Class 7: Machine Learning 1

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Clustering

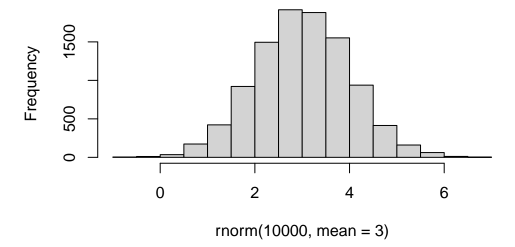
We will start with k-means clustering, one of the most prevelent of all clustering methods.

K-means clustering

K-means clustering is the commonest algorithm for doing clustering.

```
hist(rnorm(10000, mean = 3))
```

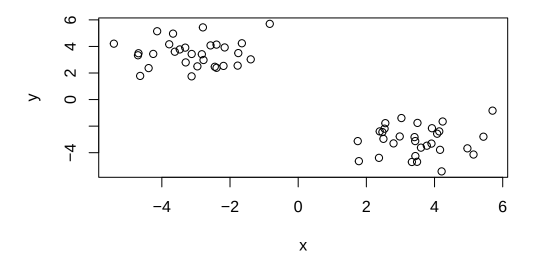
Histogram of rnorm(10000, mean = 3)



```
tmp \leftarrow c(rnorm(30, 3), rnorm(30, -3)) # we generate two bunches of points a <- cbind(x = tmp, y = rev(tmp)) # for coordinate. two buckles of points around (3, -3) a head(a)
```

```
x y
[1,] 3.437485 -4.252542
[2,] 4.077042 -2.571675
[3,] 3.412754 -2.826503
[4,] 2.369859 -4.387497
[5,] 3.499400 -1.757220
[6,] 3.334941 -4.699861
```



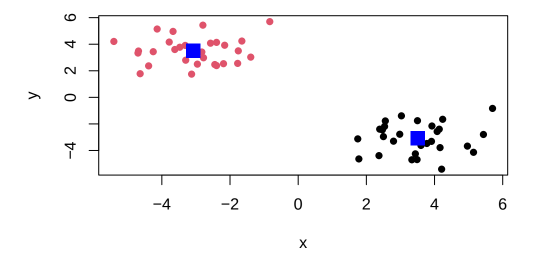


The primitive of calling k-means is just kmeans()

```
k <- kmeans(a, centers = 2, nstart = 20)
k</pre>
```

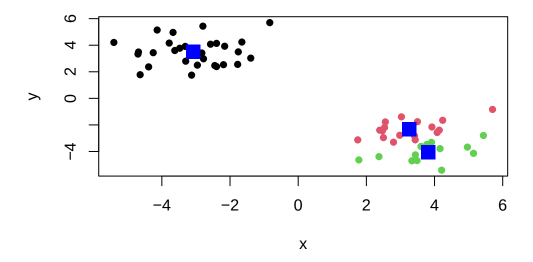
K-means clustering with 2 clusters of sizes 30, 30

```
Cluster means:
      Х
1 3.501267 -3.084773
2 -3.084773 3.501267
Clustering vector:
Within cluster sum of squares by cluster:
[1] 65.05629 65.05629
(between_SS / total_SS = 90.9 %)
Available components:
[1] "cluster"
            "centers"
                      "totss"
                                "withinss"
                                           "tot.withinss"
[6] "betweenss"
            "size"
                      "iter"
                                "ifault"
Let's check some of the properties:
 k$size
[1] 30 30
 k$cluster
k$centers
      X
1 3.501267 -3.084773
2 -3.084773 3.501267
 plot(a, col=k$cluster, pch=16)
 points(k$centers, col="blue", pch=15, cex=2)
```



What if we chance the number of centers?

```
k3 <- kmeans(a, centers=3, nstart=20)
plot(a, col=k3$cluster, pch=16)
points(k3$centers, col="blue", pch=15, cex=2)</pre>
```



Hierarchical clustering

HC can reveal multi-level structural information rather than just imposing the structure (k-means).

Funciton in R is hclust(), which requires a distance matrix as the input.

```
hc <- hclust(dist(a))
hc</pre>
```

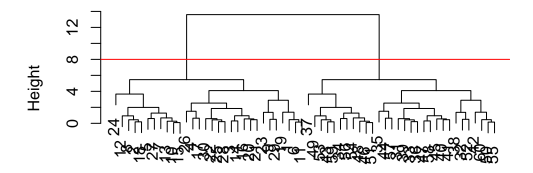
```
Call:
hclust(d = dist(a))
```

Cluster method : complete
Distance : euclidean

Number of objects: 60

```
plot(hc)
abline(h=8, col='red')
```

Cluster Dendrogram

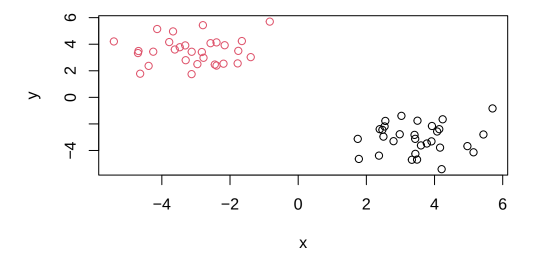


dist(a) hclust (*, "complete")

We can 'cut the tree' using cutree:

```
groups <- cutree(hc, h = 8)
groups</pre>
```

plot(a, col=groups)



Seems cool!

PCA

Now we use the UK food dataset for the PCA exploration.

```
#load the dataset
url <- "https://tinyurl.com/UK-foods"
x <- read.csv(url)
head(x)</pre>
```

```
X England Wales Scotland N.Ireland
                       105
                             103
                                       103
                                                   66
1
          Cheese
2
                             227
   Carcass_meat
                       245
                                       242
                                                  267
3
     Other_meat
                      685
                             803
                                       750
                                                  586
4
            Fish
                             160
                       147
                                       122
                                                   93
5 Fats_and_oils
                       193
                             235
                                       184
                                                  209
          Sugars
                                       147
                       156
                             175
                                                  139
```

```
dim(x)
```

Q1. How many rows and columns are in your new data frame named x? What R functions could you use to answer this questions?

17 and 4. dim

Let's set the index to be the first column:

```
rownames(x) <- x[,1]
x <- x[,-1]
head(x)
```

	England	Wales	${\tt Scotland}$	${\tt N.Ireland}$
Cheese	105	103	103	66
Carcass_meat	245	227	242	267
Other_meat	685	803	750	586
Fish	147	160	122	93
Fats_and_oils	193	235	184	209
Sugars	156	175	147	139

Alternatively, we can use the parameter in read.csv

```
x <- read.csv(url, row.names=1)
head(x)</pre>
```

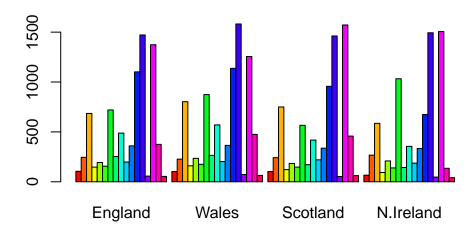
	England	Wales	Scotland	N.Ireland
Cheese	105	103	103	66
Carcass_meat	245	227	242	267
Other_meat	685	803	750	586
Fish	147	160	122	93
Fats_and_oils	193	235	184	209
Sugars	156	175	147	139

Q2. Which approach to solving the 'row-names problem' mentioned above do you prefer and why? Is one approach more robust than another under certain circumstances?

I prefer the latter method, since it preserve when running multiple times.

Spotting major differences and trends

```
barplot(as.matrix(x), beside=T, col=rainbow(nrow(x)))
```



Q3: Changing what optional argument in the above barplot() function results in the following plot?

We can use ${\tt beside=FALSE}$

```
barplot(as.matrix(x), beside=F, col=rainbow(nrow(x)))
```



Q5: Generating all pairwise plots may help somewhat. Can you make sense of the following code and resulting figure? What does it mean if a given point lies on the diagonal for a given plot?

```
pairs(x, col=rainbow(10), pch=16)
```



Points on the diagonal means two countries have the same value on a certain category.

Q6. What is the main differences between N. Ireland and the other countries of the UK in terms of this data-set?

They are further away from the diagonal.

PCA to the rescue.

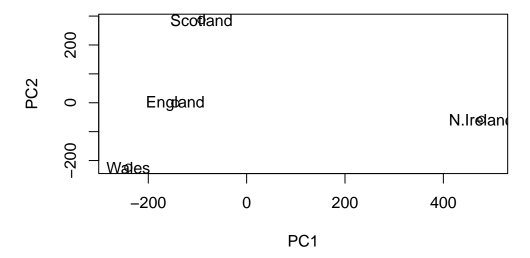
```
# Use the prcomp() PCA function
pca <- prcomp( t(x) )
summary(pca)</pre>
```

Importance of components:

	PC1	PC2	PC3	PC4
Standard deviation	324.1502	212.7478	73.87622	3.176e-14
Proportion of Variance	0.6744	0.2905	0.03503	0.000e+00
Cumulative Proportion	0.6744	0.9650	1.00000	1.000e+00

Q7. Complete the code below to generate a plot of PC1 vs PC2. The second line adds text labels over the data points.

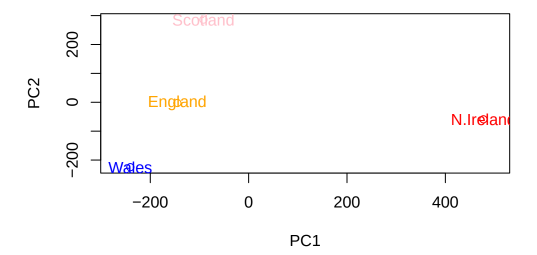
```
# Plot PC1 vs PC2
plot(pca$x[,1], pca$x[,2], xlab="PC1", ylab="PC2", xlim=c(-270,500))
text(pca$x[,1], pca$x[,2], colnames(x))
```



Q8. Customize your plot so that the colors of the country names match the colors in our UK and Ireland map and table at start of this document.

```
color <- c("orange", "blue", "pink", "red")

plot(pca$x[,1], pca$x[,2], xlab="PC1", ylab="PC2", xlim=c(-270,500), col=color)
text(pca$x[,1], pca$x[,2], colnames(x), col=color)</pre>
```



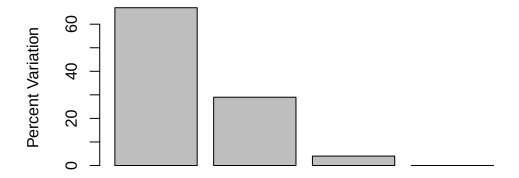
We can further analyze the standard deviation

```
v <- round( pca$sdev^2/sum(pca$sdev^2) * 100 )
v

[1] 67 29 4 0

## or the second row here...
z <- summary(pca)
z$importance</pre>
```

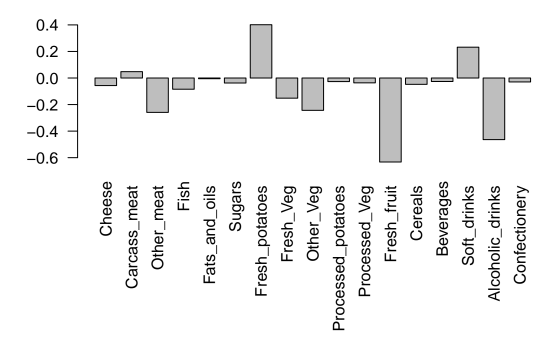
```
barplot(v, xlab="Principal Component", ylab="Percent Variation")
```



Principal Component

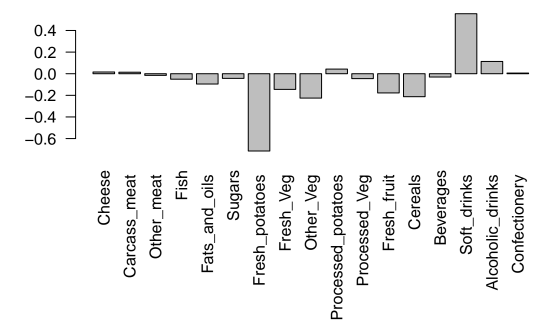
Digging deeper (variable loadings)

```
## Lets focus on PC1 as it accounts for > 90% of variance
par(mar=c(10, 3, 0.35, 0))
barplot( pca$rotation[,1], las=2 )
```



Q9: Generate a similar 'loadings plot' for PC2. What two food groups feature prominantely and what does PC2 maninly tell us about?

```
## Lets focus on PC1 as it accounts for > 90% of variance
par(mar=c(10, 3, 0.35, 0))
barplot( pca$rotation[,2], las=2 )
```



The two dominant factors are Fresh_potatoes and Soft_drinks. From the previous plot, we can see that Wales and Scotland are at two ends of the PC2, therefore, that means those two factors can explain their difference.

Using ggplot for these figures

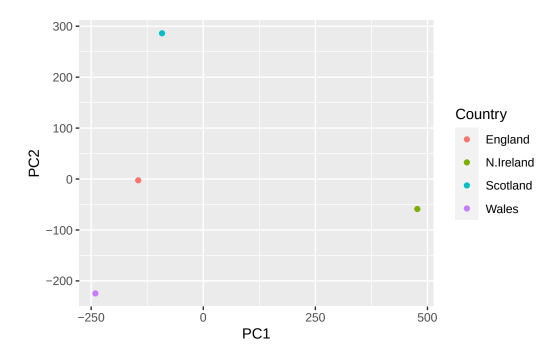
```
library(ggplot2)

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

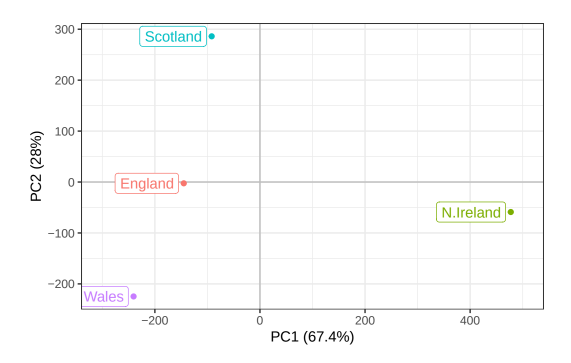
df_lab <- tibble::rownames_to_column(df, "Country")

# Our first basic plot

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

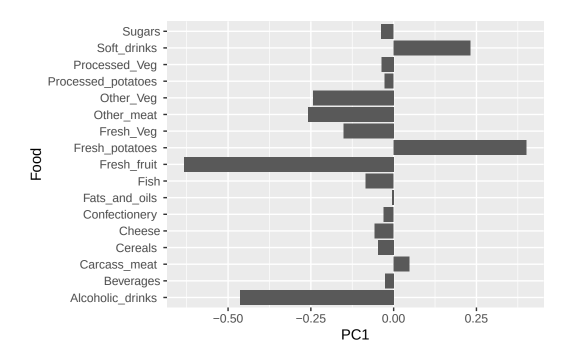


```
ggplot(df_lab) +
  aes(PC1, PC2, col=Country, label=Country) +
  geom_hline(yintercept = 0, col="gray") +
  geom_vline(xintercept = 0, col="gray") +
  geom_point(show.legend = FALSE) +
  geom_label(hjust=1, nudge_x = -10, show.legend = FALSE) +
  expand_limits(x = c(-300,500)) +
  xlab("PC1 (67.4%)") +
  ylab("PC2 (28%)") +
  theme_bw()
```

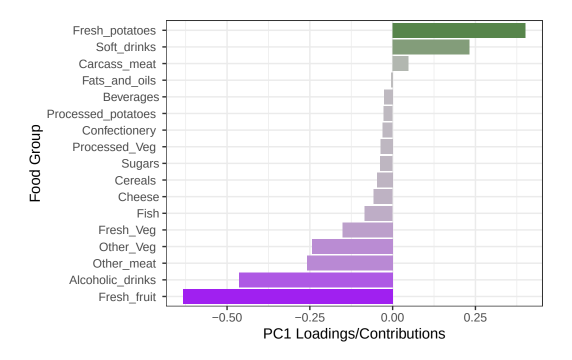


```
ld <- as.data.frame(pca$rotation)
ld_lab <- tibble::rownames_to_column(ld, "Food")

ggplot(ld_lab) +
  aes(PC1, Food) +
  geom_col()</pre>
```

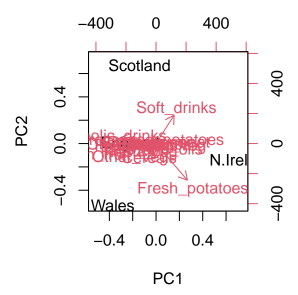


```
ggplot(ld_lab) +
  aes(PC1, reorder(Food, PC1), bg=PC1) +
  geom_col() +
  xlab("PC1 Loadings/Contributions") +
  ylab("Food Group") +
  scale_fill_gradient2(low="purple", mid="gray", high="darkgreen", guide=NULL) +
  theme_bw()
```



Biplot

The inbuilt biplot() can be useful for small datasets biplot(pca)



2. PCA of RNA-seq data

Again load the dataset:

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

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

Q10: How many genes and samples are in this data set?

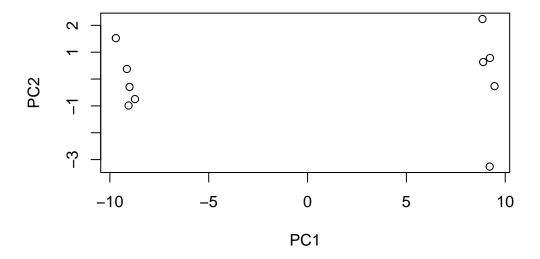
```
dim(rna.data)
```

[1] 100 10

10 samples, 100 genes.

```
## 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:

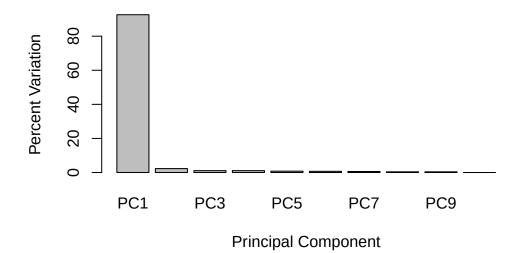
PC2 PC3 PC4 PC5 PC1 PC6 PC7 Standard deviation 9.6237 1.5198 1.05787 1.05203 0.88062 0.82545 0.80111 Proportion of Variance 0.9262 0.0231 0.01119 0.01107 0.00775 0.00681 0.00642 Cumulative Proportion 0.9262 0.9493 0.96045 0.97152 0.97928 0.98609 0.99251 PC8 PC9 PC10 Standard deviation 0.62065 0.60342 3.457e-15 Proportion of Variance 0.00385 0.00364 0.000e+00 Cumulative Proportion 0.99636 1.00000 1.000e+00

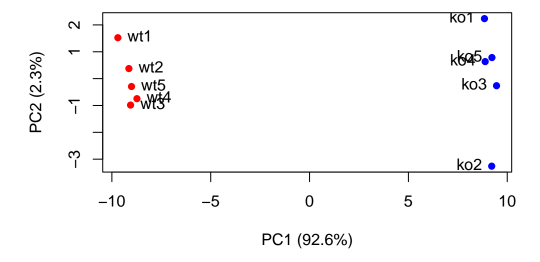
```
plot(pca, main="Quick scree plot")
```

Quick scree plot



Scree Plot

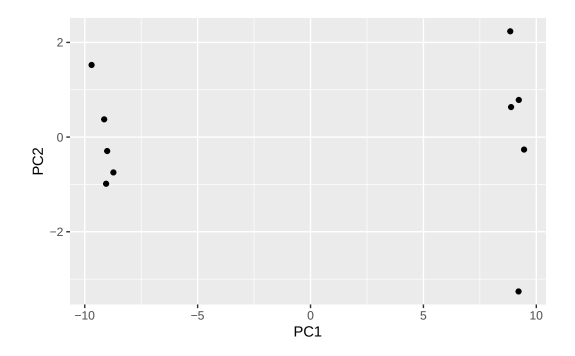


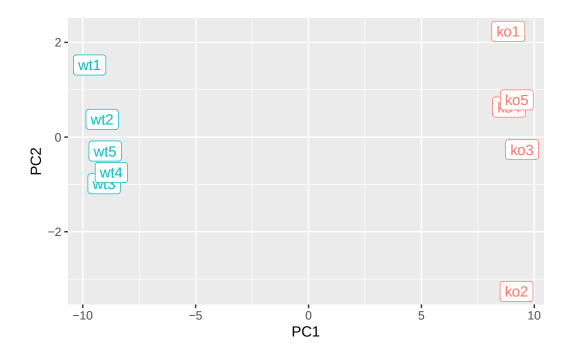


```
library(ggplot2)

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

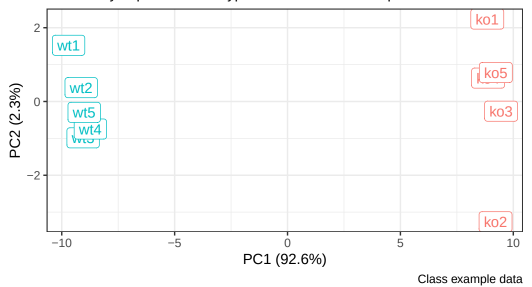
# Our first basic plot
ggplot(df) +
   aes(PC1, PC2) +
   geom_point()</pre>
```





PCA of RNASeq Data

PC1 clealy seperates wild-type from knock-out samples



(Optional) Gene loading:

```
loading_scores <- pca$rotation[,1]

## Find the top 10 measurements (genes) that contribute
## most to PC1 in either direction (+ or -)
gene_scores <- abs(loading_scores)
gene_score_ranked <- sort(gene_scores, decreasing=TRUE)

## show the names of the top 10 genes
top_10_genes <- names(gene_score_ranked[1:10])
top_10_genes

[1] "gene100" "gene66" "gene45" "gene68" "gene98" "gene60" "gene21"
[8] "gene56" "gene10" "gene90"</pre>
```