```
title: "Class 07 Machine Learning 1"
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# First up, kmeans()
Demo of using kmeans() function in R. FIrst make up some data with a known structure.
```{r}
We will generate random numbers
tmp <- c(rnorm(30, -3), rnorm(30, 3))
tmp
This creates two columns - Points that are (3,-3) and (-3, 3)
x <- cbind(x=tmp, y=rev(tmp))</pre>
Plot a graph of the two columns
plot(x)
Now that we have made up data in 'x', we will see what kmeans() does to the data
```{r}
# We will use the function kmeans() to create 2 clusters, with 20 iterations (i.e. it is
running through the algorithim 20 diff times)
k \leftarrow kmeans(x, centers = 2, nstart = 20)
k
When we print k, 2 clusters are created, with each cluster containing 30 points each. It
also tells us our cluster means, which are the (x, y) coordinates of the means of the 2
clusters.
There are different available components that can help us learn more about the cluster.
> Q. How many points are in each cluster?
```{r}
k$size
> Q. How do we get to the cluster membership/assignment?
```{r}
k$cluster
> Q. What about cluster centers?
```{r}
k$centers
Now that we have returned the main results of our clusters, we will use them to generate a
plot of our kmeans().
To color the plot according to the cluster
plot(x, col=k$cluster)
Ti distinguish the center value of each cluster with different color, in this case, blue
points(k$centers, col="blue", pch=15)
Now for hclust()
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We will cluster the same data 'x' with the `hclust()` function. In this case `hclust()
requires a distance matrix as an input.
```{r}
hc <- hclust(dist(x))</pre>
hc
Let's plot our hclust result!
```{r}
plot(hc)
To get our cluster membership vector we need to "cut" the tree with `cutree()`
```{r}
groups <- cutree(hc, h=8)
groups
Now plot our data with the hclust() results.
```{r}
plot(x, col=groups)
Hands on with Principle Component Analysis (PCA)
First, we need to import our data.
Q1. How many rows and columns are in your new data frame named x? What R functions could
you use to answer this questions?
```{r}
url <- "https://tinyurl.com/UK-foods"</pre>
x <- read.csv(url)</pre>
```{r}
nrow(x)
ncol(x)
Checking your data. It is always a good idea to examine your imported data to make sure it
meets your expectations.
```{r}
#To view the entire data frame
View(x)
#To view the first 6 rows of the data frame
head(x)
#To view the last 6 rows of the data frame
tail(x)
It appears that the data is not set properly, as the first column is labeled as 'X',
giving us 5 variables not 4. To fix this we use the function rownames().
```{r}
#To class for the first column
rownames(x) <- x[,1]
#To remove the first column
x < -x[,-1]
head(x)
```

```
Another way to do it is by calling read.csv()
x <- read.csv(url, row.names=1)</pre>
head(x)
```{r}
#To find out the dimensions (x, y) of the data frame: dim()
dim(x)
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 think the solution for the 'row-names problem' that I prefer is the: 'x <- read.csv(url,
row.names=1)' approach, as you have more control as to which column that you are changing.
I think using the x[,-1] method would work if you only had to adjust the first column,
because you might continue to erase more variables.
Spotting major differences and trends
```{r}
barplot(as.matrix(x), beside=T, col=rainbow(nrow(x)))
```{r}
cols <- rainbow(nrow(x))</pre>
barplot( as.matrix(x), col=cols)
#It is hard to compare the data in this format.
```{r}
pairs(x, col=cols)
PCA to the rescue!
The main R PCA function is called `prcomp()`. We will need to give it the transpose of our
input data.
```{r}
pca <- prcomp(t(x))</pre>
pca
```{r}
#Like kmeans(), there are different attributes for prcomp()
attributes(pca)
#Example: calling the center values.
pca$center
To make our new PCA plot (PCA score plot) we access `pca$x`
```{r}
pca$x
country_cols <- c("orange", "red", "blue", "green")
plot(pca$x[,1], pca$x[,2], xlab = "PC1", ylab = "PC2")</pre>
text(pca$x[,1], pca$x[,2], colnames(x), col = country_cols)
# PCA of RNA-seq Data
```

. . .

First, we have to read our data.

```
"\{r\}
url2 <- "https://tinyurl.com/expression-CSV"
rna.data <- read.csv(url2, row.names=1)
head(rna.data)
"\{r\}
pca <- prcomp(t(rna.data))
summary(pca)
plot(pca)
"\{r\}
plot(pca\$x[,1], pca\$x[,2], xlab = "PC1", ylab = "PC2")
text(pca\$x[,1], pca\$x[,2], colnames(rna.data))</pre>
```