



Vall d'Hebron  
Institut de Recerca

VHIR

# Vall d'Hebron Institut de Recerca

**Practical session:  
Running a differential expression  
analysis on RNA-seq data**

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2021/02/22

# HANDS ON: Introduction to RNA-Seq Differential Expression Analysis

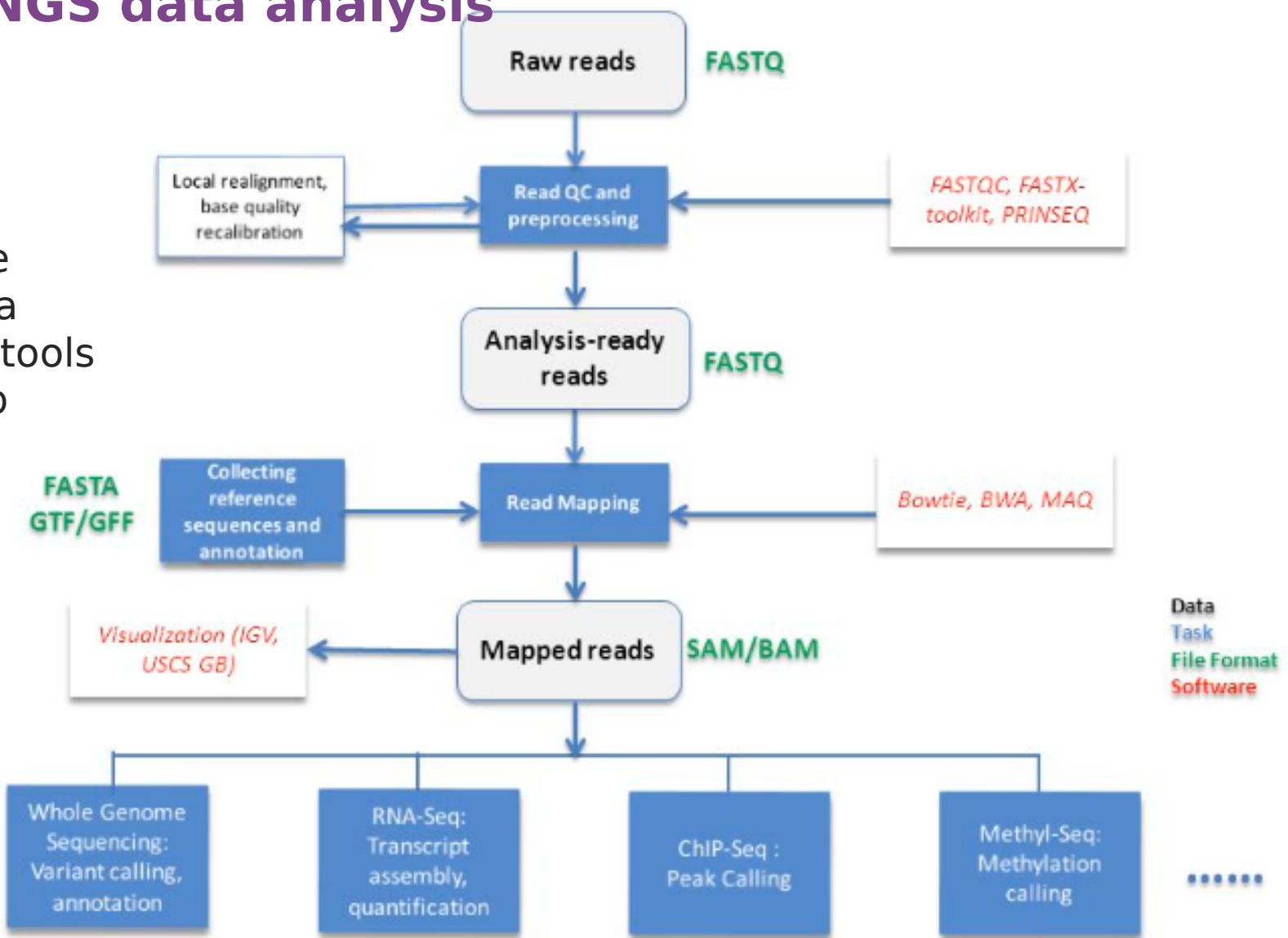
Bioinformàtica per a la Recerca Biomèdica

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Esther Camacho<sup>1</sup>, Angel Blanco<sup>1,2</sup>**

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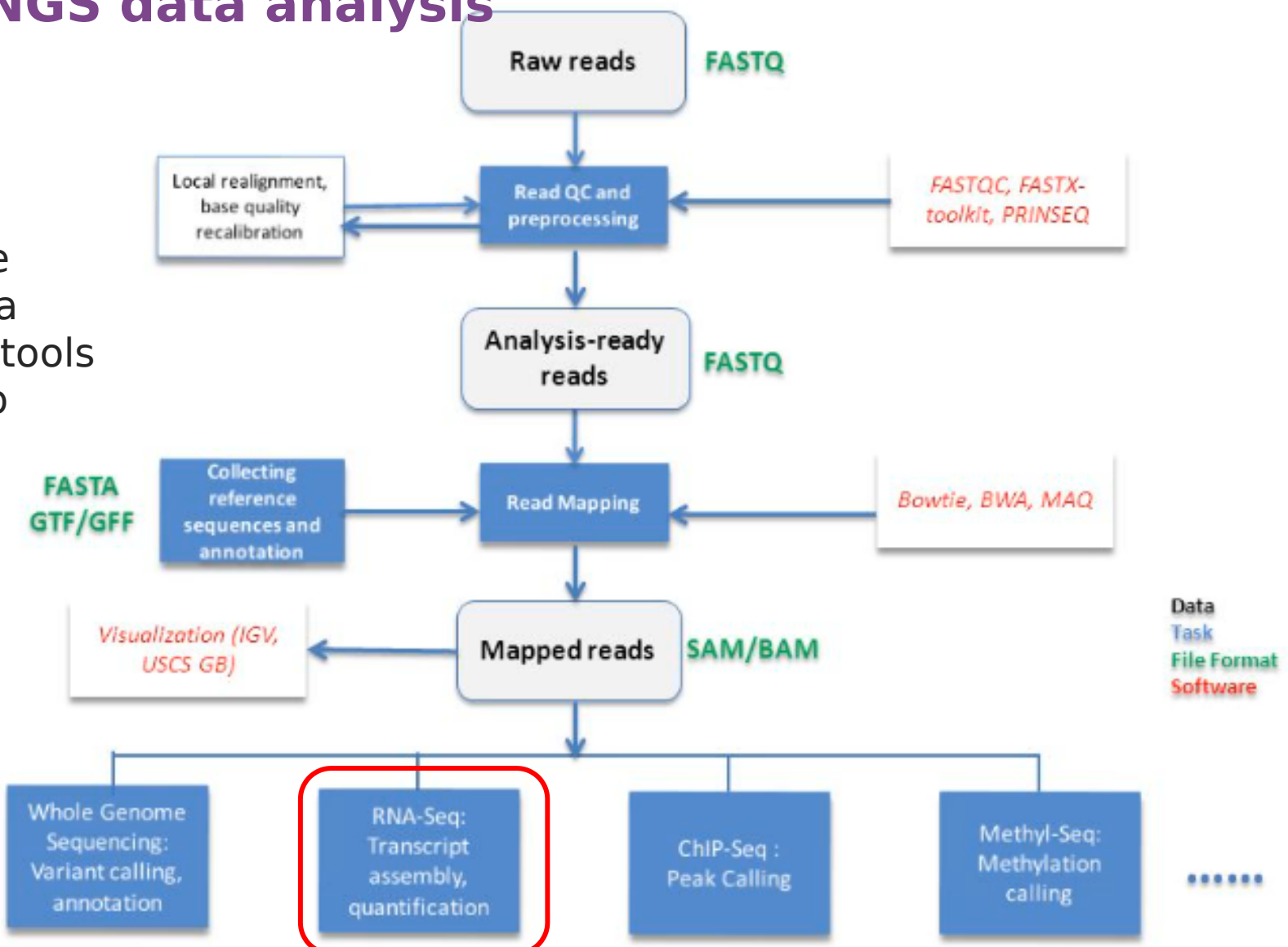
# Steps in NGS data analysis

- We will have different data formats and tools for each step

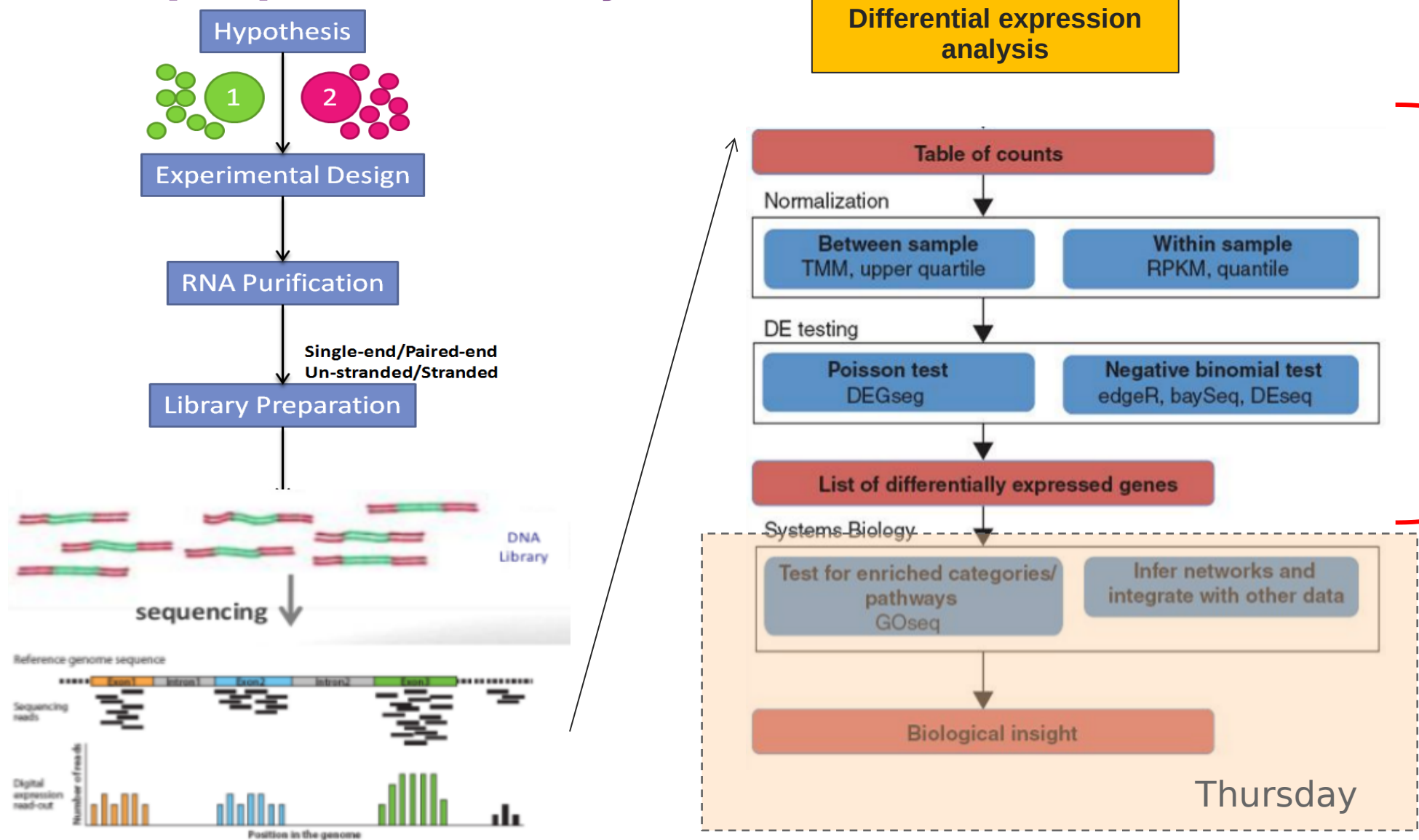


# Steps in NGS data analysis

- We will have different data formats and tools for each step

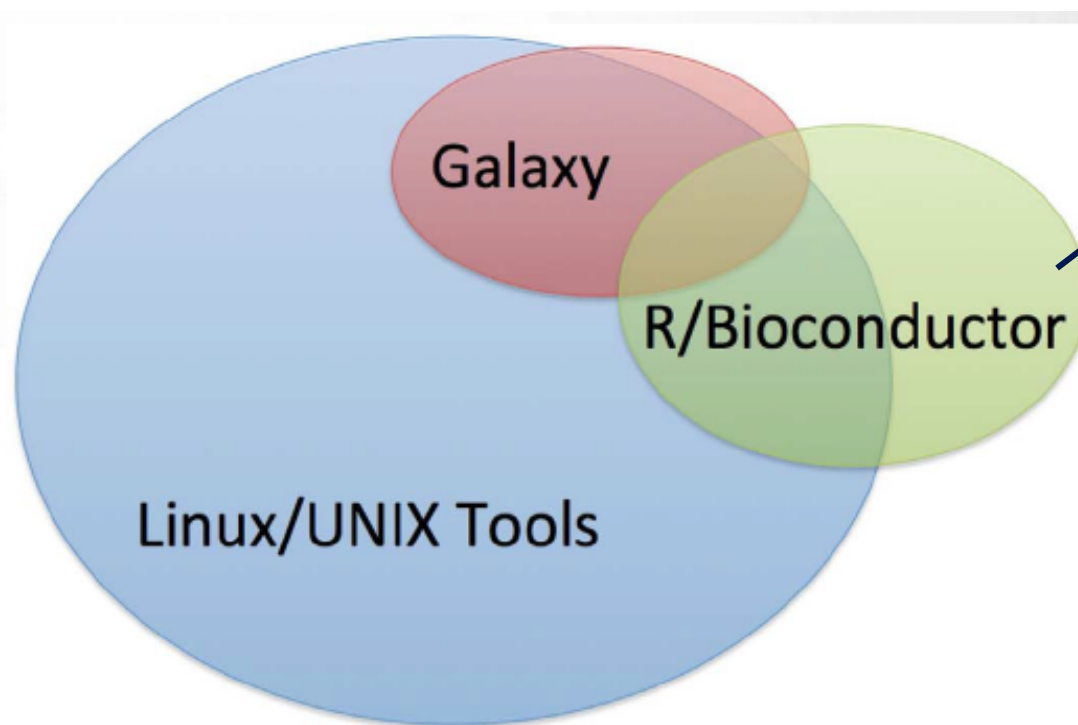


# RNA-seq Expression Analysis Workflow



## Tools for NGS data analysis

Highly efficient and fast processing tools are required to handle large volume of datasets



### About Bioconductor

Bioconductor provides tools for the analysis and comprehension of high-throughput genomic data. Bioconductor uses the R statistical programming language, and is open source and open development. It

# Today's practicum

## Steps covered by the tutorial:

- Start with the FASTQ files (how they are aligned to the genome)
- 1. Summarization/Quantification of aligned reads: obtaining the counts matrix
- 2. Data pre-processing and exploratory analysis
- 3. Differential gene expression analysis
- 4. Visually explore the results

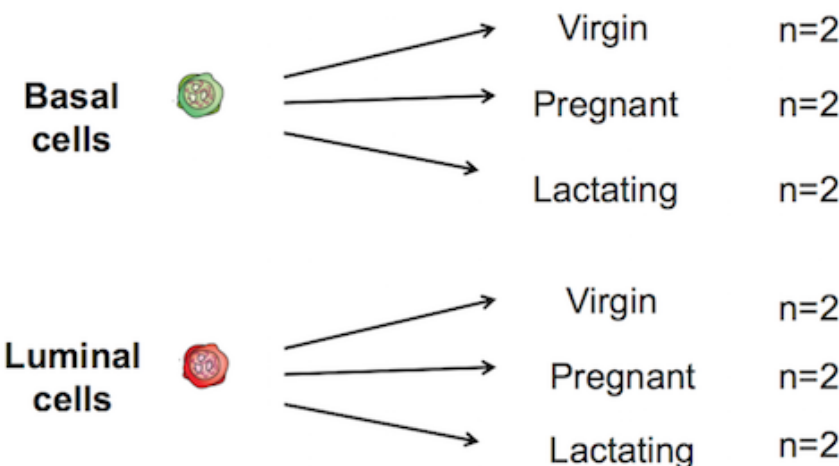
This material has been created using the following sources:

- <https://combine-australia.github.io/RNAseq-R/>
- <https://bioconductor.org/packages/release/workflows/vignettes/RNAseq123/install/doc/limmaWorkflow.html>

## Study overview

- Objective: We want to identify genes differentially expressed in the lactating versus pregnant mammary gland

Dataset: RNA-seq data of mouse mammary gland ([GSE60450](#))



Fu *et al.* (2015) 'EGF-mediated induction of Mcl-1 at the switch to lactation is essential for alveolar cell survival' Nat Cell Biol

Note: two biological replicates are used here, however three replicates is usually recommended as a minimum requirement for RNA-seq.

GSE60450

Scope: Self Format: HTML Amount: Quick GEO accession: GSE60450

## Series GSE60450

Query DataSets for GSE60450

Status	Public on Jan 19, 2015
Title	Transcriptome analysis of luminal and basal cell subpopulations in the lactating versus pregnant mammary gland
Organism	<a href="#">Mus musculus</a>
Experiment type	Expression profiling by high throughput sequencing
Summary	To identify genes specifically expressed in lactating mammary glands, the gene expression profiles of luminal and basal cells from different developmental stages were compared.
Overall design	Comparison of gene expression in luminal and basal cells harvested from the mammary glands of virgin, 18.5 day pregnant and 2 day lactating mice (2 mice per stage).
Contributor(s)	<a href="#">Fu NY</a> , <a href="#">Lun A</a> , <a href="#">Smyth GK</a> , <a href="#">Visvader JE</a>
Citation(s)	Fu NY, Rios AC, Pal B, Soetanto R et al. EGF-mediated induction of Mcl-1 at the switch to lactation is essential for alveolar cell survival. <i>Nat Cell Biol</i> 2015 Apr;17(4):365-75. PMID: <a href="#">25730472</a>
Platforms (1)	<a href="#">GPL13112</a> Illumina HiSeq 2000 (Mus musculus)
Samples (12)	<a href="#">GSM1480291</a> Luminal virgin #1 <a href="#">GSM1480292</a> Luminal virgin #2 <a href="#">GSM1480293</a> Luminal 18.5 dP #1
Relations	
BioProject	<a href="#">PRJNA258286</a>
SRA	<a href="#">SRP045534</a>

**Organism:** [Mus musculus](#)
**Library:****Instrument:** Illumina HiSeq 2000**Strategy:** RNA-Seq**Source:** TRANSCRIPTOMIC**Selection:** cDNA**Layout:** SINGLE**Construction protocol:** FACS-sorted cells were for sequencing using standard Illumina protocol

# Getting started

Hands on!

... follow **sections 1-2 of the Rmd file**

## Steps covered in this tutorial:

- 1. Summarization/Quantification of aligned reads: obtaining the counts matrix**
2. Data pre-processing and exploratory analysis
3. Differential gene expression analysis
4. Visually explore the results

# Step 1: Summarization/Quantification of reads: obtaining the counts matrix

- The alignment produces a set of **BAM** files, where each file contains the read alignments for each sample.
- In the BAM file, there is a chromosomal location for every read that was

**Headers**

First Record

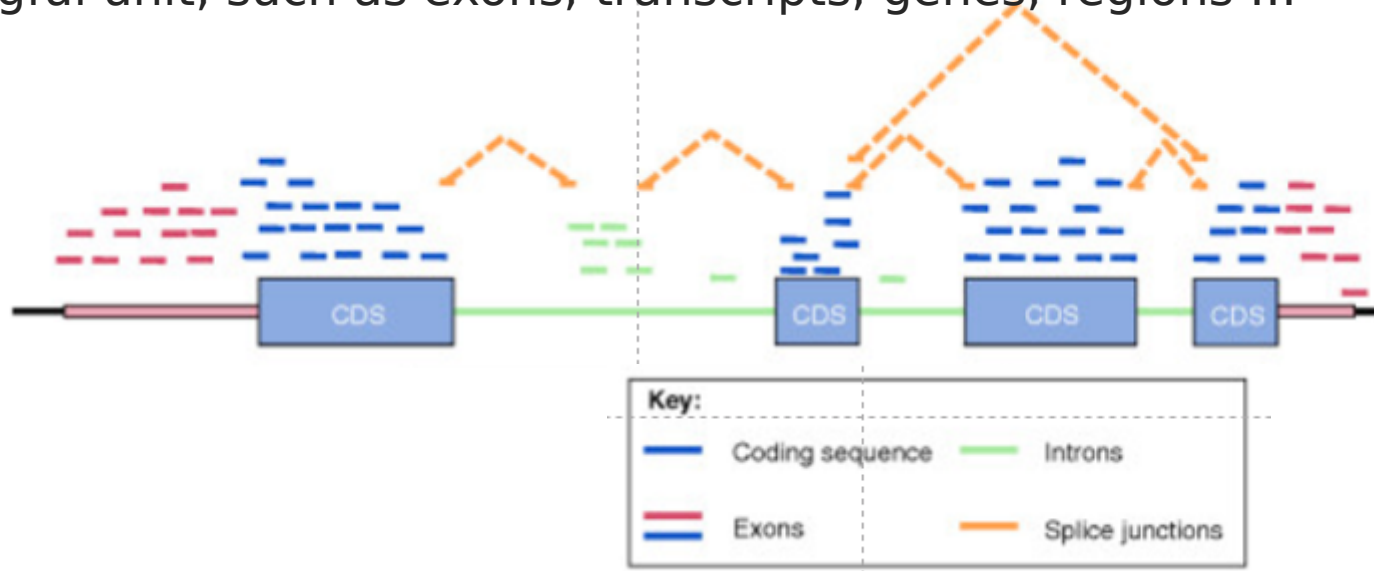
```

@HD VN:1.3 SO:coordinate
@SQ SN:22 LN:61304566 AS:NCBI37 M5:a718acaa6135fdca8357d5bfe94211dd UR
:file:/home/mktrost/seqshop/example/ref22/human.g1k.v37.chr22.fa
@RG ID:0 SM:HG00551 LB:HG00551 CN:unknown PL:ILLUMINA
@PG ID:bwa PN:bwa VN:0.7.10-r960-dirty CL:/home/mktrost/seqshop/gotcloud/bin/bwa
mem -t 1 -M -R @RG\@ID:0\@SM:HG00551\@LB:HG00551\@CN:unknown\@PL:ILLUMINA /home/mktrost/seqshop/example/ref22/human.g1k.v37.chr22.fa /home/mktrost/seqshop/example/fastq/HG00551.SRR
190851.fastq
@RG ID:1 SM:HG00551 LB:HG00551 CN:unknown PL:ILLUMINA
SRR190851.48112415 113 22 16918656 3 23M78S = 31650772 1
4732127 TCCTCGACCTCCAAAGTCTGGTATTAGCCCTTAGAGCCACCGCACCCAGCCAGTTATCTTTTTAAAATGTTATTTA
ATACATTATTTTTATACT ##### Chromosome/position :0A22 NM:i:1 OQ:Z:#####
##### :0A22 NM:i:1 OQ:Z:##### R
G:Z:1 XS:i:21 XA:Z:22,-38586564,7521M73S,0;
SRR190851.103013373 121 22 16936847 2 37S18M2D46M = 16
936847 0 CACAAGTTCAAAGTCCACAGATCTCAAAGGCAGGTACAAAATCCCACCAAGTCTCTGCTAAAGCATAGCAAGAG
TGACCTTACTCCAGTCCAAACAAGT ##### :560523/+131335234238-2241+7-,+/-+89,6 AS:i:46 MD:Z:6G11^TT30T15 NM:i:4 OQ
:Z:##### BC>D?@?>DB?CDEB?CDAE6D@C
C>A8976=?;?A98 RG:Z:1 XS:i:44 XA:Z:22,-36435775,56M1I2M3I39M,12;

```

# Step 1: Summarization/Quantification of reads: obtaining the counts matrix

- We need to summarize and aggregate reads over some biologically meaningful unit, such as exons, transcripts, genes, regions ...



- Many methods available

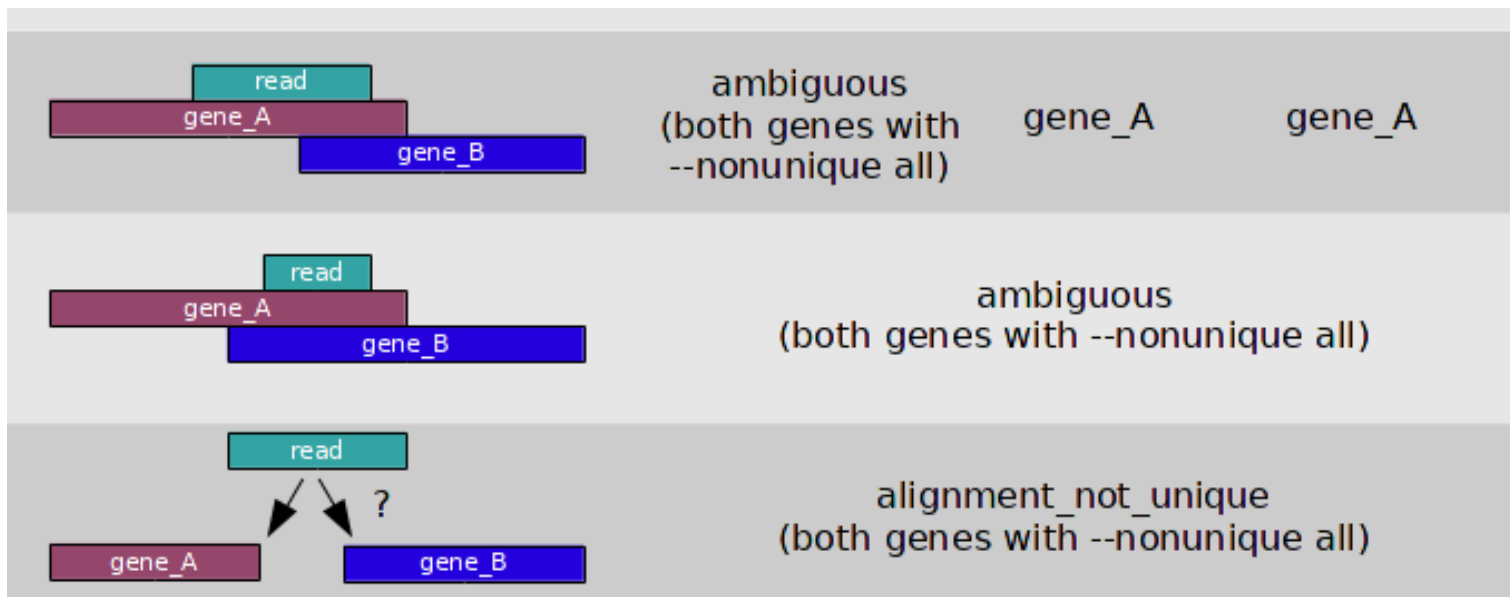
# Step 1: Summarization/Quantification of reads: obtaining the counts matrix

- Requires gene annotation specifying the genomic start and end position of each exon of each gene . Usually this is contained in a data frame in *GTF* format for each organism.

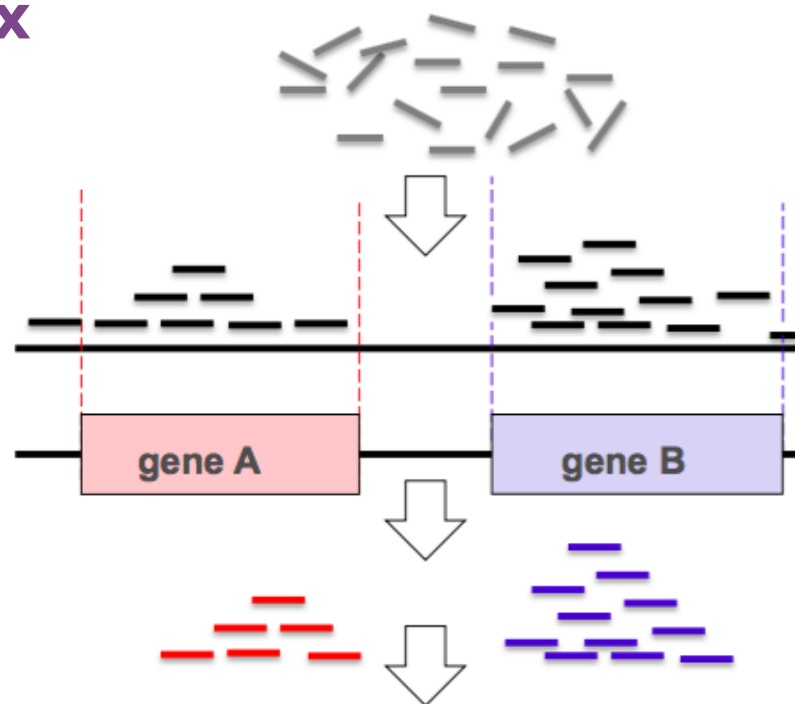
Seqname	Source	Feature	Start	End	Score	Strand	Frame	Attributes
chr4	protein_coding	CDS	24053	24477	.	+	0	exon_number "1"; gene_id "FBgn00400
chr4	protein_coding	exon	24053	24477	.	+	.	exon_number "1"; gene_id "FBgn00400
chr4	protein_coding	CDS	24979	25153	.	+	1	exon_number "2"; gene_id "FBgn00400
chr4	protein_coding	exon	24979	25153	.	+	.	exon_number "2"; gene_id "FBgn00400
chr4	protein_coding	CDS	25218	25450	.	+	0	exon_number "3"; gene_id "FBgn00400
chr4	protein_coding	exon	25218	25450	.	+	.	exon_number "3"; gene_id "FBgn00400
chr4	protein_coding	CDS	25501	25618	.	+	1	exon_number "4"; gene_id "FBgn00400
chr4	protein_coding	exon	25501	25621	.	+	.	exon_number "4"; gene_id "FBgn00400
chr4	protein_coding	stop_codon	25619	25621	.	+	0	exon_number "4"; gene_id "FBgn00400
chr4	pseudogene	exon	26994	27101	.	-	.	exon_number "7"; gene_id "FBgn00520
chr4	pseudogene	exon	27167	27349	.	-	.	exon_number "6"; gene_id "FBgn00520
chr4	pseudogene	exon	28371	28609	.	-	.	exon_number "5"; gene_id "FBgn00520

# Step 1: Summarization/Quantification of reads: obtaining the counts matrix

- Reads that map to exons of genes are added together to obtain the count for each gene
- There may be some ambiguities



# Step 1: Summarization/Quantification of reads: obtaining the counts matrix



- The summarized RNA-seq data is widely known as a ***count matrix***

	Control 1	Control 2	Control 3	Sample 1	Sample 2	Sample 3
Gene A	6	5	7	17	10	11
Gene B	11	11	10	3	4	2
Gene C	200	150	355	50	1	3
Gene D	0	1	0	2	0	1

# Step 1: Summarization/Quantification of reads: obtaining the counts matrix

Hands on!

... follow **sections 3-4 of the Rmd file**

## Steps covered in this tutorial:

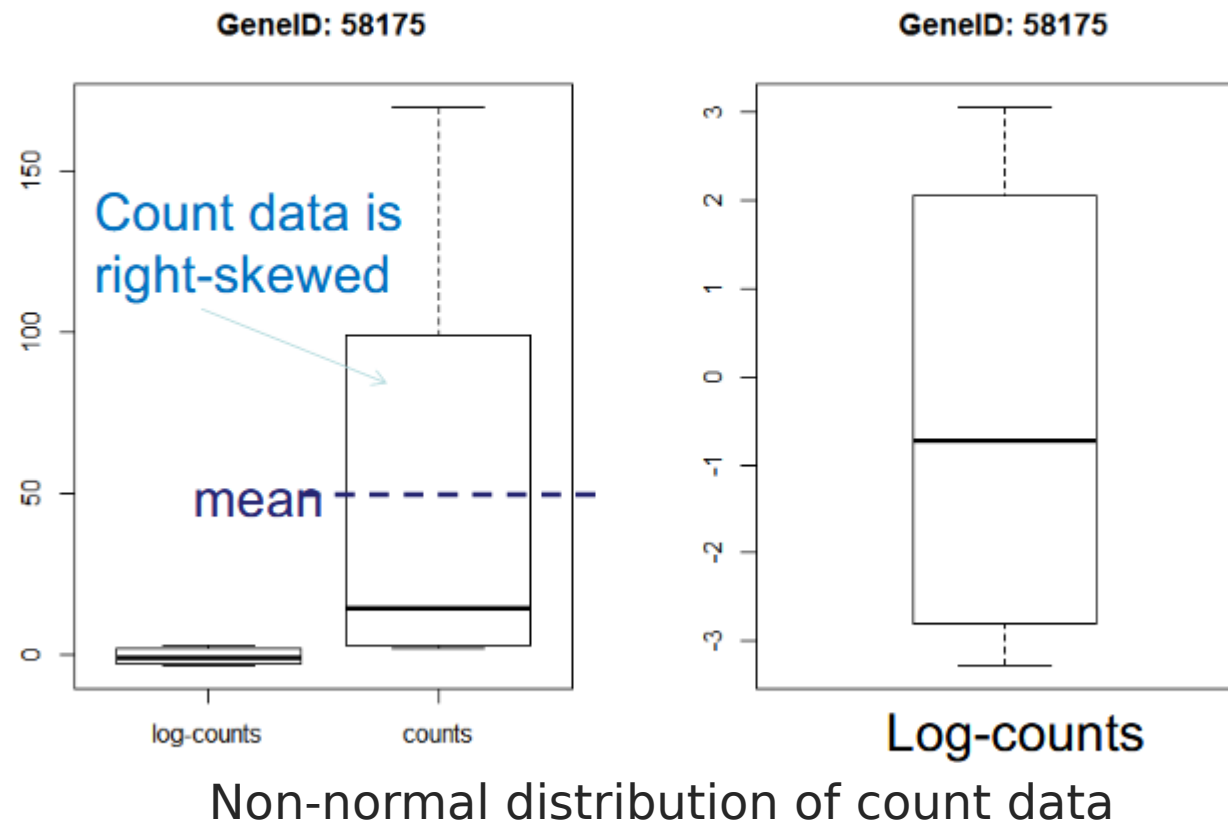
1. Summarization/Quantification of aligned reads: obtaining the counts matrix
- 2. Data pre-processing and exploratory analysis**
3. Differential gene expression analysis
4. Visually explore the results

## Step 2: Data pre-processing and exploratory analysis

- How my data looks like?
- Is it of enough quality for analysis?
- Are there some outlier samples that should be removed?
- Are samples grouped according to the experimental conditions?
- What are the main sources of variability in the data?

## Step 2: Data pre-processing and exploratory analysis

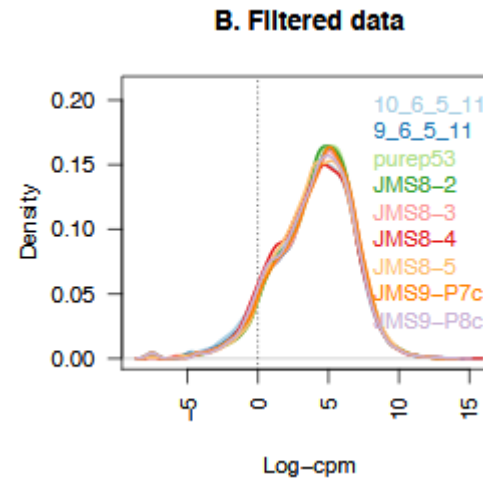
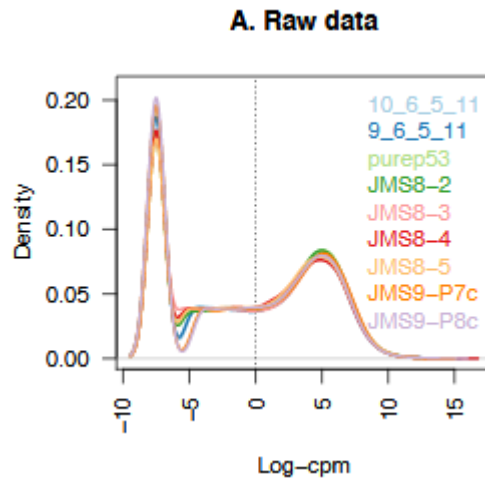
- RNA-seq data is
  - discrete
  - has non-constant mean-variance trend



## Step 2: Data pre-processing and exploratory analysis

### Filtering

- It is recommended to filter for lowly expressed genes before differential expression testing.
  - provide little evidence for differential expression and they interfere with some of the statistical approximations used
  - Add to the multiple testing burden when estimating detect differential expressed

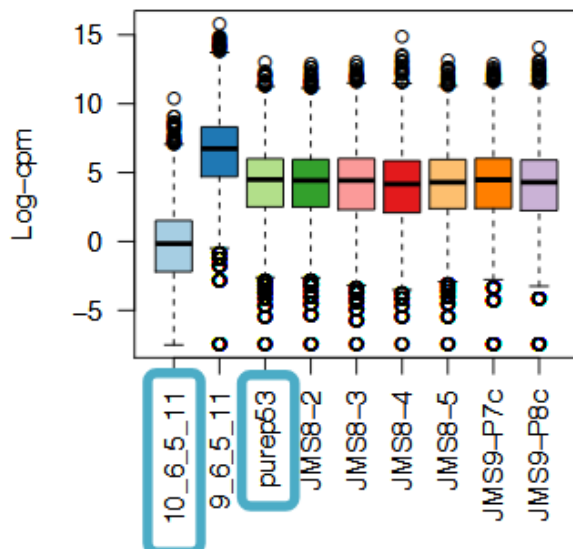


# Step 2: Data pre-processing and exploratory analysis

## Scaling and normalization

- The counts of mapped reads for each gene is proportional to the expression of RNA (“interesting”) in addition to many other factors (“uninteresting”).
- Normalization is the process of scaling raw count values to account for the “uninteresting” factors.

A. Example: Unnormalised data



If we ran a DE analysis on  
Sample 1 and Sample 3, almost  
all genes will be down-  
regulated in Sample 1!!

# Step 2: Data pre-processing and exploratory analysis

## Scaling and normalization

Main factors often considered during normalization are:

1. **Sequencing depth:** total number of reads mapped to the genome



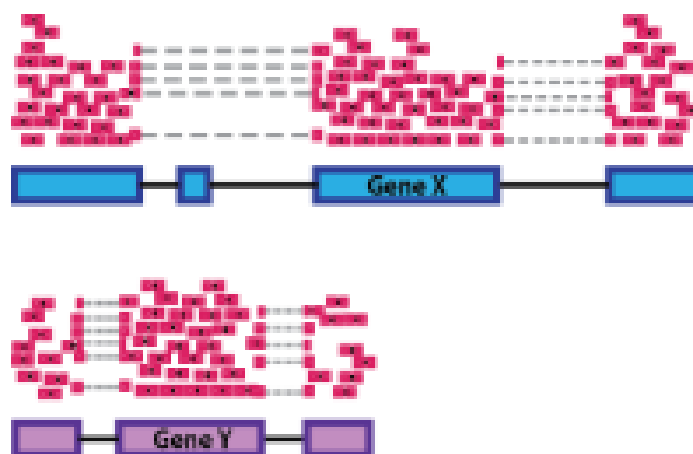
# Step 2: Data pre-processing and exploratory analysis

## Scaling and normalization

Main factors often considered during normalization are:

2. **Gene length:** Accounting for gene length is necessary for comparing expression between different genes within the same sample.

**Sample A Reads**

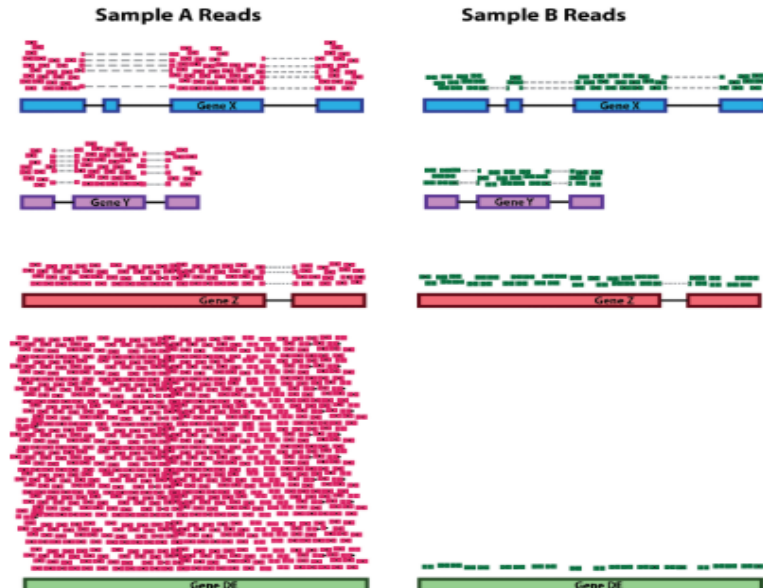


## Step 2: Data pre-processing and exploratory analysis

### Scaling and normalization

Main factors often considered during normalization are:

3. **RNA composition:** A few highly differentially expressed genes between samples, differences in the number of genes expressed between samples, or presence of contamination can skew some types of normalization methods.



[https://hbctraining.github.io/DGE\\_workshop/lessons/02\\_DGE\\_count\\_normalization.html](https://hbctraining.github.io/DGE_workshop/lessons/02_DGE_count_normalization.html)

# Step 2: Data pre-processing and exploratory analysis

## Scaling and normalization

Normalization method	Description	Accounted factors	Recommendations for use
CPM (counts per million)	counts scaled by total number of reads	sequencing depth	gene count comparisons between replicates of the same samplegroup; <b>NOT for within sample comparisons or DE analysis</b>
TPM (transcripts per kilobase million)	counts per length of transcript (kb) per million reads mapped	sequencing depth and gene length	gene count comparisons within a sample or between samples of the same sample group; <b>NOT for DE analysis</b>
RPKM/FPKM (reads/fragments per kilobase of exon per million reads/fragments mapped)	similar to TPM	sequencing depth and gene length	gene count comparisons between genes within a sample; <b>NOT for between sample comparisons or DE analysis</b>
DESeq2's median of ratios [1]	counts divided by sample-specific size factors determined by median ratio of gene counts relative to geometric mean per gene	sequencing depth and RNA composition	gene count comparisons between samples and for <b>DE analysis</b> ; <b>NOT for within sample comparisons</b>
EdgeR's trimmed mean of M values (TMM) [2]	uses a weighted trimmed mean of the log expression ratios between samples	sequencing depth, RNA composition, and gene length	gene count comparisons between and within samples and for <b>DE analysis</b>

[https://hbctraining.github.io/DGE\\_workshop/lessons/02\\_DGE\\_count\\_normalization.html](https://hbctraining.github.io/DGE_workshop/lessons/02_DGE_count_normalization.html)

## **Step 2: Data pre-processing and exploratory analysis**

### **Exploratory analysis**

Unsupervised-separation methods based on data, without prior knowledge of experimental design, can be used to get an overview of the data

- Do samples separate by experimental groups?
- Where the greatest sources of variation in the data come from?
- Are there any outliers?

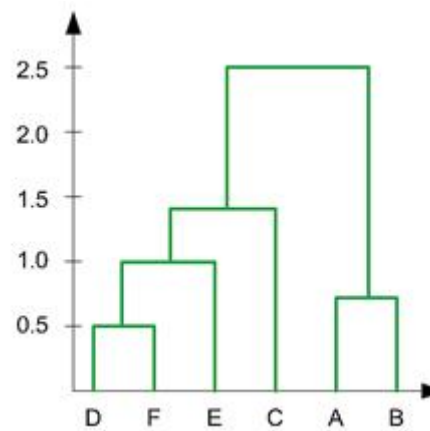
# Step 2: Data pre-processing and exploratory analysis

## Exploratory analysis

### Hierarchical clustering

- Hierarchical clustering is typically based on pairwise comparisons of individual samples, which are grouped into “neighborhoods” of similar samples. The basis of hierarchical clustering is therefore a matrix of similarity metrics.

Dist	A	B	C	D	E	F
A	0.00	0.71	5.66	3.61	4.24	3.20
B	0.71	0.00	4.95	2.92	3.54	2.50
C	5.66	4.95	0.00	2.24	1.41	2.50
D	3.61	2.92	2.24	0.00	1.00	0.50
E	4.24	3.54	1.41	1.00	0.00	1.12
F	3.20	2.50	2.50	0.50	1.12	0.00

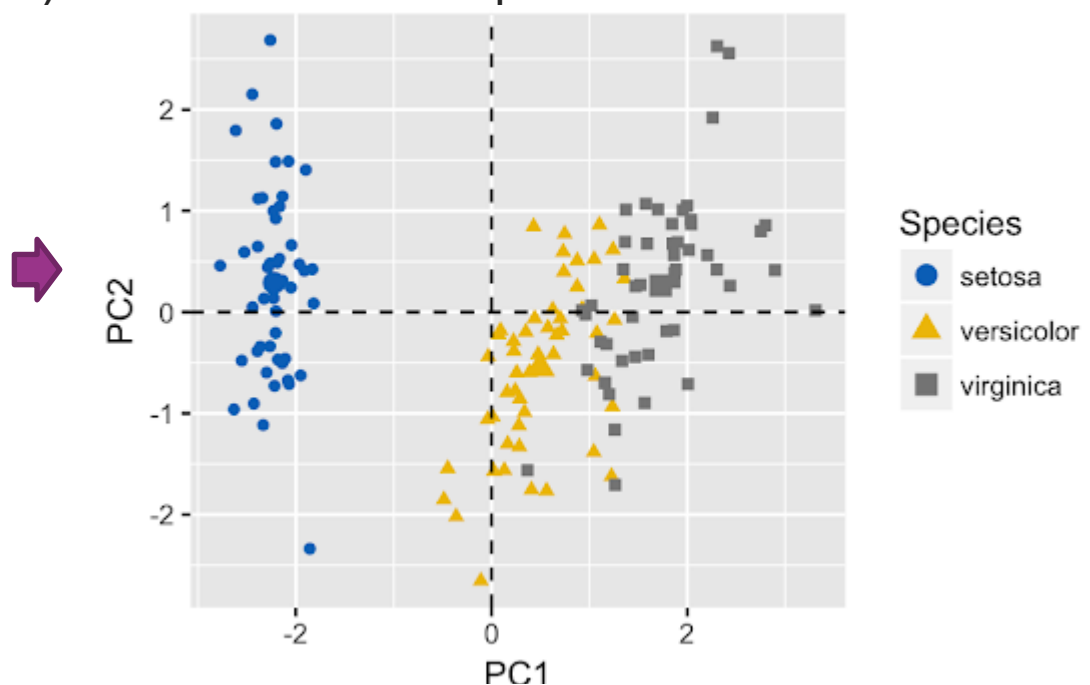
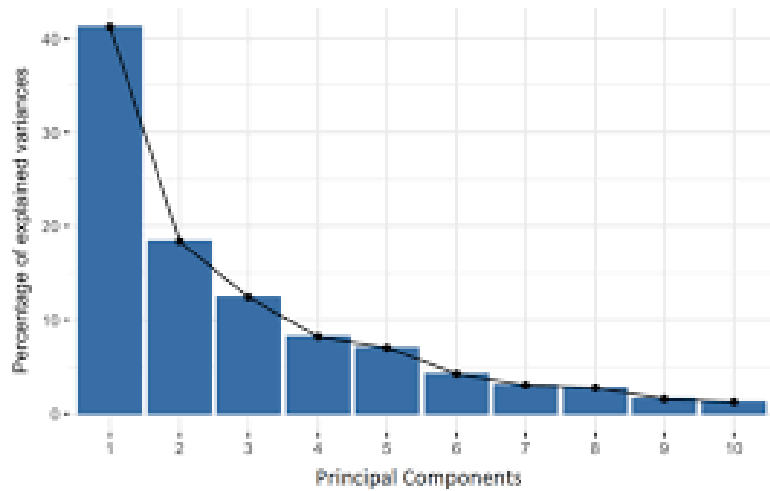


# Step 2: Data pre-processing and exploratory analysis

## Exploratory analysis

### Principal Components Analysis (PCA)

- A dimensionality reduction approach that aims to find groups of features (e.g., genes) that have something in common (e.g., certain patterns of expression across different samples)
- Few dimensions (components) can be used to represent the information from thousands of mRNAs.

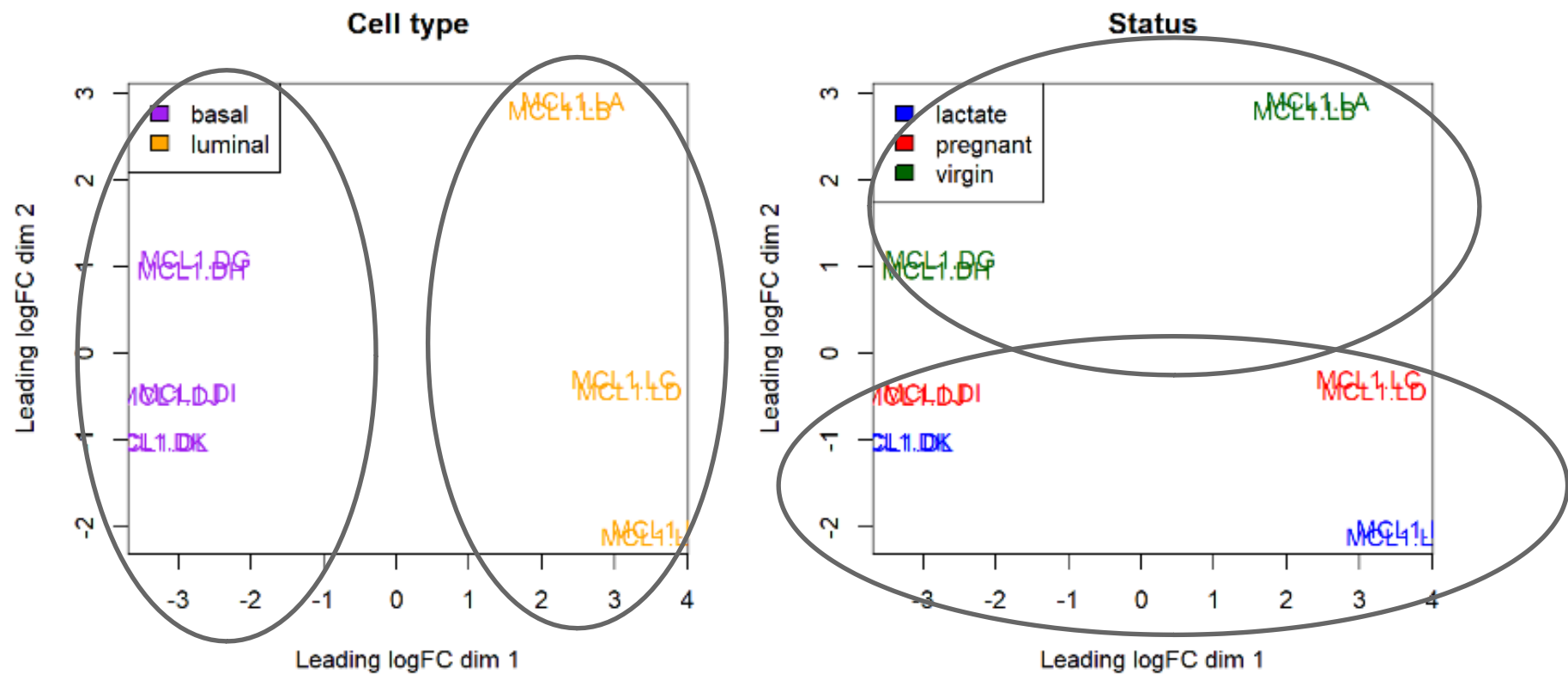


## Step 2: Data pre-processing and exploratory analysis

Hands on!

... follow **section 5 of the Rmd file**

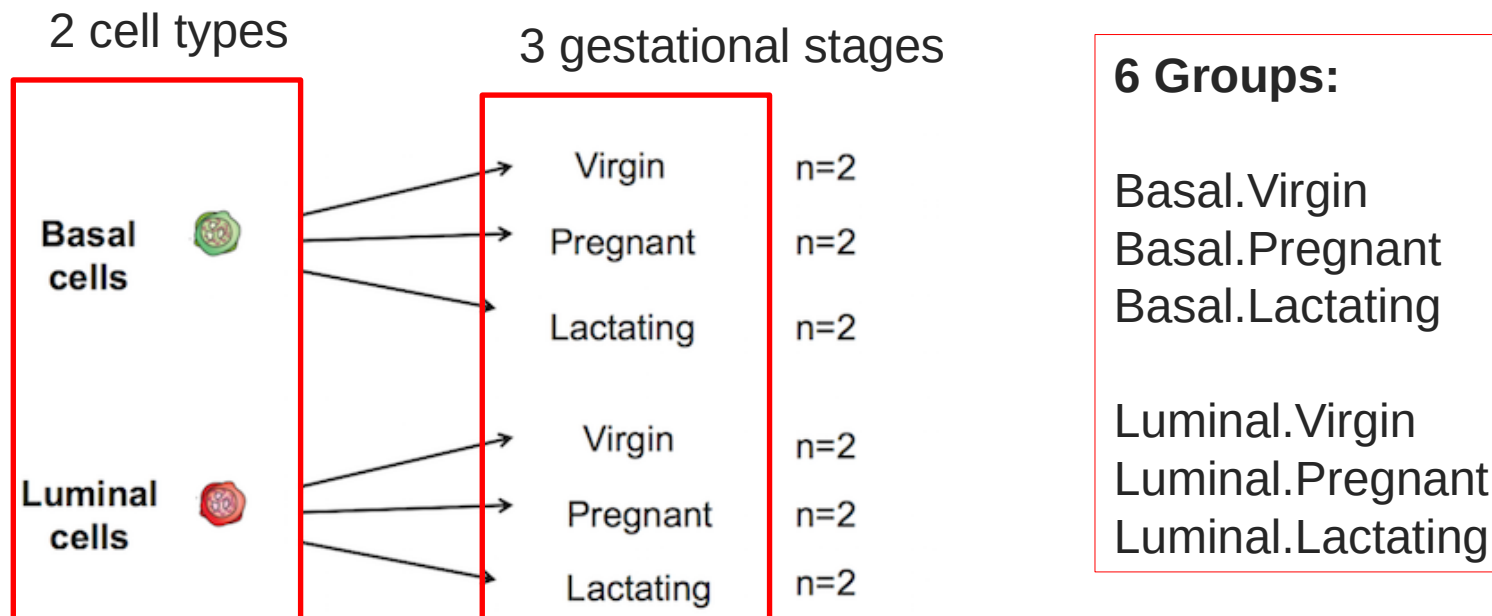
## Step 2: Data pre-processing and exploratory analysis



## Steps covered in this tutorial:

1. Summarization/Quantification of aligned reads: obtaining the counts matrix
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- 3. Differential gene expression analysis**
4. Visually explore the results

## Step 3: Differential expression analysis



Fu et al. (2015) 'EGF-mediated induction of Mcl-1 at the switch to lactation is essential for alveolar cell survival' Nat Cell Biol

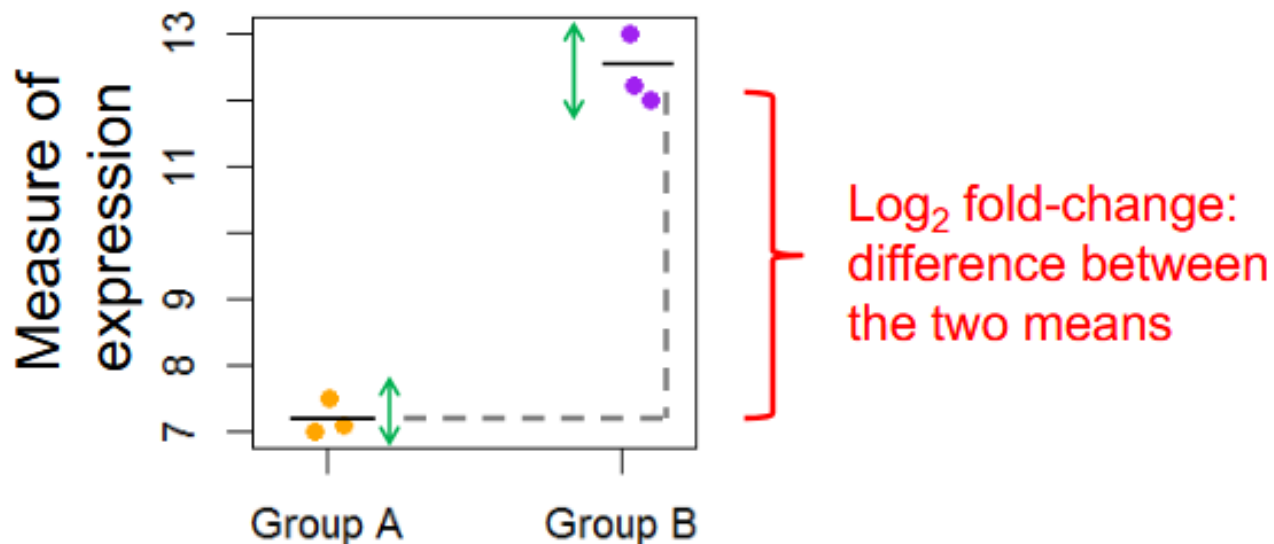
We will analyze the following comparisons:

- Basal.Lactating vs. Basal.Pregnant
- Luminal.Lactating vs. Luminal.Pregnant

## Step 3: Differential expression analysis

What do we need to perform a statistical test?

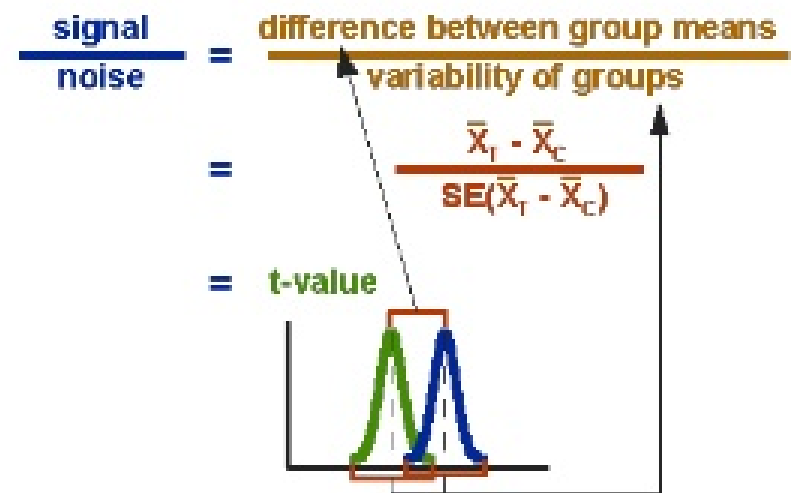
- Measure of average expression
- Measure of variability



- Finally we will be assigning a p-value for each test/gene.

## Step 3: Differential expression analysis

Couldn't we just use a Student's t test for each gene?

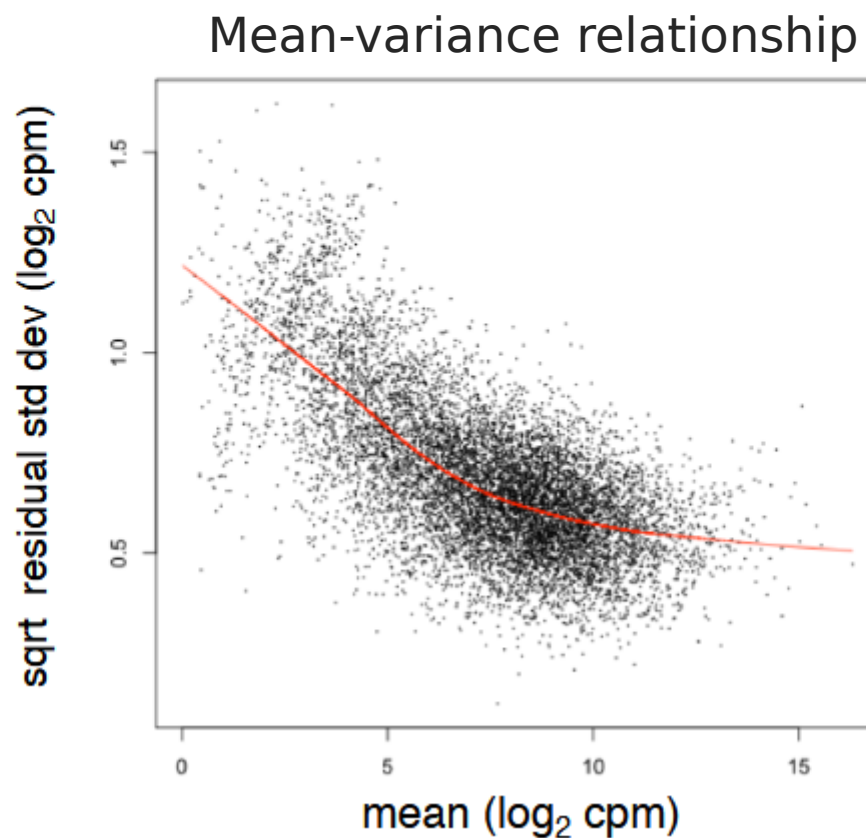
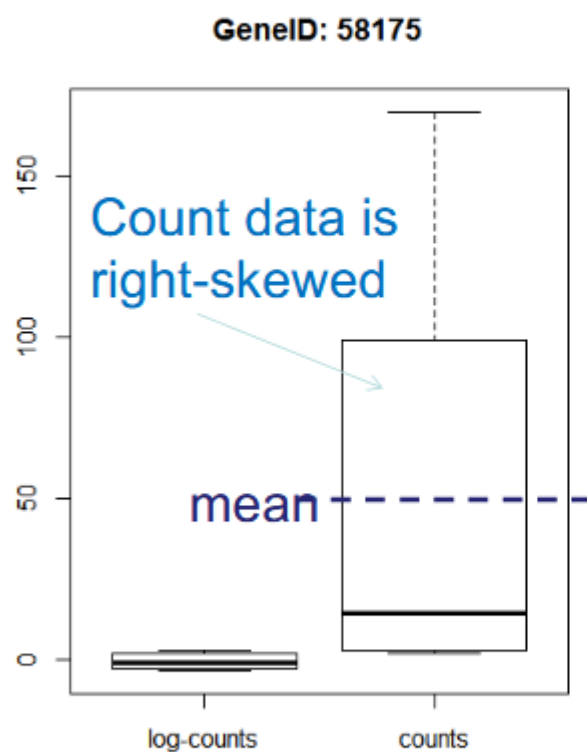


Problems with this approach:

- May have **few replicates**
- Distribution is **not normal**
- **Multiple testing** issues

[http://www.socialresearchmethods.net/kb/stat\\_t.php](http://www.socialresearchmethods.net/kb/stat_t.php)

- RNA-seq data is
  - discrete
  - has non-constant mean-variance trend



Non-normal distribution of count data

## Step 3: Differential expression analysis

Different software use different approaches to deal with the “t test issues”

*Distributional issue:* Solved by variance stabilizing transform in limma – voom() function

edgeR and DESeq model the count data using a *negative binomial distribution* and use their own modified statistical tests based on that.

*Multiple testing issue:* All of these packages report false discovery rate (corrected p values). For SAMseq based on resampling, for others usually Benjamini-Hochberg corrected p values.

*Variance estimation issue:* edgeR, DESeq2 and limma (in slightly different ways) “borrow” information across genes to get a better variance estimate. One says that the estimates “shrink” from gene-specific estimates towards a common mean value.

# Step 3: Differential expression analysis

Feature	DESeq2	edgeR	limmaVoom	Cuffdiff
<b>Seq. depth normalization</b>	Sample-wise size factor	Gene-wise trimmed median of means (TMM)	Gene-wise trimmed median of means (TMM)	FPKM-like or DESeq-like
<b>Dispersion estimate</b>	Cox-Reid approximate conditional inference with focus on maximum <i>individual</i> dispersion estimate	Cox-Reid approximate conditional inference moderated towards the <i>mean</i>	squeezes gene-wise residual variances towards the global variance	
<b>Assumed distribution</b>	Neg. binomial	Neg. binomial	<i>log</i> -normal	Neg. binomial
<b>Test for DE</b>	Wald test (2 factors); LRT for multiple factors	exact test for 2 factors; LRT for multiple factors	<i>t</i> -test	<i>t</i> -test
<b>False positives</b>	Low	Low	Low	High
<b>Detection of differential isoforms</b>	No	No	No	Yes
<b>Support for multi-factored experiments</b>	Yes	Yes	Yes	No
<b>Runtime (3-5 replicates)</b>	Seconds to minutes	Seconds to minutes	Seconds to minutes	Hours

<https://chagall.med.cornell.edu/RNASEQcourse/Intro2RNAseq.pdf>

# Step 3: Differential expression analysis

Feature	DESeq2	edgeR	limmaVoom	Cuffdiff
<b>Seq. depth normalization</b>	Sample-wise size factor	Gene-wise trimmed median of means (TMM)	Gene-wise trimmed median of means (TMM)	FPKM-like or DESeq-like
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<b>Assumed distribution</b>	Neg. binomial	Neg. binomial	<i>log</i> -normal	Neg. binomial
<b>Test for DE</b>	Wald test (2 factors); LRT for multiple factors	exact test for 2 factors; LRT for multiple factors	<i>t</i> -test	<i>t</i> -test
<b>False positives</b>	Low	Low	Low	High
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<https://chagall.med.cornell.edu/RNASEQcourse/Intro2RNAseq.pdf>

## Step 3: Differential expression analysis

Hands on!

... follow **sections 6.1-6.4 of the Rmd file**

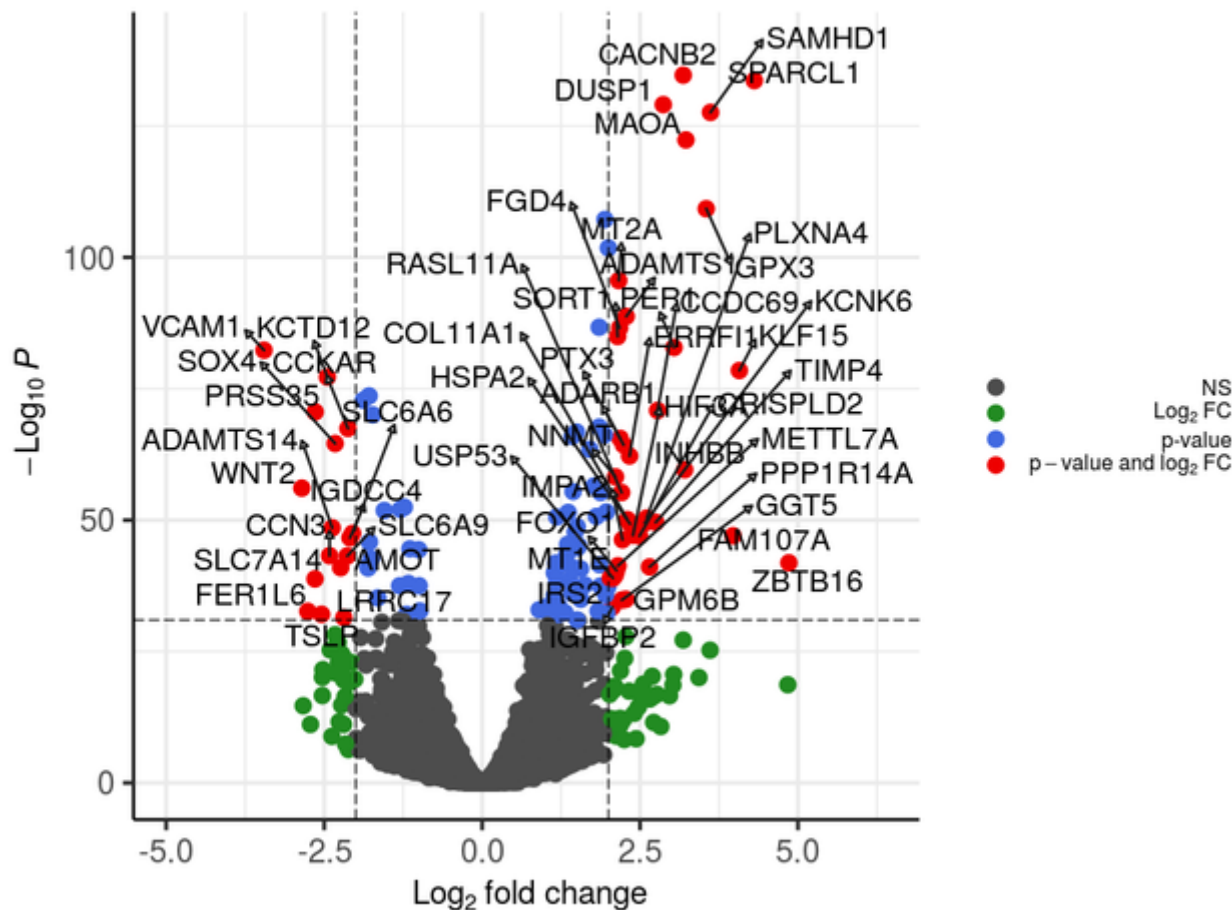
## Steps covered in this tutorial:

1. Summarization/Quantification of aligned reads: obtaining the counts matrix
2. Data pre-processing and exploratory analysis
3. Differential gene expression analysis
- 4. Visually explore the results**



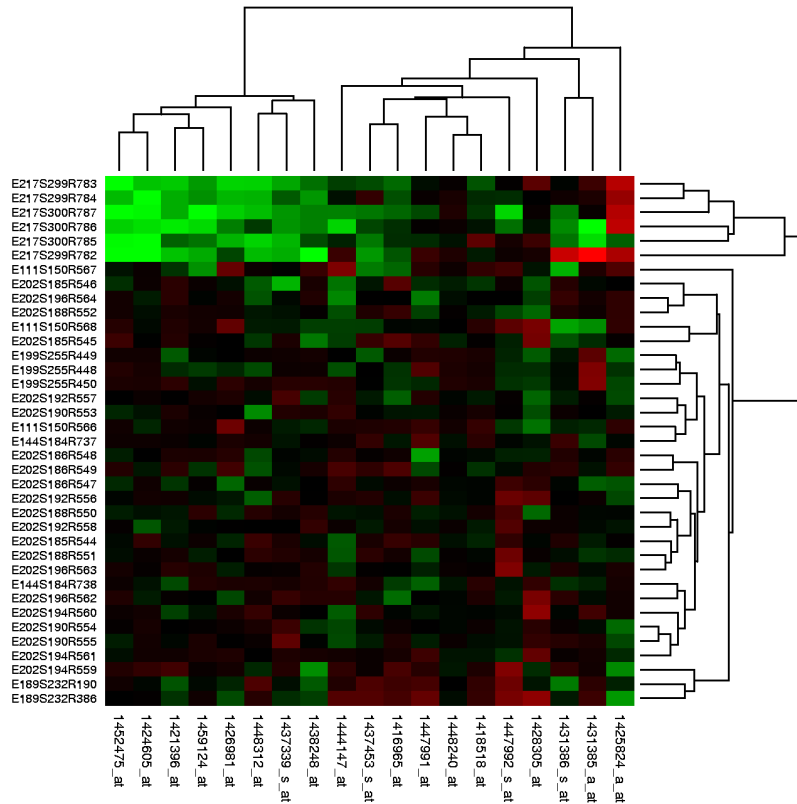
## Step 4: Visual exploration of results

- A common visualisation is the **Volcano Plot** which display a measure of significance on the y-axis and fold-change on the x-axis



## Step 4: Visual exploration of results

- Heatmap is a very useful tool for quick representation of quantitative differences in expression levels of biological data.
  - Each gene is represented as a row and is color-coded to represent the intensity of its variation (either positive or negative) relative to a reference value.
  - Biological samples are represented as columns in the grid.
- Heatmap representations are also combined with clustering methods to group genes and/or samples based on their expression patterns.



## Step 4: Visual exploration of results

Hands on!

... follow **section 6.5 of the Rmd file**

## Next step...

- We have performed analysis on gene level
- It would be interesting to look for how these genes act together



Analysis of biological significance