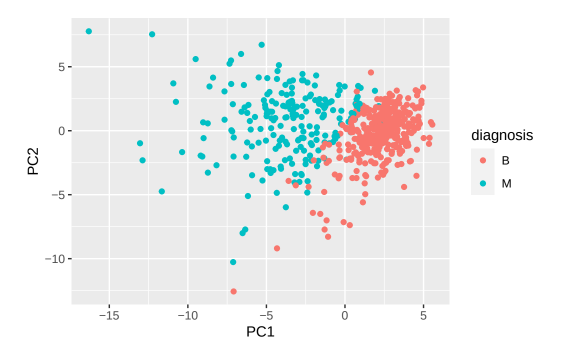
Cluster Dendrogram



d hclust (*, "complete")

Come back here. later to see how our cluster grps correspond to M or B groups.

```
ggplot(pc)+
aes(PC1, PC2, col = diagnosis)+
geom_point()
```



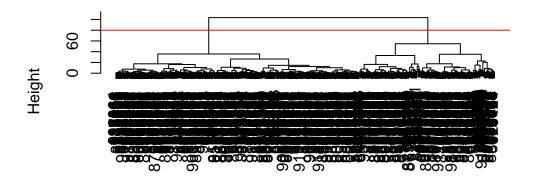
##5. Combining methods

Here we will perform clustering on our PCA results rather than the original data.

In other words we will cluster using wisc.pr\$x - our new better variables or PCs. We can choose as many or as few PCs to use as we like.It is your call!

```
d.pc <- dist(wisc.pr$x[,1:3])
wisc.pr.hclust <- hclust(d.pc,method = "ward.D2")
plot(wisc.pr.hclust)
abline(h=80, col = "red")</pre>
```

Cluster Dendrogram



d.pc hclust (*, "ward.D2")

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

Average method is my favorite results since it is more straightforward to understand

```
grps<- cutree(wisc.pr.hclust, h =80)
table(grps)</pre>
```

grps 1 2 203 366

We can use table() finction to male a cross table as well as just a count table

```
table(diagnosis)
```

diagnosis

B M

357 212

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

```
2
```

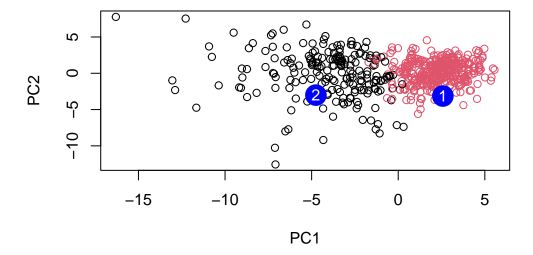
```
table(grps, diagnosis)

diagnosis
grps B M
1 24 179
2 333 33
```

Write a note here about how to read this cross table result. The results indicate that our cluster 1 mostly captures cancer(M) and our cluster 2 mostly captures healthy (B) samples/individuals.

7. Prediction

```
#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
          PC1
                                                   PC5
                    PC2
                              PC3
                                         PC4
                                                             PC6
                                                                        PC7
     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
                                                  PC12
           PC8
                              PC10
                                        PC11
                                                            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
[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
[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=grps)
  points(npc[,1], npc[,2], col="blue", pch=16, cex=3)
```



Q18. Which of these new patients should we prioritize for follow up based on your results? patient 2