

# RNA-seq to study HIV Infection in cells

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Uku-Kaspar Uustalu Data Science Specialist

- ✓ Consultation on Projects and Grants
- ✓ High Performance Compute Cluster
- ✓ Workshops

https://it.tufts.edu/research-technology

### Outline

Bulk and single cell RNA sequencing

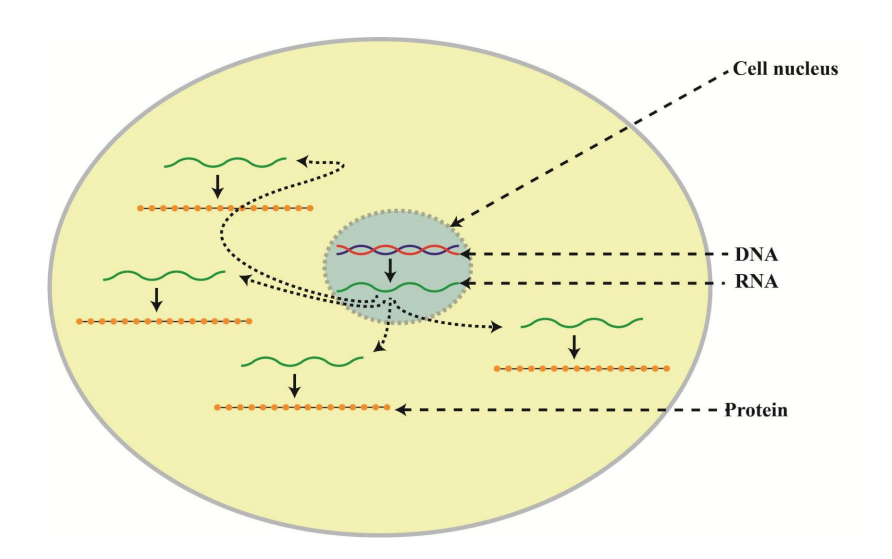
Intro to Galaxy Platform for Bioinformatics (Tufts network or VPN required)

https://galaxy.cluster.tufts.edu/

Work through RNAseq example together on Galaxy

https://rbatorsky.github.io/in tro-to-rnaseq-with-galaxy/ 2 days!

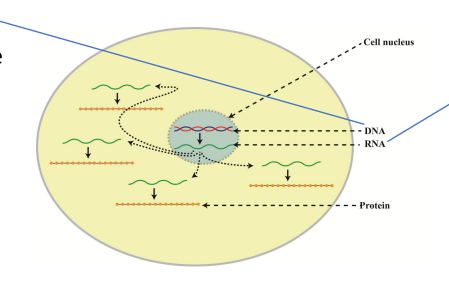
#### DNA and RNA in a cell



#### Two common analyses

#### **DNA Sequencing**

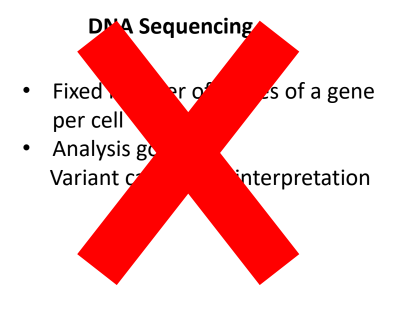
- Fixed number of copies of a gene per cell
- Analysis goal:
   Variant calling and interpretation

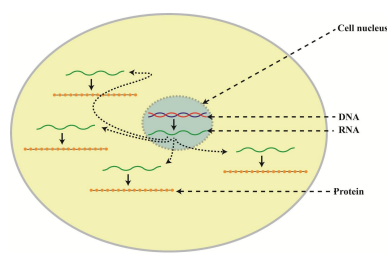


#### **RNA Sequencing**

- Number of copies of a gene transcript per cell depends on gene expression
- Analysis goal:
  - Bulk : Differential expression
  - Single cell : Quantify different cell populations

#### Today we will cover RNA sequencing





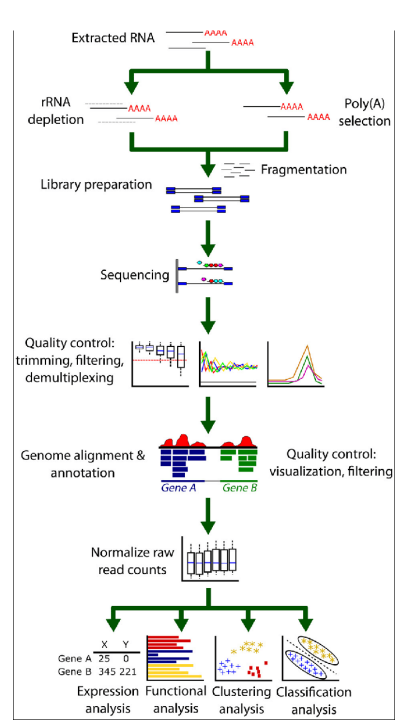
#### **RNA Sequencing**

- Number of copies of a gene transcript per cell depends on gene expression
- Analysis goal:
  - Bulk : Differential expression
  - Single cell : Quantify different cell populations

#### "Bulk" RNA seq workflow

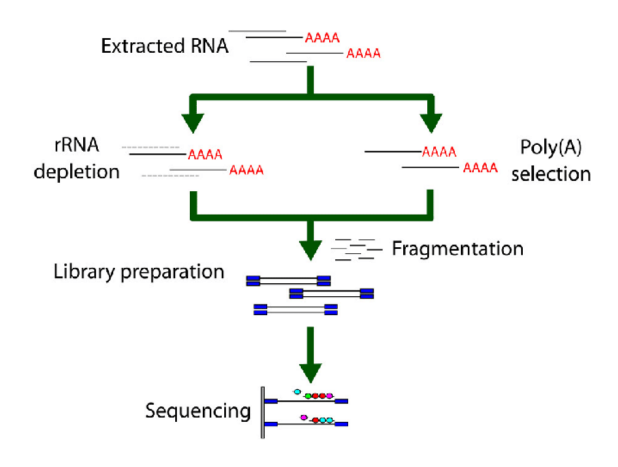
Library prep and sequencing

**Bioinformatics** 



Good resource: Griffiths et al Plos Comp Bio 2015

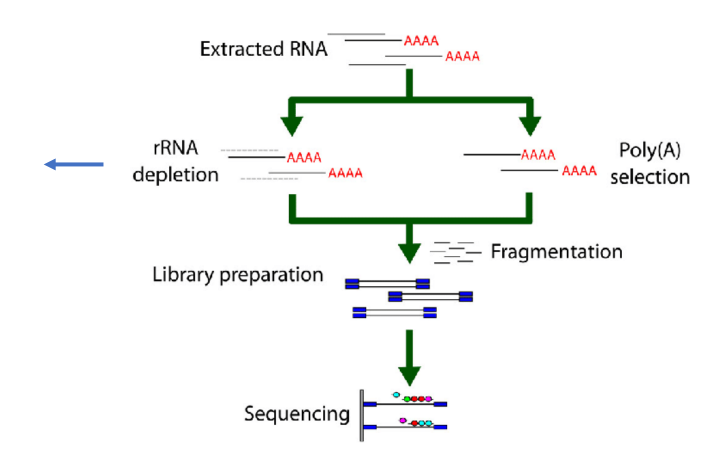
### RNA seq library prep and sequencing



Good resource: <u>Griffiths et al Plos Comp Bio 2015</u>

### RNA seq library prep and sequencing

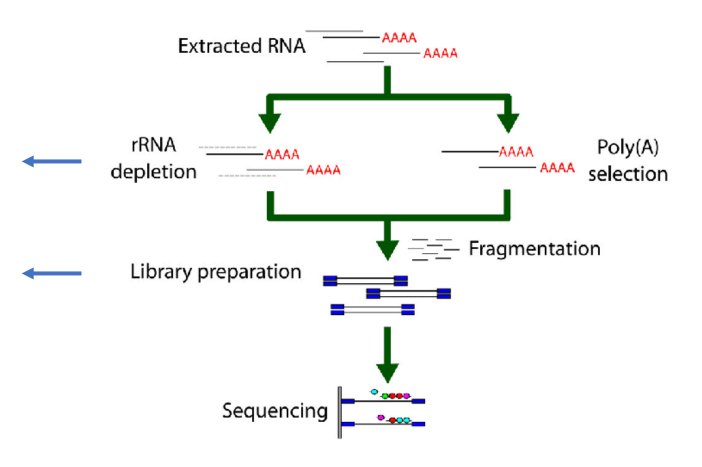
- Enrichment for mRNA, two options
- In humans, ~95%–98% of all RNA molecules are rRNAs



Good resource: <u>Griffiths et al Plos Comp Bio 2015</u>

### RNA seq library prep and sequencing

- Enrichment for mRNA, two options
- In humans, ~95%–98% of all RNA molecules are rRNAs
- Random priming and reverse transcription
- Double stranded cDNA synthesis
- Sequencing adapter ligation



#### **Resources:**

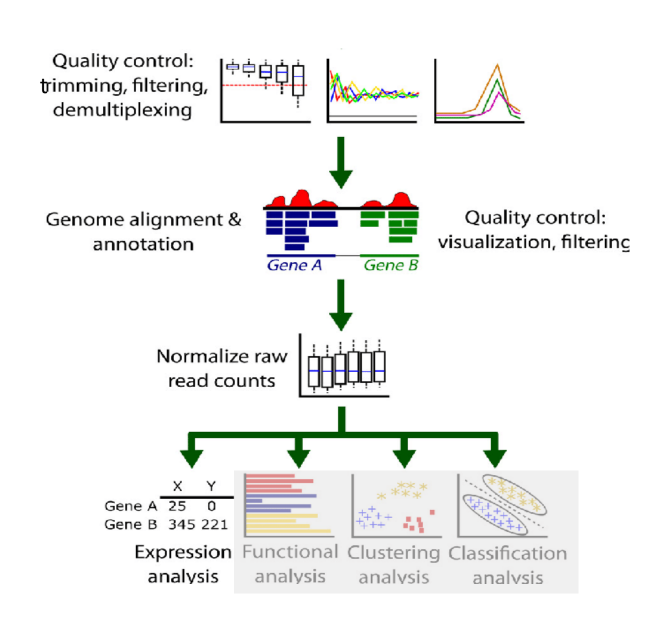
Illumina Sequencing by Synthesis
Griffiths et al Plos Comp Bio 2015

#### RNA seq bioinformatics

Goal of Differential Expression

"How can we detect genes for which the counts of reads change between conditions **more systematically** than as expected by chance"

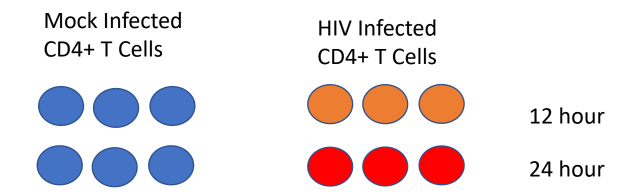
Oshlack et al. 2010. From RNA-seq reads to differential expression results. Genome Biology 2010, 11:220



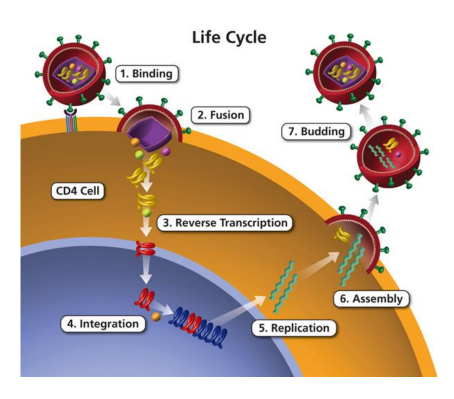
#### Our dataset

## Next-Generation Sequencing Reveals HIV-1-Mediated Suppression of T Cell Activation and RNA Processing and Regulation of Noncoding RNA Expression in a CD4<sup>+</sup> T Cell Line

Stewart T. Chang, Pavel Sova, Xinxia Peng, Jeffrey Weiss, G. Lynn Law, Robert E. Palermo, Michael G. Katze



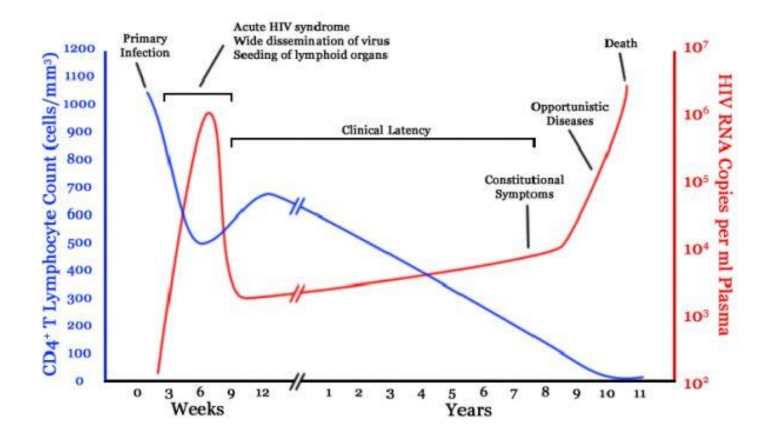
### HIV lifecycle



### HIV lifecycle

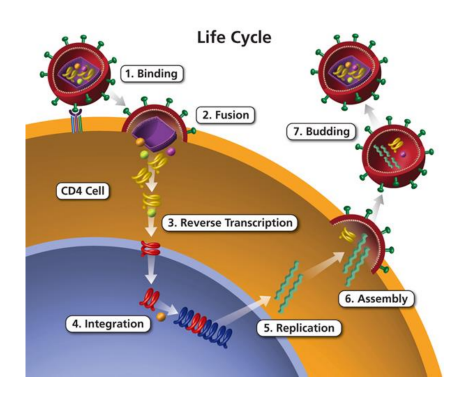
#### Life Cycle 1. Binding 2. Fusion 7. Budding CD4 Cell 3. Reverse Transcription 6. Assembly 4. Integration 5. Replication

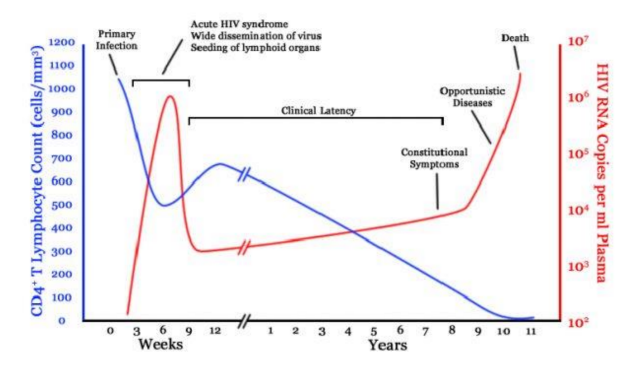
#### HIV infection in a human host



#### The study question

What changes take place in the first 12-24 hours of HIV infection in terms of gene expression of host cell and viral replication levels?

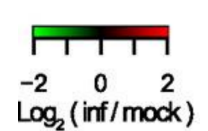


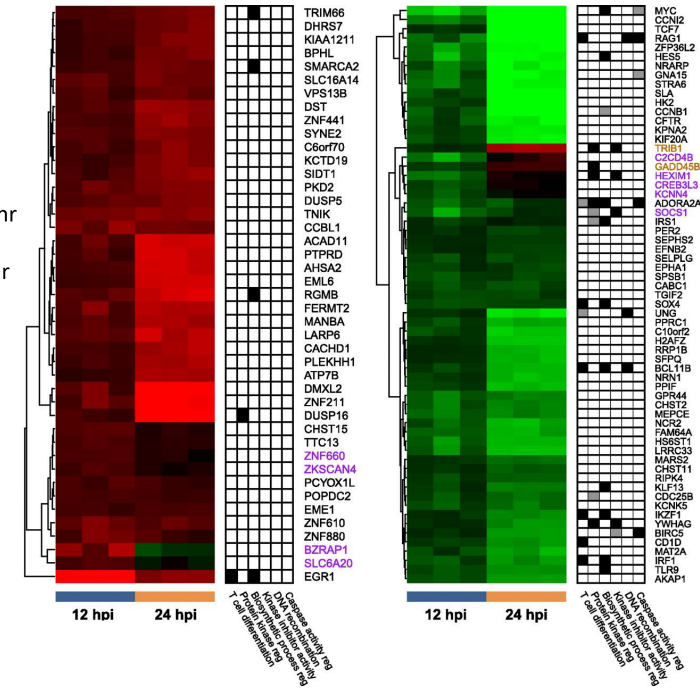


### Study findings

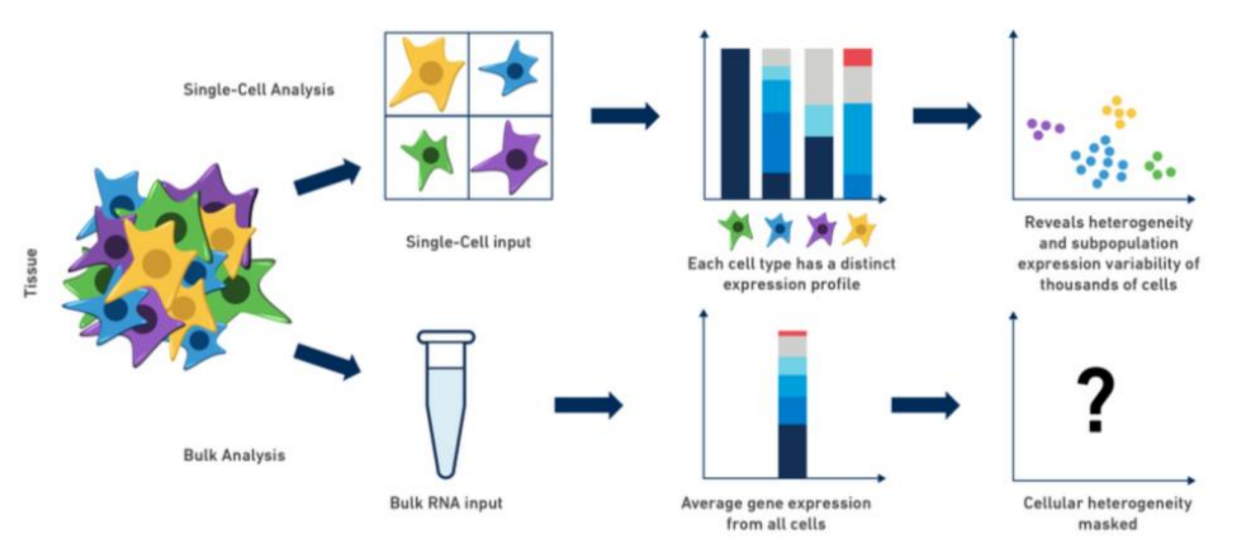
Using RNAseq, authors demonstrate:

- 20% of reads mapped to HIV at 12 hr, 40% at 24hr
- Downregulation of T cell differentiation genes at 12hr
- 'Large-scale disruptions to host transcription' at 24hr

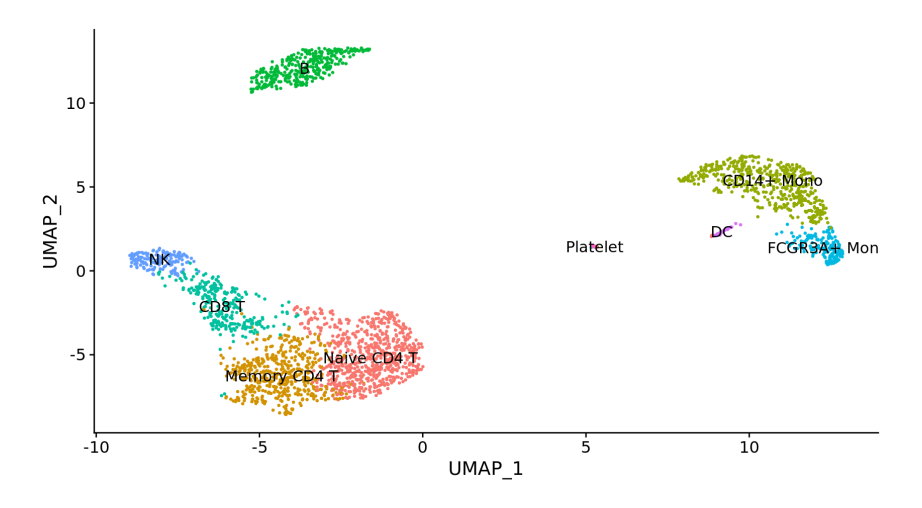




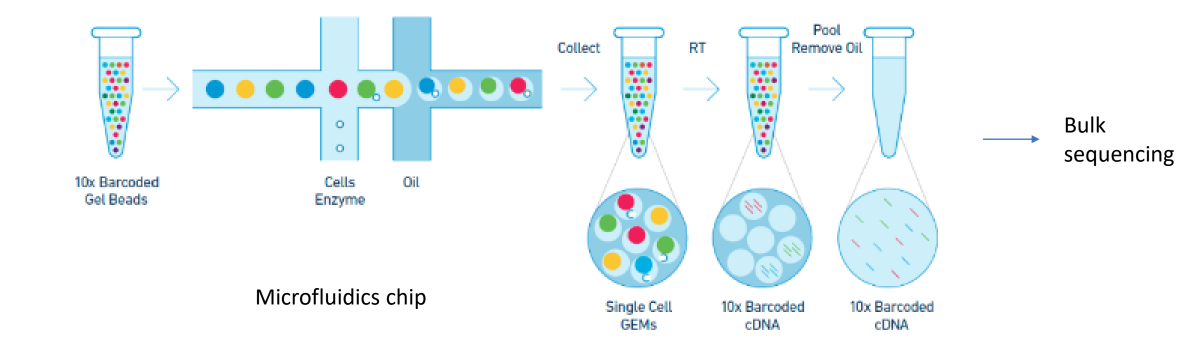
### Bulk vs Single Cell RNA Sequencing



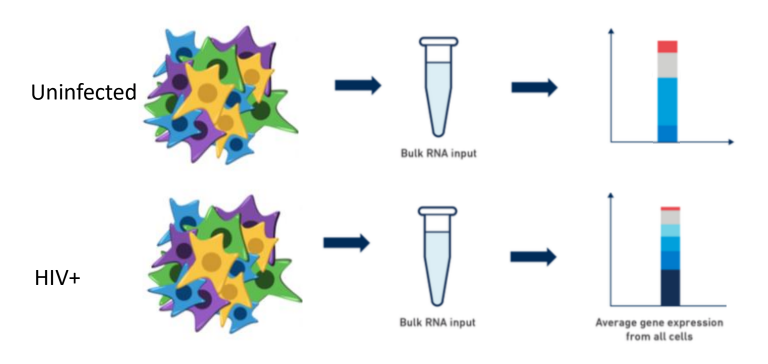
#### scRNA cell subsets in PBMC



### 10x single cell technology

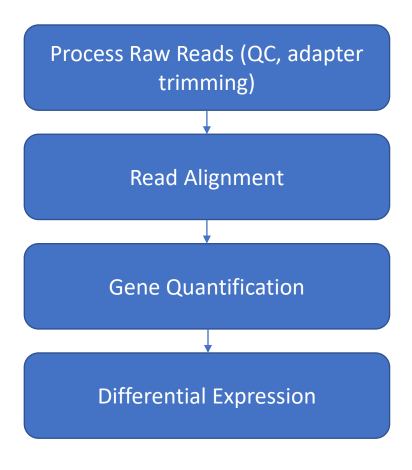


### Bulk RNAseq for Differential Expression is OK!

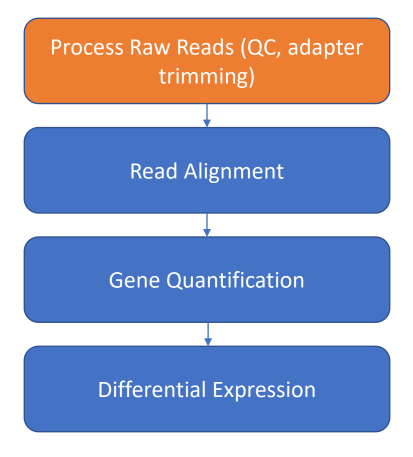


Compare relative gene expression between conditions

### Our (bulk) RNAseq Workflow



### Quality control on Raw Reads



#### Raw reads in Fastq format

@SRR098401.109756285

GACTCACGTAACTTTAAACTCTAACAGAAATATACTA...

+

CAEFGDG?BCGGGEEDGGHGHGDFHEIEGGDDDD...

- 1. Sequence identifier
- 2. Sequence
- 3. + (optionally lists the sequence identifier again)
- 4. Quality string

#### Base Quality Scores

The symbols we see in the read quality string are an encoding of the quality score:

```
Quality encoding: !"#$%&'()*+,-./0123456789:;<=>?@ABCDEFGHI
```

A quality score is a prediction of the probability of an error in base calling:

Quality Score	Probability of Incorrect Base Call	Inferred Base Call Accuracy
10 (Q10)	1 in 10	90%
20 (Q20)	1 in 100	99%
30 (Q30)	1 in 1000	99.9%

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30 (Q30)	1 in 1000	99.9%

#### Back to our read:

```
@SRR098401.109756285
GACTCACGTAACTTTAAACTCTAACAGAAATATACTA...
+
CAEFGDG?BCGGGEEDGGHGHGDFHEIEGGDDDD...
```

 $C \rightarrow Q = 34 \rightarrow Probability < 1/1000 of an error$ 

#### Raw read quality control

#### Fastq File

@SRR497699.30343179.1 HWI-EAS39X\_10175\_FC61MK0\_4\_117\_4812\_10346 length=75 CAGATGGCCGCAGAGGGAAGCCATGAAGGCCCTGCATGGGGAAGACCGGTTCAGCAGGAATGCCGAGAC

+

+

+

-

HHHHHHHHHHGGGGGHHHGHGEBEEGGEDGGGGGGHHHHHGGEGBDGGGDDGBGGC<EADBEBE<GGGGBEEDGD

• • •

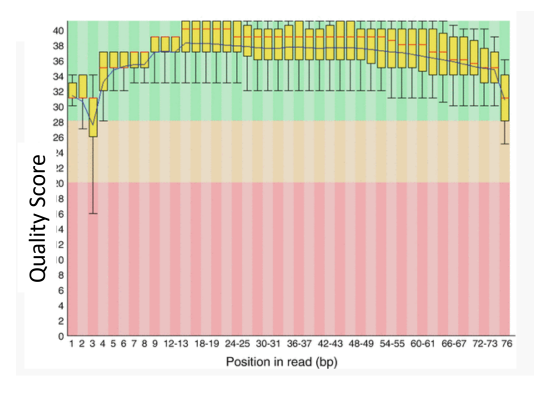
#### <u>Metrics</u>

- Sequence Quality
- GC content

FastQC Tool

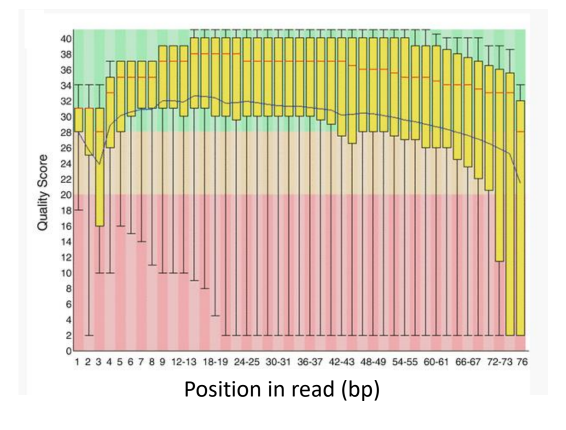
- Per base sequence content
- Adapters in Sequence

#### FastQC: Sequence Quality Histogram

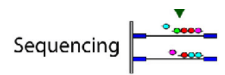


Position in read (bp)

GOOD
High quality over the length of the read

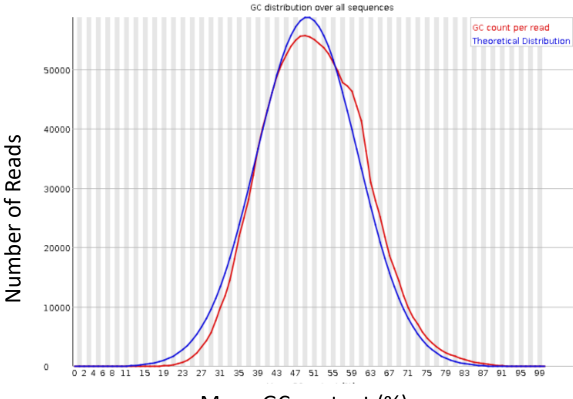


BAD Read quality drops at the beginning and end

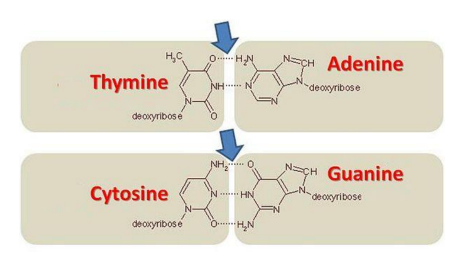


#### FastQC: Per sequence GC content

#### Per sequence GC content

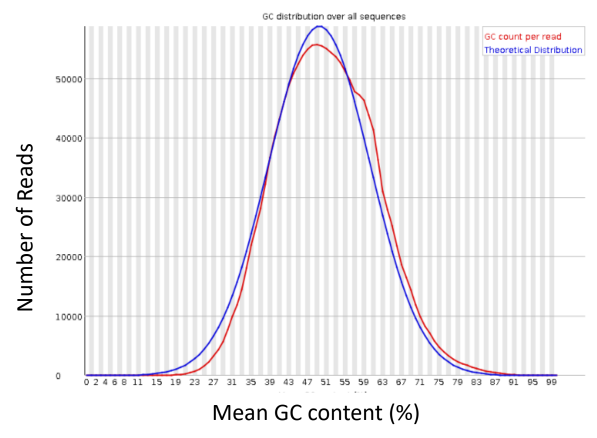


Mean GC content (%)



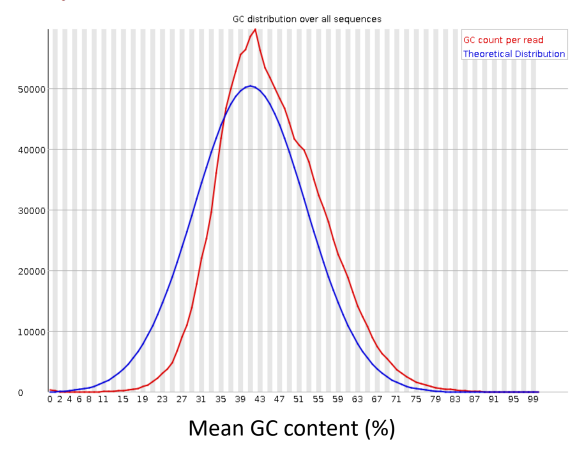
#### FastQC: Per sequence GC content

#### Per sequence GC content



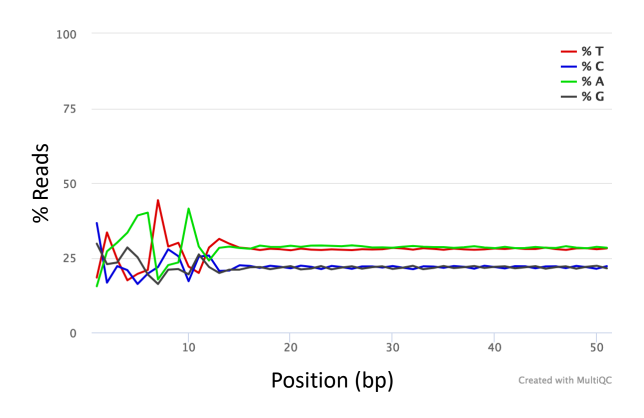
GOOD: follows normal distribution (sum of deviations is < 15% of reads)

#### **Per sequence GC content**



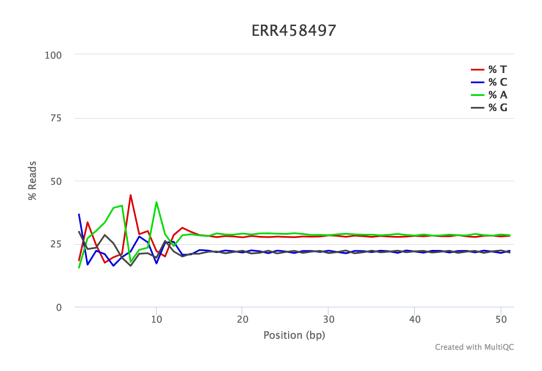
BAD: can indicate contamination with adapter dimers, or another species

#### FastQC: Per Base Sequence Content

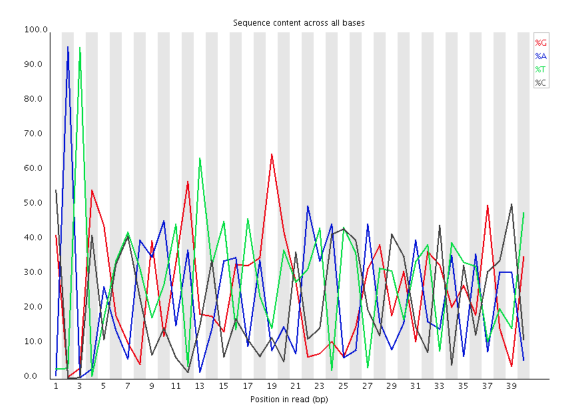


- Proportion of each position for which each DNA base has been called
- RNAseq data tends to show a positional sequence bias in the first ~12 bases
- The "random" priming step during library construction is not truly random and certain hexamers are more prevalent than others

#### FastQC: Per Base Sequence Content



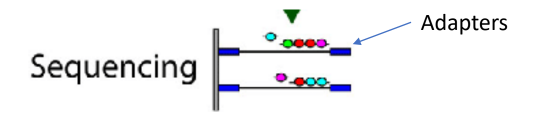
**EXPECTED** for RNAseq



BAD:

Shows a strong positional bias throughout the reads, which in this case is due to the library having a certain sequence that is overrepresented

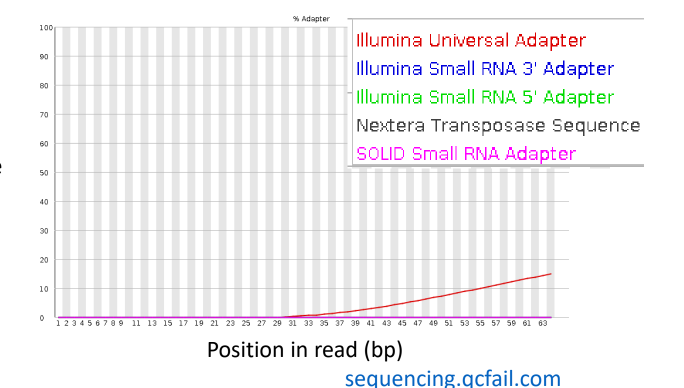
#### FastQC: Adapter content



FastQC will scan each read for the presence of known adapter sequences

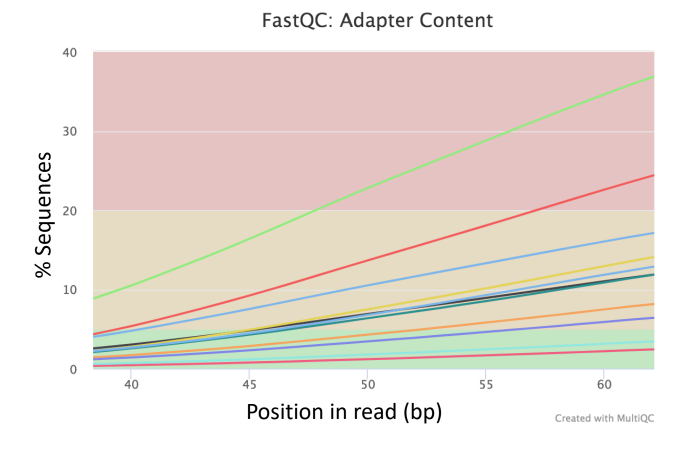
The plot shows that the adapter content rises over the course of the read

Solution – Adapter trimming!



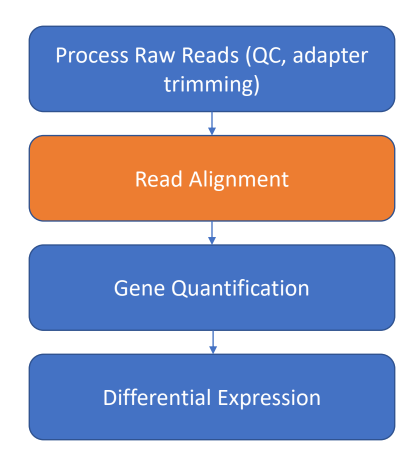
#### FastQC -> MultiQC

Should view all samples at once to notice abnormalities for our dataset.



We'll use a tool called "Trim Galore!" to trim adapters and remove low quality bases/reads.

#### Workflow



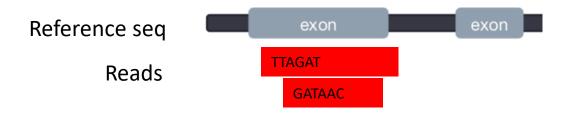
#### Read Alignment

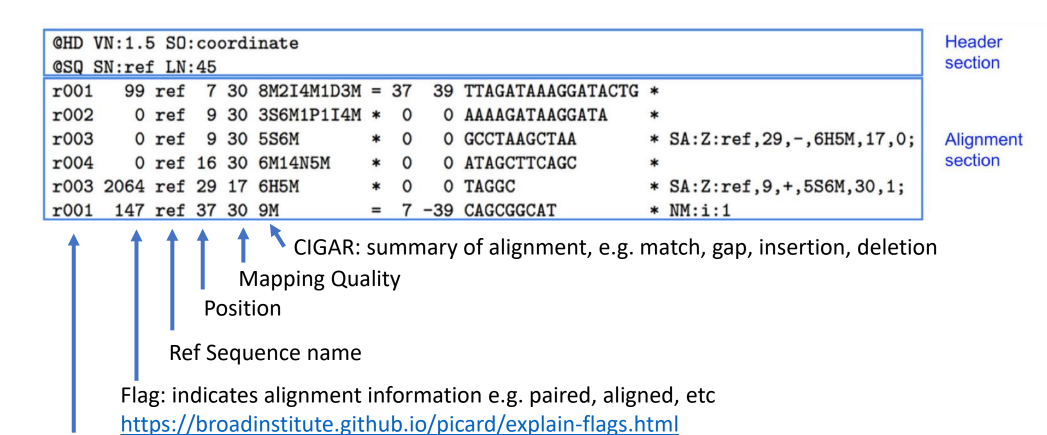
- RNAseq data originates from spliced mRNA (no introns)
- When aligning to the genome, our aligner must find a spliced alignment for reads
- We use a tool called STAR (Spliced Transcripts Alignment to a Reference) that has a exon-aware mapping algorithm.

full-length mRNA AAAAAA 3' **cDNA** fragments sequencing aligned reads exon

Reference sequence

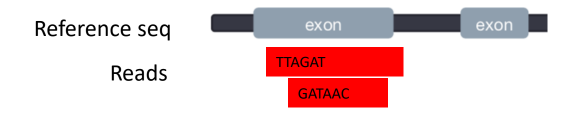
### Sequence Alignment Map (SAM)

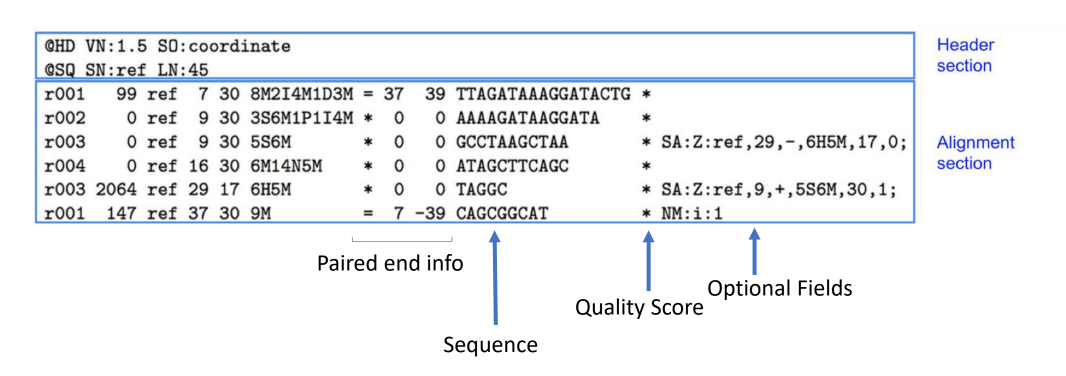




Read ID

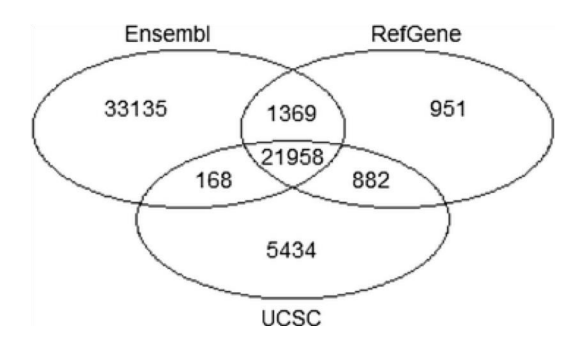
## Sequence Alignment Map (SAM)





#### Genome Annotation Standards

- STAR can use an annotation file gives the location and structure of genes in order to improve alignment in known splice junctions
- Annotation is dynamic and there are at least three major sources of annotation
- The intersection among RefGene, UCSC, and Ensemble annotations shows high overlap. RefGene has the fewest unique genes, while more than 50% of genes in Ensemble are unique
- Be consistent with your choice of annotation source!



### Gene Annotation Format (GTF)

In order to count genes, we need to know where they are located in the reference sequence STAR uses a Gene Transfer Format (GTF) file for gene annotation

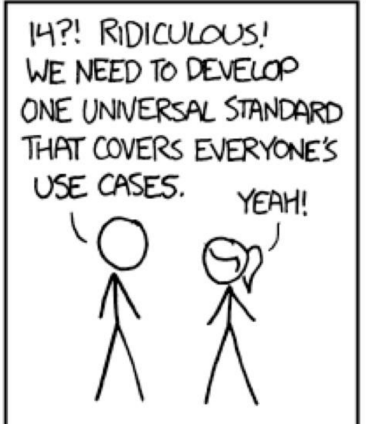
Frame

						Stra	and	
Chrom	Source	Feature type	Start	Stop	(Sco	re)		Attribute
chr5	hg38_refGene	exon	138465492	138466068		+		gene_id "EGR1";
chr5	hg38_refGene	CDS	138465762	138466068		+	0	gene_id "EGR1";
chr5	hg38_refGene	start_codon	138465762	138465764		+		gene_id "EGR1";
chr5	hg38_refGene	CDS	138466757	138468078		+	2	gene_id "EGR1";
chr5	hg38_refGene	exon	138466757	138469315		+		gene_id "EGR1";
chr5	hg38_refGene	stop_codon	138468079	138468081		+		gene_id "EGR1";

#### A note on standards

HOW STANDARDS PROLIFERATE: (SEE: A/C CHARGERS, CHARACTER ENCODINGS, INSTANT MESSAGING, ETC.)

SITUATION: THERE ARE 14 COMPETING STANDARDS.

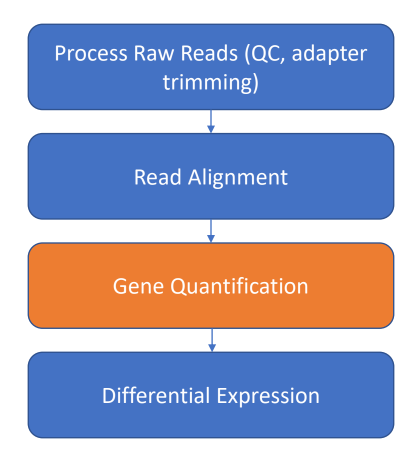




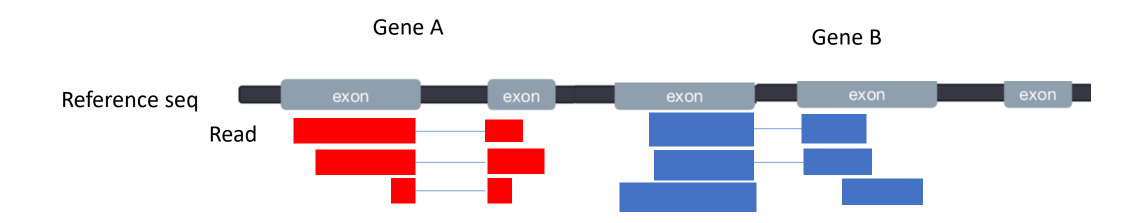
## Visualizing reads with JBrowse



### Workflow

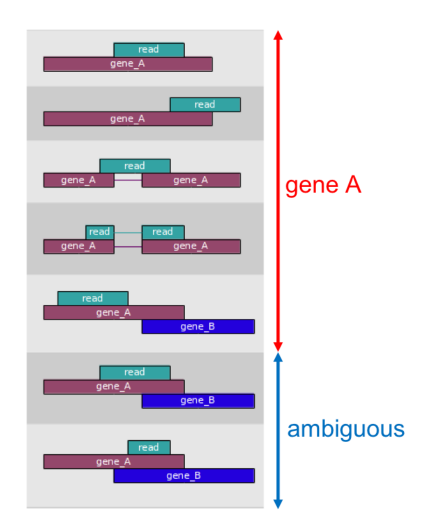


# Counting reads for each gene



# Counting reads: featurecounts

- The mapped coordinates of each read are compared with the features in the GTF file
- Reads that overlap with a gene by >=1 bp are counted as belonging to that feature
- Ambiguous reads will be discarded

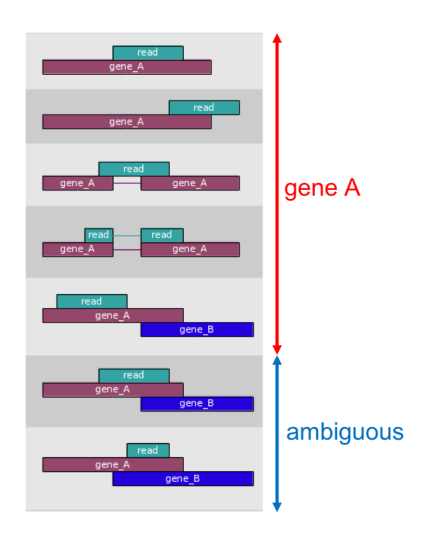


# Counting reads: featurecounts

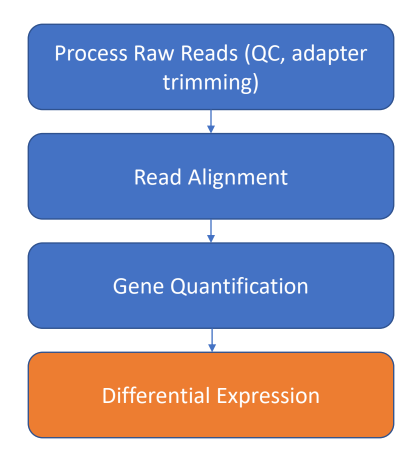
- The mapped coordinates of each read are compared with the features in the GTF file
- Reads that overlap with a gene by >=1 bp are counted as belonging to that feature
- Ambiguous reads will be discarded

#### Result is a gene count matrix:

Gene	Sample 1	Sample 2	Sample 3	Sample 4
Α	1000	1000	100	10
В	10	1	5	6
С	10	1	10	20

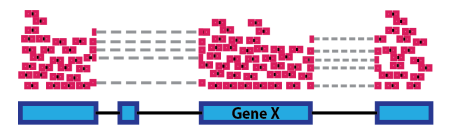


### Workflow



- Raw Count != Expression strength
- Normalization:
  - Eliminates factors that are not of interest for our experiment
  - Enables accurate comparison between samples or genes

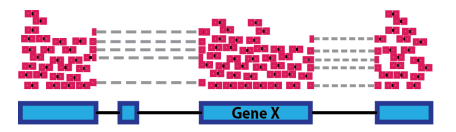
#### **Sample A Reads**

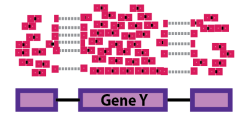


The number of reads mapped to a gene depends on

Gene Length

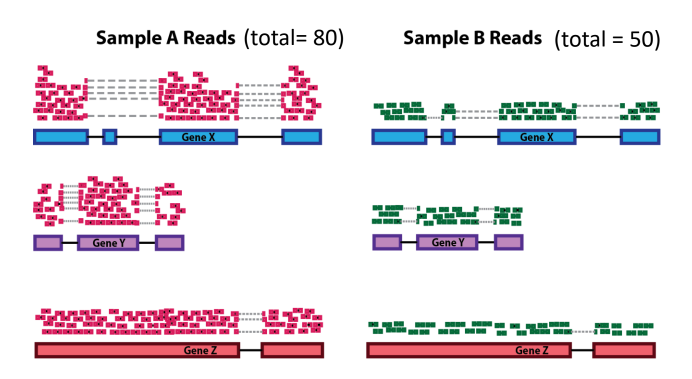
#### **Sample A Reads**





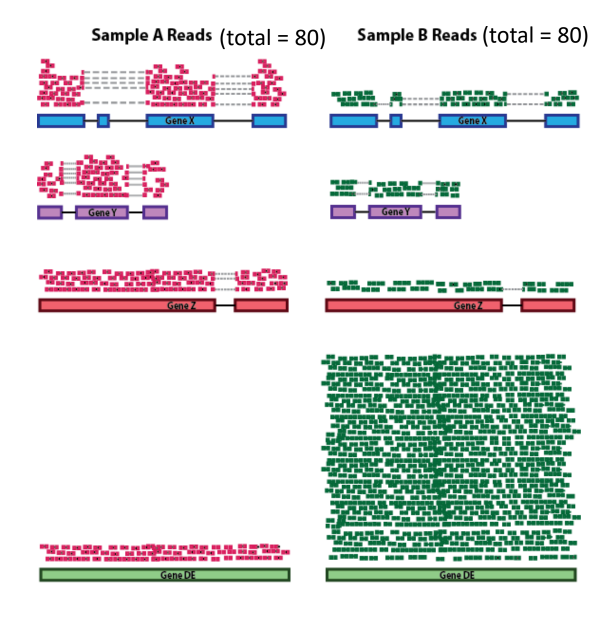
The number of reads mapped to a gene depends on

- Gene Length
- Sequencing depth



The number of reads mapped to a gene depends on

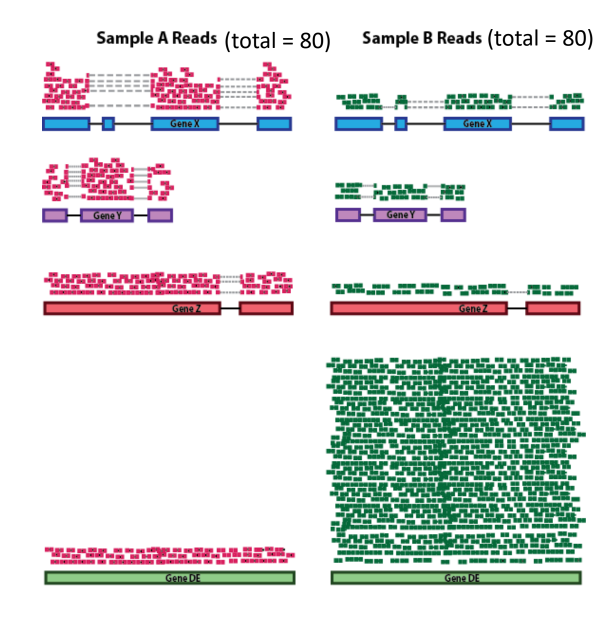
- Gene Length
- Sequencing depth
- The expression level of other genes in the sample (RNA Composition)



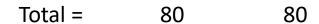
The number of reads mapped to a gene depends on

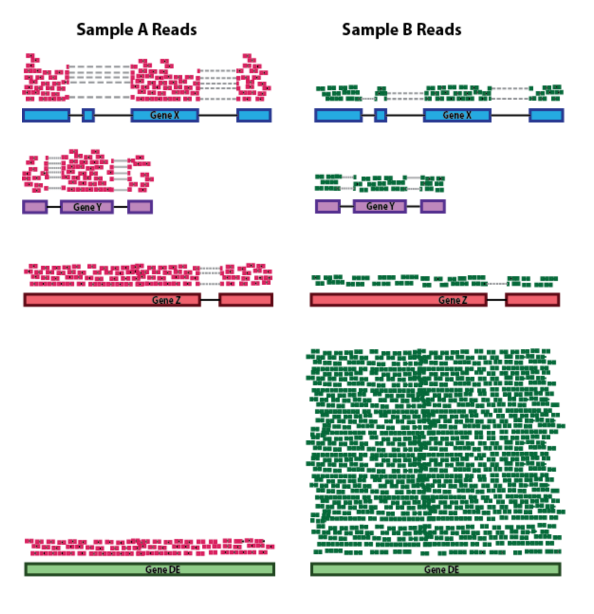
- Gene Length
- Sequencing depth
- The expression level of other genes in the sample (RNA Composition)

**DESeq2 Median of Ratios** 



Gene	Sample A	Sample B
X	26	10
Υ	26	10
Z	26	10
DE	2	50





1. Take a row-wise average to produce an average sample (geometric mean)

$\sqrt[n]{x_1x_2}$	•	•	•	$\overline{x_n}$

Gene	Sample A	Sample B	Avg. Sample
Х	26	10	 16
Υ	26	10	16
Z	26	10	16
DE	2	50	10

1. Take a row-wise average to produce an average sample (geometric mean)

an)	$\sqrt[n]{x_1x_2}$	•	•	•	$x_n$	

Gene	Sample A	Sample B	Avg. Sample
Χ	26	10	 16
Υ	26	10	16
Z	26	10	16
DE	2	50	10

2. Divide all rows by the Average Sample for that gene (Ratio)

Gene	Sample A/Avg.	Sample B /Avg.
X	26/16 = 1.6	10/16 = 0.6
Υ	1.6	0.6
Z	1.6	0.6
DE	0.2	5

1. Take a row-wise average to produce an average sample (geometric mean)  $\sqrt[n]{x}$ 

$$\sqrt[n]{x_1x_2\cdots x_n}$$

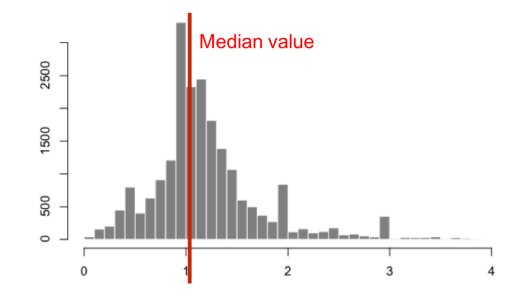
Gene	Sample A	Sample B	Avg. Sample
Х	26	10	 16
Υ	26	10	16
Z	26	10	16
DE	2	50	16

2. Divide all rows by the Average Sample for that gene (Ratio)

Gene	Sample A/Avg.	Sample B /Avg.
Χ	26/16 = 1.6	10/16 = 0.6
Υ	1.6	0.6
Z	1.6	0.6
DE	0.2	5

3. Take the **median** of each column. Should be ~1 for all

Size factor	1.6	0.6
-------------	-----	-----



1. Take a row-wise average to produce an average sample (geometric mean)

$$\sqrt[n]{x_1x_2\cdots x_n}$$

Gene	Sample A	Sample B	Avg. Sample
Χ	26	10	 16
Υ	26	10	16
Z	26	10	16
DE	2	50	16

2. Divide all rows by the Average Sample for that gene (Ratio)

Gene	Sample A/Avg.	Sample B /Avg.
X	26/16 = 1.6	10/16 = 0.6
Υ	1.6	0.6
Z	1.6	0.6
DE	0.2	5

4. Divide all counts by sample specific size factor

Gene	Sample A / S <sub>A</sub>	Sample B / S <sub>B</sub>
X	16.3	16.7
Υ	16.3	16.7
Z	16.3	16.7
DE	1.3	83.3

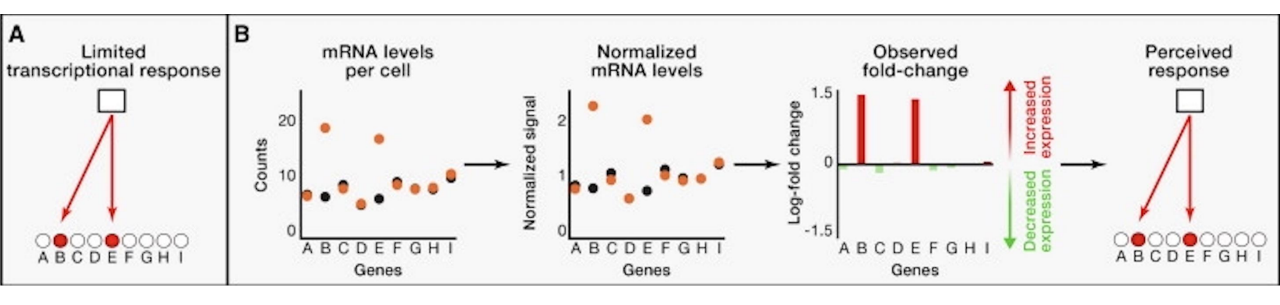
Normalized counts for non-DE genes are similar!

3. Take the **median** of each column. Should be ~1 for all

Size factor	1.6	0.6
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# Assumption of DESeq2 Median of Ratios

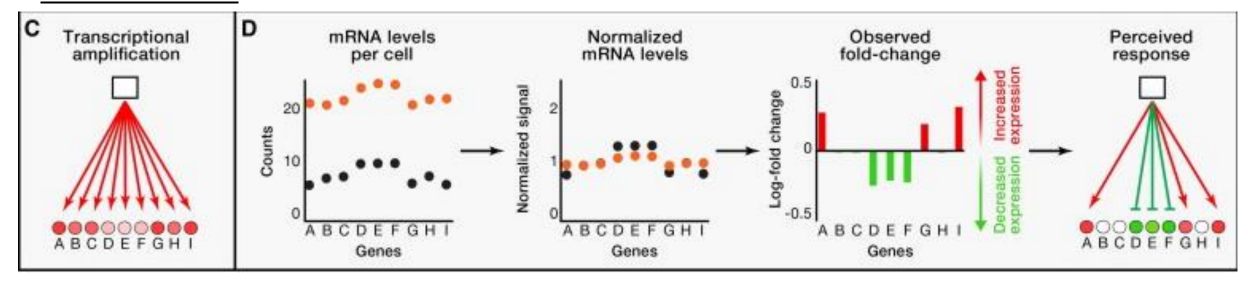
Median of Ratios method assumes that most genes are not Differentially Expressed between samples.



# Assumption of DESeq2 Median of Ratios

Median of Ratios method assumes that most genes are not Differentially Expressed between samples.

#### **COUNTER EXAMPLE**



NOTE: add back full picture or remove

- Late stage cell death (total RNA DOWN)
- High c-Myc cells (total RNA UP)

Known quantity spike-in transcripts (ERCC) can be used to normalize in these cases.

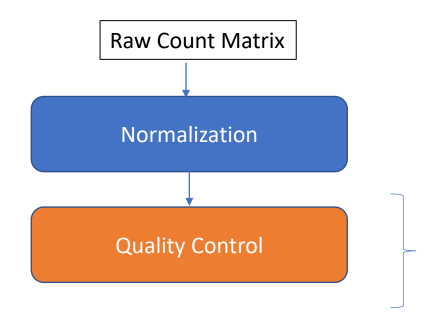
Loven et al "Revisiting Global Gene Expression Analysis" Cell 2012 <a href="https://doi.org/10.1016/j.cell.2012.10.012">https://doi.org/10.1016/j.cell.2012.10.012</a>

#### Normalization methods

Normalization method	Description	Accounted factors	Recommended use
CPM (counts per million)	K <sub>i</sub> Total Reads per Sample/10 <sup>6</sup>	sequencing depth	Comparison between replicates of the sample group
<b>R/FPKM</b> (reads/fragments per kilobase of exon per million reads/fragments mapped)	$\frac{K_{i}}{\textit{Gene Length}/10^{3}*\;\textit{Total Reads per Sample}/10^{6}}$	sequencing depth and gene length	Comparison between genes in a sample
DESeq2's <b>median of ratios</b> [1]	$K_i$ divided by sample-specific size factors		Differential Expression between samples

Similar to DESeq2: EdgeR, limma-voom

## Quality Control Visualizations



Examine sources of variation in the data

- Principal Component Analysis
- Hierarchical Clustering

(Log2 + 1) Transformed, Normalized Count Table

Gene	Sample A	Sample B	Sample C
1	1	1.6	0.5
2	2.2	-0.2	1
3	-1	1	3.1

### Principle Component Analysis

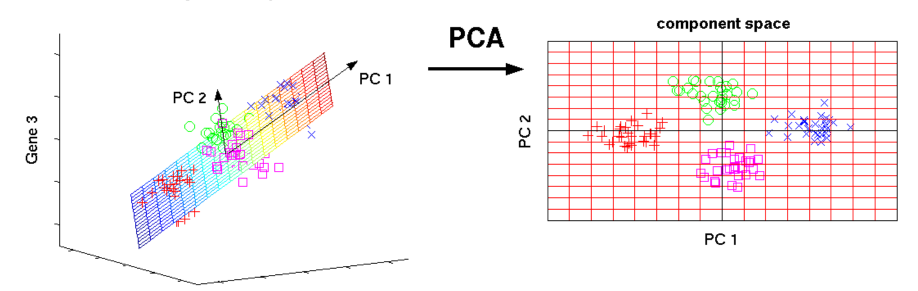
Dimension reduction technique Example: 3 gene dimensions -> 2 PC

Gene	Mock_12h	Mock_12h	Mock_24h	Mock_24h	HIV_12h	HIV_12h	HIV_24h	HIV_24h
Gene 1	8.9	8.9	8.9	9.0	8.9	8.9	9.0	6.8
Gene 2	0.6	-1.0	0.6	-1.0	0.6	-1.0	0.6	3.8
Gene 3	4.1	11.9	4.1	-0.5	4.1	8.7	4.0	4.4

#### original data space

Gene 1

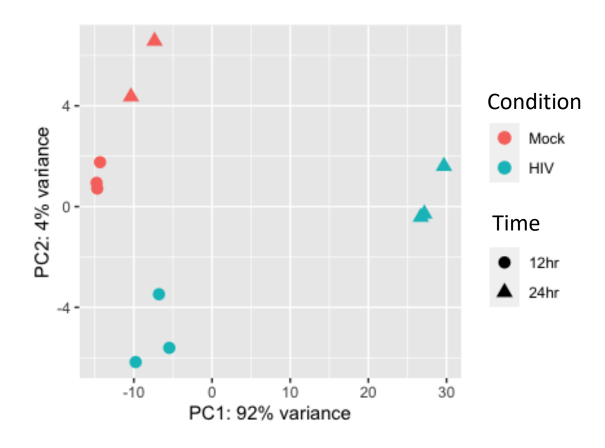
Gene 2



Do your samples cluster as expected?

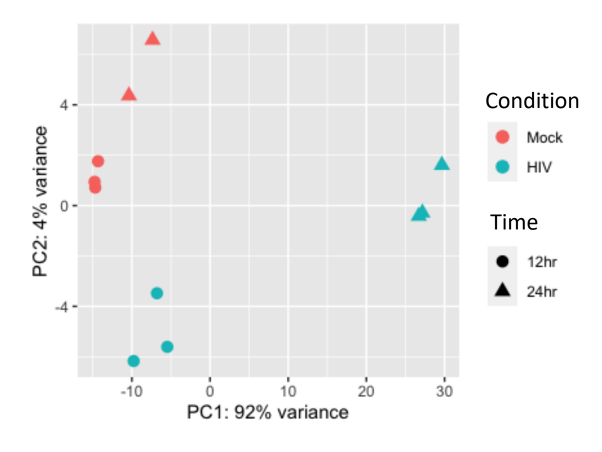
What are the major sources of variation in the data?

# Principle Component Analysis



- ✓ Do your samples cluster as expected?
- ✓ What are the major sources of variation in the data?

## Principle Component Analysis



- ✓ Do your samples cluster as expected?
- ✓ What are the major sources of variation in the data?
- ✓ Is there a batch effect?

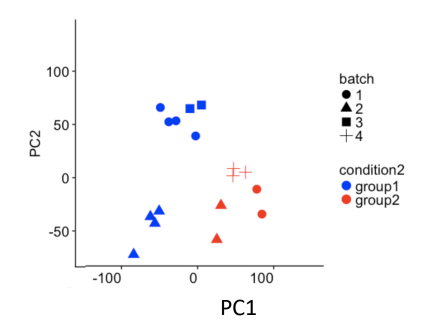
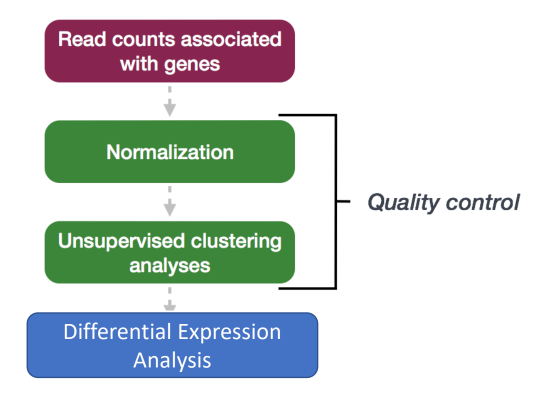
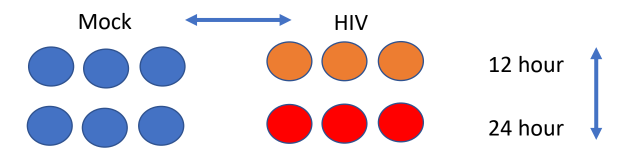


Image https://support.bioconductor.org/p/111491/

# Differential Expression with DESeq2



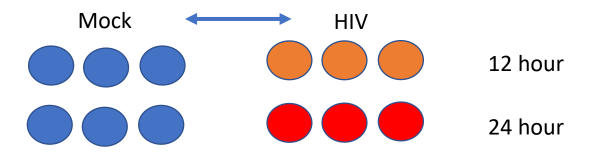
# Multi-factor experiment design



Factor 1: Infection status (Mock or HIV)

Factor 2: Time (12 or 24 hr)

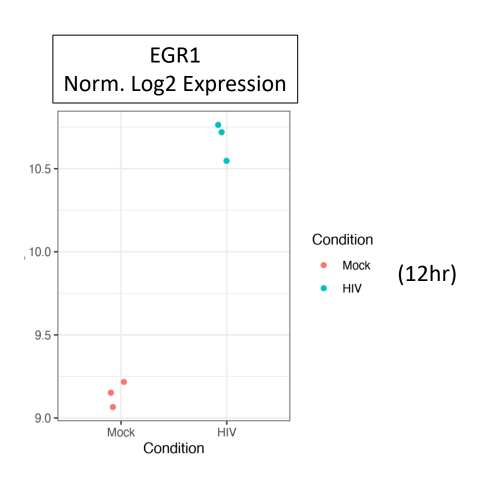
## Multi-factor experiment design



- Differential Expression compares two conditions
- We'll choose Infection status at 12 hr (Mock or HIV) for comparison
- We could also choose time, or a combination of multiple factors

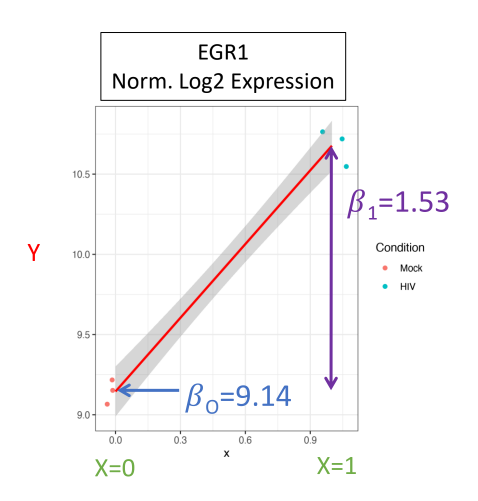
# Step 1: Modeling gene expression values

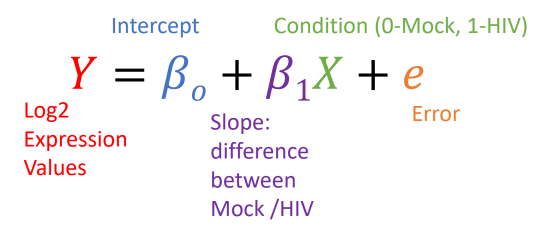
All leading DE tools use regression models to estimate the fold change between conditions for each gene



# Step 1: Modeling gene expression values

All leading DE tools use **regression models** to estimate the fold change between conditions for **each gene** Example, simple linear regression:

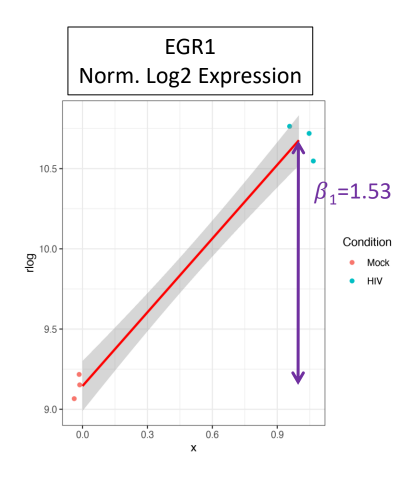




DESeq2 uses a Generalized Linear Model with a Negative Binomial error Distribution, which has been shown to be best fit for RNAseq data.

Love, M.I., Huber, W. & Anders, S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biol* **15**, 550 (2014).

#### Step 2: Hypothesis Testing



Is EGR1 differentially expressed?

$$H_0: \beta_1 = 0$$
 vs.  $H_A: \beta_1 \neq 0$ 

H<sub>o</sub>: there is no systematic difference between the average read count values for Mock vs. HIV

- Statistical test Wald test (similar to t-test) on  $\beta_1$
- $Z = \beta_1/SE_{\beta_1}$
- Z-statistic is compared to the normal distribution and probability of getting a statistic at least as extreme is computed

### DESeq2 Results table

GeneID	Base mean	log2FoldChange	StdErr	P-value	P-adj
EGR1	1273	1.55	0.13	1.19e-77	1.52e-73
MYC	5226	-1.53	0.14	1.63e-36	1.03e-32

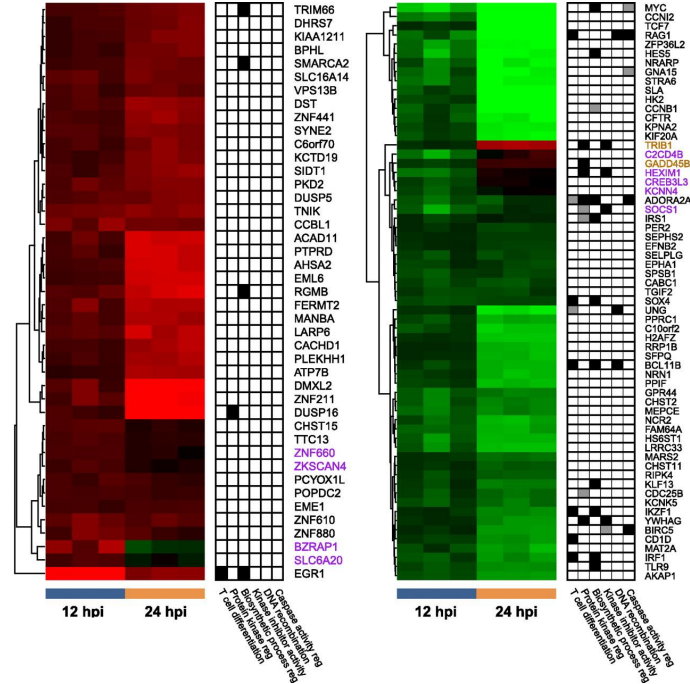
- Mean of normalized counts averaged over all samples from two conditions (HIV, Mock)
- Log of the fold change between two conditions
- StdErr Standard error of coefficient (e.g. b<sub>1</sub>)
- P-value the probability that the Wald statistic is as extreme as observed if H<sub>O</sub> were true
- P-adj accounting for multiple testing correction

# Study findings

 T cell differentiation-related genes were overrepresented in the DEG at 24hr

'Large-scale disruptions to host transcription' at 24hr





#### References

#### DESeq2 vignette (R/Rstudio):

http://www.bioconductor.org/packages/release/bioc/vignettes/DESeq2/inst/doc/DESeq2.html#dif ferential-expression-analysis

HBC Training (Command line/R):

https://hbctraining.github.io/DGE workshop

Galaxy Training:

https://galaxyproject.org/tutorials/rb\_rnaseq/

Next: Introduction To Galaxy