Class 08: Machine Learning

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Breast Cancer Project

Today we are going to explore some data from the University of Wisconsin Cancer Center on Breast biopsy data.

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

	diagnosis	radius_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		
	smoothnes	s_mean compa	ctness_mean co	oncavity_mean co	oncave.poi	nts_mean	
842302	0	.11840	0.27760	0.3001		0.14710	
842517	0	.08474	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	.12780	0.17000	0.1578		0.08089	
symmetry_mean fractal_dimension_mean radius_se texture_se perimeter_se							
842302	0.	2419	0.0787	1.0950	0.9053	8.589	
842517	0.	1812	0.0566	0.5435	0.7339	3.398	
84300903	0.	2069	0.0599	0.7456	0.7869	4.585	
84348301	0.	2597	0.0974	14 0.4956	1.1560	3.445	
84358402	0.	1809	0.0588	3 0.7572	0.7813	5.438	
843786	0.	2087	0.0761	0.3345	0.8902	2.217	
	area_se s	moothness_se	compactness_s	se concavity_se	concave.p	oints_se	
842302	153.40	0.006399	0.0490	0.05373		0.01587	

842517	74.08	0.005225	0.01308	0.01860	0.01340
84300903	94.03	0.006150	0.04006	0.03832	0.02058
84348301	27.23	0.009110	0.07458	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	fractal_dime	nsion_se rad	ius_worst text	ure_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_wo	rst area_wor	st smoothnes	s_worst compac	tness_worst
842302	184	.60 2019	0.0	0.1622	0.6656
842517	158	.80 1956	3.0	0.1238	0.1866
84300903	152	.50 1709	0.0	0.1444	0.4245
84348301	98	.87 567	7.7	0.2098	0.8663
84358402	152	.20 1575	5.0	0.1374	0.2050
843786	103	.40 741	6	0.1791	0.5249
	concavity_wo	rst concave.	points_worst	symmetry_wors	t
842302	0.7	119	0.2654	0.460	1
842517	0.2	416	0.1860	0.275	0
84300903	0.4	504	0.2430	0.361	3
84348301	0.6	869	0.2575	0.663	8
84358402	0.4	000	0.1625	0.236	4
843786	0.5	355	0.1741	0.398	5
	<pre>fractal_dime</pre>	nsion_worst			
842302		0.11890			
842517		0.08902			
84300903		0.08758			
84348301		0.17300			
84358402		0.07678			
843786		0.12440			

Q. How many patient samples are in this dataset?

nrow(wisc.data)

[1] 569

There are 569 patients in this dataset.

Q. How many cancer (M) and non cancer (B) samples are there?

```
table(wisc.data$diagnosis)
```

```
B M
357 212
```

Save the diagnosis for later use as a reference to compare how well we do with PCA etc.

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

```
[75] В М В М М В В В М М В М М В В В М В В М М В В В М М В В В М В В В М В В
[482] B B B B B B B M B M B B B B B B B B M M B M B B B B B B M B B M B M B M M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B M B 
[556] B B B B B B B M M M M M B
Levels: B M
```

Now exclude diagnosis column from the data

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

Q. How many "dimensions", "variables", "columns" are there in this dataset?

ncol(wisc)
```

[1] 30

```
dim(wisc)
```

[1] 569 30

Principal Component Analysis (PCA)

To perform PCA in R we can use the prcomp() function. It takes as input a numeric dataset and optional scale=FALSE/TRUE argument.

We general always want to set scale=TRUE but let's make sure by checking if the mean and standard deviation values are different across these 30 columns.

round(colMeans(wisc))

radius_mean	toxturo moon	perimeter_mean
-	texture_mean	•
14	19	92
area_mean	${\tt smoothness_mean}$	compactness_mean
655	0	0
${\tt concavity_mean}$	concave.points_mean	symmetry_mean
0	0	0
fractal_dimension_mean	radius_se	texture_se
0	0	1
perimeter_se	area_se	smoothness_se
3	40	0
compactness_se	concavity_se	concave.points_se
0	0	0
symmetry_se	fractal_dimension_se	radius_worst
0	0	16
texture_worst	perimeter_worst	area_worst
26	107	881
smoothness_worst	compactness_worst	concavity_worst
0	0	0
concave.points_worst	symmetry_worst	<pre>fractal_dimension_worst</pre>
0	0	0

```
pca <- prcomp(wisc, scale =TRUE)
summary(pca)</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
                                  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
```

attributes(pca)

```
$names
[1] "sdev"          "rotation" "center"          "x'
$class
[1] "prcomp"

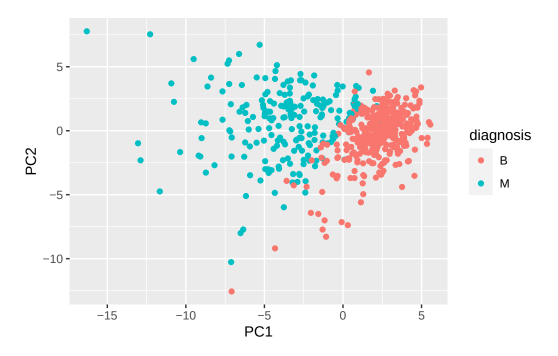
plot(pca$x[,1], pca$x[,2], col=diagnosis)
```

Cumulative Proportion 1.00000 1.00000

```
library(ggplot2)

x <- as.data.frame(pca$x)

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



Q. How much variance is captured in the top 3 PCs.

They capture 76% of the total variance.

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? This tells us how much this original feature contributes to the first PC.

```
pca$rotation["concave.points_mean",1]
```

[1] -0.2608538

attributes(pca)

```
$names
```

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

\$class

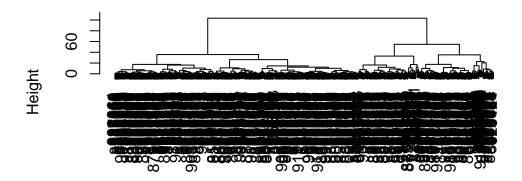
[1] "prcomp"

Combine PCA results with clustering.

We can use our new PCA variables (i.e. the scores along the PCs contained in the pca\$x) as input for other methods such as clustering.

```
# hclust needs a distance matrix as an input
d <-dist( pca$x[, 1:3] )
hc <- hclust(d, method = "ward.D2")
plot(hc)</pre>
```

Cluster Dendrogram



d hclust (*, "ward.D2")

To get our cluster membership vector we can use the cutree() function and specify a height (h) or number of groups (k).

```
grps <- cutree(hc, h =80)
table(grps)

grps
    1      2
203 366</pre>
```

I want to find out how many diagnosis "M" and "B" are in each group?

```
table(diagnosis)
```

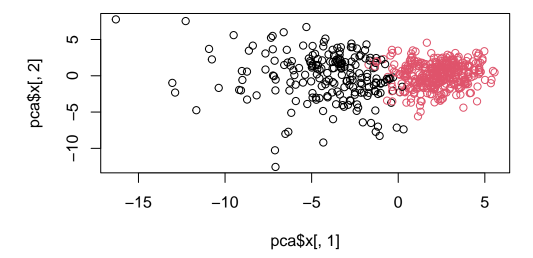
```
diagnosis
B M
357 212
```

table(diagnosis, grps)

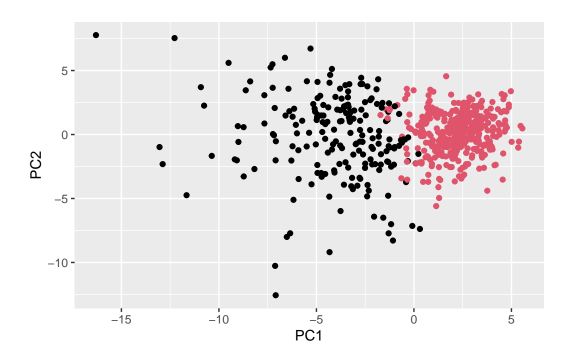
```
grps
diagnosis 1 2
B 24 333
M 179 33
```

We can also plot our results using out clustering vector grps.

```
plot(pca$x[,1], pca$x[,2], col = grps)
```



```
ggplot(x) +
  aes( PC1, PC2) +
  geom_point( col=grps)
```



Q15. what is the specificity and sensitivity?

```
tbl <- table(diagnosis, grps)
tbl</pre>
```

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

```
# TP: grps 1,M (179) [2,1]
TP <- tbl[2,1]
# FP grps 1, B (24) [1,1]
FP <- tbl[1,1]
# TN: grps 2, B (333) [1,2]
TN <- tbl[1,2]
# FN: grps 2, M (33) [2,2]</pre>
```

```
# Sensitivity TP/(TP+FN)
sens <- TP/(TP+FN)
sens

[1] 0.8443396

# Specificity TN/(TN+FN)
spec <- TN/(TN+FN)
spec

[1] 0.9098361

The sensitivity is 84% and the specificity is 91%.

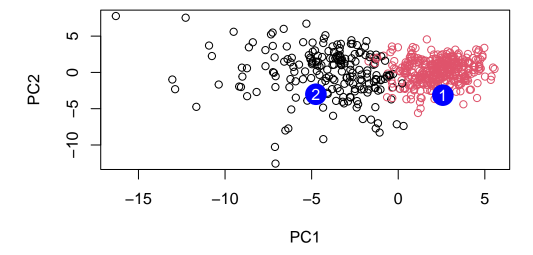
Prediction
```

url <- "https://tinyurl.com/new-samples-CSV"</pre>

#url <- "new_samples.csv"</pre>

```
new <- read.csv(url)</pre>
  npc <- predict(pca, newdata=new)</pre>
  npc
          PC1
                   PC2
                             PC3
                                        PC4
                                                 PC5
                                                            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
           PC8
                    PC9
                             PC10
                                       PC11
                                                PC12
                                                          PC13
                                                                  PC14
[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
[1,] 0.3216974 -0.1743616 -0.07875393 -0.11207028 -0.08802955 -0.2495216
PC21
                    PC22
                              PC23
                                         PC24
                                                    PC25
                                                                PC26
[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
                       PC28
                                               PC30
            PC27
[1,] 0.220199544 -0.02946023 -0.015620933 0.005269029
[2,] -0.001134152  0.09638361  0.002795349 -0.019015820
```

```
plot(pca$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")
```



Q16 Which of these new patients should we prioritize for follow up based on your results?

Patient 2 should be prioritized since in the previous graph, that group predicted malignant cancer.