

Recap of Lecture 7

- Reviewed the major steps for R function writing
- Introduced the CRAN & Bioconductor repositories for packages that extend R functionality
- Explored installing and using the ggplot2, bio3d, rgl, rentrez, igraph, blogdown, shiny, and circlize packages
- Discussed how to judge the utility, usability and development status of R packages.

[Muddy Point Feedback Link]

Today's Menu

- Introduction to machine learning
 - Unsupervised, supervised and reinforcement learning
- Clustering
 - K-means clustering
 - Hierarchical clustering
 - Heatmap representations
- Dimensionality reduction, visualization and 'structure' analysis
 - Principal Component Analysis (PCA)
- Hands-on application to cell classification

Types of machine learning

- Unsupervised learning
 - Finding structure in unlabeled data
- Supervised learning
 - Making predictions based on labeled data
 - Predictions like regression or classification
- Reinforcement learning
 - Making decisions based on past experience

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Here your eyes can clearly see 3 natural groupings

- Breaks observations into k pre-defined number of clusters
- You define k the number of clusters!
 - Imagine you had data that you could plot along a line and you knew you had to put them into k=3 "clusters" (e.g. data from three types of tumor cells)



Here your eyes can clearly see 3 natural groupings How does k-means attempt to define this grouping?

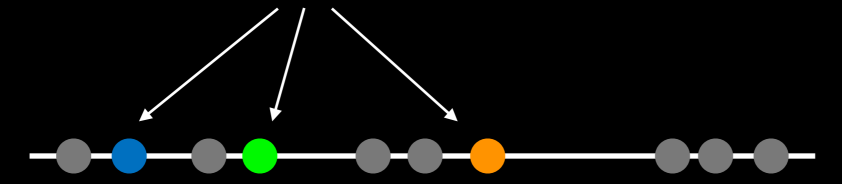
Step 1.
Select *k* (the number of clusters)



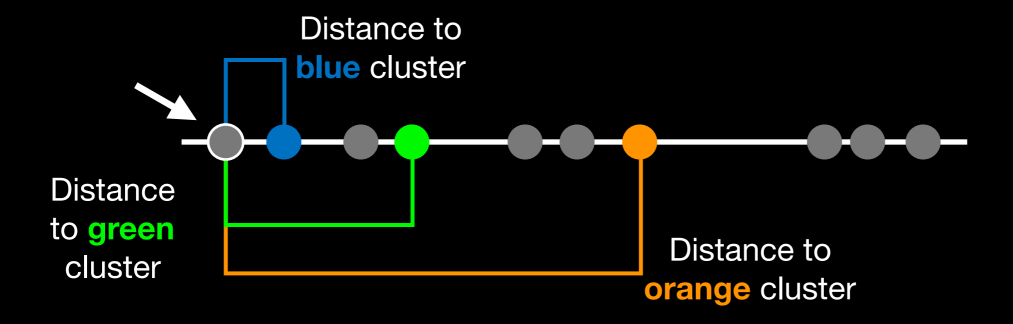
Step 2.

Select **k=3** distant data points at random

These are the initial clusters



Step 3. Measure distance between the 1st point and the k=3 initial clusters



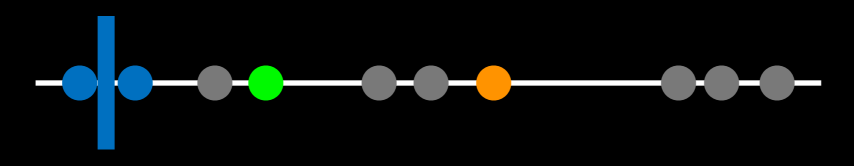
Step 4.
Assign the 1st point to the nearest cluster



Step 5.

Update cluster centers

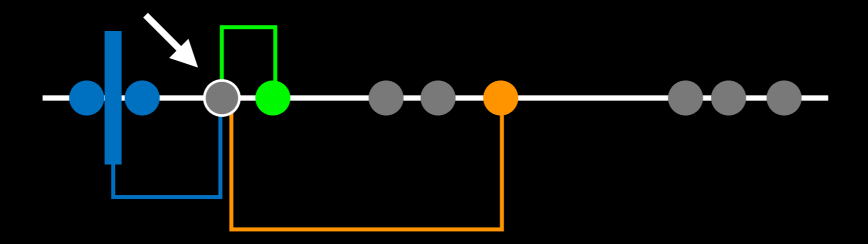
Calculate the mean value for the blue cluster including the new point



Step 6.

Assign next point to closest cluster

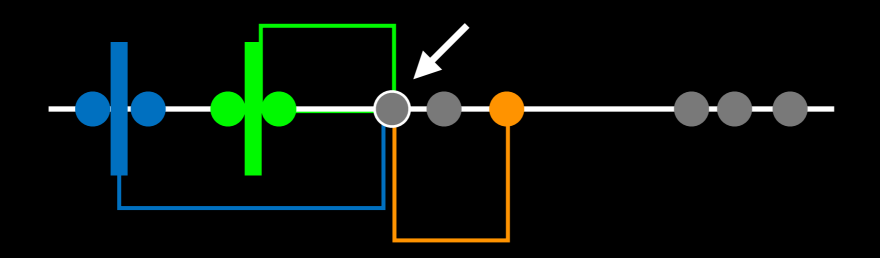
Use updated cluster centers for distance calculation



Step 7.

Update cluster centers and move to next point

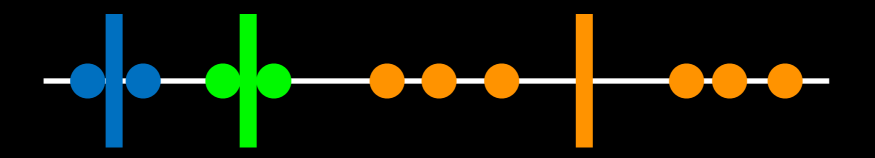
Use updated cluster centers for distance calculation



Step 8.

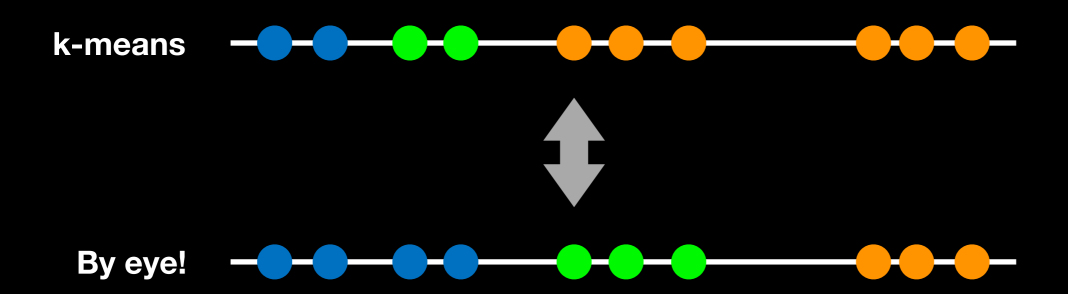
Repeat for each point

Each time updating cluster centers



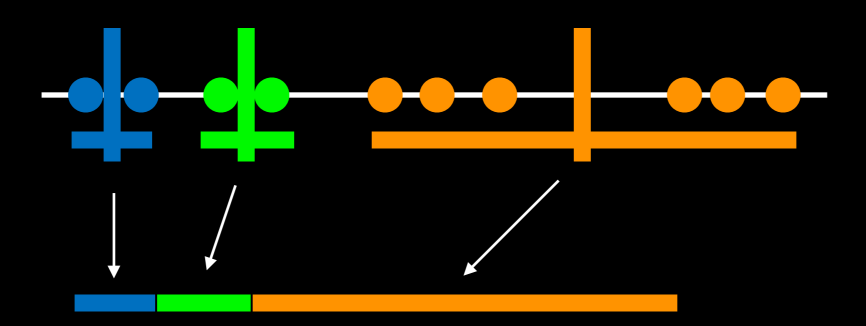
Hmm....

Here the k-means result does not look as good as what we were able to do by eye!



Step 9.

Assess the quality of the clustering by adding up the variation within each cluster



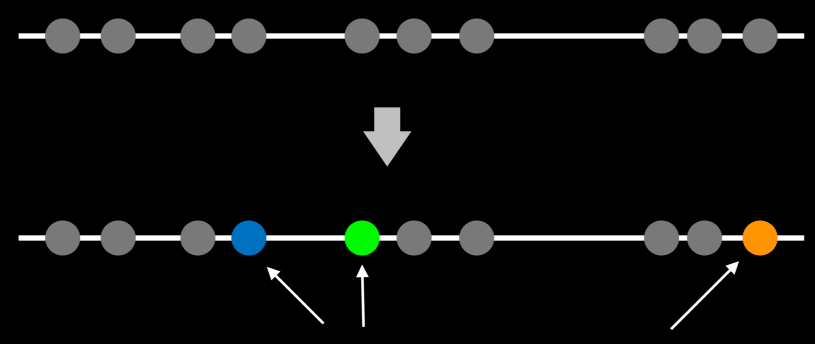
The total variation within clusters

K-means keeps track of these clusters and their total **variance** and then does the whole thing over again with different starting points

Step 10.

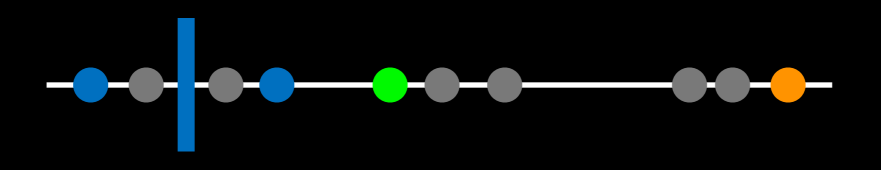
Repeat with different starting points

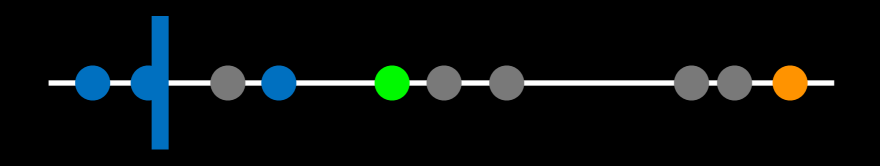
Back to the beginning and do all steps over again...

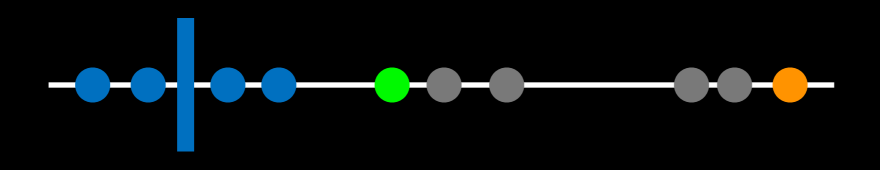


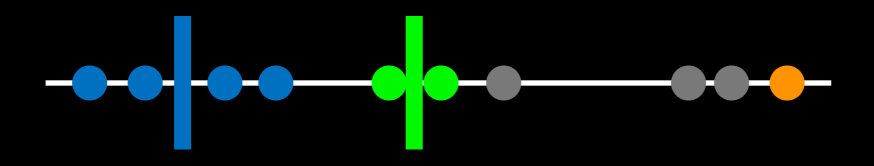
Pick new points as "initial" clusters

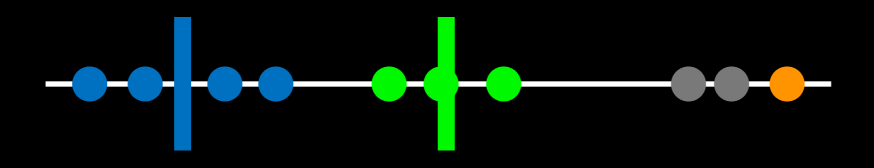


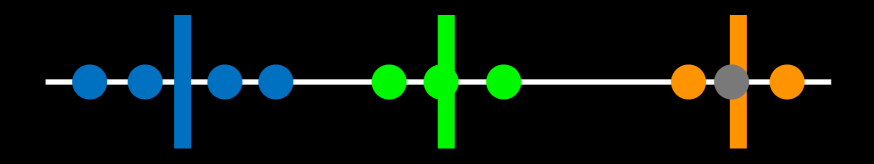


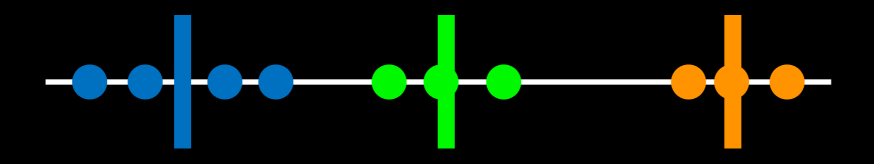




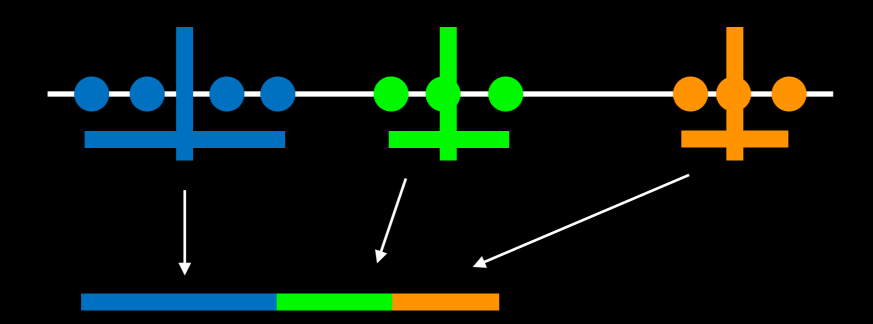








Now the data are all assigned to clusters, we again sum the variation within each cluster



The total variation within clusters

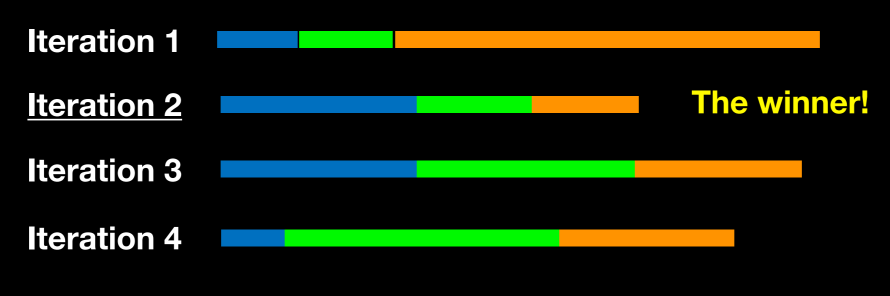
Step 10.

Repeat again with different starting points



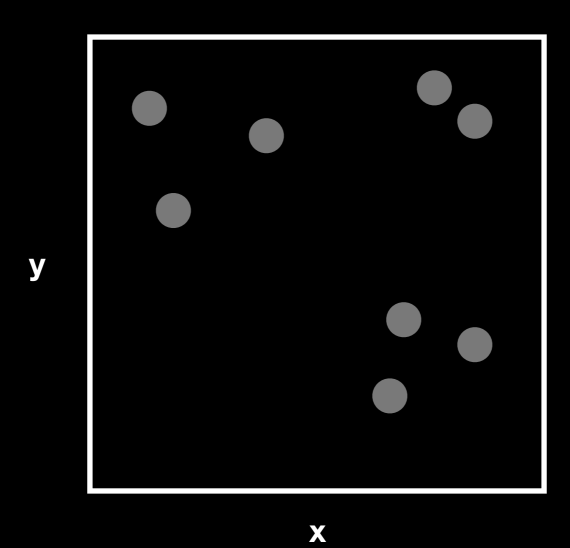
After several iterations k-means clustering knows it has the best clustering so-far based on the smallest total variation with clusters.

However, it does not know if it has found the <u>best overall</u>. So it will perform several more iterations with different starting points...

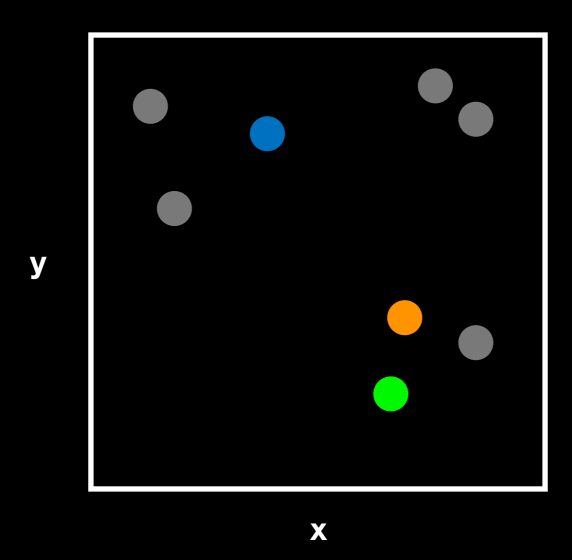


The total variation within clusters

What if we have more dimensions?

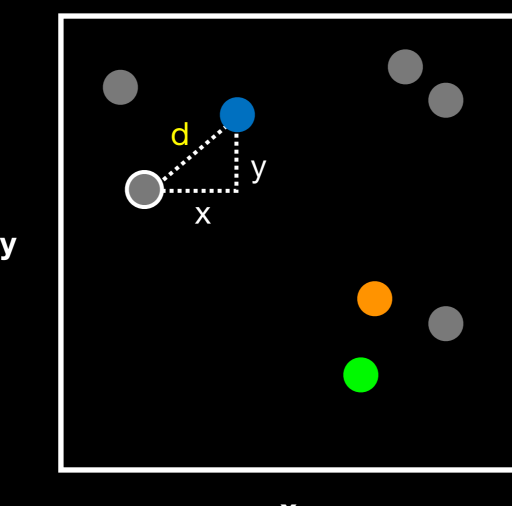


Just like before, we pick 3 random points...

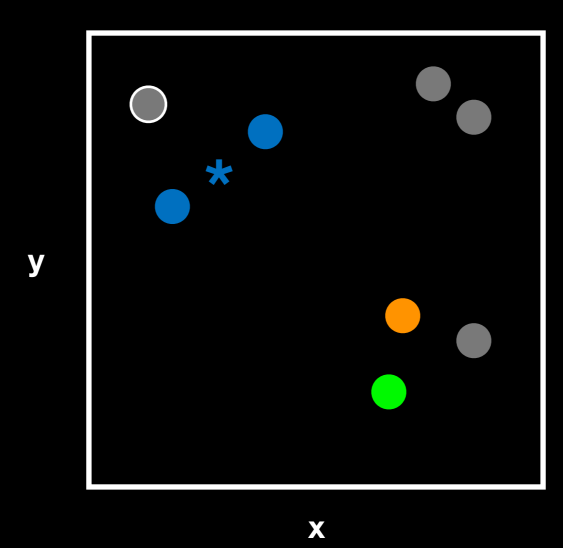


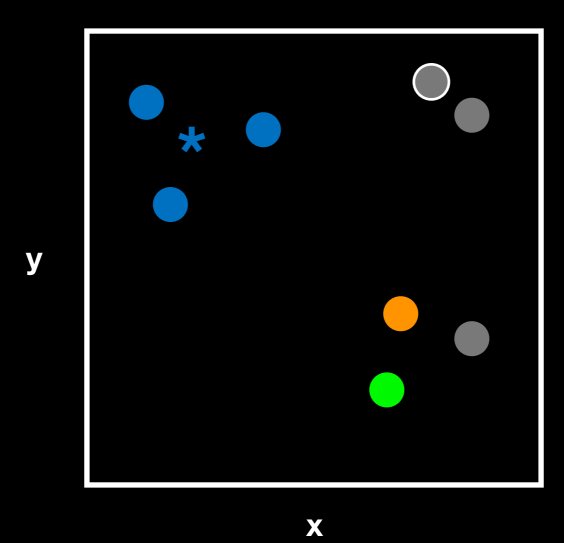
...and use the Euclidean distance.

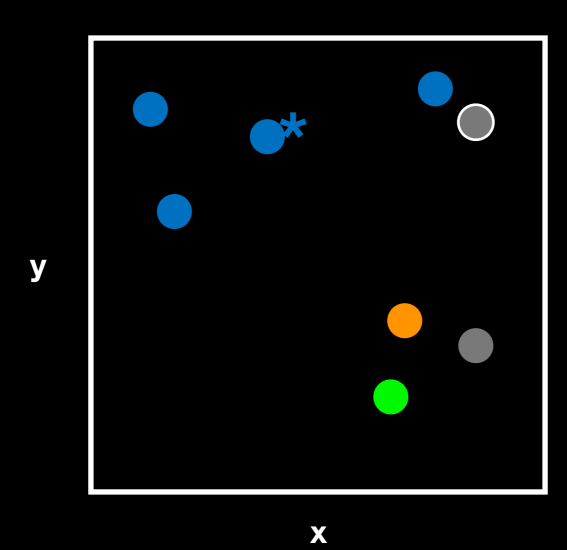
In 2 dimensions the Euclidean distance is the same as the Pythagorean theorem $d = sqrt(x^2 + y^2)$



...assign point to nearest cluster and update cluster center *

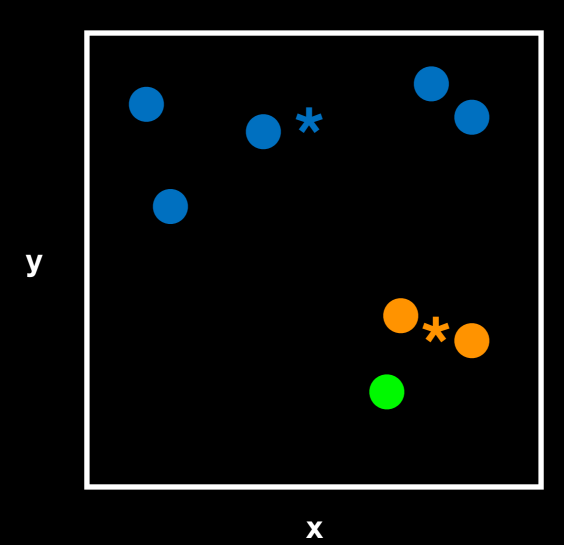




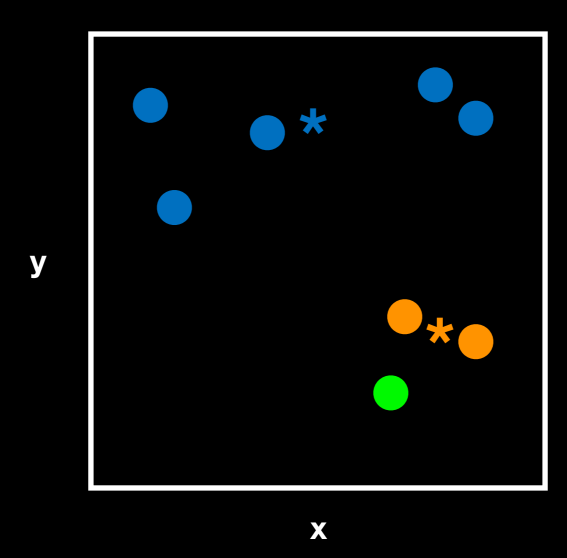




X



Again we have to use a number of different starting conditions before deciding on a good clustering!



What if we have even more dimensions?

Cell Samples

	#1	#2	#3
Gene 1	12	6	-13
Gene 2	-7	13	10
Gene 3	8	6	-9
Gene 4	9	5	-11
Gene 5	-3	1	6
Gene 6	10	4	-8

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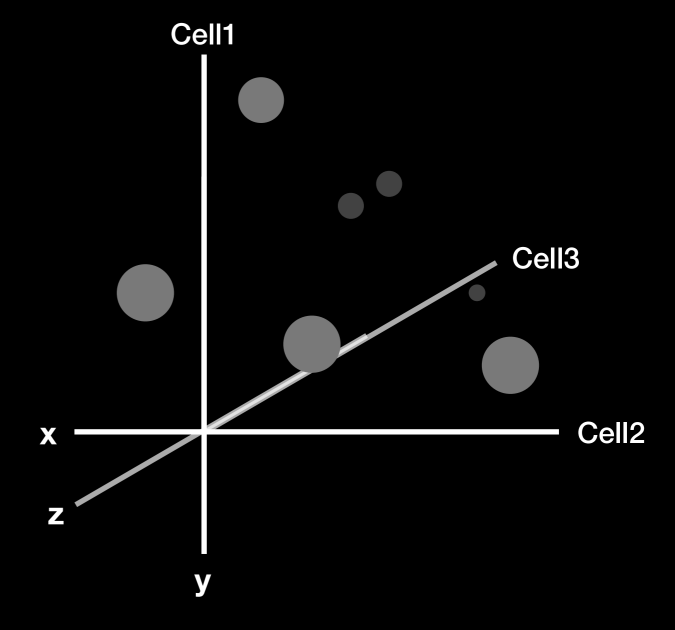
We could simply plot them by relabeling the cell samples as x, y, and z (i.e. a 3D plot)

x y z

What if we have even more dimensions?

Cell Samples

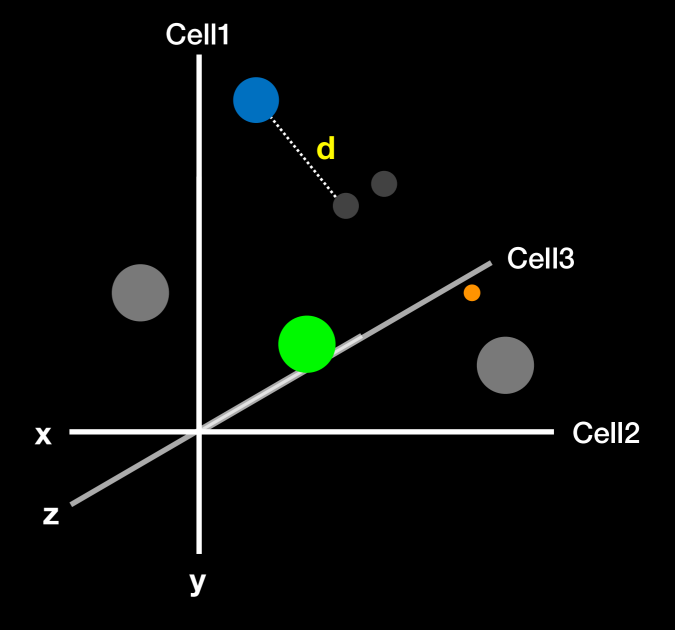
	#1	#2	#3
Gene 1	12	6	-13
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...and go through exactly the same procedure with initial cluster assignment followed by distance calculation etc...

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$$d = sqrt(x^2 + y^2 + z^2)$$

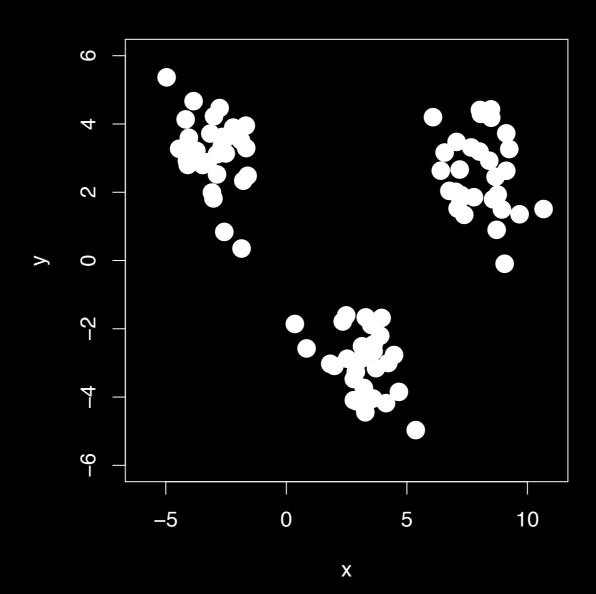
Of course we don't actually need to plot anything.

We can just calculate the Euclidean distance along any number of dimensions and perform our k-means clustering in the same way.

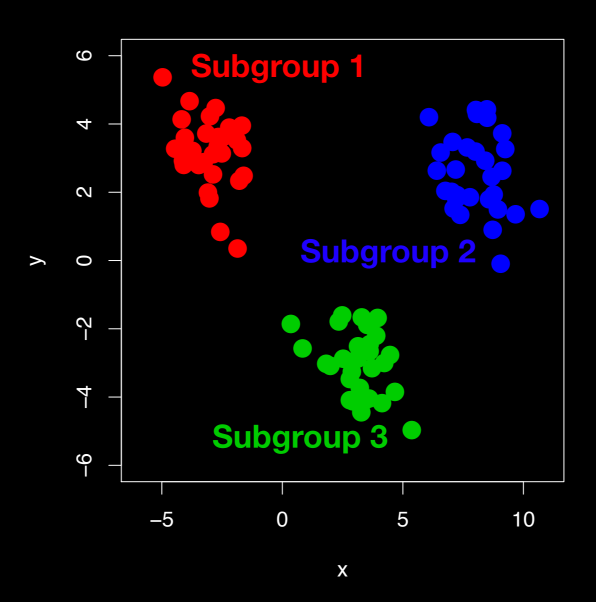
k-means in R

```
# k-means algorithm with 3 centers, run 20 times
kmeans(x, centers= 3, nstart= 20)
```

- Input x is a numeric matrix, or data.frame, with one observation per row, one feature per column
- k-means has a random component
- Run algorithm multiple times to improve odds of the best model



3 Groups



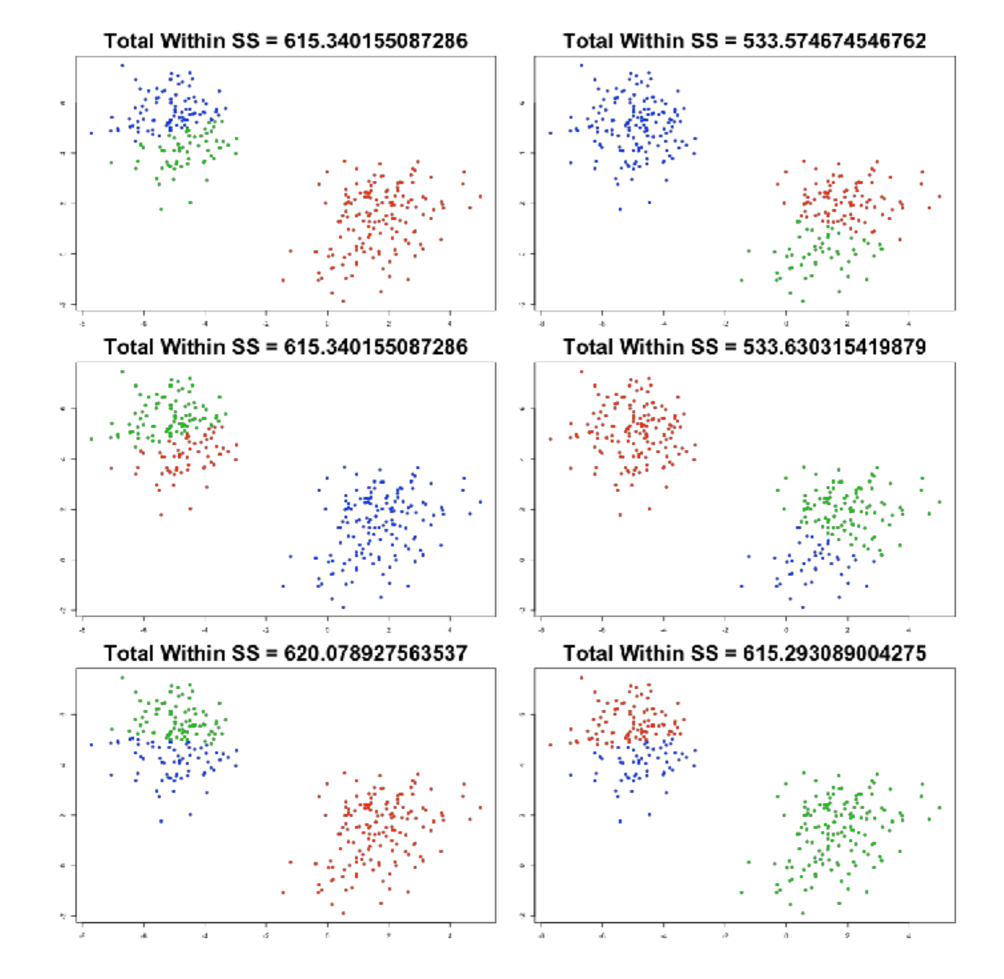
Model selection

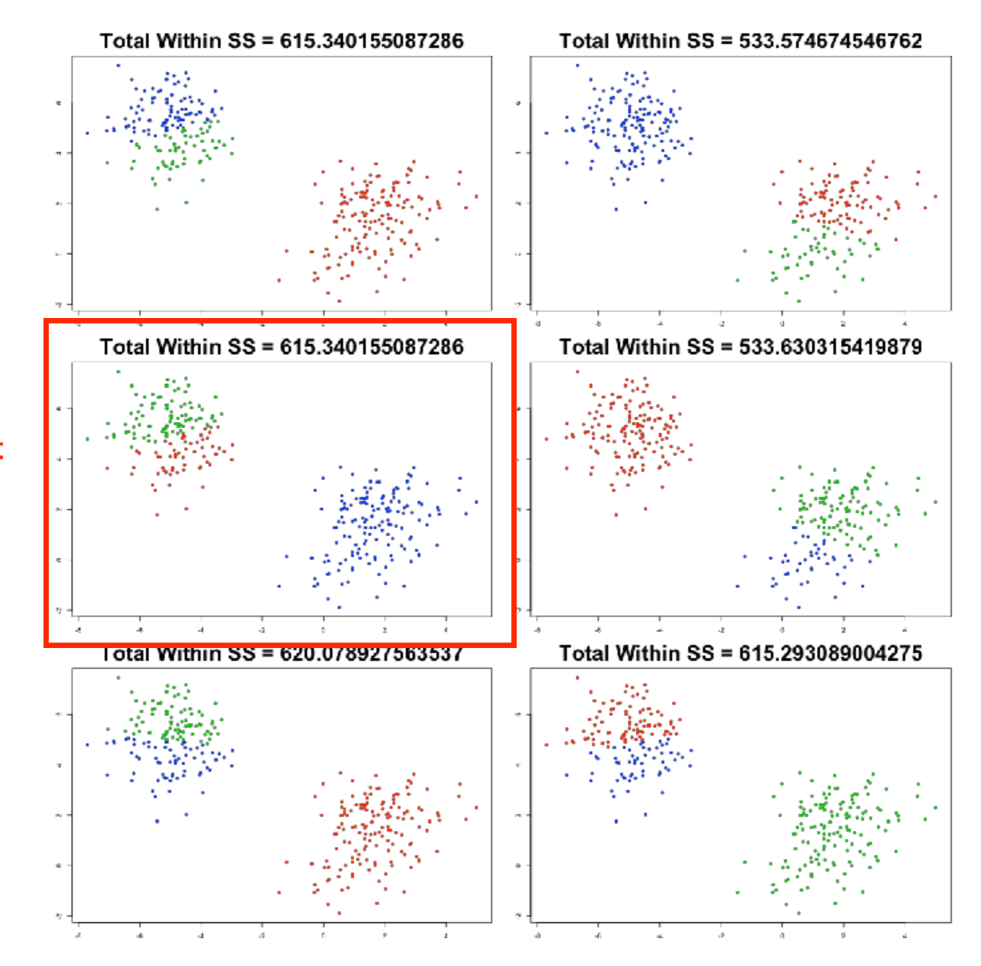
- Recall k-means has a random component
- Best outcome is based on total within cluster sum of squares:
 - For each cluster
 - For each observation in the cluster
 - Determine squared distance from observation to cluster center
 - Sum all of them together

Model selection

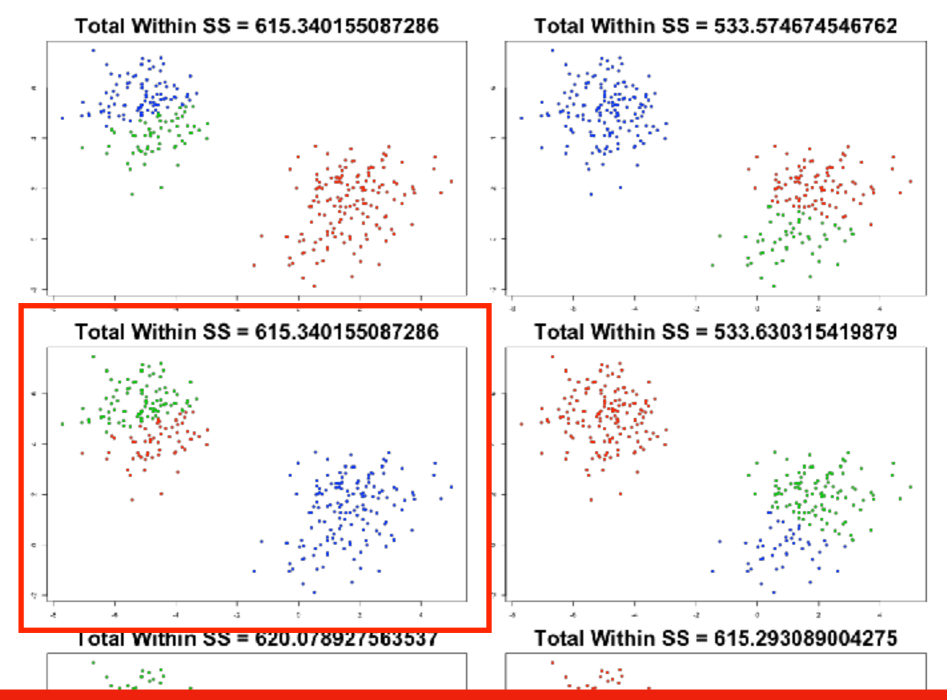
```
# k-means algorithm with 5 centers, run 20 times
kmeans(x, centers=5, nstart=20)
```

- Running algorithm multiple times (i.e. setting nstart) helps find the global minimum total within cluster sum of squares
- Increasing the default value of nstart is often sensible

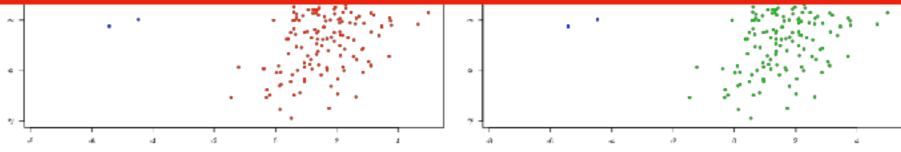




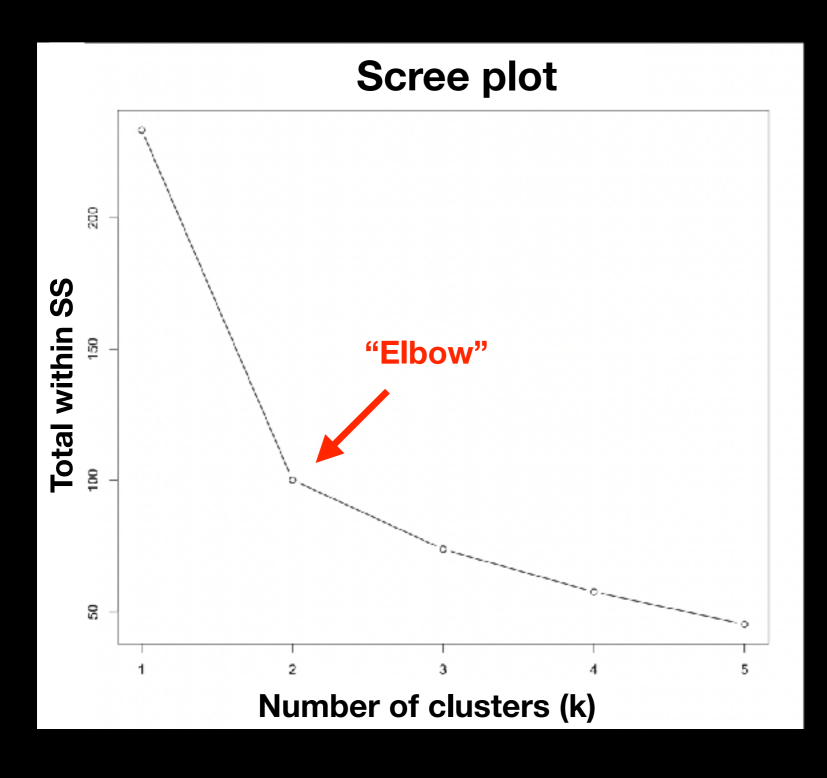
Winner has the smallest within cluster SS



Note. k-means will always give you the renumber of clusters you request! (Here for example, k=2 may be better but we asked for k=3)



Determining number of clusters



Trial and error is not the best approach

Systematically try a range of different k values and plot a "scree plot".

Here there is a large reduction in SS with **k=2** but after that the values do not go down as quickly!

Your Turn!

```
tmp <- c(rnorm(30,-3), rnorm(30,3))
x <- cbind(x=tmp, y=rev(tmp))
plot(x)
```

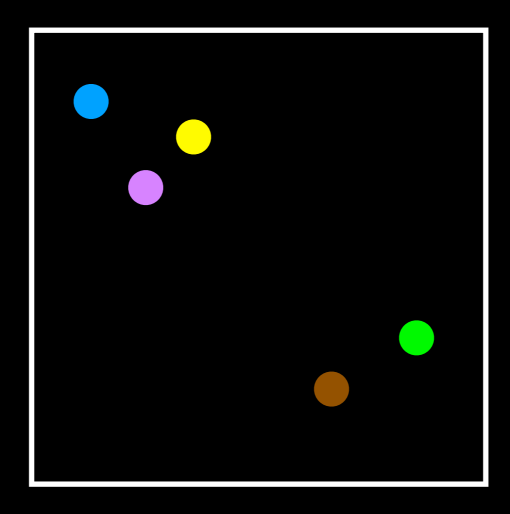
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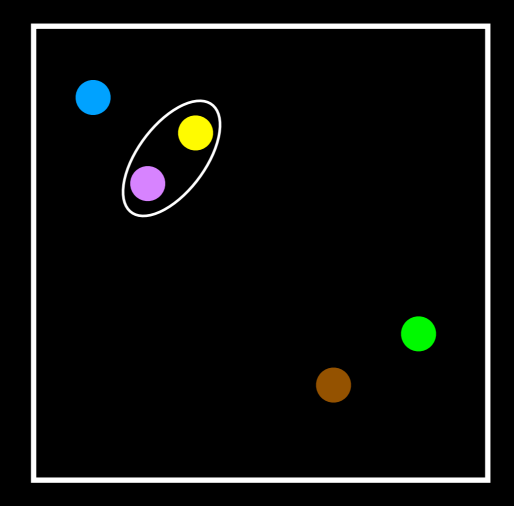
- Number of clusters is not known ahead of time
- Two kinds of hierarchical clustering:
 - bottom-up
 - top-down

Simple example:

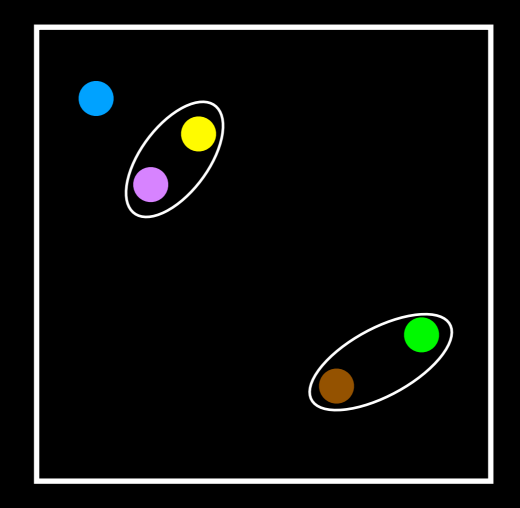
5 clusters: Each point starts as it's own "cluster"!



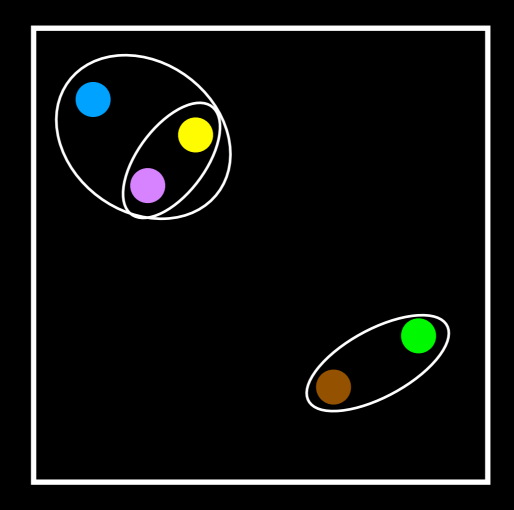
4 clusters



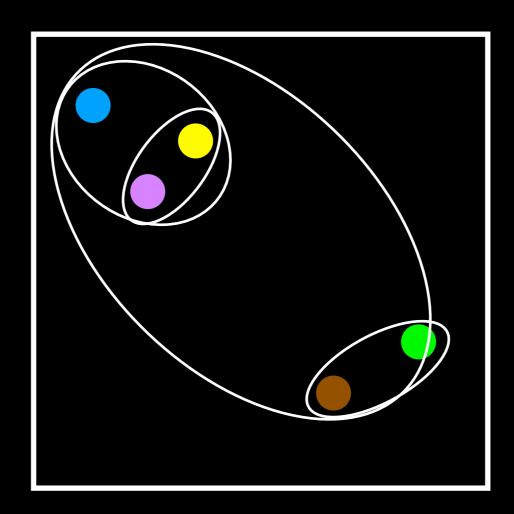
3 clusters



2 clusters



End: 1 cluster



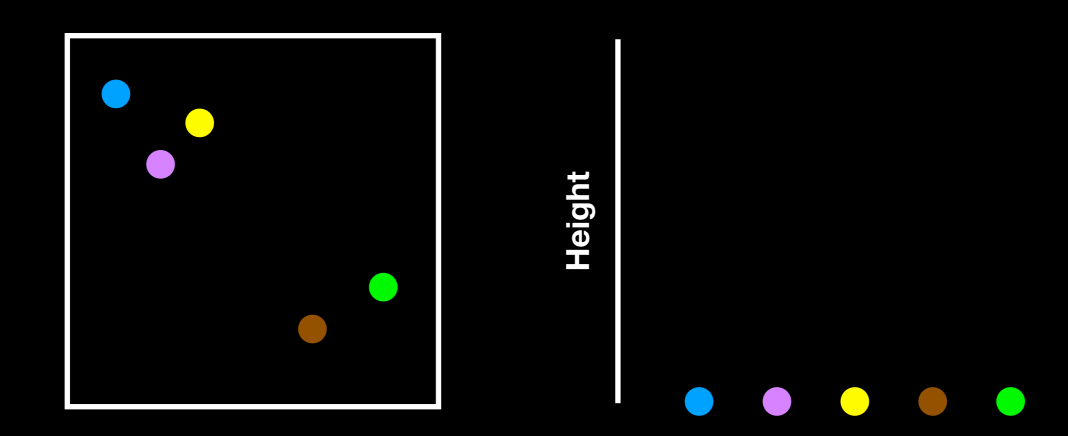
```
dist matrix <- dist(x)</pre>
hc <- hclust(d = dist matrix)</pre>
hc
Call:
hclust(d = dist matrix)
Cluster method : complete
Distance : euclidean
Number of objects: 60
```

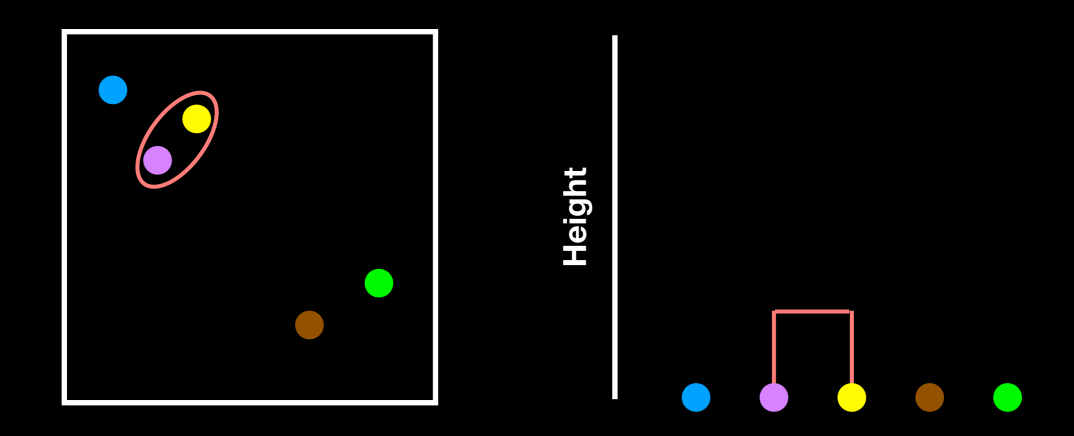
Lets have a closer look...

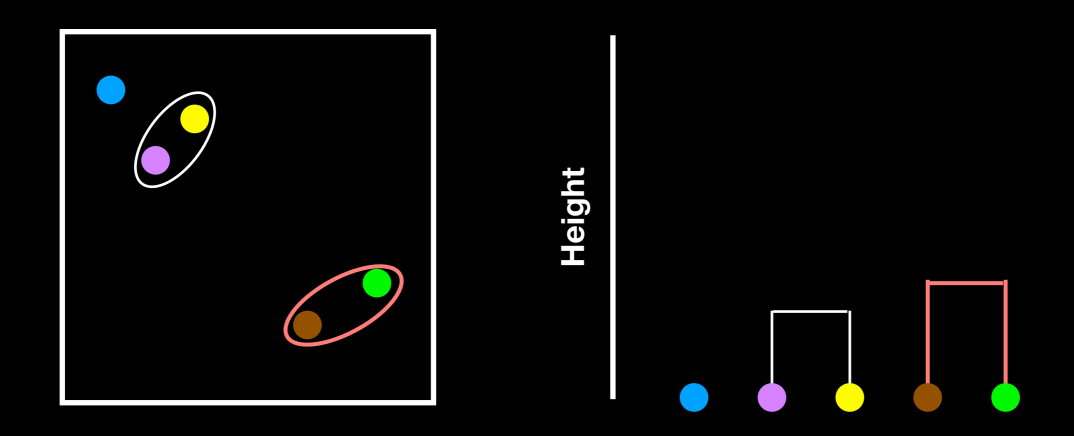
```
dist matrix <- dist(x)</pre>
dim(dist matrix)
NULL
View( as.matrix(dist matrix) )
dim(x)
[1] 60 2
dim( as.matrix(dist_matrix) )
[1] 60 60
```

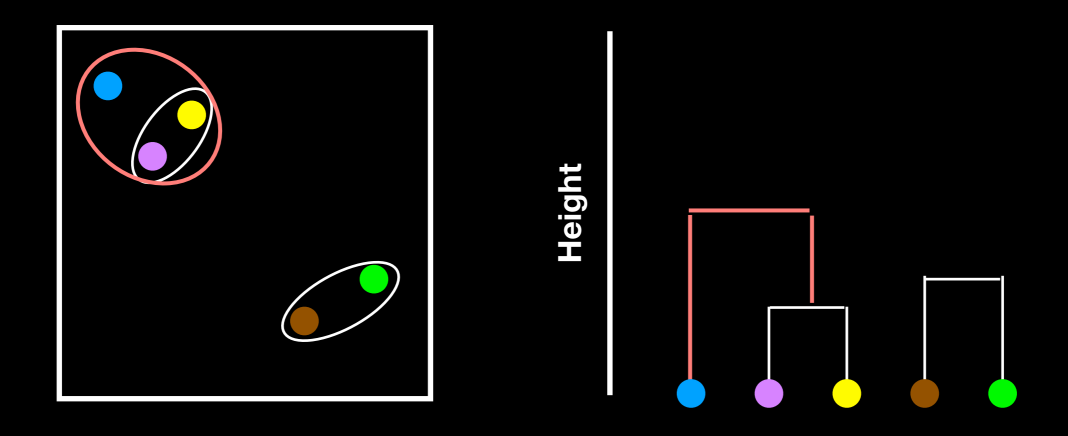
Interpreting results

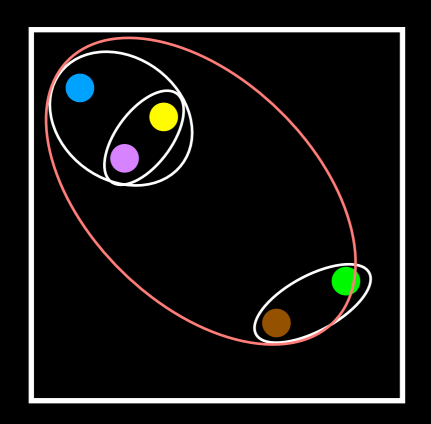
```
hc <- hclust(dist(x))</pre>
plot(hc)
```

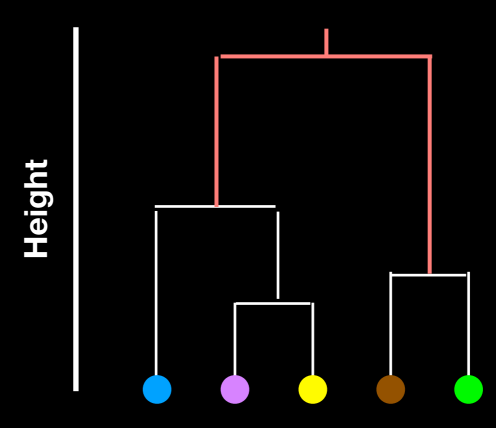


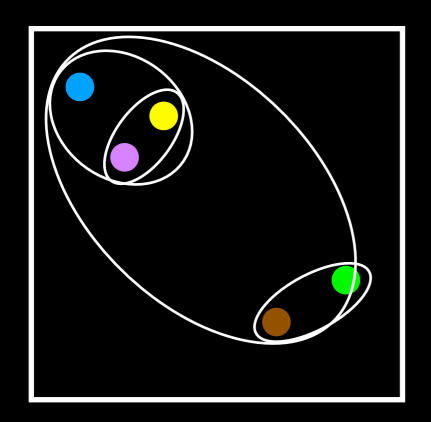


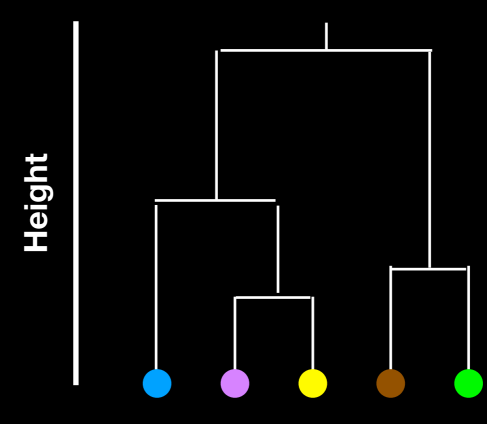




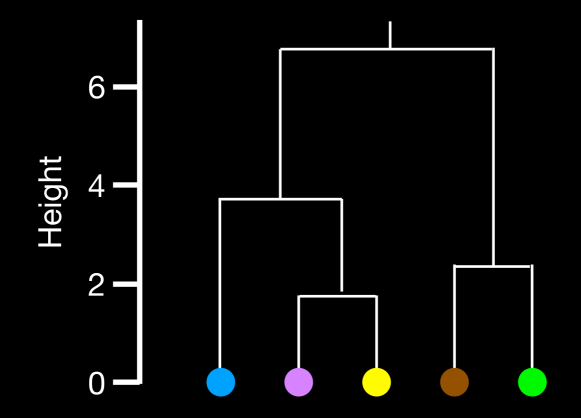




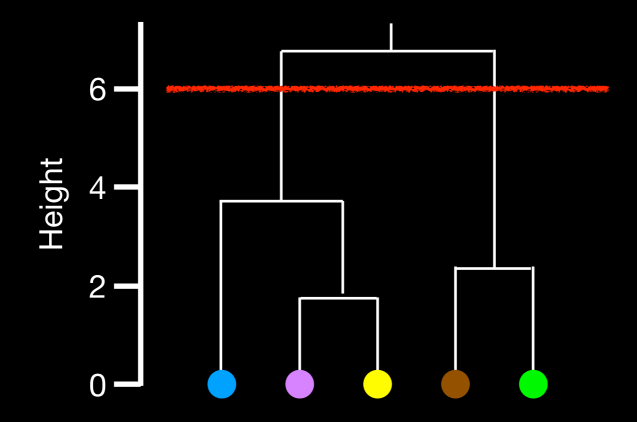




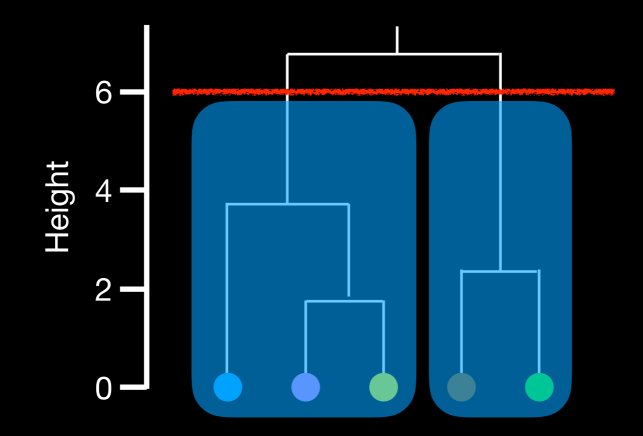
```
# Draws a dendrogram
plot(hc)
```



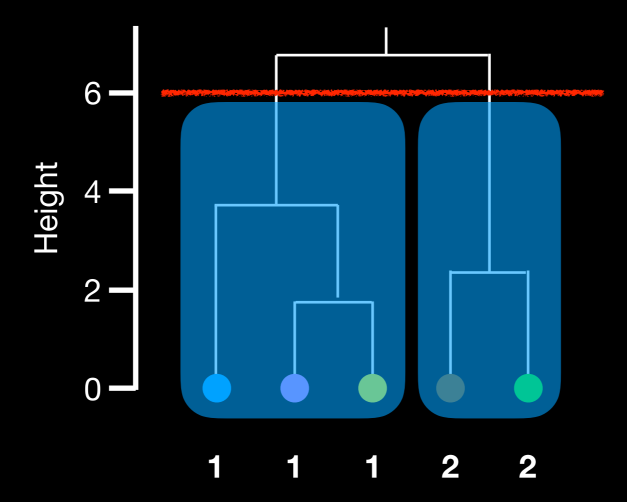
```
# Draws a dendrogram
plot(hc)
abline(h=6, col="red")
```



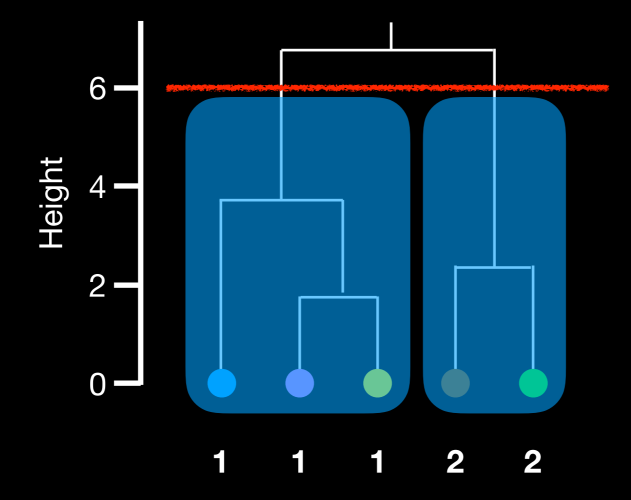
```
# Draws a dendrogram
plot(hc)
abline(h=6, col="red")
```



```
# Draws a dendrogram
plot(hc)
abline(h=6, col="red")
cutree(hclust.out, h=6) # Cut by height h
[1] 1,1,1,2,2
```



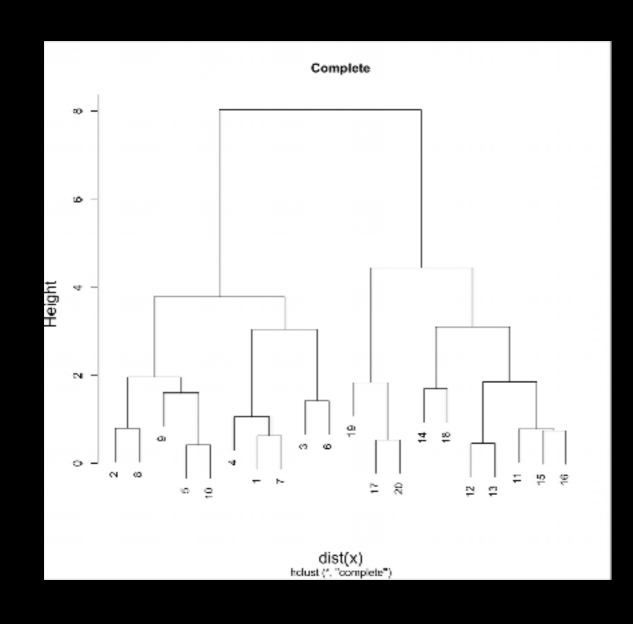
```
# Draws a dendrogram
plot(hclust.out)
abline(h=6, col="red")
cutree(hclust.out, k=2)  # Cut into k grps
[1] 1,1,1,2,2
```

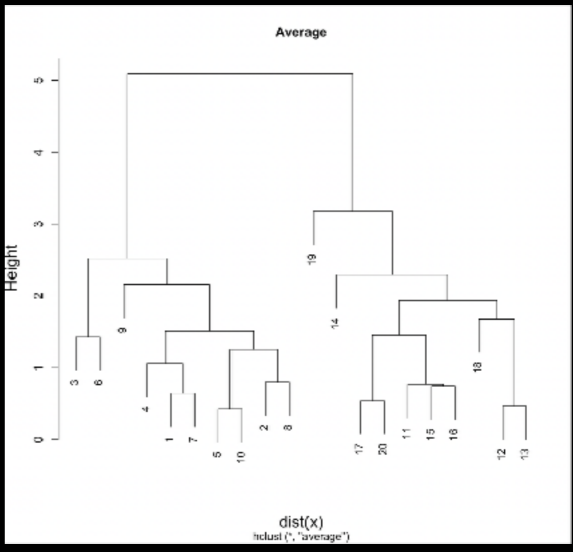


Linking clusters in hierarchical clustering

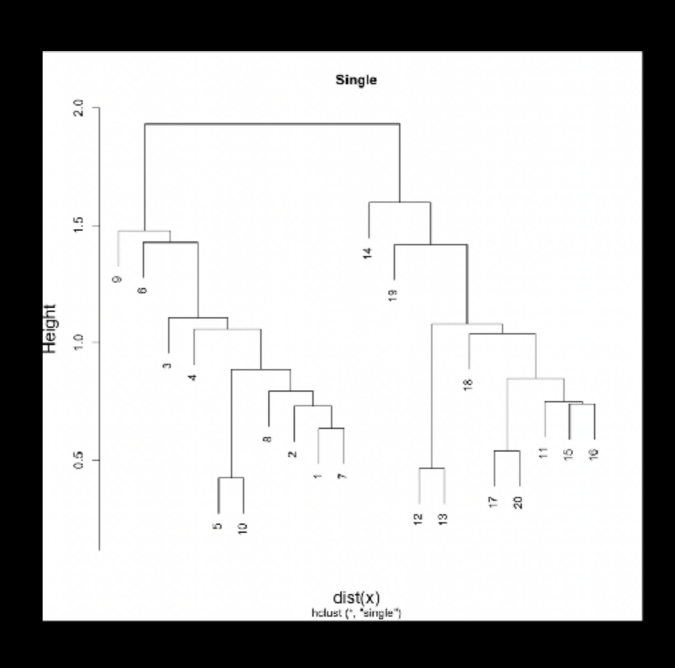
- How is distance between clusters determined?
- There are four main methods to determine which cluster should be linked:
 - Complete: pairwise similarity between all observations in cluster 1 and cluster 2, and uses largest of similarities
 - Single: same as above but uses smallest of similarities
 - Average: same as above but uses average of similarities
 - → Centroid: finds centroid of cluster 1 and centroid of cluster 2, and uses similarity between two centroids

Linking methods: complete and average

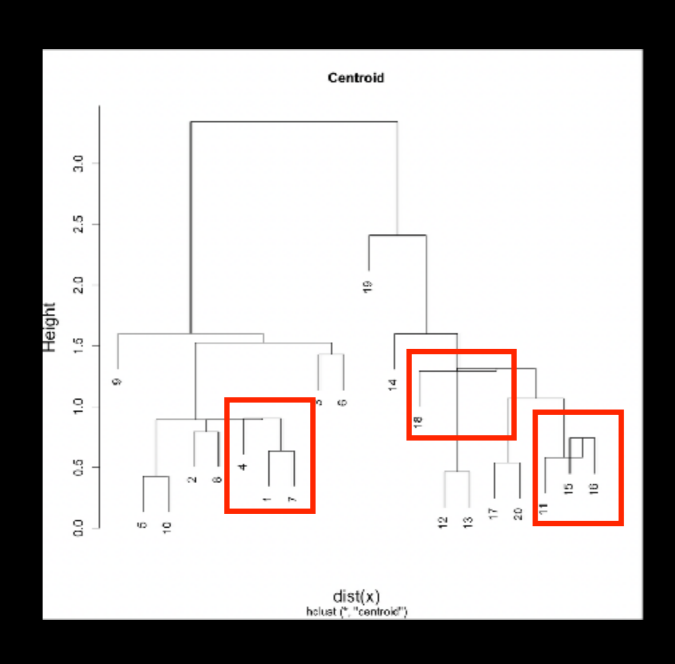




Linking method: single



Linking method: centroid



Linkage in R

```
# Using different hierarchical clustering methods
hc.complete <- hclust(d, method="complete")
hc.average <- hclust(d, method="average")
hc.single <- hclust(d, method="single")</pre>
```

Side-Note: Practical matters

- Data on different scales can cause undesirable results in clustering methods
- Solution is to scale data so that features have same mean and standard deviation
 - Subtract mean of a feature from all observations
 - Divide each feature by the standard deviation of the feature
 - Normalized features have a mean of zero and a standard deviation of one

Side-Note: Practical matters

```
colMeans(x) # x is a data matrix
[1] -0.1337828 0.0594019
apply(x, 2, sd)
[1] 1.974376 2.112357
scaled x <- scale(x)</pre>
colMeans(scaled x)
[1] 2.775558e-17 3.330669e-17
apply(scaled x, 2, sd)
```

Your Turn!

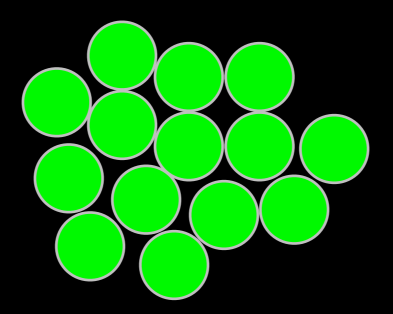
```
x <- rbind(
  matrix(rnorm(100, mean=0, sd = 0.3), ncol = 2), # c1
  matrix(rnorm(100, mean = 1, sd = 0.3), ncol = 2), # c2
  matrix(c(rnorm(50, mean = 1, sd = 0.3),
                                                     # c3
           rnorm(50, mean = 0, sd = 0.3)), ncol = 2))
colnames(x) <- c("x", "y")
plot(x)
col <- as.factor( rep(c("c1", "c2", "c3"), each=50) )
plot(x, col=col)
```

Today's Menu

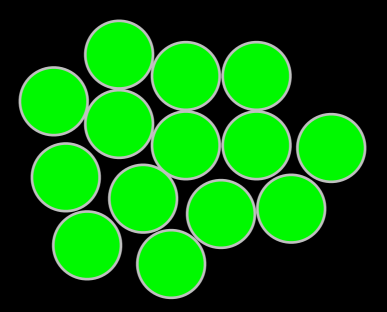
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PCA: The absolute basics

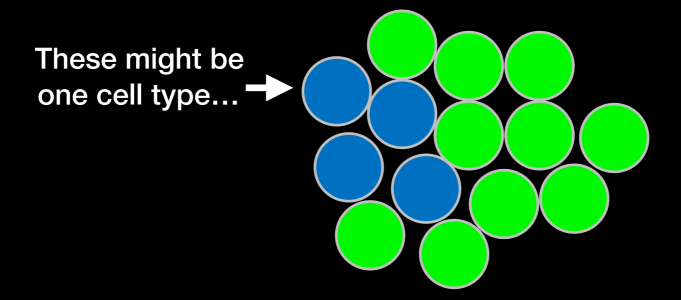
Bunch of "normal" cells

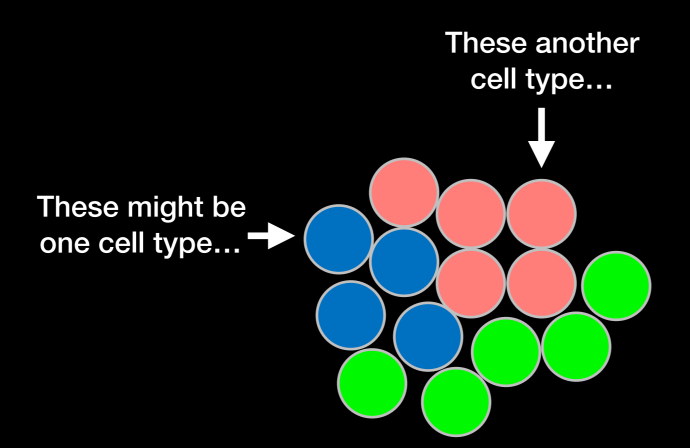


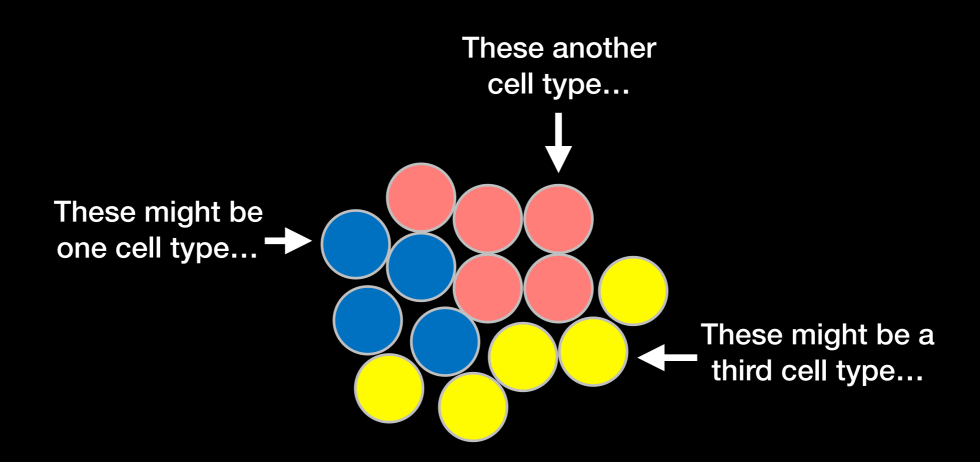
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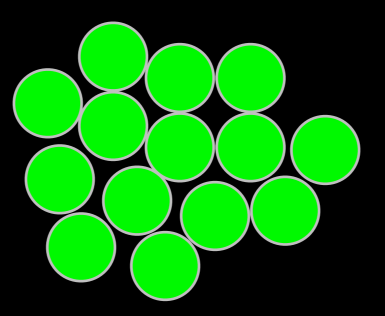


Even though they look the same we suspect that there are differences...

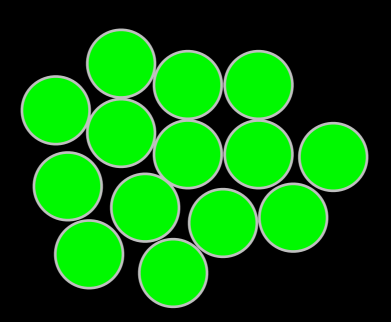








Unfortunately we can't observe the differences visually



Unfortunately we can't observe the differences visually

So we sequence the mRNA in each cell to identify which genes are active and at what levels.

Here is the data...

	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

Each column shows how much each gene is transcribed in each cell

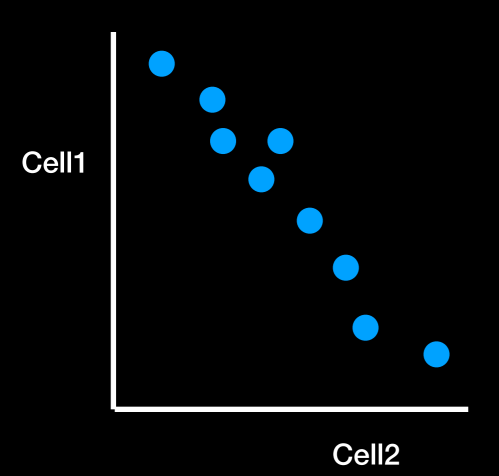
Here is the data...

	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

For now lets consider only two cells

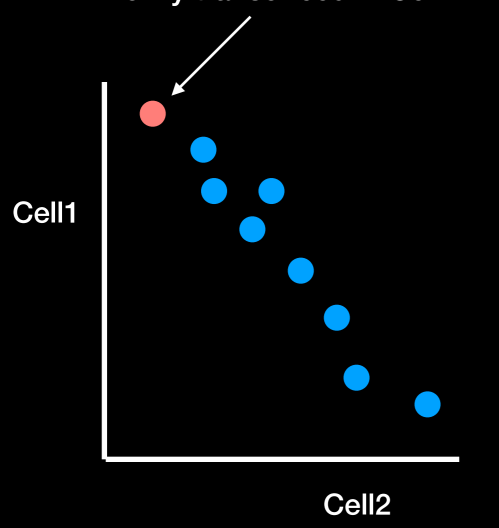
	Cell1	Cell2
Gene1	3	0.25
Gene2	2.9	8.0
Gene3	2.2	1
Gene4	2	1.4
Gene5	1.3	1.6
Gene6	1.5	2
Gene7	1.1	2.2
Gene8	1	2.7
Gene9	0.4	3

We have just 2 cells so we can plot the measurements for each gene



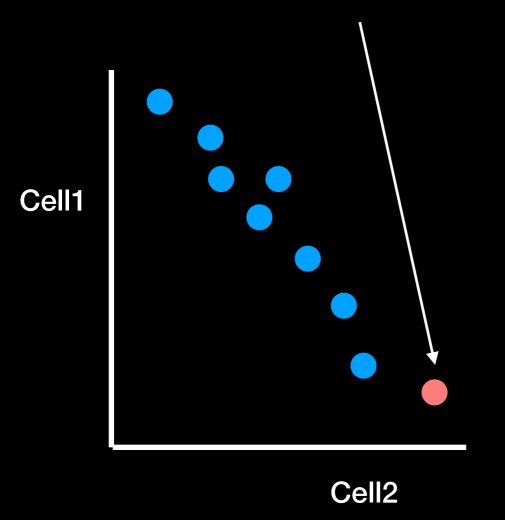
	Cell1	Cell2
Gene1	3	0.25
Gene2	2.9	8.0
Gene3	2.2	1
Gene4	2	1.4
Gene5	1.3	1.6
Gene6	1.5	2
Gene7	1.1	2.2
Gene8	1	2.7
Gene9	0.4	3

This gene (Gene1) is highly transcribed in Cell1 and lowly transcribed in Cell2...



	Cell1	Cell2
Gene1	3	0.25
Gene2	2.9	8.0
Gene3	2.2	1
Gene4	2	1.4
Gene5	1.3	1.6
Gene6	1.5	2
Gene7	1.1	2.2
Gene8	1	2.7
Gene9	0.4	3

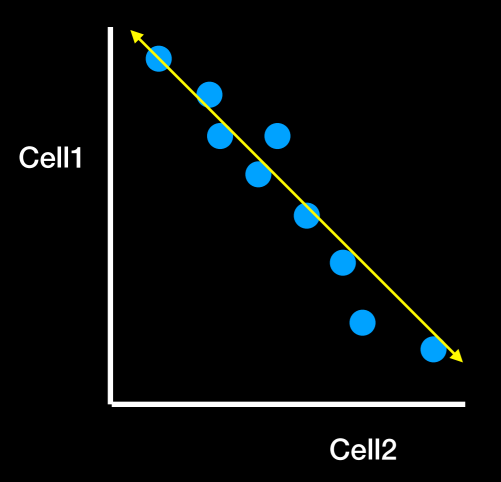
This gene (Gene9) is lowly transcribed in Cell1 and highly transcribed in Cell2...



	Cell1	Cell2
Gene1	3	0.25
Gene2	2.9	8.0
Gene3	2.2	1
Gene4	2	1.4
Gene5	1.3	1.6
Gene6	1.5	2
Gene7	1.1	2.2
Gene8	1	2.7
Gene9	0.4	3

In generel, Cell1 and Cell2 have an inverse correlation.

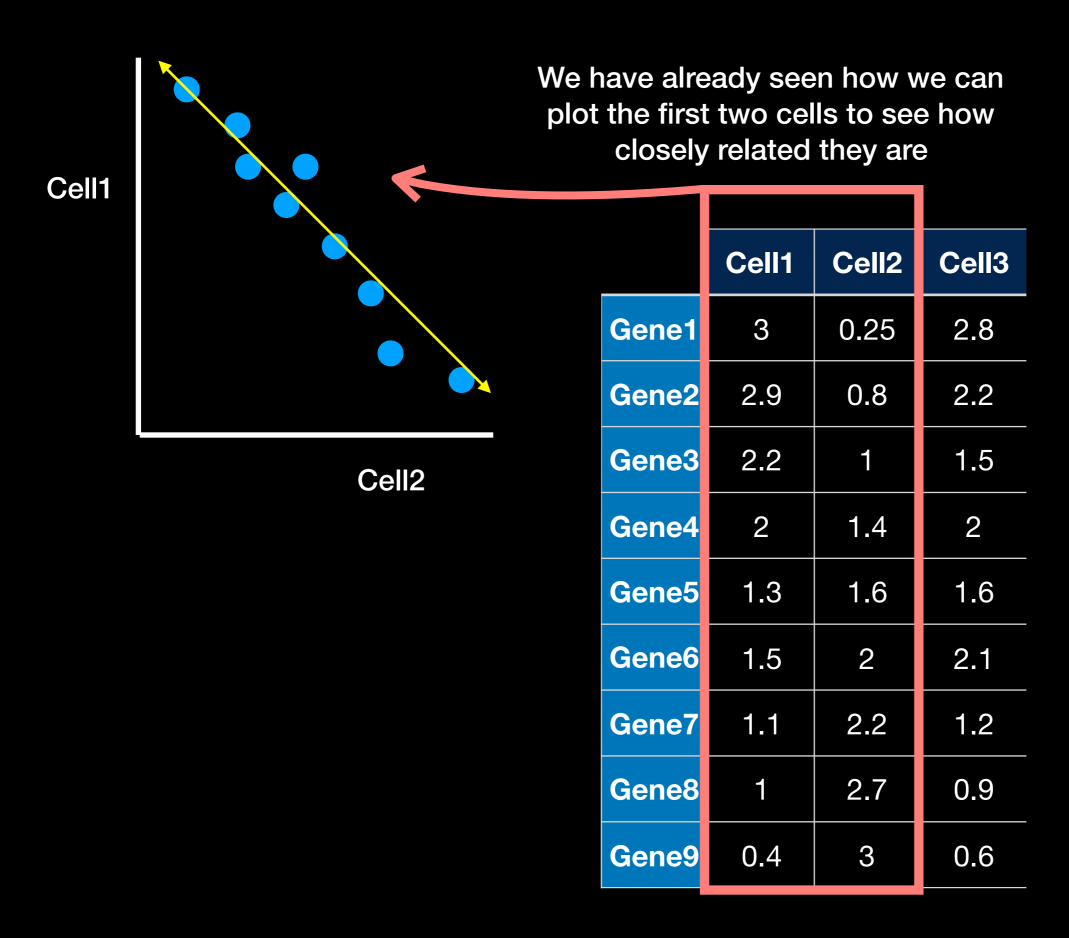
This suggests that they may be two different types of cells as they are using different genes



	Cell1	Cell2
Gene1	3	0.25
Gene2	2.9	8.0
Gene3	2.2	1
Gene4	2	1.4
Gene5	1.3	1.6
Gene6	1.5	2
Gene7	1.1	2.2
Gene8	1	2.7
Gene9	0.4	3

Now lets imagine there are three cells

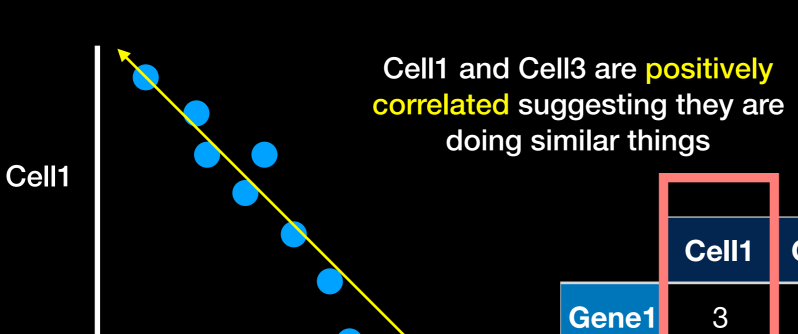
	Cell1	Cell2	Cell3
Gene1	3	0.25	2.8
Gene2	2.9	8.0	2.2
Gene3	2.2	1	1.5
Gene4	2	1.4	2
Gene5	1.3	1.6	1.6
Gene6	1.5	2	2.1
Gene7	1.1	2.2	1.2
Gene8	1	2.7	0.9
Gene9	0.4	3	0.6



Cell1 Cell2 Cell1 Cell3

Now we can also compare Cell1 to Cell3

	Cell1	Cell2	Cell3
Gene1	3	0.25	2.8
Gene2	2.9	8.0	2.2
Gene3	2.2	1	1.5
Gene4	2	1.4	2
Gene5	1.3	1.6	1.6
Gene6	1.5	2	2.1
Gene7	1.1	2.2	1.2
Gene8	1	2.7	0.9
Gene9	0.4	3	0.6

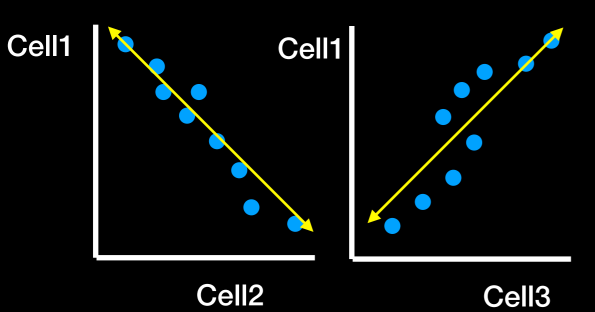


Cell2

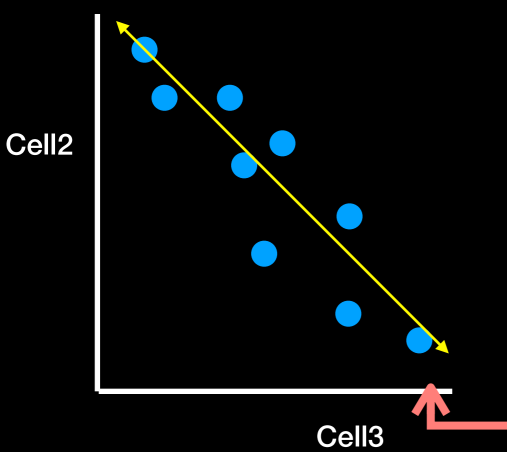
Cell1	

	Cell1	Cell2	Cell3
Gene1	3	0.25	2.8
Gene2	2.9	8.0	2.2
Gene3	2.2	1	1.5
Gene4	2	1.4	2
Gene5	1.3	1.6	1.6
Gene6	1.5	2	2.1
Gene7	1.1	2.2	1.2
Gene8	1	2.7	0.9
Gene9	0.4	3	0.6

Cell3



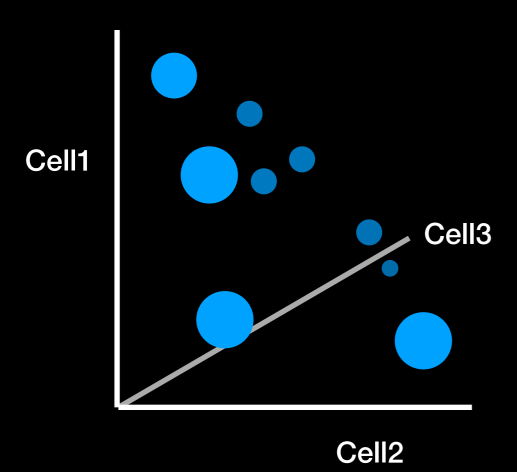
The inverse correlation suggests that Cell2 is doing something different from Cell3



We can also compare Cell2 to Cell3...

	Cell1	Cell2	Cell3
Gene1	3	0.25	2.8
Gene2	2.9	0.8	2.2
Gene3	2.2	1	1.5
Gene4	2	1.4	2
Gene5	1.3	1.6	1.6
Gene6	1.5	2	2.1
Gene7	1.1	2.2	1.2
Gene8	1	2.7	0.9
Gene9	0.4	3	0.6

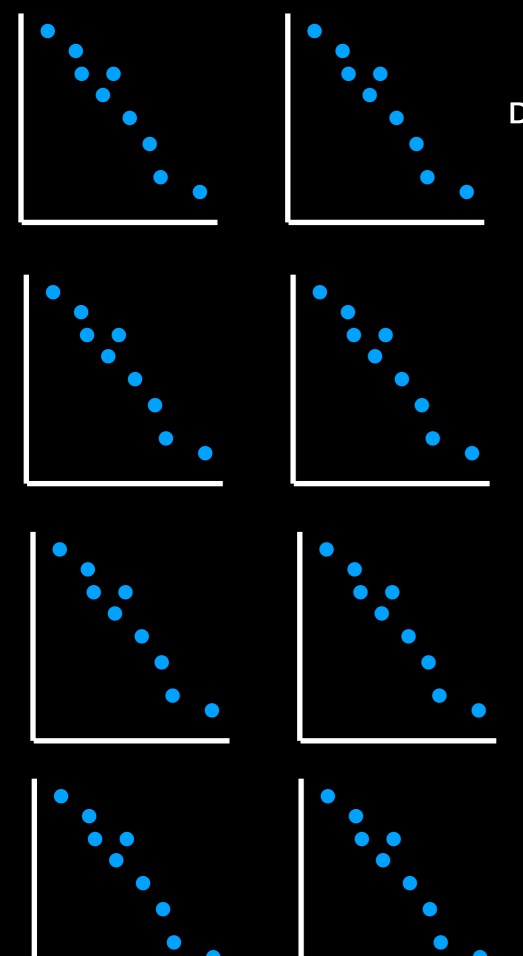
Alternatively, we could try to plot all 3 cells at once on a 3-dimensional graph.



	Cell1	Cell2	Cell3	
Gene1	3	0.25	2.8	
Gene2	2.9	8.0	2.2	
Gene3	2.2	1	1.5	
Gene4	2	1.4	2	
Gene5	1.3	1.6	1.6	
Gene6	1.5	2	2.1	
Gene7	1.1	2.2	1.2	
Gene8	1	2.7	0.9	
Gene9	0.4	3	0.6	

But what if we have 4 or more Cells?

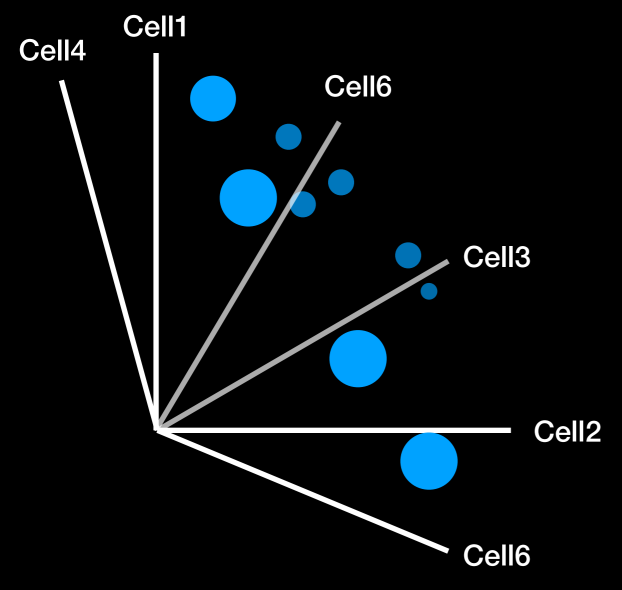
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	



Draw lots of 2 cell plots and try to make sense of them all?

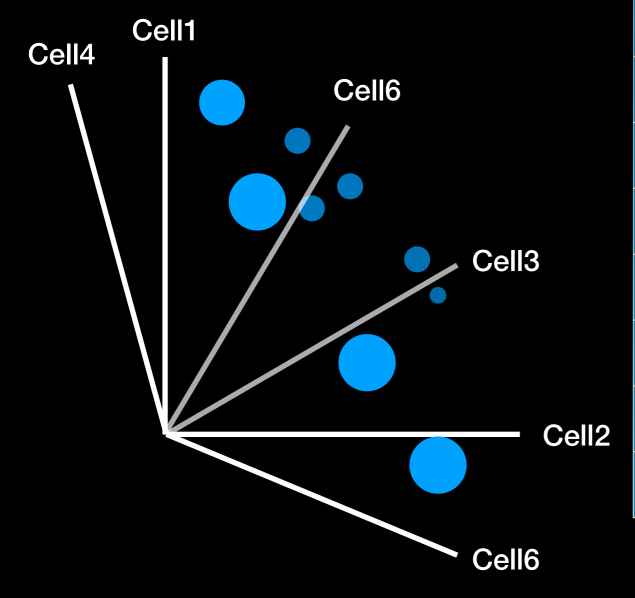
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

Or draw some crazy graph that has an axis for each cell and makes or brains hurt!



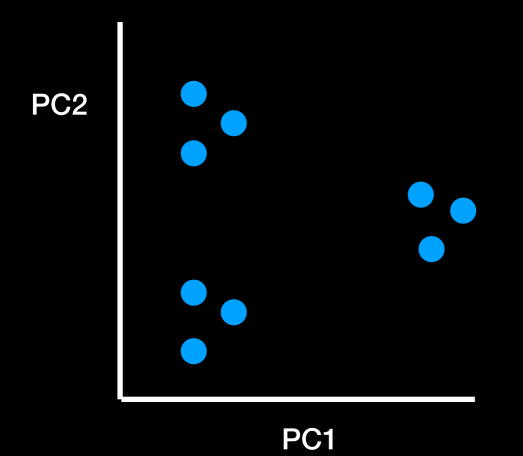
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	





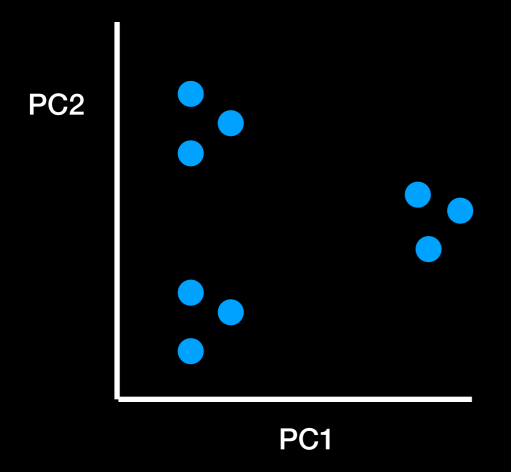
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

Enter Principal Component Analysis (PCA)



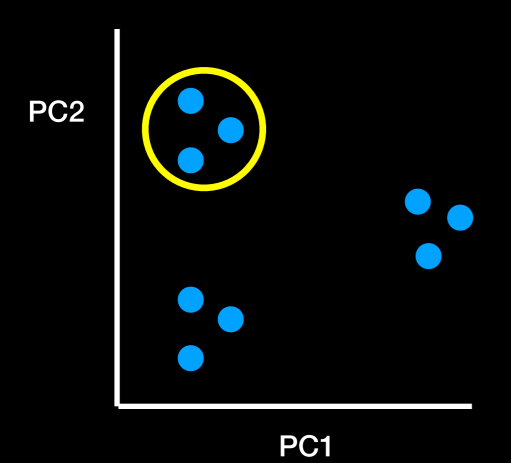
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

PCA converts the correlations (or lack there of) among all cells into a representation we can more readily interpret (e.g. a 2D graph!)



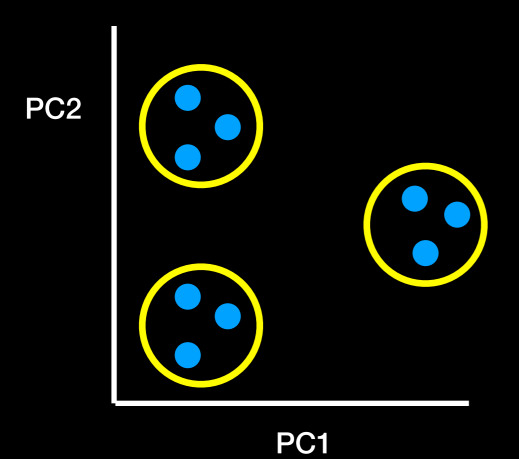
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

Cells that are highly correlated cluster together



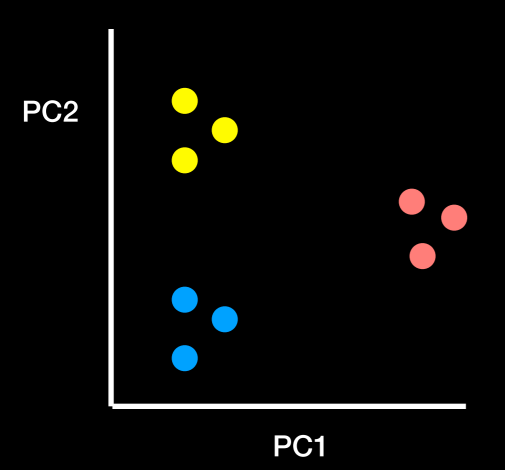
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

Cells that are highly correlated cluster together



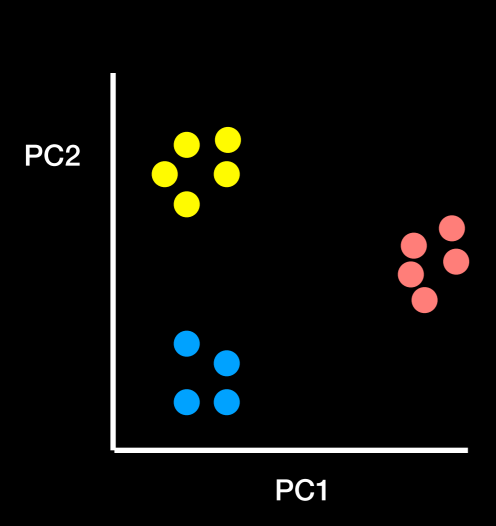
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

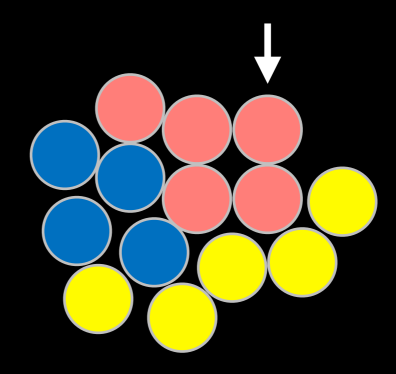
To make the clusters easier to see we can color code them...



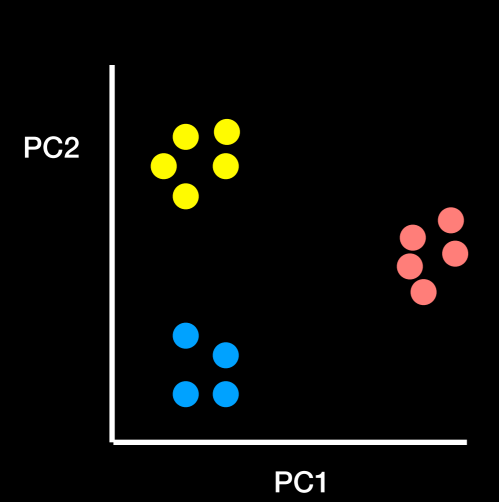
	Cell1	Cell2	Cell3	Cell4	
Gene1	3	0.25	2.8	0.1	
Gene2	2.9	8.0	2.2	1.8	
Gene3	2.2	1	1.5	3.2	
Gene4	2	1.4	2	0.3	
Gene5	1.3	1.6	1.6	0	
Gene6	1.5	2	2.1	3	
Gene7	1.1	2.2	1.2	2.8	
Gene8	1	2.7	0.9	0.3	
Gene9	0.4	3	0.6	0.1	

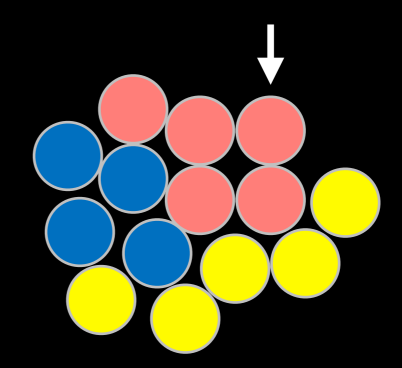
Once we have identified the clusters from our PCA results, we can go back to or original cells...





Once we have identified the clusters from our PCA results, we can go back to or original cells...





...and see they represent three different types of cells doing three different things with their genes!

PC2

PC1

Some key points:

The PCs (i.e. new plot axis) are ranked by their importance

So PC1 is more important than PC2 which in turn is more important than PC3 etc.

PC2

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PC2 (11%) PC1 (44%)

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The PCs (i.e. new plot axis) are ranked by the amount of variance in the original data (i.e. gene expression values) that they "capture"

PC2 (11%) PC1 (44%)

Some key points:

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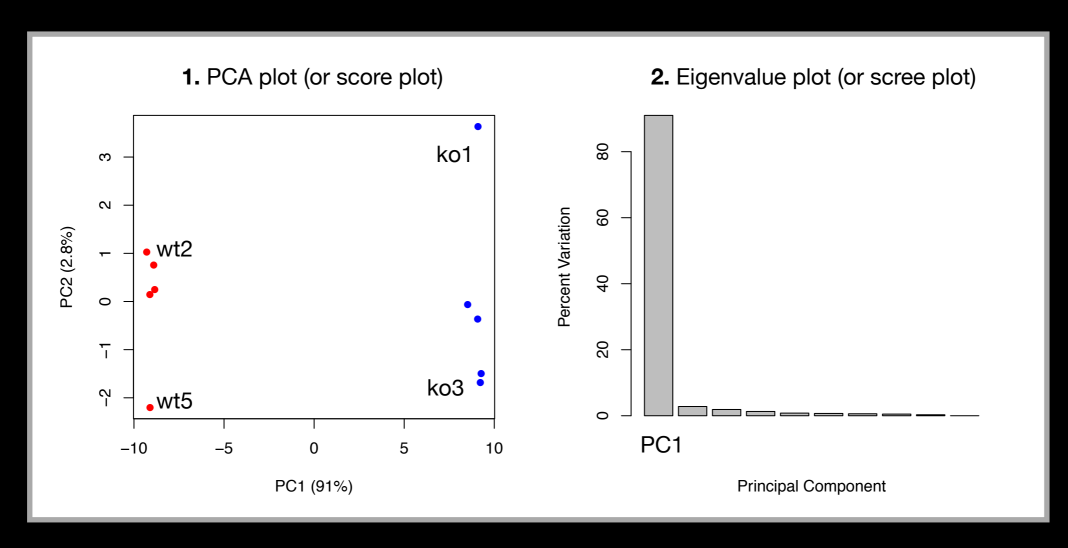
So PC1 is more important than PC2 which in turn is more important than PC3 etc.

So the red and blue cluster are more dissimilar than the yellow and blue clusters

The PCs (i.e. new plot axis) are ranked by the amount of variance in the original data (i.e. gene expression values) that they "capture"

In this example PC1 'captures' 4x more of the original variance than PC2 (44/11 = 4)

- We actually get two main things out of a typical PCA
 - The new axis (called PCs or Eigenvectors) and
 - Eigenvalues that detail the amount of variance captured by each PC



- Another cool thing we can get out of PCA is a quantitive report on how the original variables contributed to each PC
 - In other words, which were the most important genes that lead to the observed clustering in PC-space
 - These are often called the loadings and we can plot them to see which are the most important genes for the observed separation as well as outputting ranked lists of genes that act to discriminate the samples

gene64 gene39 0.1047968 0.1047629

gene7 gene65 -0.1047629 -0.1047443

Hands-on time!

http://setosa.io/ev/principal-component-analysis/

Outline: How to do PCA in R

- How to use the prcomp() function to do PCA.
- How to draw and interpret PCA plots
- How to determine how much variation each principal component accounts for and the the "intrinsic dimensionality" useful for further analysis
- How to examine the loadings (or loading scores) to determine what variables have the largest effect on the graph and are thus important for discriminating samples.

```
## Initialize a blank 100 row by 10 column matrix
mydata <- matrix(nrow=100, ncol=10)</pre>
```

```
## Initialize a blank 100 row by 10 column matrix
mydata <- matrix(nrow=100, ncol=10)</pre>
## Lets label the rows gene1, gene2 etc. to gene100
rownames (mydata) <- paste("gene", 1:100, sep="")</pre>
```

```
## Initialize a blank 100 row by 10 column matrix
mydata <- matrix(nrow=100, ncol=10)</pre>
## Lets label the rows gene1, gene2 etc. to gene100
rownames (mydata) <- paste("gene", 1:100, sep="")</pre>
## Lets label the first 5 columns wt1, wt2, wt3, wt4 and wt5
     and the last 5 kol, ko2 etc. to ko5 (for "knock-out")
colnames(mydata) <- c( paste("wt", 1:5, sep=""),</pre>
                        paste("ko", 1:5, sep="") )
```

```
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mydata <- matrix(nrow=100, ncol=10)</pre>
## Lets label the rows gene1, gene2 etc. to gene100
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## Lets label the first 5 columns wt1, wt2, wt3, wt4 and wt5
     and the last 5 kol, ko2 etc. to ko5 (for "knock-out")
colnames(mydata) <- c( paste("wt", 1:5, sep=""),</pre>
                        paste("ko", 1:5, sep="") )
## Fill in some fake read counts
for(i in 1:nrow(mydata)) {
  wt.values <- rpois(5, lambda=sample(x=10:1000, size=1))</pre>
  ko.values <- rpois(5, lambda=sample(x=10:1000, size=1))</pre>
  mydata[i,] <- c(wt.values, ko.values)</pre>
head (mydata)
```

NOTE: the samples are columns, and the genes are rows!

```
mydata <- matrix(nrow=100, ncol=10)</pre>
rownames (mydata) <- paste("gene", 1:100, sep="")</pre>
colnames(mydata) <- c( paste("wt", 1:5, sep=""),</pre>
                       paste("ko", 1:5, sep="") )
for(i in 1:nrow(mydata)) {
  wt.values <- rpois(5, lambda=sample(x=10:1000, size=1))</pre>
  ko.values <- rpois(5, lambda=sample(x=10:1000, size=1))</pre>
  mydata[i,] <- c(wt.values, ko.values)</pre>
head (mydata)
            wt1 wt2 wt3 wt4 wt5 ko1 ko2 ko3 ko4 ko5
     genel 147 171 160 175 187
                                    63 57
     gene2 151 134 148 126 134 838 831 894 847 830
     gene3 702 672 653 681 701 593 579 644 596 610
     gene4 319 297 310 296 304 754 807 734 750 774
```

- Now we have our data we call prcomp() to do PCA
 - NOTE: prcomp() expects the samples to be rows and genes to be columns so we need to first transpose the matrix with the t() function!

```
## lets do PCA
pca <- prcomp(t(mydata), scale=TRUE)</pre>
```

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 - NOTE: prcomp() expects the samples to be rows and genes to be columns so we need to first transpose the matrix with the t() function!

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## lets do PCA
pca <- prcomp(t(mydata), scale=TRUE)

## See what is returned by the prcomp() function
attributes(pca)

# $names
#[1] "sdev" "rotation" "center" "scale" "x"

# $class
#[1] "prcomp"</pre>
```

- The returned pca\$x here contains the principal components (PCs) for drawing our first graph.
 - Here we will take the first two columns in pca\$x
 (corresponding to PC1 and PC2) to draw a 2-D plot

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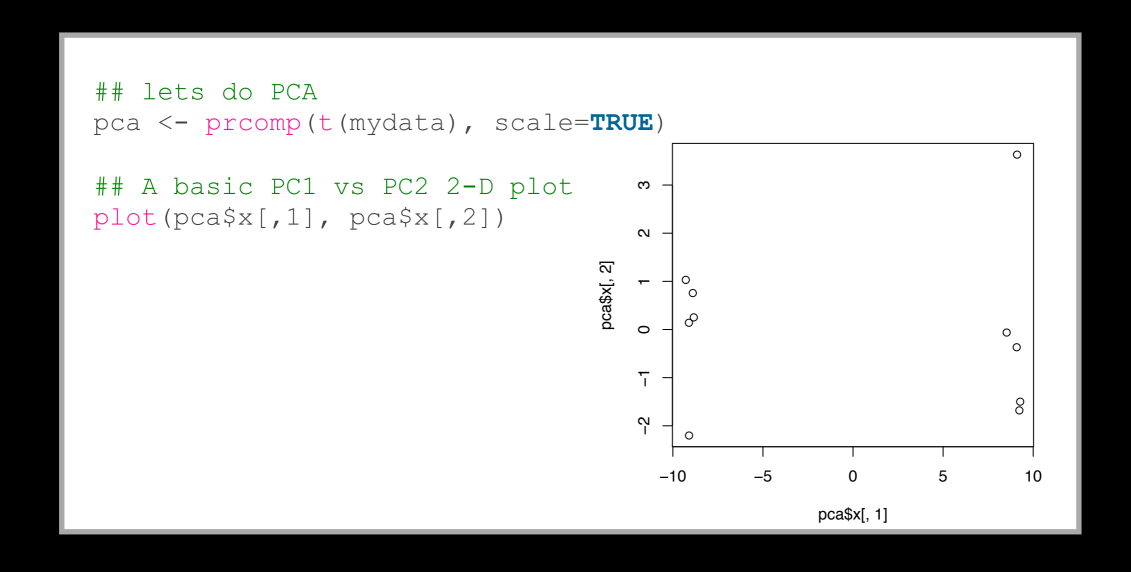
# $names
#[1] "sdev" "rotation" "center" "scale" "x"

# $class
#[1] "prcomp"</pre>
```

- The returned pca\$x here contains the principal components (PCs) for drawing our first graph.
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## A basic PC1 vs PC2 2-D plot
plot (pca$x[,1], pca$x[,2])
```

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- Looks interesting with a nice separation of samples into two groups of 5 samples each
 - Now we can use the square of pca\$sdev, which stands for "standard deviation", to calculate how much variation in the original data each PC accounts for

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## Precent variance is often more informative to look at
pca.var <- pca$sdev^2
pca.var.per <- round(pca.var/sum(pca.var)*100, 1)</pre>
```

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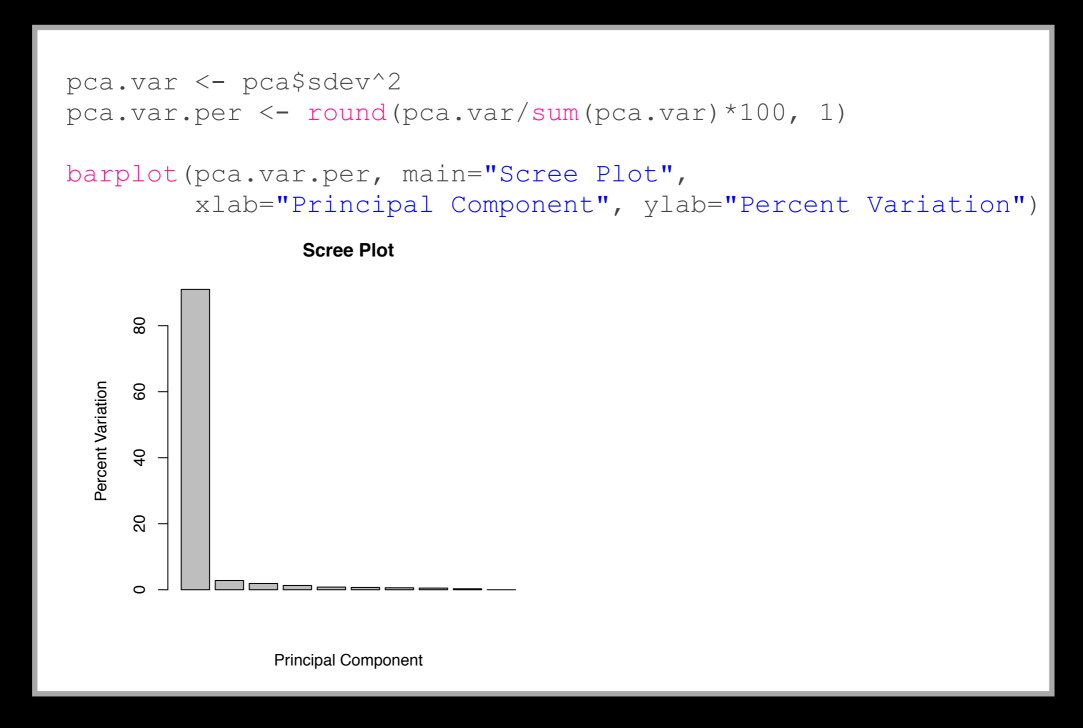
## Precent variance is often more informative to look at
pca.var <- pca$sdev^2
pca.var.per <- round(pca.var/sum(pca.var)*100, 1)

pca.var.per

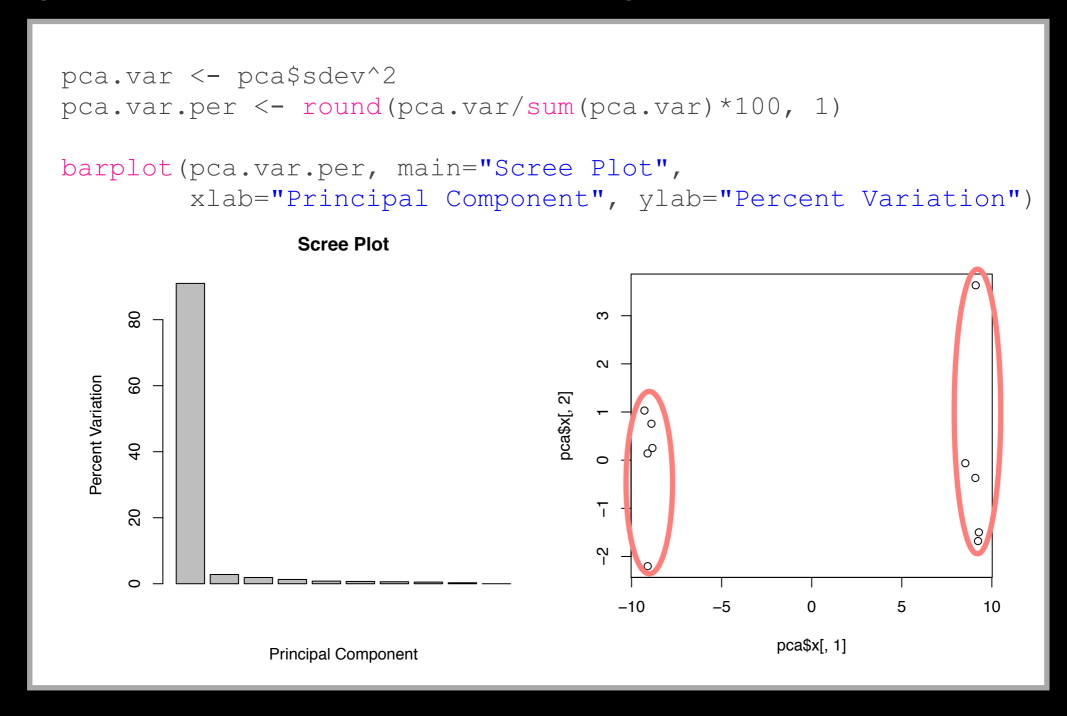
[1] 91.0 2.8 1.9 1.3 0.8 0.7 0.6 0.5 0.3 0.0</pre>
```

- Looks interesting with a nice separation of samples into two groups of 5 samples each
 - Now we can use the square of pca\$sdev, which stands for "standard deviation", to calculate how much variation in the original data each PC accounts for

 From the "scree plot" it is clear that PC1 accounts for almost all of the variation in the data!



 Which means there are big differences between these two groups that are separated along the PC1 axis...



Lets make our plot a bit more useful...

```
## A vector of colors for wt and ko samples
colvec <- colnames (mydata)</pre>
colvec[grep("wt", colvec)] <- "red"</pre>
colvec[grep("ko", colvec)] <- "blue"</pre>
plot(pca$x[,1], pca$x[,2], col=colvec, pch=16,
      xlab=paste0("PC1 (", pca.var.per[1], "%)"),
      ylab=paste0("PC2 (", pca.var.per[2], "%)"))
                                       က
                                       \alpha
                                    PC2 (2.8%)
                                       0
                                       T
                                       7
                                                              5
                                                -5
                                         -10
                                                       0
                                                                    10
                                                    PC1 (91%)
```

And add some labels...

```
plot(pca$x[,1], pca$x[,2], col=colvec, pch=16,
     xlab=paste0("PC1 (", pca.var.per[1], "%)"),
     ylab=paste0("PC2 (", pca.var.per[2], "%)"))
## Click to identify which sample is which
identify(pca$x[,1], pca$x[,2], labels=colnames(mydata))
## Press ESC to exit...
                                                              ko1
                                     \alpha
                                   PC2 (2.8%)
                                     0
                                      T
                                                             ko3
                                     7
                                        wt5
                                                           5
                                       -10
                                              -5
                                                    0
                                                                 10
                                                  PC1 (91%)
```

- Finally, lets look at how to use the loading scores to determine which genes have the largest effect on where samples are plotted in the PCA plot
 - The prcomp() function calls loading scores \$rotation

```
## Lets focus on PC1 as it accounts for > 90% of variance loading_scores <- pca$rotation[,1]
```

- Finally, lets look at how to use the loading scores to determine which genes have the largest effect on where samples are plotted in the PCA plot
 - The prcomp() function calls loading scores \$rotation

```
## Lets focus on PC1 as it accounts for > 90% of variance
loading_scores <- pca$rotation[,1]

summary(loading_scores)
    Min. 1st Qu. Median Mean 3rd Qu. Max.
-0.104763 -0.104276 -0.068784 -0.005656 0.103926 0.104797

## We are interested in the magnitudes of both plus
## and minus contributing genes
gene_scores <- abs(loading_scores)</pre>
```

- Finally, lets look at how to use the loading scores to determine which genes have the largest effect on where samples are plotted in the PCA plot
 - The prcomp() function calls loading scores \$rotation

```
loading_scores <- pca$rotation[,1]
gene_scores <- abs(loading_scores)

## Sort by magnitudes from high to low
gene_score_ranked <- sort(gene_scores, decreasing=TRUE)</pre>
```

- Finally, lets look at how to use the loading scores to determine which genes have the largest effect on where samples are plotted in the PCA plot
 - The prcomp() function calls loading scores \$rotation

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loading_scores <- pca$rotation[,1]

gene_scores <- abs(loading_scores)

## Sort by magnitudes from high to low
gene_score_ranked <- sort(gene_scores, decreasing=TRUE)

## Find the names of the top 5 genes
top_5_genes <- names(gene_score_ranked[1:5])</pre>
```

- Finally, lets look at how to use the loading scores to determine which genes have the largest effect on where samples are plotted in the PCA plot
 - The prcomp() function calls loading scores \$rotation

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

gene_scores <- abs(loading_scores)

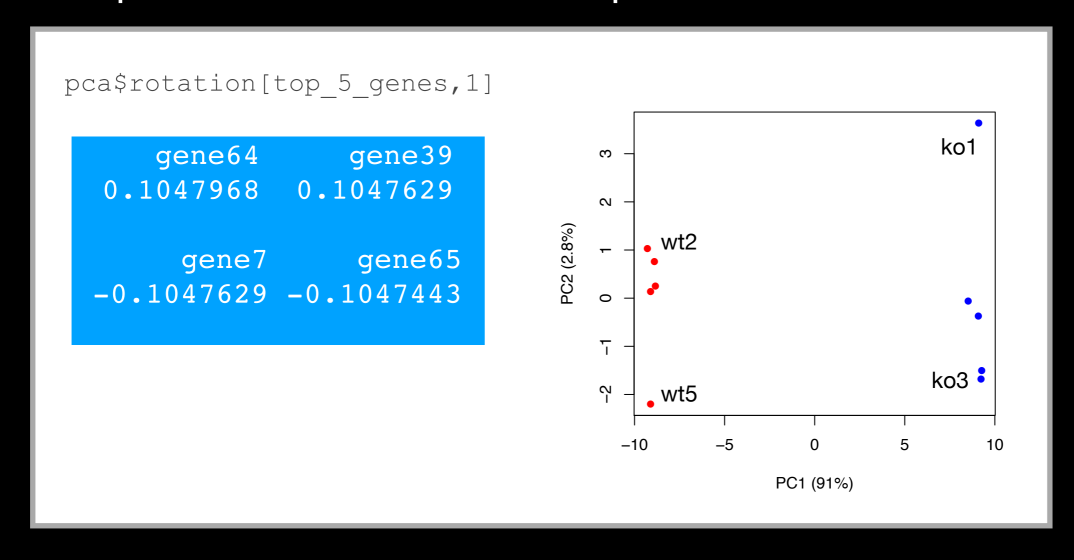
## Sort by magnitudes from high to low
gene_score_ranked <- sort(gene_scores, decreasing=TRUE)

## Find the names of the top 5 genes
top_5_genes <- names(gene_score_ranked[1:5])

## Show the scores (with +/- sign)
pca$rotation[top_5_genes,1]</pre>
```

- Here we see genes with the largest positive loading scores that effectively 'push' the "ko" samples to the right positive side of the plot.
- And the genes with high negative scores that push "wt" samples to the left side of the plot.

- Here we see genes with the largest positive loading scores that effectively 'push' the "ko" samples to the right positive side of the plot.
- And the genes with high negative scores that push "wt" samples to the left side of the plot.



PCA Recap

- PCA is classic "multivariate statistical technique" used to reduce the dimensionality of a complex data set to a more manageable number (typically 2D or 3D)
- For a matrix of m genes x n samples, we mean center (i.e. subtract the sample mean from each sample column), optionally rescale the values for each sample column, then calculate a new covariance matrix of size n x n
- We finally diagonalize the covariance matrix to yield our n Eigenvectors (called principal components or PCs) and n Eigenvalues.
- The top PCs (with largest Eigenvalues) retain the essential features of the original data and represent a useful subspace for further analysis (e.g. visualization, clustering, feature extraction, outlier detection etc...)

PCA objectives in a nutshell

- to reduce dimensionality
- to visualize multidimensional data
- to choose the most useful variables (features)
- to identify groupings of objects (e.g. genes/samples)
- to identify outliers

Your turn!

Perform a PCA on the UK foods dataset

UK foods.csv

Input: read, View/head,

PCA: prcomp,

Plots: PCA plot

scree plot,

loadings plot.

