



MSIB32500 Advanced Bioinformatics Spring 2018

RNAseq Data Analysis and Clinical Applications, Part II

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The University of Chicago



Outline

- ▶ Part I (05/19/2018)
 - ▶ Introduction to RNAseq technology and clinical applications
 - ▶ Hands on: From raw data to gene expression quantification
- ▶ Part II (05/26/2018)
 - ▶ Differential gene expression analysis and data visualization
 - ▶ Hands on: Identification of genes and pathways significantly changed under condition
 - ▶ **Homework assignment**
- ▶ Part III (06/02/2018)
 - ▶ How to associate gene expression data with clinical outcome
 - ▶ Hands on: Use gene expression data to discover tumor subtypes and survival analysis



Class materials

- GitHub

- <https://github.com/MScBiomedicalInformatics/MSIB32500>

- This lecture note contains the same contents as the notebook. In addition, the notebook also contains hands-on materials

- **lecture9.pdf**

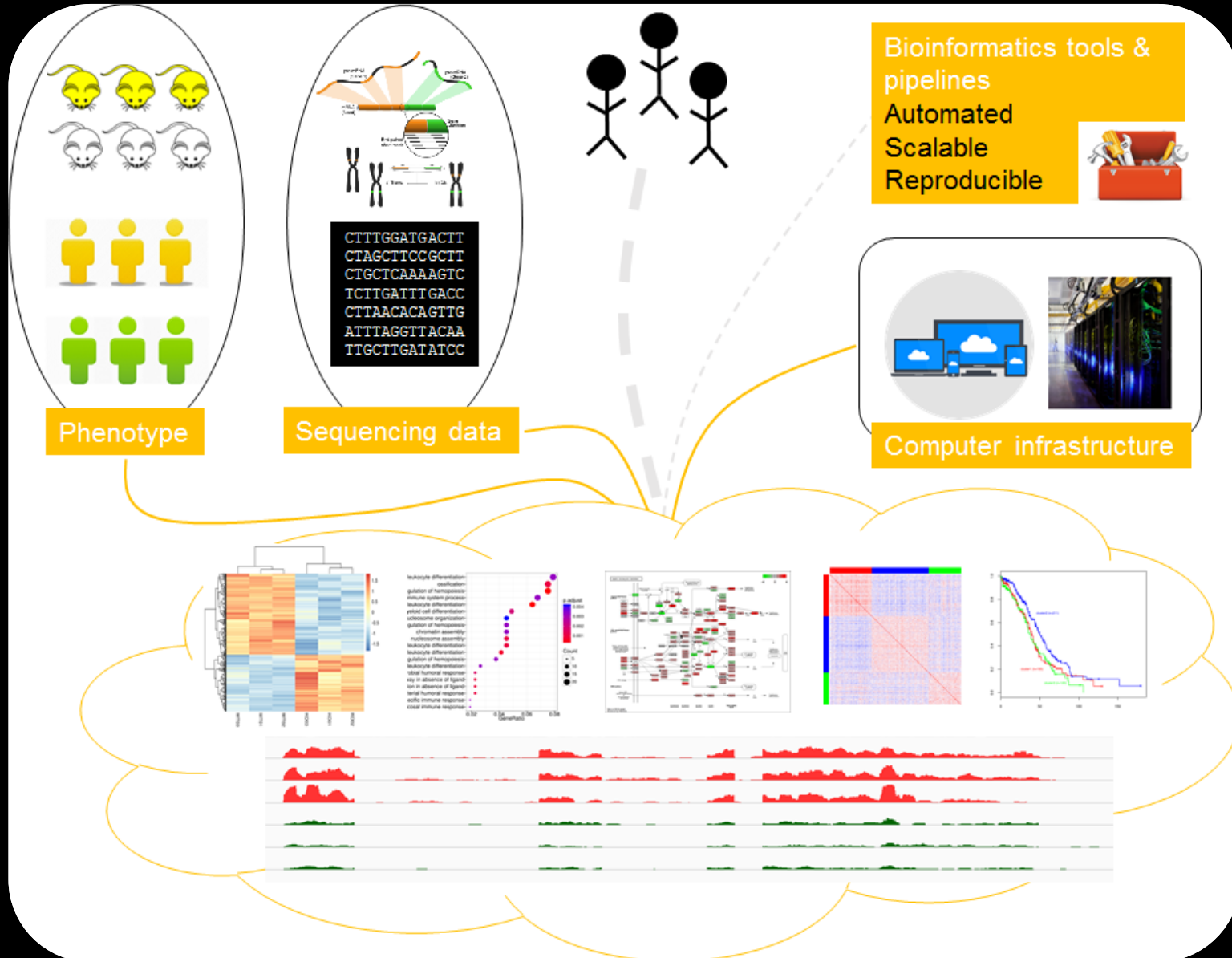
- **Handson9.Rmd**

- Rstudio (or R console) on personal computers (hands on practice)



Objective

- *(Recap from last class)*
- Detect genes differentially expressed between conditions
- Identify pathways / network enriched in genes of interest
- Generate high-quality figures for publication (PCA, heatmap, sample/gene cluster, GO/pathways, etc.)
- Become familiar with running commands in R / Rstudio

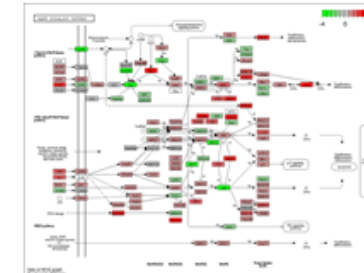
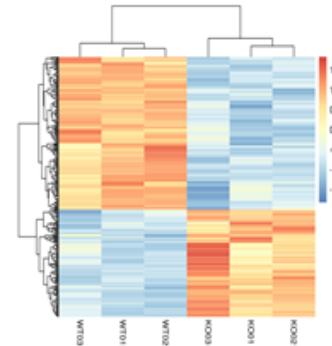
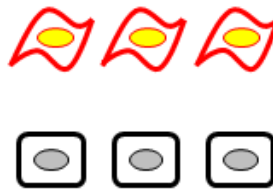


How to perform RNAseq analysis

The good-practice analysis protocol takes 8 major steps.

- **01-04:** From raw sequencing to transcript quantification
- **05-08:** DEG and pathway analysis (05/25, part II)

```
CTTTGGATGACTTCACA  
CTAGCTTCCGCTTTCTT  
CTGCTCAAAAGTCTTCA  
TCTTGATTTGACCAGTT  
CTTAACACAGTTGCATA  
ATTTAGGTTACAATTTA  
TTGCTTGATATCCACCA
```

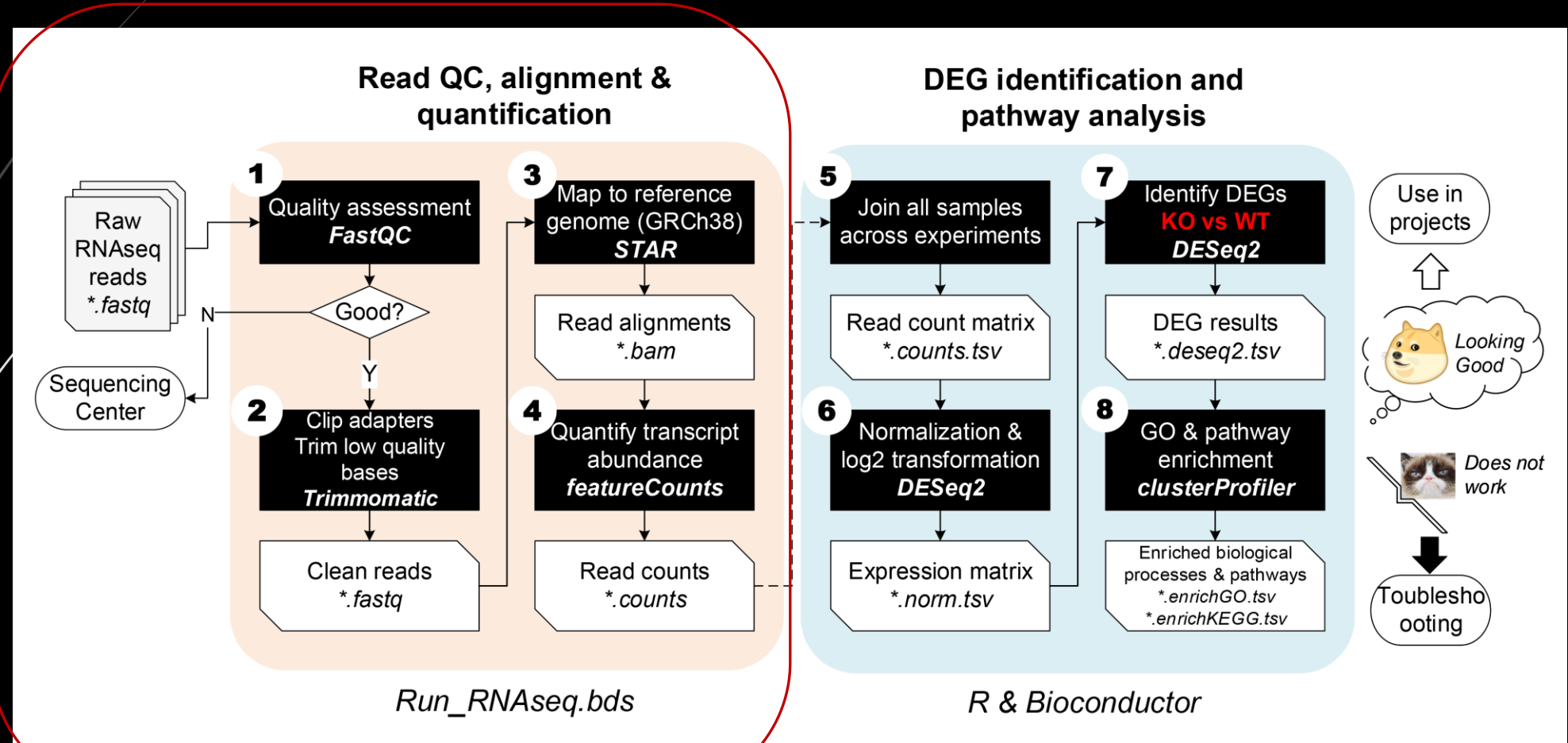


Raw sequencing data + sample group



Differentially expressed genes and pathways

How to perform RNAseq analysis



IGV (Integrative Genome Viewer)

<http://software.broadinstitute.org/software/igv/home>



- Load existing genomes, or generate custom genomes
- Visualize standard file formats
 - BAM
 - BED
 - GTF
 - ... and more!



Reference databases

- ▶ **Gene annotation database: GENCODE**
 - ▶ <https://www.gencodegenes.org/>
- ▶ **Ensembl database**
 - ▶ <https://www.ensembl.org/index.html>
- ▶ **UCSC Genome Browser**
 - ▶ <https://genome.ucsc.edu/>
- ▶ **NCBI databases**
 - ▶ <https://www.ncbi.nlm.nih.gov/guide/genomes-maps/>

[https://www.gencodegenes.org/
model organisms](https://www.gencodegenes.org/model%20organisms)



GENCODE

Data

Stats

DATA
STATISTICS



HUMAN

GENCODE 28 (05.04.18)

DATA
STATISTICS



MOUSE

GENCODE M17 (05.04.18)

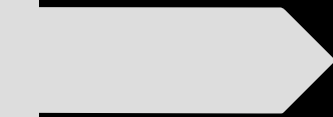
GTF / GFF3 files

Content	Regions	Description	Download
Comprehensive gene annotation	CHR	<ul style="list-style-type: none">It contains the comprehensive gene annotation on the reference chromosomes onlyThis is the main annotation file for most users	GTF GFF3
Comprehensive gene annotation	ALL	<ul style="list-style-type: none">It contains the comprehensive gene annotation on the reference chromosomes, scaffolds, assembly patches and alternate loci (haplotypes)This is a superset of the main annotation file	GTF GFF3
Comprehensive gene annotation	PRI	<ul style="list-style-type: none">It contains the comprehensive gene annotation on the primary assembly (chromosomes and scaffolds) sequence regionsThis is a superset of the main annotation file	GTF GFF3
Basic gene annotation	CHR	<ul style="list-style-type: none">It contains the basic gene annotation on the reference chromosomes onlyThis is a subset of the corresponding comprehensive annotation, including only those transcripts tagged as 'basic' in every gene	GTF GFF3
Basic gene annotation	ALL	<ul style="list-style-type: none">It contains the basic gene annotation on the reference chromosomes, scaffolds, assembly patches and alternate loci (haplotypes)This is a subset of the corresponding comprehensive annotation, including only those transcripts tagged as 'basic' in every gene	GTF GFF3
Long non-coding RNA gene annotation	CHR	<ul style="list-style-type: none">It contains the comprehensive gene annotation of lncRNA genes on the reference chromosomesThis is a subset of the main annotation file	GTF GFF3
PolyA feature annotation	CHR	<ul style="list-style-type: none">It contains the polyA features (polyA_signal, polyA_site, pseudo_polyA) manually annotated by HAVANA on the reference chromosomesThis dataset does not form part of the main annotation file	GTF GFF3
Consensus pseudogenes predicted by the Yale and UCSC pipelines	CHR	<ul style="list-style-type: none">2-way consensus (retrotransposed) pseudogenes predicted by the Yale and UCSC pipelines, but not by HAVANA, on the reference chromosomesThis dataset does not form part of the main annotation file	GTF GFF3
Predicted tRNA genes	CHR	<ul style="list-style-type: none">tRNA genes predicted by ENSEMBL on the reference chromosomes using tRNAscan-SEThis dataset does not form part of the main annotation file	GTF GFF3

Version 28 (November 2017 freeze, GRCh38) - Ensembl 92

General stats

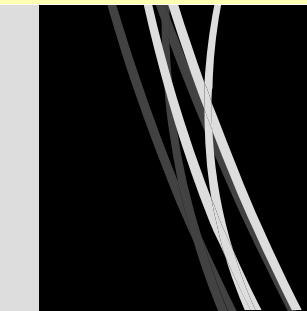
Total No of Genes	58381	Total No of Transcripts	203835
Protein-coding genes	19901	Protein-coding transcripts	82335
Long non-coding RNA genes	15779	- full length protein-coding:	56541
Small non-coding RNA genes	7569	- partial length protein-coding:	25794
Pseudogenes	14723	Nonsense mediated decay transcripts	14889
- processed pseudogenes:	10693	Long non-coding RNA loci transcripts	28468
- unprocessed pseudogenes:	3519		
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Immunoglobulin/T-cell receptor gene segments		Total No of distinct translations	61132
- protein coding segments:	408	Genes that have more than one distinct translations	13641
- pseudogenes:	237		

















Fasta files

Content	Regions	Description	Download
Transcript sequences	CHR	<ul style="list-style-type: none">Nucleotide sequences of all transcripts on the reference chromosomes	Fasta
Protein-coding transcript sequences	CHR	<ul style="list-style-type: none">Nucleotide sequences of coding transcripts on the reference chromosomesTranscript biotypes: protein_coding, nonsense_mediated_decay, non_stop_decay, IG_*_gene, TR_*_gene, polymorphic_pseudogene	Fasta
Protein-coding transcript translation sequences	CHR	<ul style="list-style-type: none">Amino acid sequences of coding transcript translations on the reference chromosomesTranscript biotypes: protein_coding, nonsense_mediated_decay, non_stop_decay, IG_*_gene, TR_*_gene, polymorphic_pseudogene	Fasta
Long non-coding RNA transcript sequences	CHR	<ul style="list-style-type: none">Nucleotide sequences of long non-coding RNA transcripts on the reference chromosomes	Fasta
Genome sequence (GRCh38.p12)	ALL	<ul style="list-style-type: none">Nucleotide sequence of the GRCh38.p12 genome assembly version on all regions, including reference chromosomes, scaffolds, assembly patches and haplotypesThe sequence region names are the same as in the GTF/GFF3 files	Fasta
Genome sequence, primary assembly (GRCh38)	PRI	<ul style="list-style-type: none">Nucleotide sequence of the GRCh38 primary genome assembly (chromosomes and scaffolds)The sequence region names are the same as in the GTF/GFF3 files	Fasta

- Genome-based alignment: STAR
- Transcriptome-based (pseudo)alignment: Kallisto, Salmon

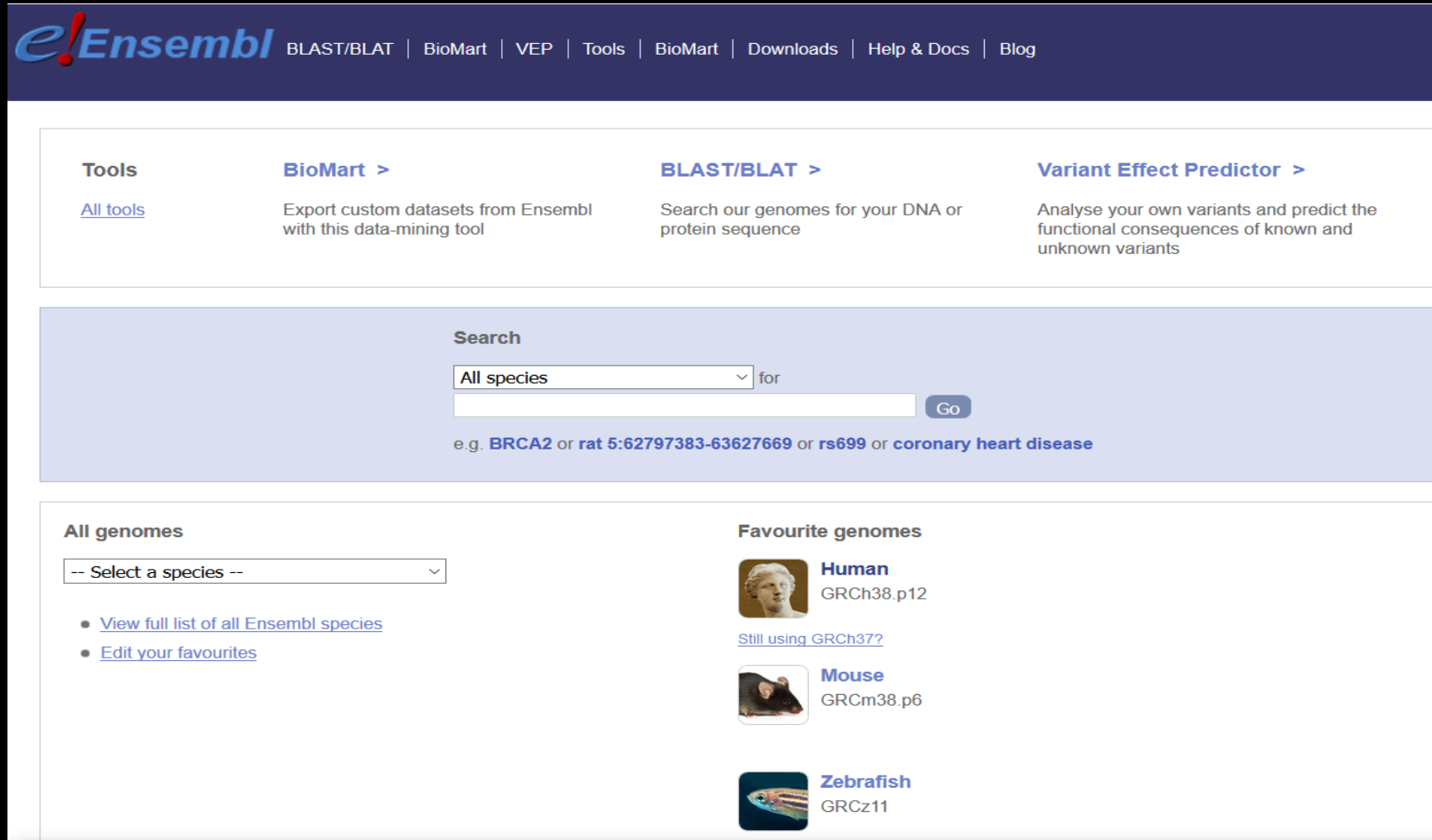


Metadata files

Content	Regions	Description	Download
Annotation remarks	ALL	<ul style="list-style-type: none">Remarks made during the manual annotation of the transcript	Metadata 
Entrez gene ids	ALL	<ul style="list-style-type: none">Entrez gene ids associated to GENCODE transcripts (from Ensembl xref pipeline)	Metadata 
Exon annotation evidence	ALL	<ul style="list-style-type: none">Piece of evidence used in the annotation of an exon (usually peptides, mRNAs, ESTs)	Metadata 
Gene source	ALL	<ul style="list-style-type: none">Source of the gene annotation (Ensembl, Havana, Ensembl-Havana merged model or imported in the case of small RNA and mitochondrial genes)	Metadata 
Gene symbol	ALL	<ul style="list-style-type: none">HGNC approved gene symbol (from Ensembl xref pipeline)	Metadata 
PDB id	ALL	<ul style="list-style-type: none">PDB entries associated to the transcript (from Ensembl xref pipeline)	Metadata 
PolyA features	ALL	<ul style="list-style-type: none">Manually annotated polyA features overlapping the transcript 3'-end	Metadata 
PubMed id	ALL	<ul style="list-style-type: none">Pubmed ids of publications associated to the transcript (from HGNC website)	Metadata 
RefSeq	ALL	<ul style="list-style-type: none">RefSeq RNA and/or protein associated to the transcript (from Ensembl xref pipeline)	Metadata 
Selenocysteine	ALL	<ul style="list-style-type: none">Amino acid position of a selenocysteine residue in the transcript	Metadata 
SwissProt	ALL	<ul style="list-style-type: none">UniProtKB/SwissProt entry associated to the transcript (from Ensembl xref pipeline)	Metadata 
Transcript source	ALL	<ul style="list-style-type: none">Source of the transcript annotation	Metadata 
Transcript annotation evidence	ALL	<ul style="list-style-type: none">Piece of evidence used in the annotation of the transcript	Metadata 
TrEMBL	ALL	<ul style="list-style-type: none">UniProtKB/TrEMBL entry associated to the transcript (from Ensembl xref pipeline)	Metadata 

ID conversion between different annotation databases (e.g. NCBI/RefSeq, Ensembl)

<https://www.ensembl.org/index.html>
all organisms

 [BLAST/BLAT](#) | [BioMart](#) | [VEP](#) | [Tools](#) | [BioMart](#) | [Downloads](#) | [Help & Docs](#) | [Blog](#)

Tools
[All tools](#)

BioMart >
Export custom datasets from Ensembl with this data-mining tool

BLAST/BLAT >
Search our genomes for your DNA or protein sequence

Variant Effect Predictor >
Analyse your own variants and predict the functional consequences of known and unknown variants

Search

All species ▾

for

Go


e.g. [BRCA2](#) or [rat 5:62797383-63627669](#) or [rs699](#) or [coronary heart disease](#)


All genomes


-- Select a species -- ▾

- [View full list of all Ensembl species](#)
- [Edit your favourites](#)

Favourite genomes

**Human**
GRCh38.p12
[Still using GRCh37?](#)

**Mouse**
GRCm38.p6

**Zebrafish**
GRCz11

Index of ftp://ftp.ensembl.org/pub/release-92/gtf/homo_sapiens/

 [Up to higher level directory](#)

Name	Size	Last Modified	
File: CHECKSUMS	1 KB	3/10/2018	10:08:00 PM
File: Homo_sapiens.GRCh38.92.abinitio.gtf.gz	3320 KB	3/9/2018	11:16:00 AM
File: Homo_sapiens.GRCh38.92.chr.gtf.gz	41909 KB	3/9/2018	11:07:00 AM
File: Homo_sapiens.GRCh38.92.chr_patch_hapl_scaff.gtf.gz	45818 KB	3/9/2018	11:12:00 AM
File: Homo_sapiens.GRCh38.92.gtf.gz	41916 KB	3/9/2018	11:07:00 AM
File: README	10 KB	3/9/2018	11:14:00 AM

Index of ftp://ftp.ensembl.org/pub/release-92/fasta/homo_sapiens/

 [Up to higher level directory](#)

Name	Size	Last Modified	
 cdna		3/10/2018	7:37:00 PM
 cds		3/10/2018	7:37:00 PM
 dna		3/10/2018	7:37:00 PM
 dna_index		3/10/2018	7:40:00 PM
 ncrna		3/10/2018	7:37:00 PM
 pep		3/10/2018	7:37:00 PM

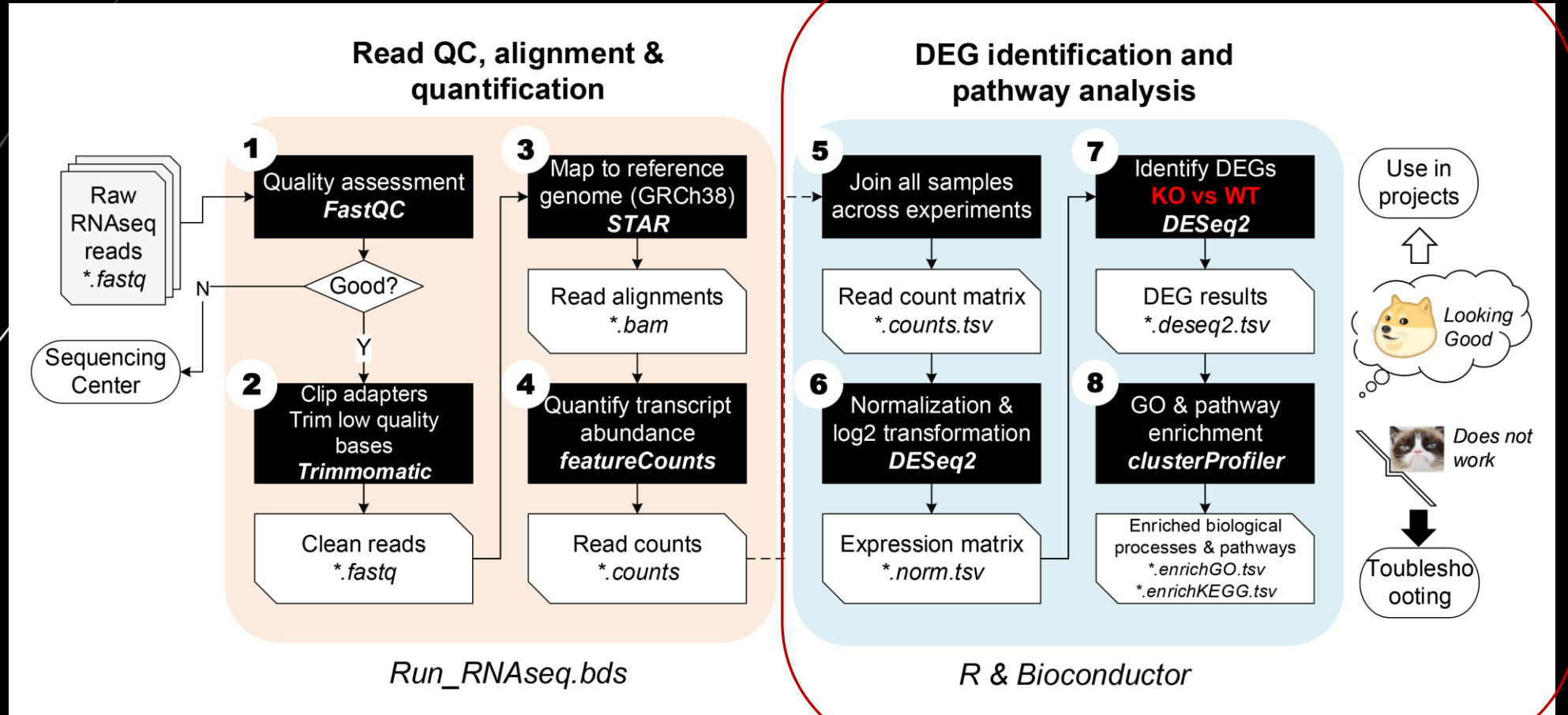
Version 28 (November 2017 freeze, GRCh38) - Ensembl 92

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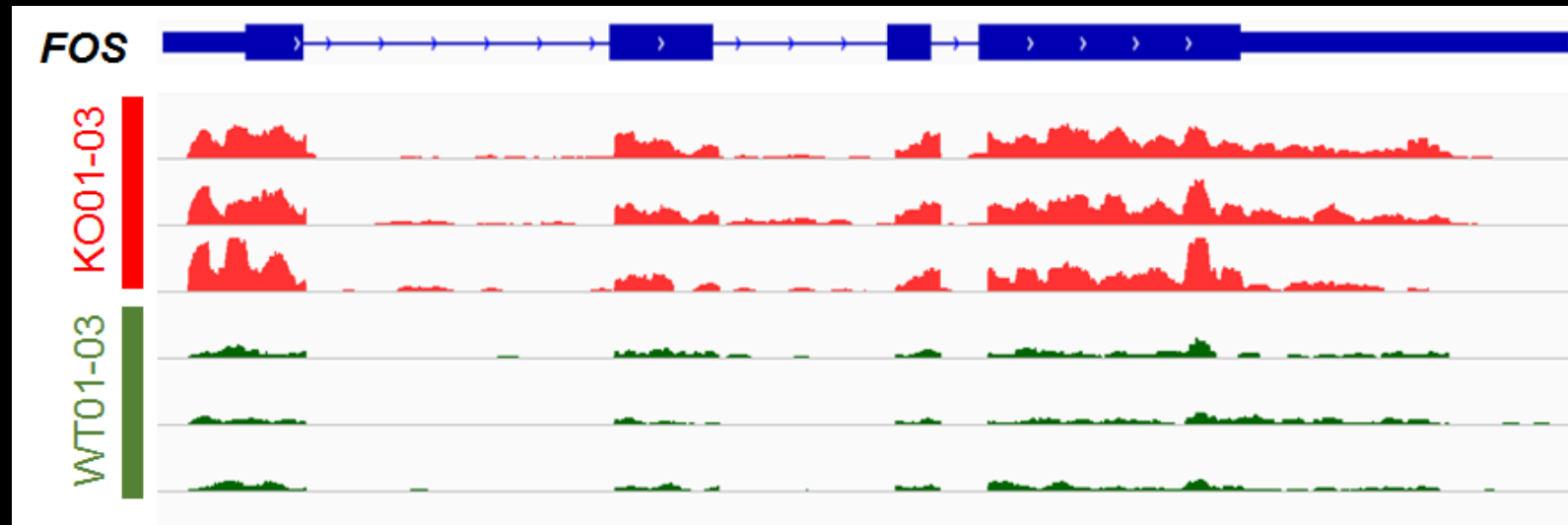
Gencode and Ensembl are generally in sync

How to perform RNAseq analysis



05-08: Identify differentially expressed genes and pathways: DESeq2, clusterProfiler

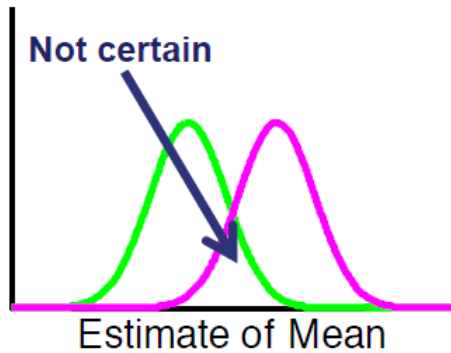
- After steps 01-04, we have generated read alignment and counts for every annotated gene on the genome
- The next step is to utilize the read counts data to detect DEGs
- For example, if we visualize *FOS* gene across 6 samples in genome browser



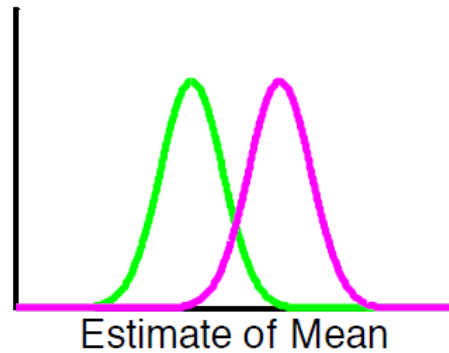
FOS = *Fos* proto-oncogene, AP-1 transcription factor subunit

DEG detection

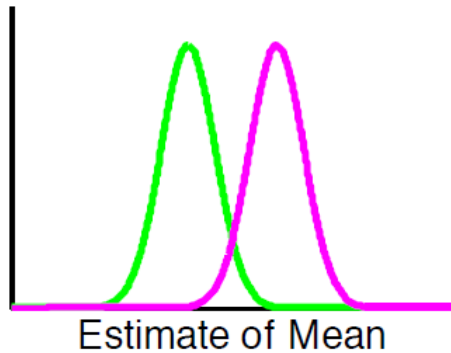
2 Replicates



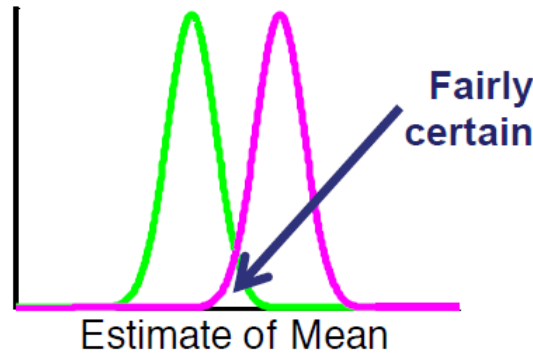
3 Replicates



4 Replicates



5 Replicates



$$SE = \frac{\sigma}{\sqrt{n}}$$

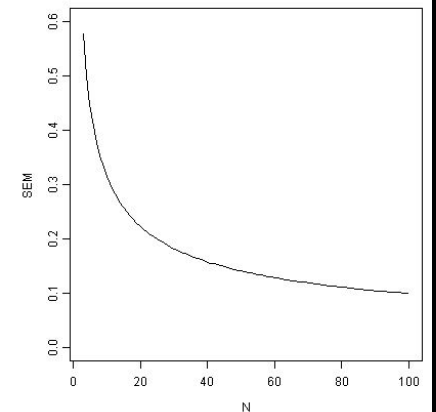
← Standard deviation

← Number of samples

Standard Error of the Mean

$$SE_{M_x} = \frac{\sigma}{\sqrt{N}}$$

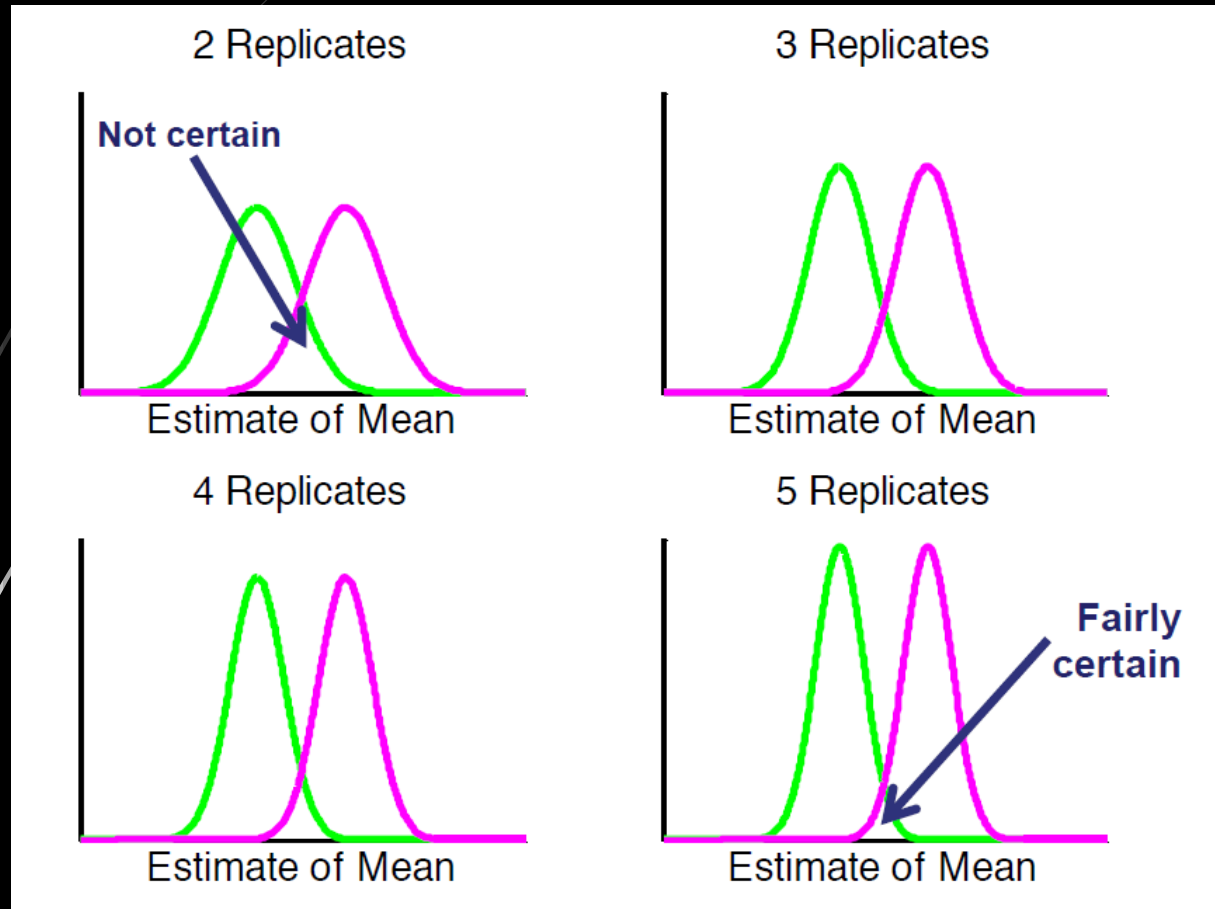
- This equation implies that sampling error decreases as sample size increases.
- This is important because it suggests that if we want to make sampling error as small as possible, we need to use as large of a sample size as we can manage.



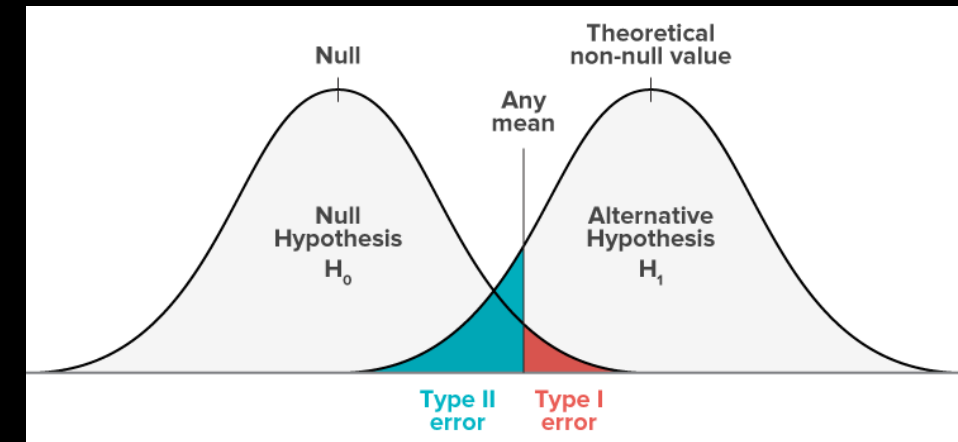
More biological replicates per group lead to higher discovery power, sensitivity and specificity.

DEG detection

Sensitivity: $TP/(TP+FN)$ Specificity: $TN/(FP+TN)$

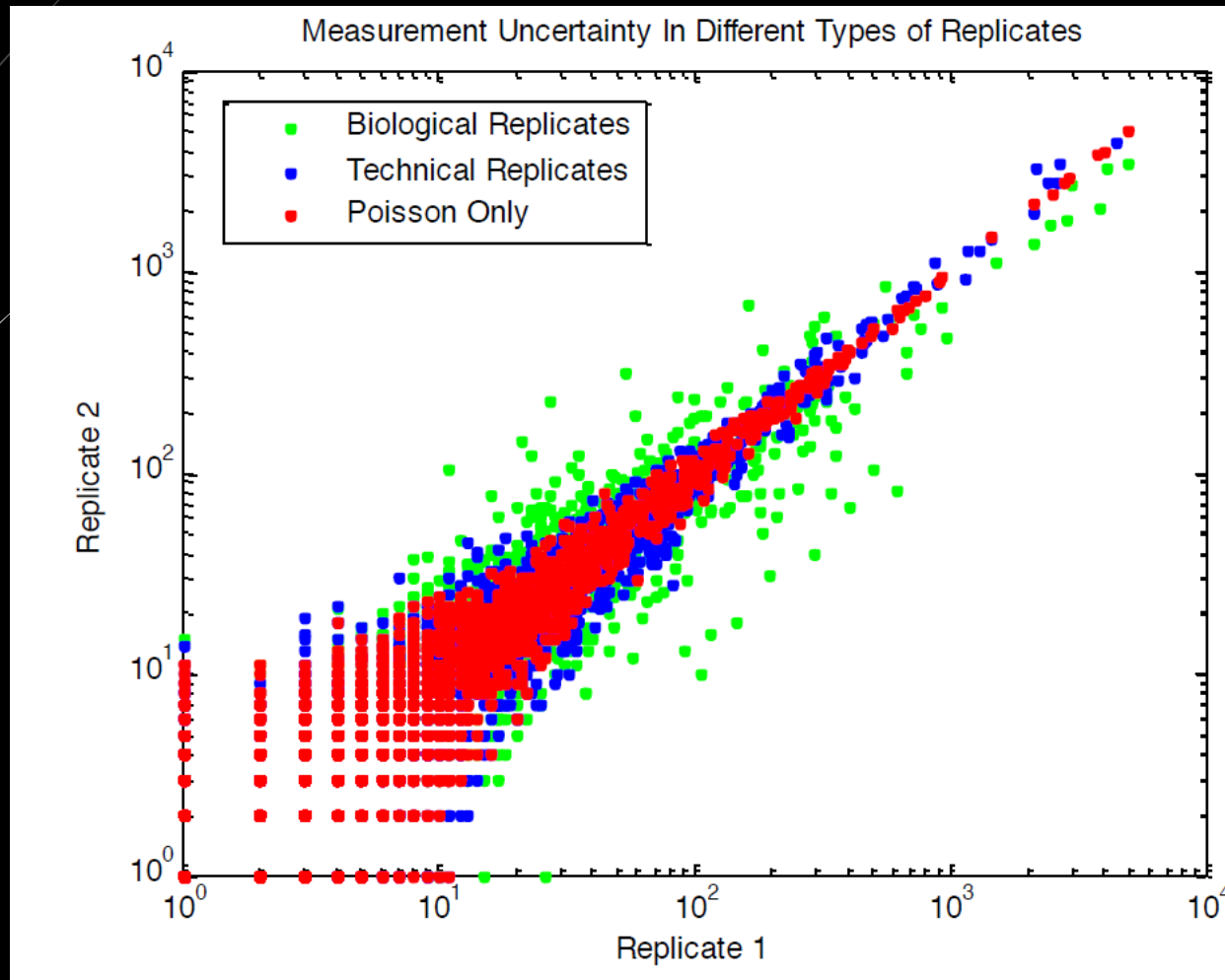


		Reality	
		Positive	Negative
Study Finding	Positive	True Positive (Power) $(1-\beta)$	FP Type I Error (α)
	Negative	FN Type II Error (β)	True Negative



More biological replicates per group lead to higher discovery power, sensitivity and specificity.

DEG detection

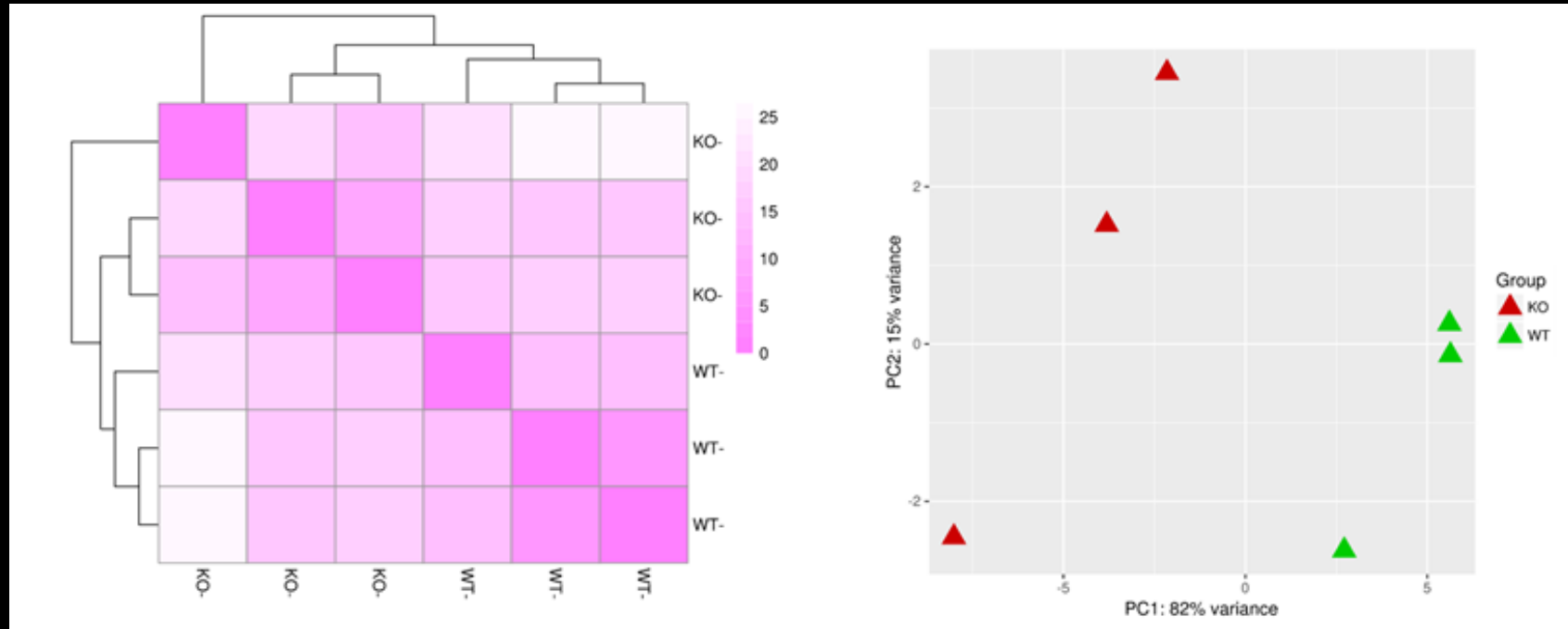


High correlation is expected between biological replicates.

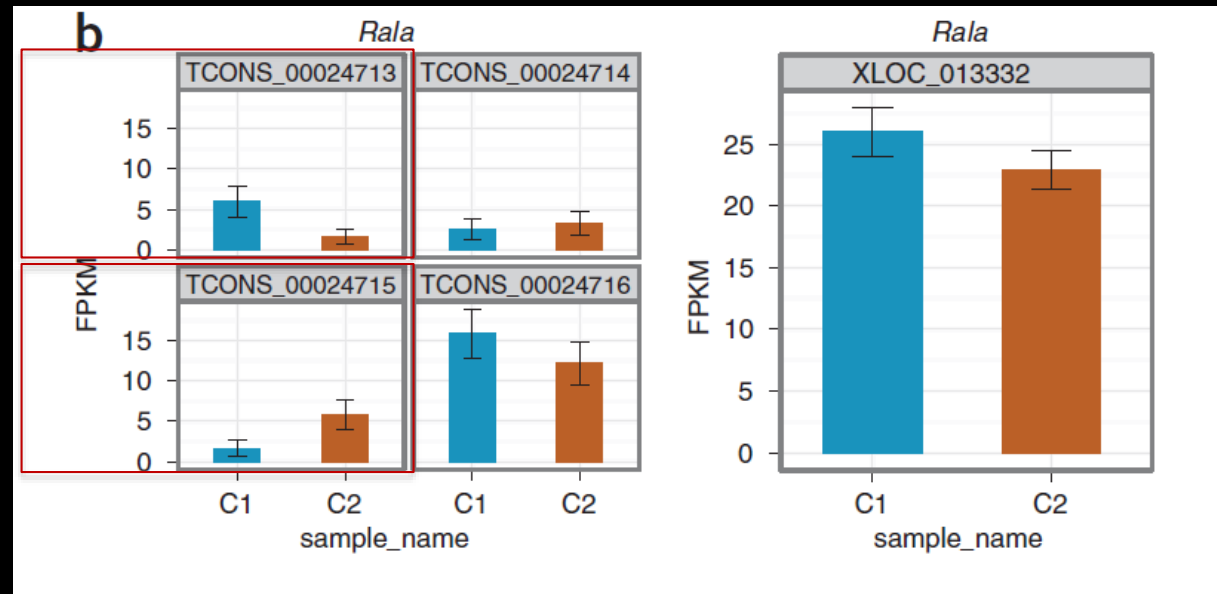
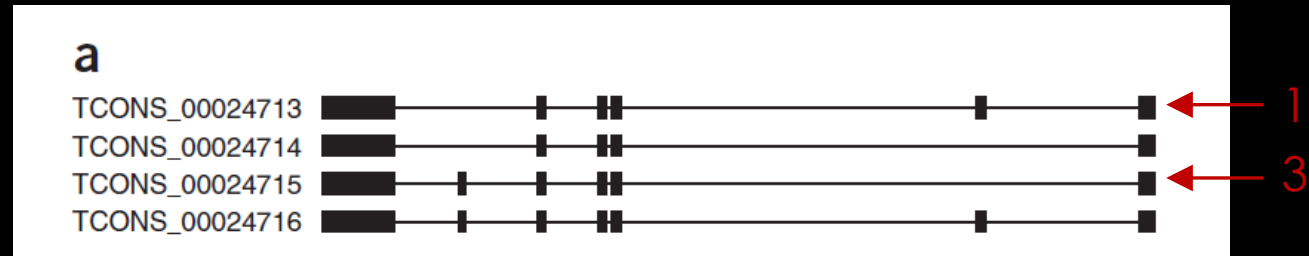
If one sample is an outlier, it can be identified if multiple replicates are included in an experiment.

How to identify an outlier?

- PCA plot (visualization)
- Unsupervised sample clustering based on all genes or *top variable genes* (e.g. 1500)



Transcript vs gene level quantification



Isoform 1 *

Isoform 3 *

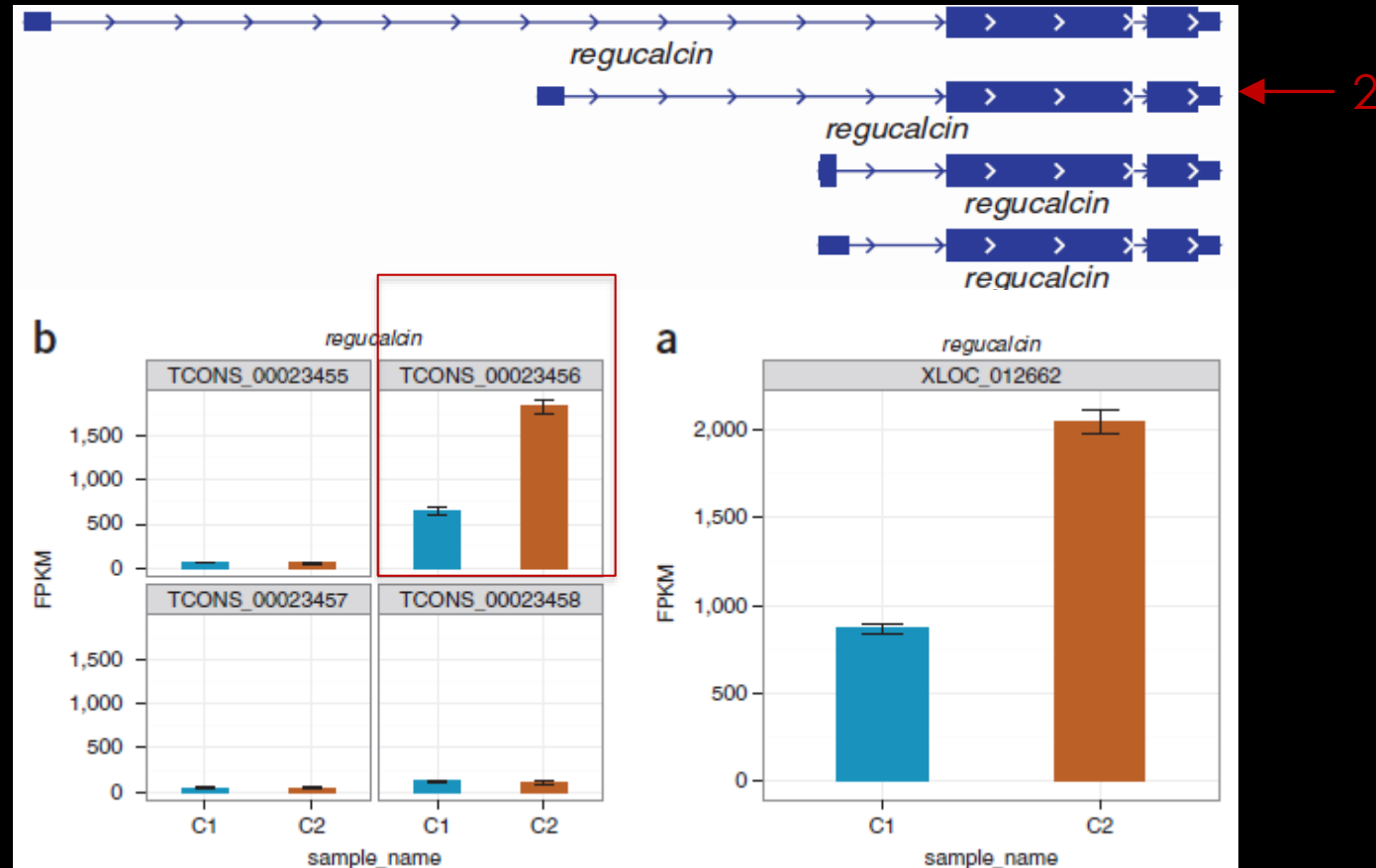
Isoforms 1 – 4

Gene

Difference in gene-level expression is not significant due to variability of isoforms

Transcript vs gene level quantification

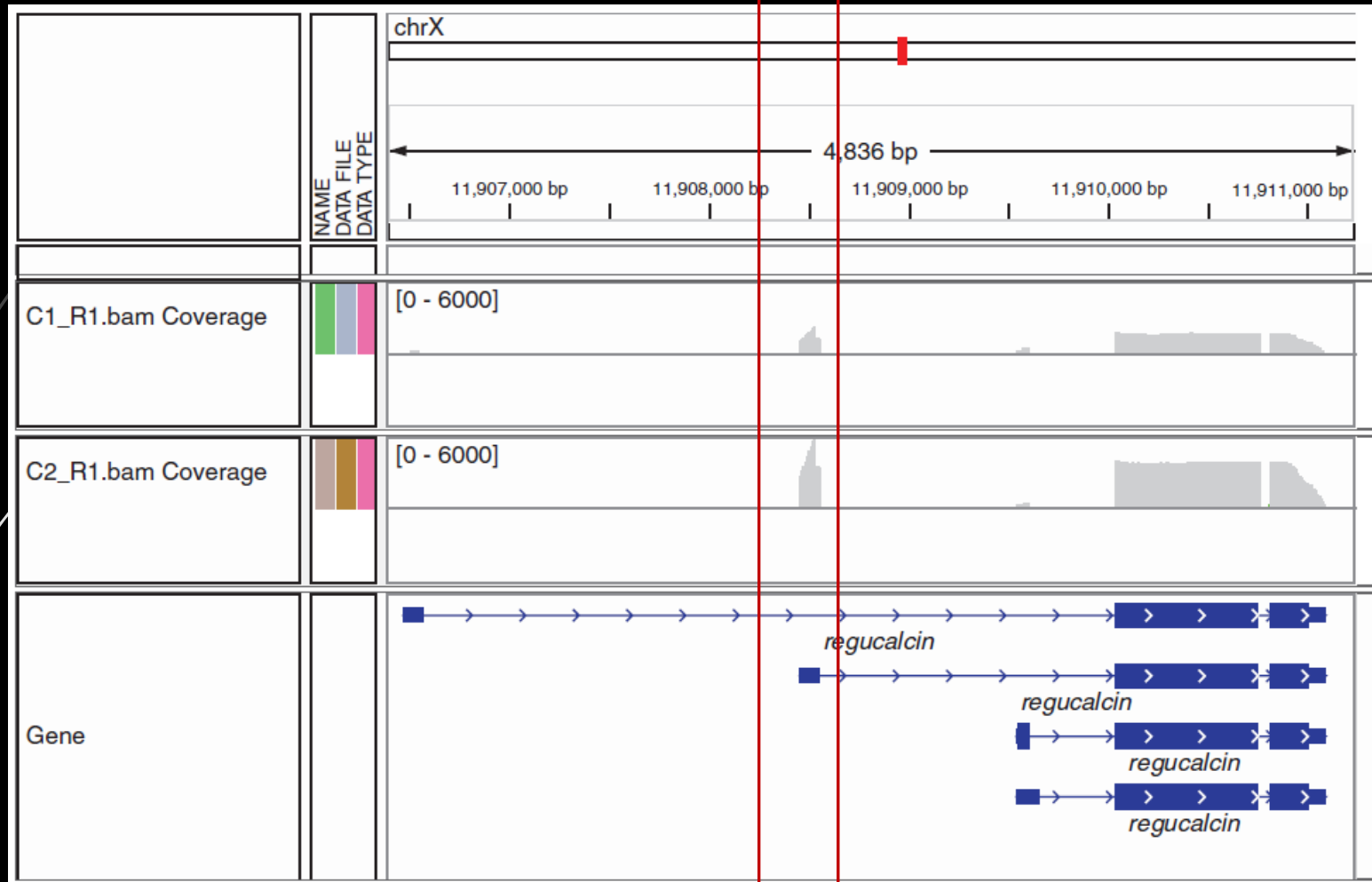
Isoform 2 *



Isoforms 1 – 4

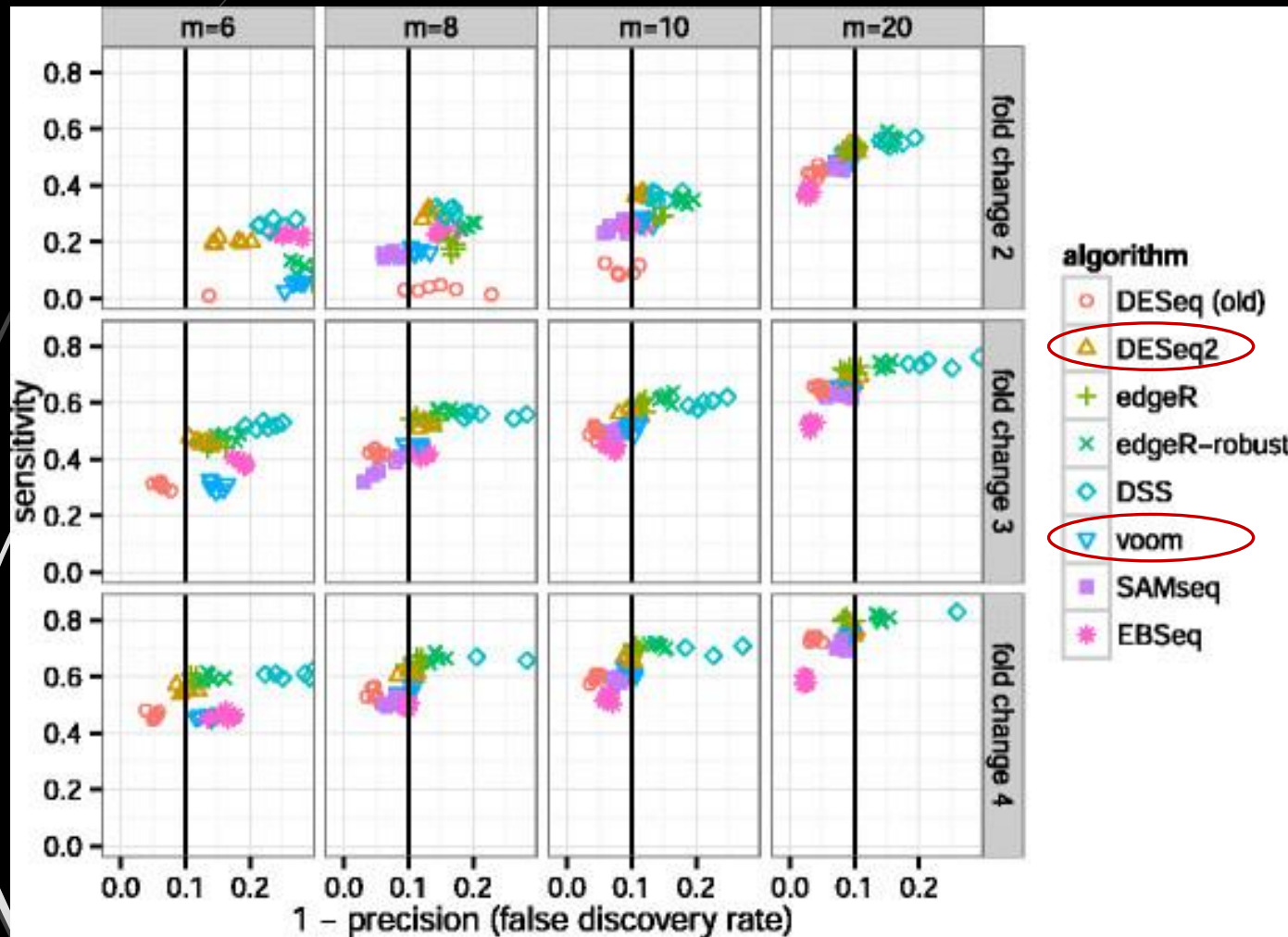
Gene

Difference in gene-level expression is significant, which is largely due to a great increase in the expression of isoform 2



Difference in gene-level expression is significant, which is largely due to a great increase in the expression of isoform 2

Comparison of different DEG identification methods



Sensitivity and precision of algorithms across combinations of sample size and effect size.

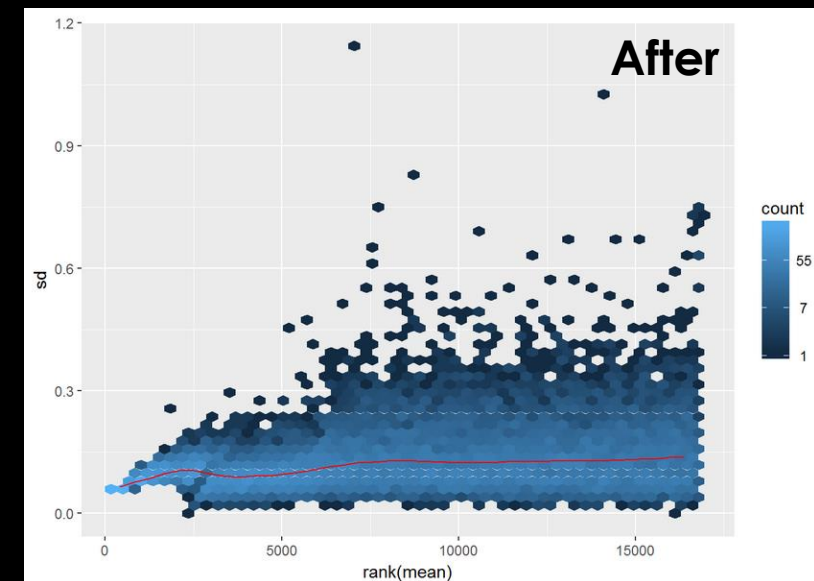
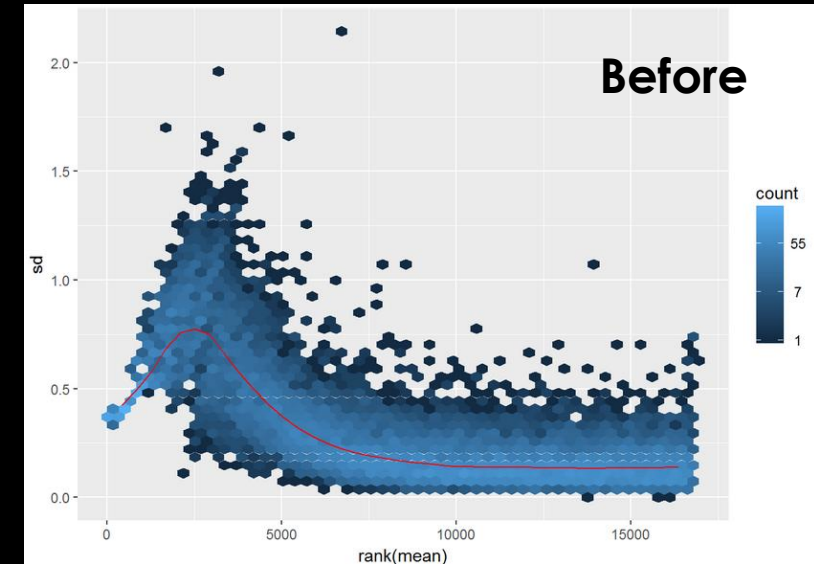
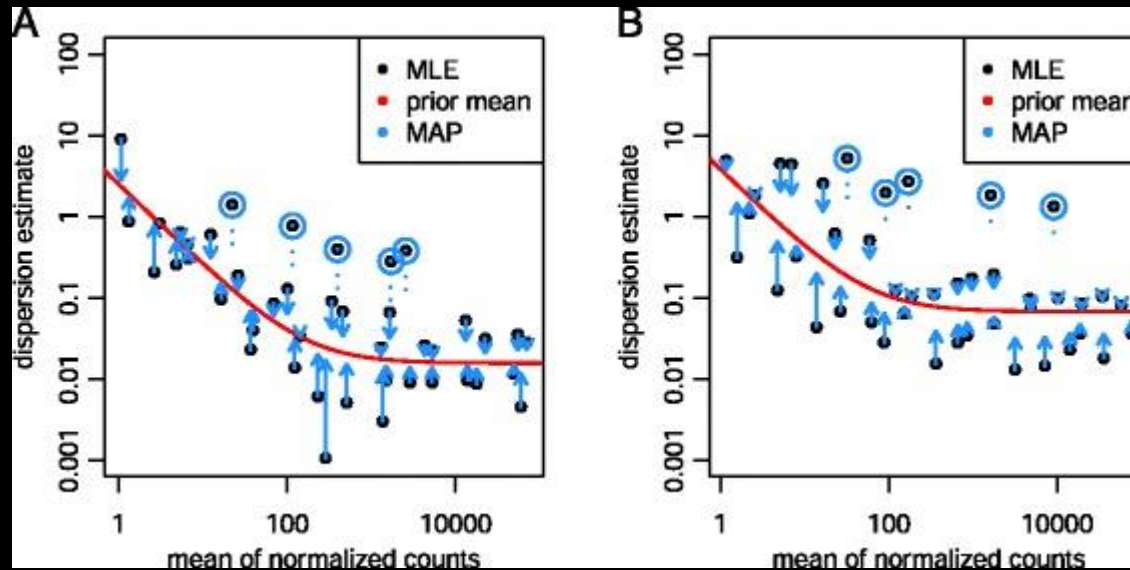
DESeq2 and edgeR often had the highest sensitivity of those algorithms that controlled the FDR, i.e., those algorithms which fall on or to the left of the vertical black line.

m: total sample size; split into two even-sized groups for comparison

DESeq2

- Count matrix data
- Assume data follow negative binomial distribution (sometimes also called a gamma-Poisson distribution) with mean (μ) and dispersion (α) parameters
- Within-group variability, i.e., the variability between replicates, is modeled by the dispersion parameter alpha, which describes the *variance* of counts
- Empirical Bayes shrinkage for dispersion estimation

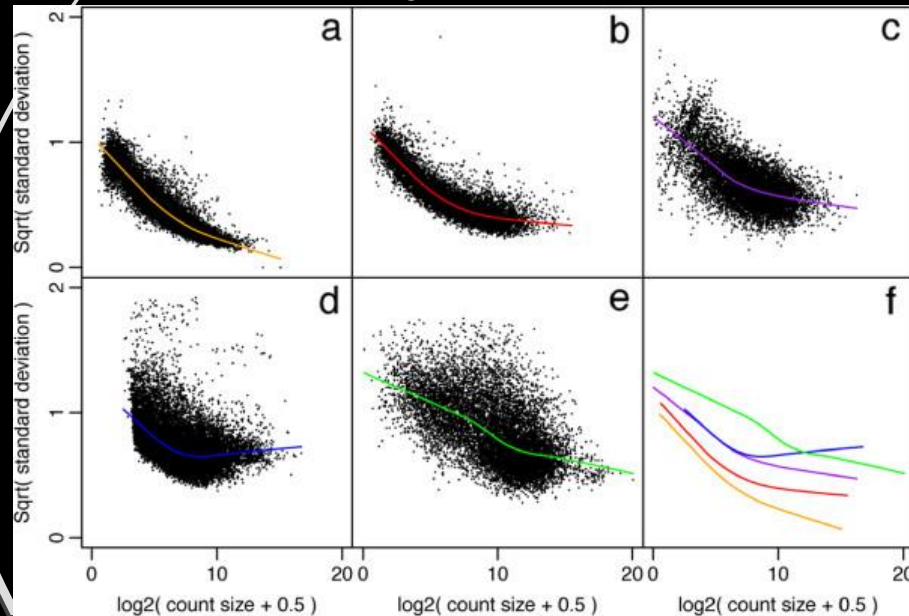
MAP, maximum a posteriori; MLE, maximum-likelihood estimate



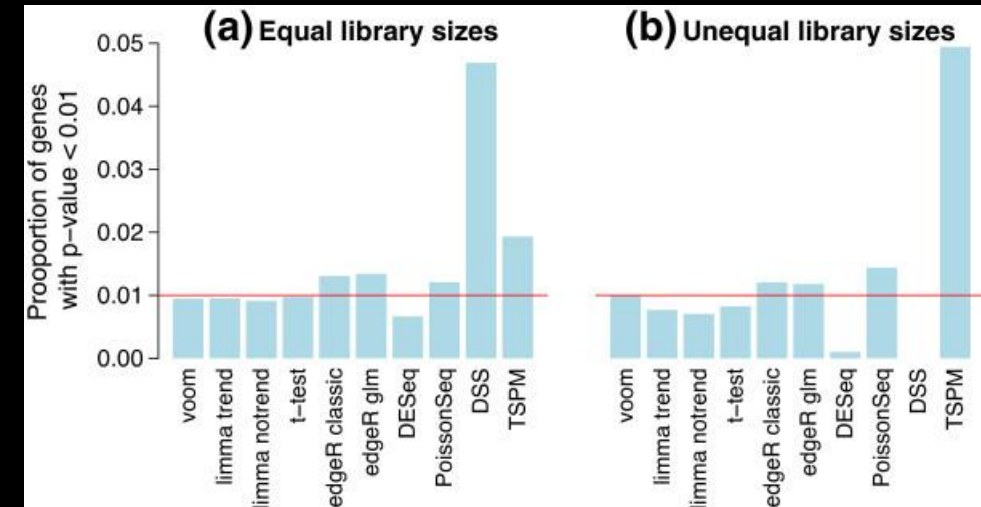
Limma voom (weighted algorithm)

- To model the mean-variance relationship than to specify the exact probabilistic distribution of the counts (e.g. NB or Poisson)
- Provide accurate Type I (alpha) and Type II error (beta) control compared to other methods, especially when sample size is small
- *Voom with sample quality weights*

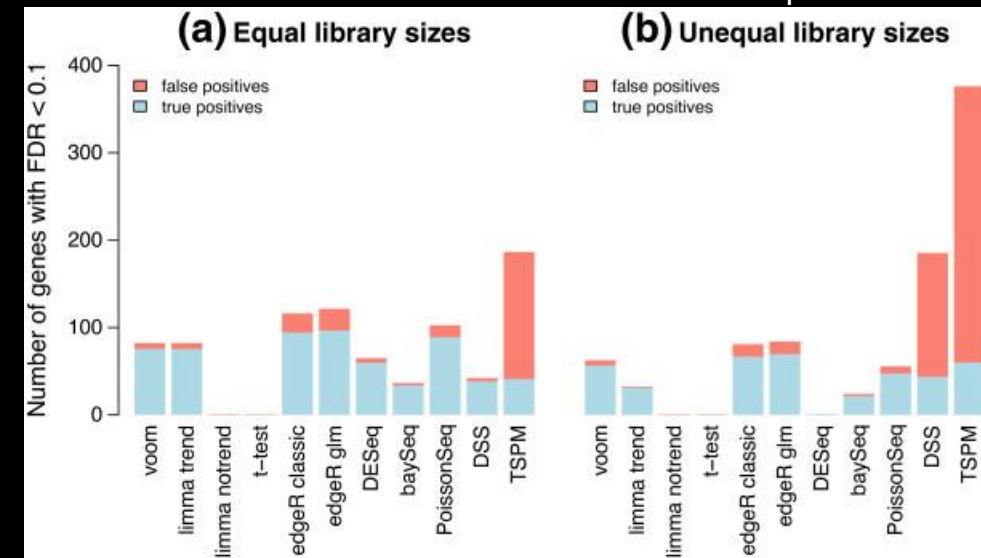
Law et al., Genome Biology 2014



Type I error rates in the absence of true differential expression



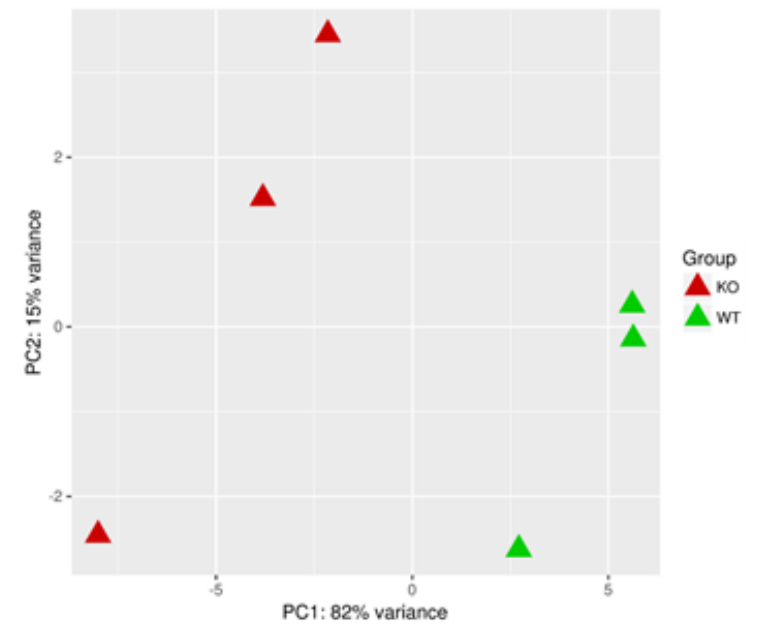
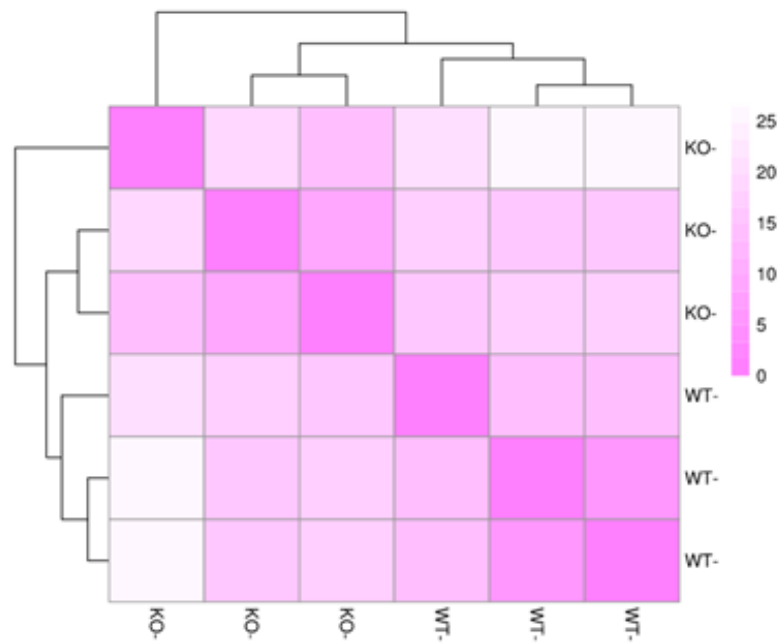
Power to detect true differential expression



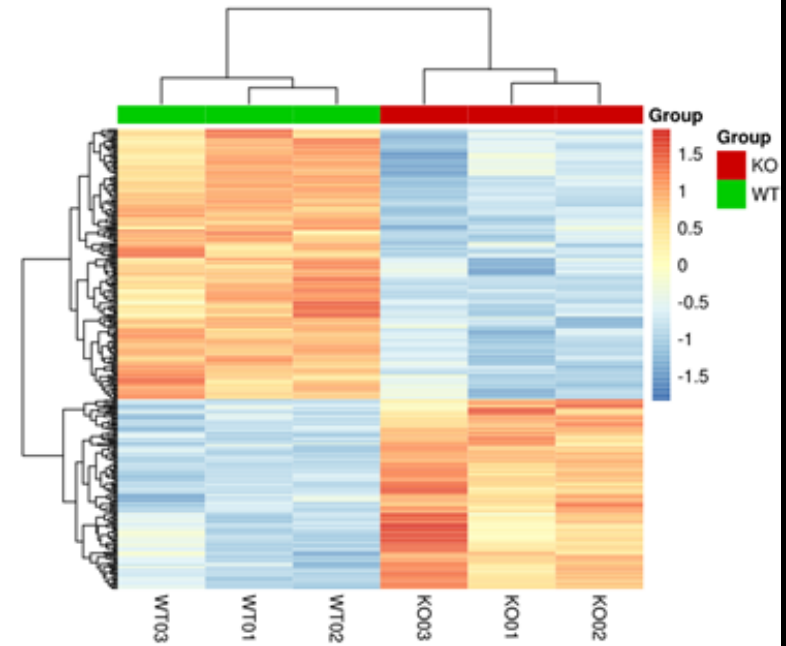
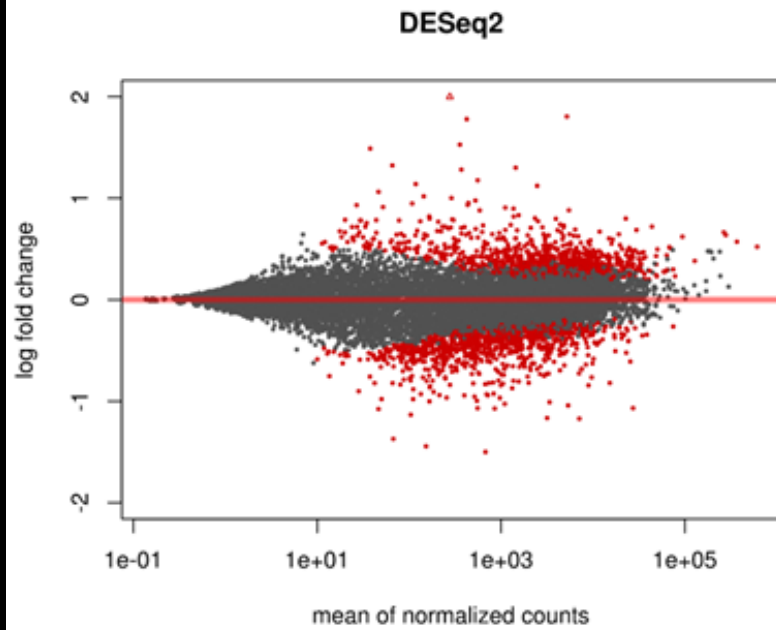


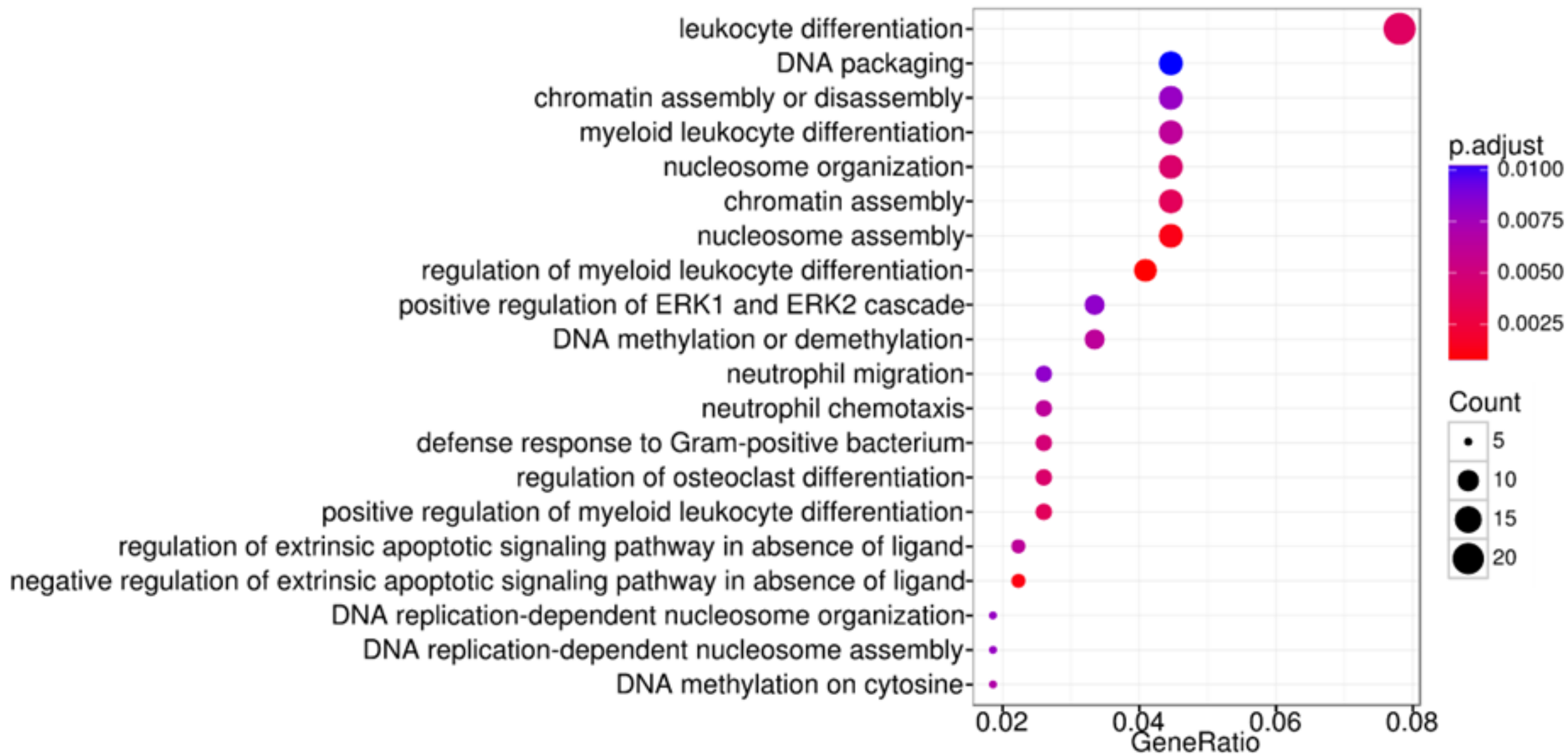
More databases!

- ▶ Gene annotation database: GENCODE
 - ▶ <https://www.gencodegenes.org/>
- ▶ Gene Ontology (GO) database: Gene Ontology Consortium
 - ▶ <http://www.geneontology.org/>
- ▶ Pathway database: KEGG
 - ▶ <http://www.genome.jp/kegg/>
- ▶ Predefined gene sets: MSigDB
 - ▶ <http://software.broadinstitute.org/gsea/msigdb/>

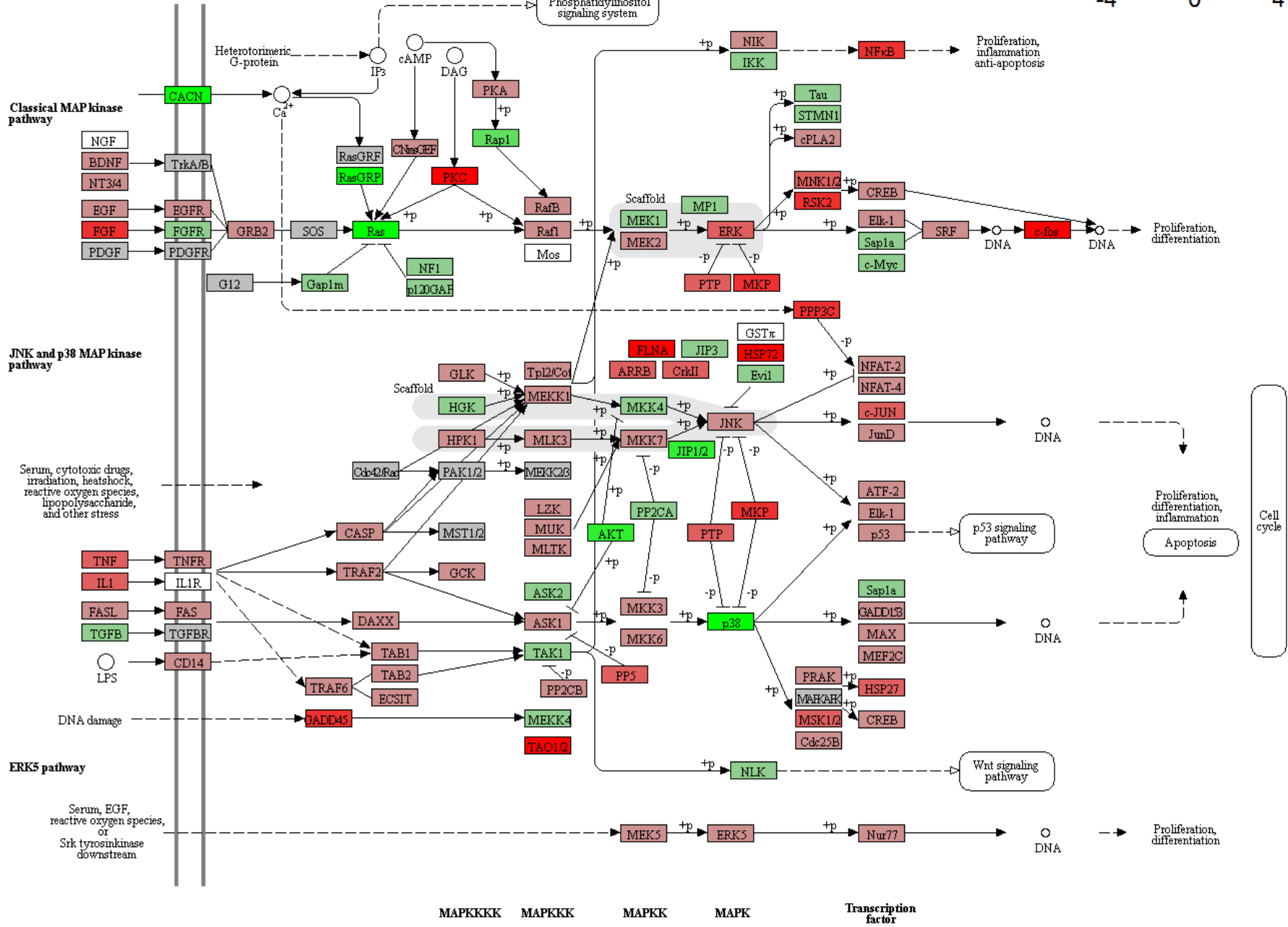
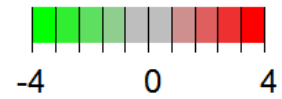


[1] "Genes significant = 296 (fc, 1.5, fdr 0.05)"
 [1] "Heatmap = 296 genes on the row, 6 samples on the column"

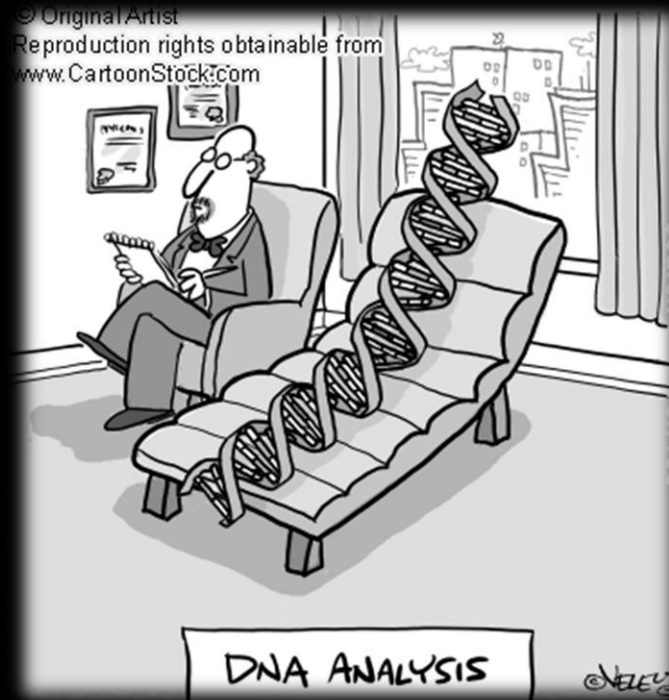




MAPK SIGNALING PATHWAY



Thank you!



Questions



Hands-on practice START

- Open your handson.Rmd on the Github or download to local computer
- <https://github.com/MScBiomedicalInformatics/MSIB32500/raw/master/lectures/handson8.html>
- Dataset: two groups (PRDM11 KO vs WT, human U2932 cells), 6 samples
- Single-end reads, unstranded libraries

Sample	Group	Sequencing File	Sequencing Data
KO01	KO	KO01.fastq.gz	74,126,025 reads
KO02	KO	KO02.fastq.gz	64,695,948 reads
KO03	KO	KO03.fastq.gz	52,972,573 reads
WT01	WT	WT01.fastq.gz	55,005,729 reads
WT01	WT	WT02.fastq.gz	61,079,377 reads
WT01	WT	WT03.fastq.gz	66,517,156 reads

Fog. et al. 2015. Loss of *PRDM11* promotes MYC-driven lymphomagenesis. Blood 125(8):1272-81

PRDM11 = PR/SET domain 11

