# Design of RNA-seq Experiments and Differential Expression Analysis

Genomic Technologies Workshop (PLPTH885)

Sanzhen Liu Plant Pathology

6/5/2024

# Schedule

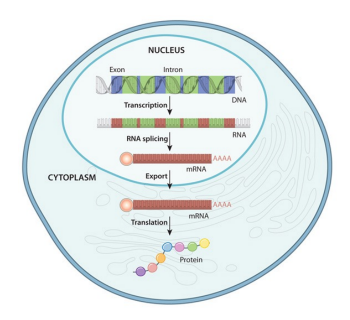
- 1:00 am Lecture DE
  Design of RNA-Seq Experiments and Differential Expression Analysis
- 2:00 am Break
- 2:10 am **Computer Lab R** Introduction to R programming
- 2:00 am **Break**
- 3:00 4:30pm Computer Lab DE RNA-Seq data analysis using R

# Outline

- RNA-seq procedure
- Experimental design
- Multiple testing correction
- Data visualization

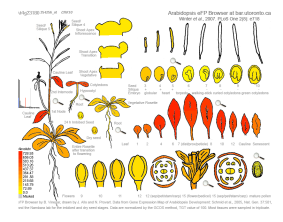
#### Gene expression

#### What is the expression level of a gene?



#### DNA to protein in eukaryote

nature.com/scitable/topicpage/gene-expression-14121669



#### Expression profiles in different tissues

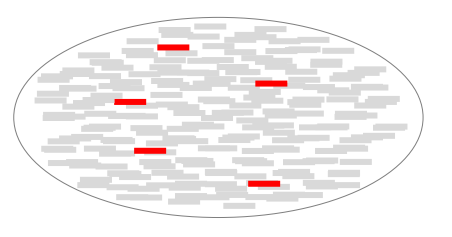


#### Adaptation to environmental change



Response to biotic stress

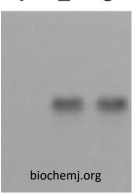
#### Approaches for quantification of gene expression



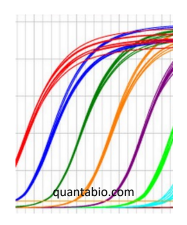
How can we measure the accumulative level of transcripts of a given gene in millions/billions of transcripts?

Northern blot

1 2 3



qRT-PCR

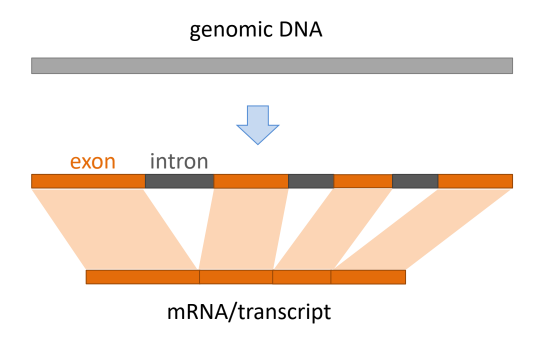


microarray



RNA-seq

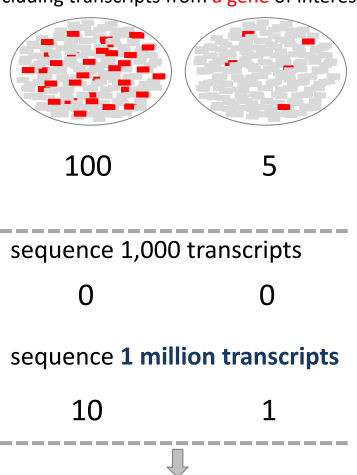
#### Rationale of RNA-seq (mRNA sequencing)



Essentially, RNA-seq is designed to measure mRNA accumulation levels of genes by

- 1) recognizing transcripts based on sequences
- 2) and quantifying transcripts of each gene

10 millions of transcripts in each sample Including transcripts from a gene of interest



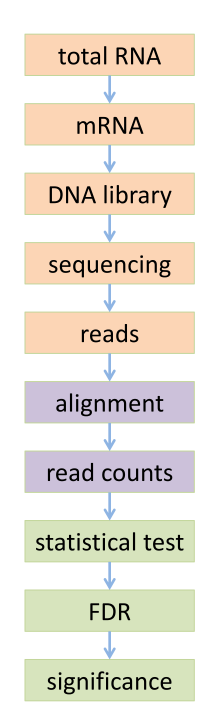
Differential expression (DE)?

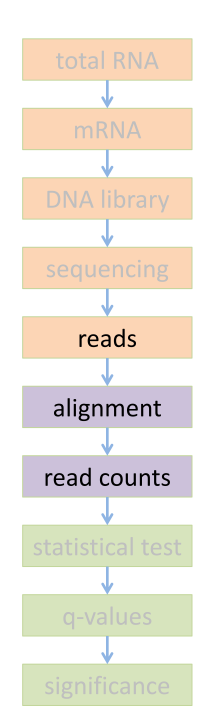
# RNA-Seq procedure

RNA to sequencing reads

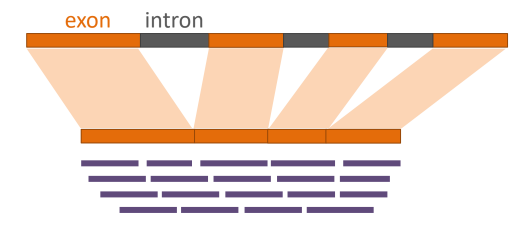
reads to read counts per gene

read counts to significant genes





#### Reads to read counts per gene



2. alignment to the reference genome (DNA sequence)



An **intron-aware** aligner is important for RNA-seq reads alignment e.g., STAR, HiSAT2

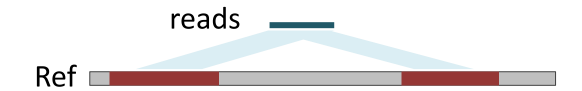
#### 3. read counts

1. reads

N = 19 if all reads can be confidently mapped to the reference genome

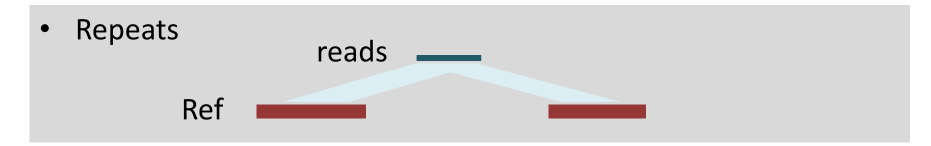
#### Alignment issues

Repeats

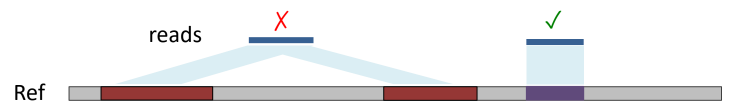


- Sequencing errors
- Polymorphisms (reference and sequenced individuals)
- Quality of reference genomes (mis-assembly and incomplete genome)

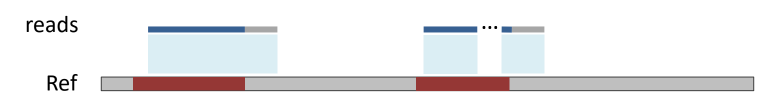
# Solutions to mitigate problems - I



Unique mapped reads



Longer reads or Paired-end reads



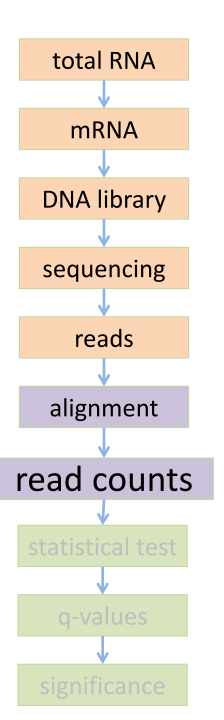
#### Solutions to mitigate problems

- Sequencing errors
- Polymorphisms (reference and sequenced individuals)

Tolerance of mismatches or gaps for each alignment



- Quality of reference genomes (mis-assembly and incomplete genome)
- Better reference genome

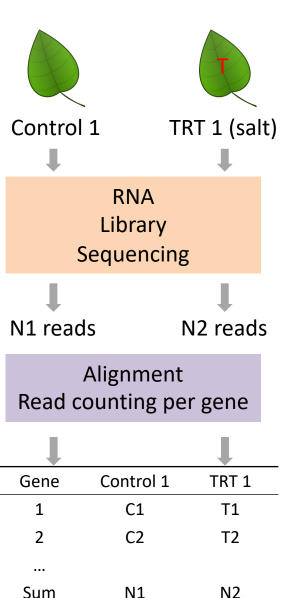


# Count matrix Read counts (Raw) per gene

Gene	sample 1	sample 2
gene 1	6,075	5,934
gene 2	295	377
•••	•••	•••

# total RNA **DNA** library read counts statistical test q-values significance

#### Read counts to significant genes



#### 2x2 Table for Gene 1

	Gene 1	Others
Control 1	C1	N1 – C1
TRT 1	T1	N2 – T1

 Fisher's Exact Test or χ² test on Gene 1

A p-value for Gene 1

- Repeat on all the genes
   p-values
- Multiple testing correction q-values
- Declaration of significance
   a significant gene set

#### Statistical test for differential expression

- Statistical test to discover differential expression (DE)
  - Count data: Generalized Linear Model (GLM) to deal with count data
  - e.g., Poisson GLM could handle count data but overdispersion exits
  - Overdispersion issue: Using negative binomial GLM to incorporate a dispersion parameter into the model

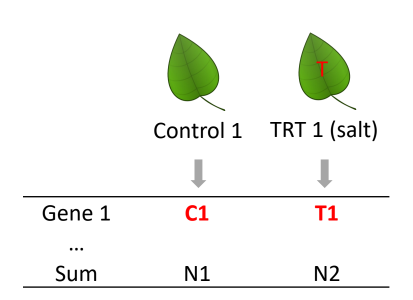
edgeR (Robinson and Smyth, 2007), **DESeq** (Anders and Huber, 2010), NBPSeq (Di et al., 2011), and QuasiSeq (Lund 2012)

Conesa et al. Genome Biology (2016) 17:13 DOI 10.1186/s13059-016-0881-8

Genome Biology

A survey of best practices for RNA-seq data analysis

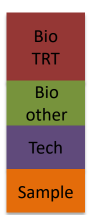
#### Source of variance in counts



Our interest: the effect of the salt treatment on gene expression **Question**: what could cause the difference between two values, C1 and T1?

- Treatment effect
- Plant difference
- RNA quality
- Library preparation
- Sequencing
- Sampling





# Sampling variance

 Sampling variance derived from the inherent nature of counting experiments

total molecules: 109

gene X: 1000 molecules

Randomly sample 10<sup>7</sup>

First sampling	6
Second sampling	13
Third sampling	8

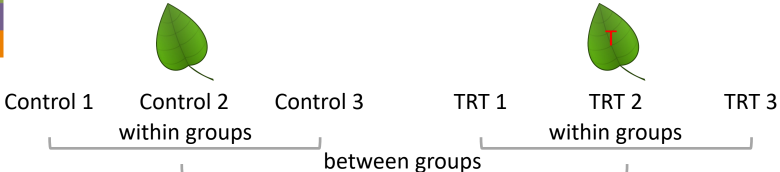
#### Randomly sample 108

First sampling	102
Second sampling	93
Third sampling	97

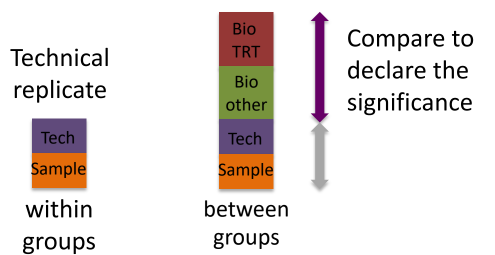
Sequence depth (sampling number) matters.



#### Technical replication



Technical replication
refers to the
sequencing of multiple
libraries derived from
the same biological
sample.



Spurious power

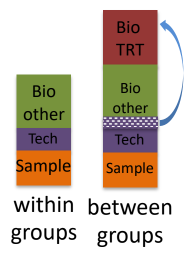
#### Question

Goal: to identify the DEs between two biological groups

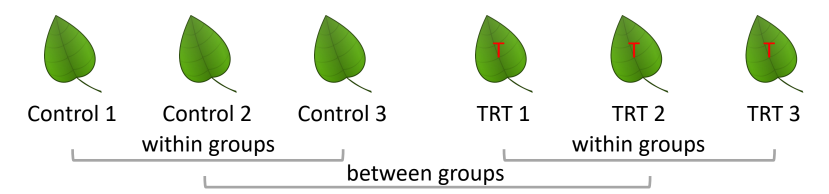
Design: Each group has five biological replicates

To avoid messing up samples across groups, the experiment of each group was conducted separately.

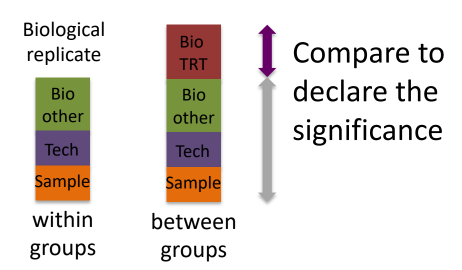
Is this a sound experimental design? Why?



#### Biological replication



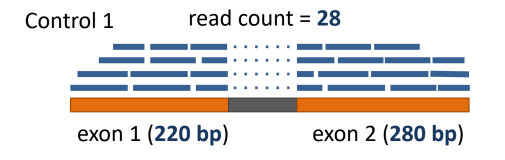
Biological replication refers to the sequencing of multiple libraries derived from different biological samples.



- 1. Use *biological replication* instead of technical replication unless you have your own interest.
- 2. More replicates increase the power to detect small effect

#### A normalization method: RPKM and FPKM

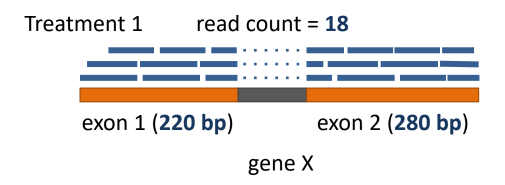
RPKM: Read number per kilobase of exons per million of total reads



total reads: 15 millions of total reads

RPKM of 
$$X =$$

= 3.7



total reads: 10 millions of total reads

RPKM of 
$$X =$$

?

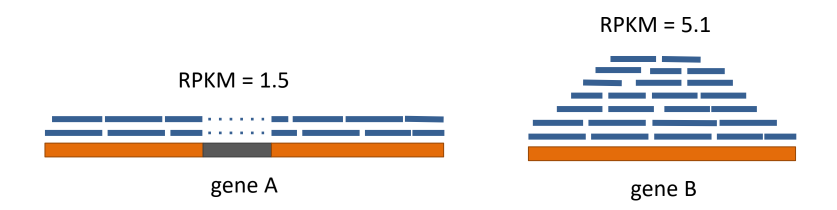
= 3.6

FPKM: Fragment number per kilobase per million of total reads.

Fragment = one pair of paired-end reads or one single-end read



#### More about RPKM



Can we say that the gene B has higher expression than the gene A?

- RPKM is not an ideal indicator to compare the expression/accumulation levels between two genes
  - 1. amplification bias
  - 2. alignment efficiency

#### **Experimental Design**

#### Sequencing depth

Increasing sequencing depth decreases sampling variance relative to the mean

#### Biological replication

A reasonable number of biological replication helps accurately estimate variances to achieve reliable statistical inference.

#### Randomization and unbiasedness

Try to avoid confounding effect

#### DE result

	DE Result		
GeneID	Log2FC*	p-value	
1	-0.40	0.037	
2	0.03	0.916	
3	-0.89	2.42E-05	
4	0.30	0.130	
5	-0.36	0.140	
6	-0.07	0.811	
•••			

<sup>\*</sup> Log2FC: log2 of fold change (trt / control)

# Outline

- RNA-seq procedure
- Experimental design
- Multiple testing correction
- Data visualization

## Single statistical test

H0: the null hypothesis

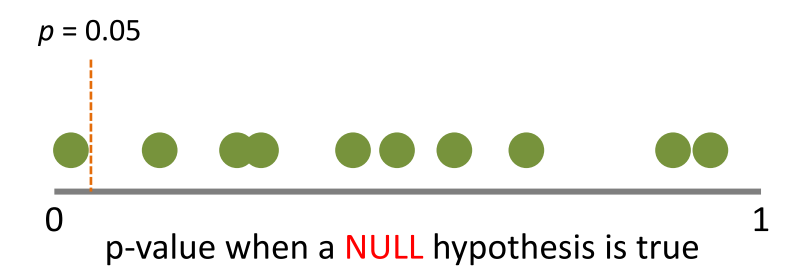


# Single statistical test

H0: the null hypothesis

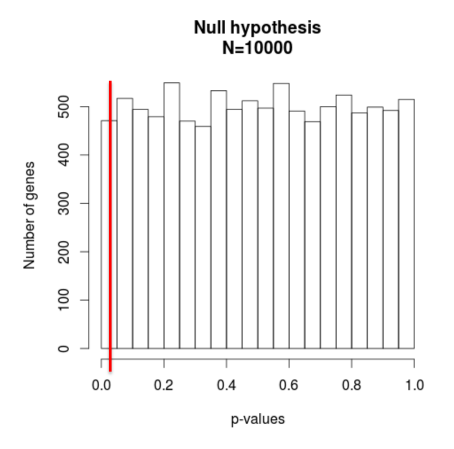


#### Multiple testing correction



"A p-value is only statistically valid when a single score is computed."

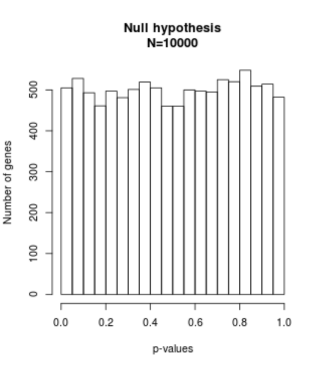
# P-value distribution under the null hypothesis (e.g., no treatment effect)



No matter how stringent the criteria are, you'll identify genes with very small p-values and the false discovery rate (FDR) is 100%.

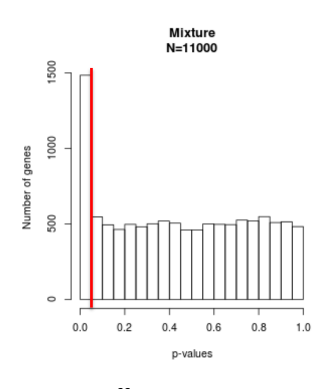
When the null hypothesis is true, the p-value is distributed uniformly from 0 to 1.

# P-value distribution under both the null and non-null hypotheses



Alternative hypothesis N=1000

Sequence of the sequence of the



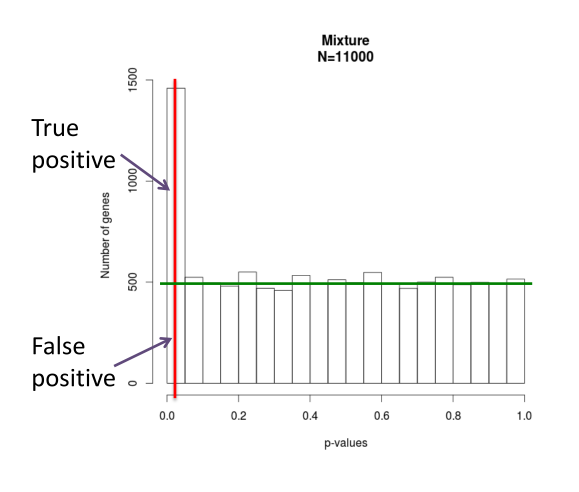
When the null hypothesis is true, the p-value is distributed uniformly.

When the null hypothesis is false, the p-value distribution is skewed toward 0.

Cutoff: p=0.05 FDR=471/(471+989)=32%

Cutoff: p=0.01 FDR=102/(102+912)=10%

#### Multiple test correction – FDR method



P-values < 0.00009

DE=992

False DE=99

**FDR 10%** 

#### False discovery rate (concept)

For example, among 10,000 tests (10,000 genes), 100 significant genes are declared, in which 10 gene is falsely rejected. In this case, the false discovery rate is 10%.

	True null hypothesis (H <sub>0</sub> )	False null hypothesis (H <sub>1</sub> )	Total
Rejected (Declared significance)	10	90	100

#### q-values (adjusted p-values)

The **q-value** is **the smallest FDR** for which we can reject the null hypothesis for that one test and all others with smaller p-values.

Gene	p-values	q-values
1	0.000	0.006
2	0.002	0.015
3	0.009	0.059
4	0.013	0.063
5	0.035	0.139
6	0.051	0.171
7	0.155	0.442
8	0.197	0.492
9	0.247	0.539
10	0.269	0.539
11	0.358	0.651
12	0.396	0.656
13	0.426	0.656
14	0.493	0.702
15	0.526	0.702
16	0.622	0.777
17	0.782	0.920
18	0.862	0.958
19	0.925	0.974
20	0.992	0.992

FDR (False Discovery Rate) method (BH) is a method to calculate q-values/adjusted p-values/corrected p-values based on p-values

5% FDR, q-values < 0.05

10% FDR, q-values < 0.1

20% FDR, q-values < 0.2

Total number of tests: m = 20

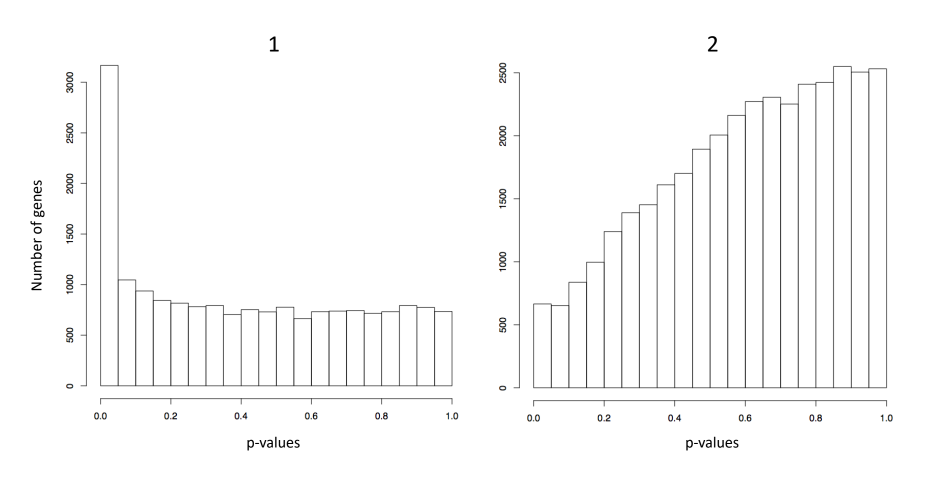
#### Question

If we identify 500 differential expression (DE) genes using the 5% FDR to account for multiple tests. Which one below is a better description?

1. I am 95% confident that 500 genes are DE.

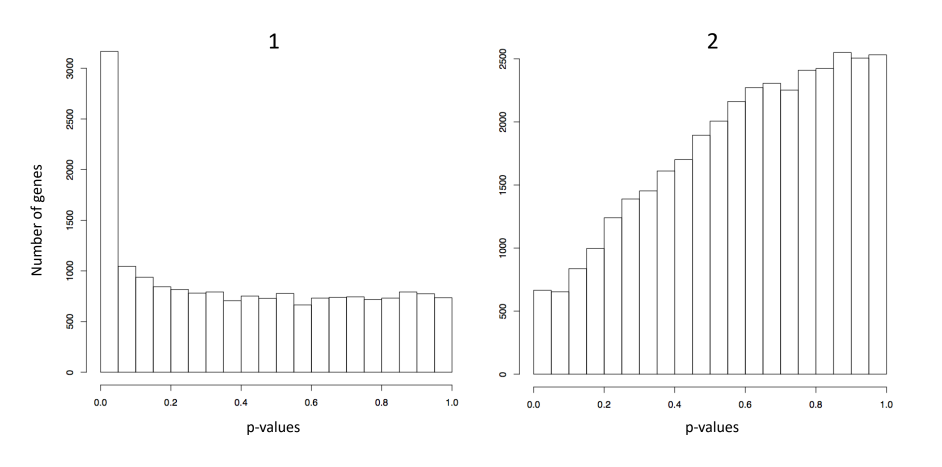
2. The 5% genes (25 genes) in the set are expected to be false DE genes.

#### P-value histograms from real studies



If you perform an RNA-Seq experiment, which one would you hope to obtain? Why?

## P-value histograms from real studies

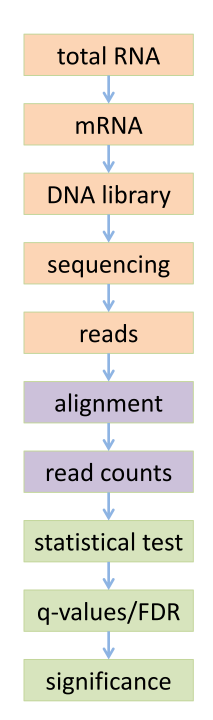


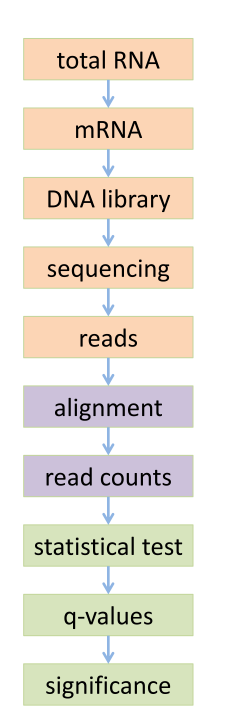
# RNA-Seq procedure

1 RNA to sequencing reads

reads to read counts per gene

read counts to significant genes





#### Keywords

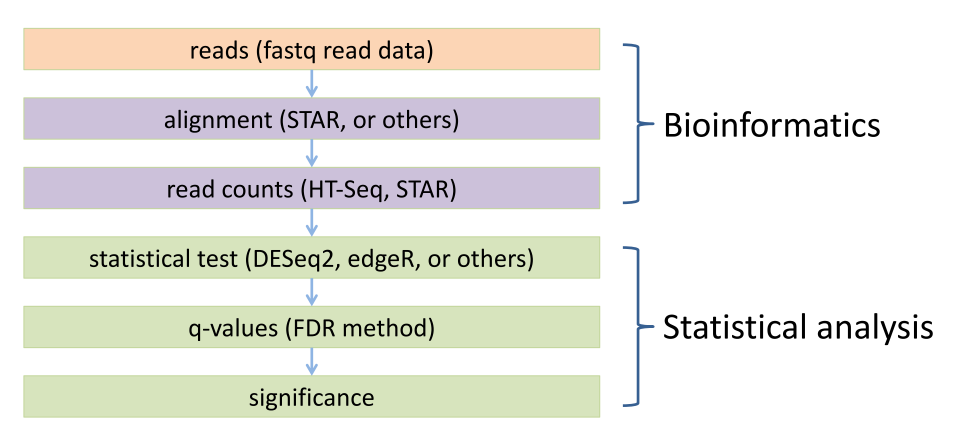
randomization, replication, RNA quality

short or long reads single- or paired-end reads, read length sequencing depths (e.g., >20 million short reads for most experiments)

intron-spanning aligners (e.g., STAR, HiSAT2)

count data statistical analysis (DESeq2 & edgeR) multiple test p-value adjustment (FDR method)

# Bioinformatics and Statistics (Illumina data)



#### **STAR** pipeline – from reads to counts

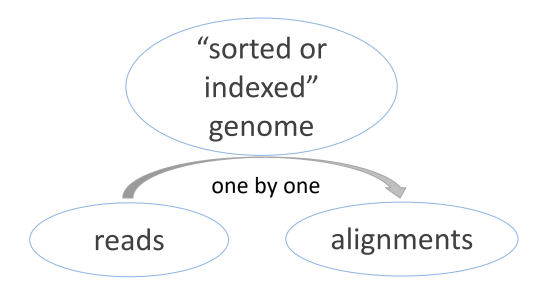
#### Required files:

- 1. Reference genome (fasta file)
- 2. Gene information (gff or gtf gene annotation)
- 3. Reads (fastq files) your own data

Many reference genomes and gff/gtf files are available at: <a href="http://ensembl.org/info/data/ftp">http://ensembl.org/info/data/ftp</a>

Species	DNA (FASTA)	cDNA (FASTA)	CDS (FASTA)	ncRNA (FASTA)	Protein sequence (FASTA)	Annotated sequence (EMBL)	Annotated sequence (GenBank)	Gene sets
<u>Human</u> Homo sapiens	FASTA ₽	FASTA ₽	FASTA ₺	FASTA ₺	FASTA ₽	<u>EMBL</u> ₽	GenBank ঐ	GTF& GFF3&
Mouse Mus musculus	FASTA ₽	FASTA ₽	FASTA ₽	FASTA ₽	FASTA ₽	<u>EMBL</u> ₽	GenBank®	GTF® GFF3®
Zebrafish Danio rerio	FASTA ₽	FASTA ₽	FASTA ₽	FASTA ₽	FASTA ₪	<u>EMBL</u> ₽	GenBank &	GTF® GFF3₽

#### Reads to counts - reference indexing



```
STAR --runMode genomeGenerate \
--genomeDir . \
--genomeFastaFiles reference.fas \
--sjdbGTFfile genes.gtf \
--runThreadN 4
```

#### Reads to counts – alignment and read counting

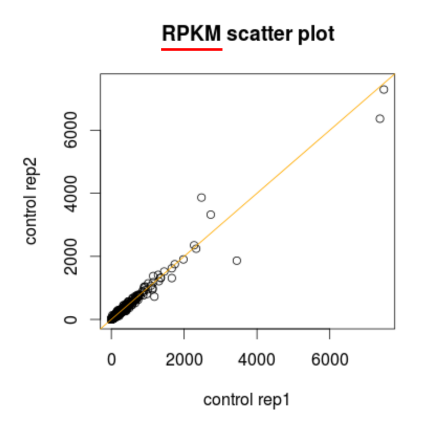
```
STAR --genomeDir reference.fas \
  --readFilesIn read1.fq read2.fq \
  --alignIntronMax 100000 \
  --alignMatesGapMax 100000 \
  --outFileNamePrefix output \
  --outSAMattrlHstart 0 \
  --outSAMmultNmax 1 \
  --outSAMstrandField intronMotif \
  --outFilterIntronMotifs RemoveNoncanonicalUnannotated \
  --outSAMtype BAM SortedByCoordinate \
  --quantMode GeneCounts \
  --outFilterMismatchNmax 5 \
  --outFilterMismatchNoverLmax 0.05 \
  --outFilterMatchNmin 50 \
  --outSJfilterReads Unique \
  --outFilterMultimapNmax 1 \
  --outSAMmapqUnique 60 \
  --outFilterMultimapScoreRange 2
```

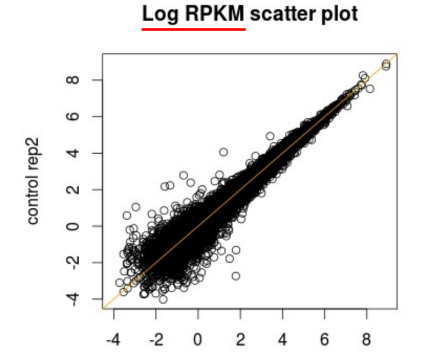
### Count matrix: Read counts (Raw) per gene

Gene	sample 1	sample 2	sample 3
gene 1	6,075	5,934	3,370
gene 2	295	377	169
•••	•••	•••	•••

## Overall comparisons of read counts among samples

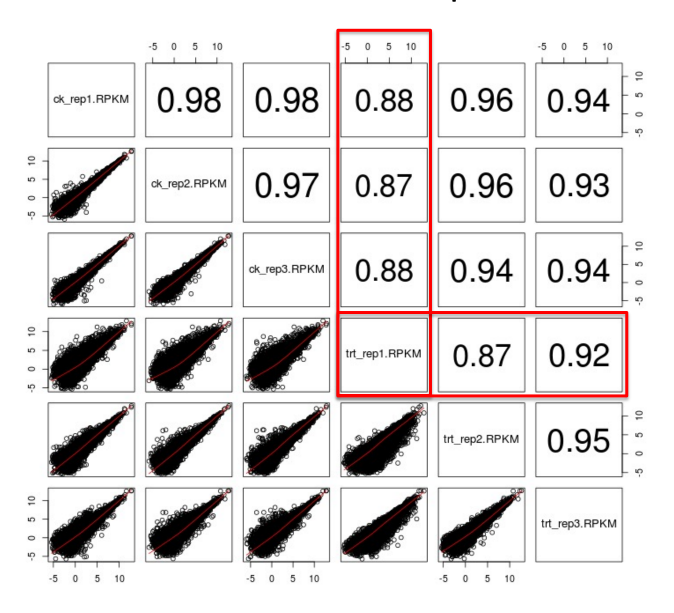
#### Scatter plot





control rep1

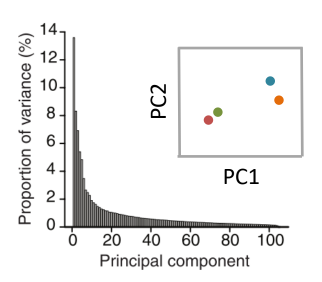
#### Pair-wise scatter plot



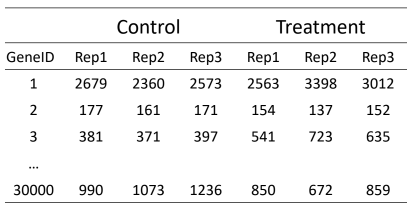
#### Principal Component Analysis (PCA)

PCA is a mathematical algorithm that reduces the dimensionality of the data while retaining most of the variation in the data set.

Feature/vari able	John	Mike	Jack	Justin
Weight (lb)	150	243	186	128
Height (cm)	171	190	178	175
•••				

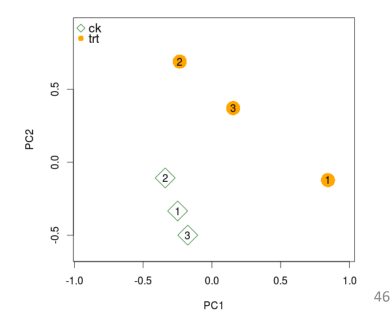


Nature Biotech, 2008, 26:303-4



#### Normalized and standardized data

PCA - full gene set



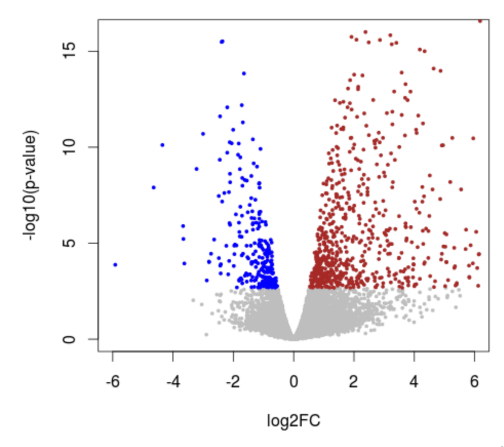
## Overview of differential expression

## Volcano plot



#### Volcano plot

	DE Result				
GeneID	Log2FC	p-value	-log10(pvalue)		
1	-0.40	0.037	1.43		
2	0.03	0.916	0.04		
3	-0.89	2.42E-05	4.62		
4	0.30	0.130	0.89		
5	-0.36	0.140	0.85		
6	-0.07	0.811	0.09		
•••					

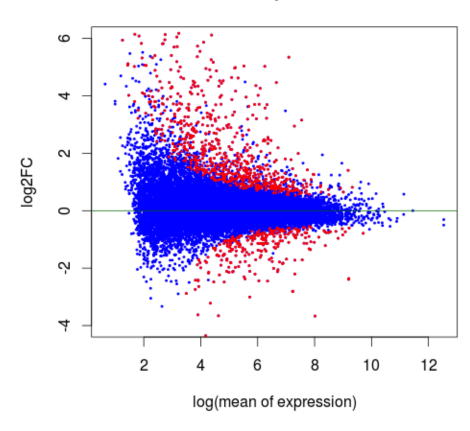


#### MA plot

#### M (log ratios) and A (mean average)

GeneID	Mean RPKM	log mean	log2FC
1	0.51	-0.29	-0.40
2	1.25	0.10	0.03
3	3.52	0.55	-0.89
4	0.19	-0.72	0.30
5	2.34	0.37	-0.36
6	6.14	0.79	-0.07
•••			

#### MA plot



#### Summary

- Biological replication rather than technical replication are typically needed for an RNA-Seq experiment.
- P-values need to be corrected to account for multiple tests. The FDR method is a reliable approach for the correction.
- Many bioinformatics pipelines and statistical methods have been developed. Most methods work fine but parameters in each method need to be carefully selected.

#### REFERENCES

- 1. Benjamini Y, et al. 1995. Controlling the False Discovery Rate a Practical and Powerful Approach to Multiple Testing. Journal of the Royal Statistical Society Series B-Methodological 57:289-300.
- Conesa A, et al. 2016. A survey of best practices for RNA-seq data analysis. Genome Biol 17:13.
- Love MI, et al. 2014. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol 15:550.
- 4. Robinson MD, et al. 2010. edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics 26:139-140.