Class 16 Mini Project

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Downstream analysis

Install tximport and rhdf5 in console with BiocManager

```
# BiocManager::install("tximport")
# BiocManager::install("rhdf5")
library(tximport)
library(rhdf5)
```

Setup the folder and filenames to read

```
folders <- dir(pattern="SRR21568*")
samples <- sub("_quant", "", folders)
files <- file.path( folders, "abundance.h5" )
names(files) <- samples

txi.kallisto <- tximport(files, type = "kallisto", txOut = TRUE)</pre>
```

1 2 3 4

head(txi.kallisto\$counts)

SRR2156848 SRR2156849 SRR2156850 SRR2156851 ENST00000539570 0.00000 ENST00000576455 0 2.62037 0 ENST00000510508 0 0.00000 0 0 ENST00000474471 1 1.00000 0 ENST00000381700 0 0.00000 0 ENST00000445946 0.00000

We now have our estimated transcript counts for each sample in R. We can see how many transcripts we have for each sample:

```
colSums(txi.kallisto$counts)

SRR2156848 SRR2156849 SRR2156850 SRR2156851
2563611 2600800 2372309 2111474
```

And how many transcripts are detected in at least one sample:

```
sum(rowSums(txi.kallisto$counts)>0)
```

```
[1] 94561
```

Before subsequent analysis, we might want to filter out those annotated transcripts with no reads:

```
to.keep <- rowSums(txi.kallisto$counts) > 0
kset.nonzero <- txi.kallisto$counts[to.keep,]</pre>
```

And those with no change over the samples:

```
keep2 <- apply(kset.nonzero,1,sd)>0
x <- kset.nonzero[keep2,]</pre>
```

Principal Component Analysis

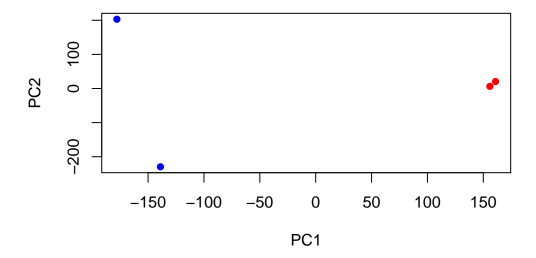
We compute the principal components, centering and scaling each transcript's measured levels so that each feature contributes equally to the PCA:

```
pca <- prcomp(t(x), scale=TRUE)
summary(pca)</pre>
```

Importance of components:

```
PC1 PC2 PC3 PC4
Standard deviation 183.6379 177.3605 171.3020 1e+00
Proportion of Variance 0.3568 0.3328 0.3104 1e-05
Cumulative Proportion 0.3568 0.6895 1.0000 1e+00
```

We can use the first two principal components as a co-ordinate system for visualizing the summarized transcriptomic profiles of each sample:



Q. Use ggplot to make a similar figure of PC1 vs PC2 and a seperate figure PC1 vs PC3 and PC2 vs PC3.

```
library(ggplot2)
library(ggrepel)
```

Make metadata object for the samples

```
colData <- data.frame(condition = factor(rep(c("control", "treatment"), each = 2)))
rownames(colData) <- colnames(txi.kallisto$counts)</pre>
```

Make the data.frame for ggplot

```
y <- as.data.frame(pca$x)
y$Condition <- as.factor(colData$condition)</pre>
```

Plot

```
ggplot(y) +
  aes(PC1, PC2, col=Condition) +
  geom_point() +
  geom_text_repel(label=rownames(y)) +
  theme_bw()
```

