

Whole transcriptome sequencing data analysis workshop

Quality control, trimming,
alignment and quantification

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1-8/02/2019

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Introduction

- Most of the genome does not appear in RNA-seq data: in humans, genes only constitute approx. 3% of the genome.

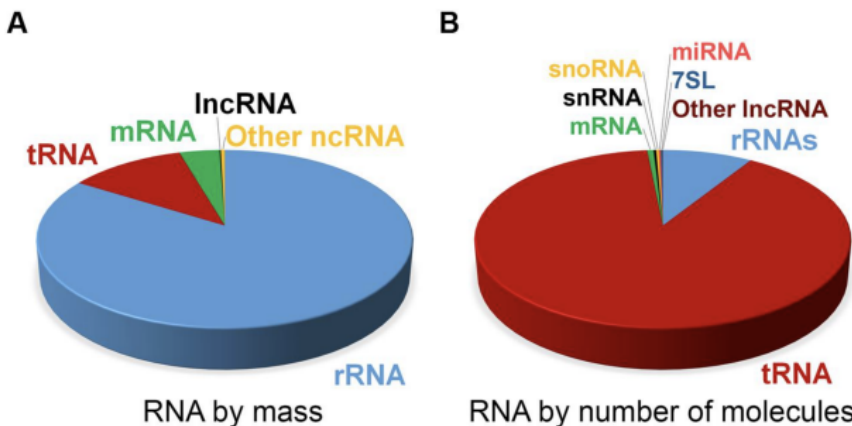


FIGURE 1. Estimate of RNA levels in a typical mammalian cell. Proportion of the various classes of RNA in mammalian somatic cells by total mass (A) and by absolute number of molecules (B). Total number of RNA molecules is estimated at roughly 10^7 per cell. Other ncRNAs in (A) include snRNA, snoRNA, and miRNA. Note that due to their relatively large sizes, rRNA, mRNA, and IncRNAs make up a larger proportion of the mass as compared to the overall number of molecules.

Overview of sequencing technologies

- First generation: microarrays (mostly replaced by RNA-seq).
- **Next generation sequencing or second generation: RNA-seq (mostly Illumina).**
- Third generation: long-reads (mostly PacBio and Oxford nanopore). It is growing in parallel, mostly useful for de-novo assembly, new transcript discoveries and transcript level analyses.
- “Recent” developments of RNA-seq: single-cell RNA-seq (scRNA-seq). Similar to “bulk RNA-seq” (standard RNA-seq) but with a single-cell resolution, i.e. observations are available for single cells instead of aggregating expression over many cells.

Protocol: Single-end vs. paired-end reads

- Single-end:
 - ▶ reads are sequenced from one end only;
 - ▶ cheaper and faster;
 - ▶ reads are approx. 100/150 base pairs (bp) long.
- Paired-end (fragments):
 - ▶ reads are sequenced from both ends;
 - ▶ more expensive but the most popular protocol nowadays;
 - ▶ fragments are approx. 250-800 (bp) long, i.e. approx. 100/150 at each end plus a gap in between (of known length);
 - ▶ more accurate as it reduces the number of multi-mapping reads.

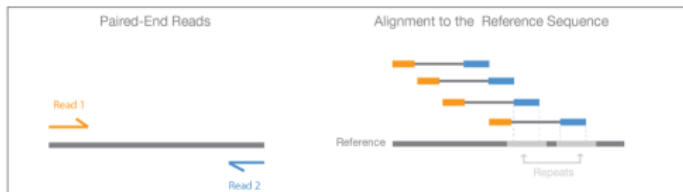


Figure 4: Paired-End Sequencing and Alignment—Paired-end sequencing enables both ends of the DNA fragment to be sequenced. Because the distance between each paired read is known, alignment algorithms can use this information to map the reads over repetitive regions more precisely. This results in better alignment of reads, especially across difficult-to-sequence, repetitive regions of the genome.

Protocol: Stranded vs. unstranded

- Unstranded:
 - ▶ we don't know what strand reads come from.
- Stranded:
 - ▶ we know what strand reads come from:
 - ▶ it decreases the number of ambiguous reads, e.g., when reads fall in regions overlapping between multiple genes.
- Ideal scenario:
 - ▶ ~150 bps long reads;
 - ▶ paired-end reads;
 - ▶ stranded reads;
 - ▶ “many” samples per condition (e.g., ≥ 6 per group).

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FASTQ file

- The raw reads are stored in a FASTQ file.
 - Each read is represented in 4 lines:
 - ▶ 1) a header for the read (starting with @);
 - ▶ 2) the sequence itself;
 - ▶ 3) a header for the quality (starting with +);
 - ▶ 4) a quality score as long as the sequence: each base of the sequence is associated to a score.

@HWI-ST1034:40:C08PJACXX:2:1101:20681:1994 1:N:0:ATCACG
CTCGNAGACTGGCAACTTGTCTGGTTACTGCACCTCTTTAAAGGCAGAAAGGC
+
CCCC#2ADHHHCHIIIIIIIIIIIIIIIBGIIIIIIIIIIIIIIIIIIIIIIIIII

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Phred score

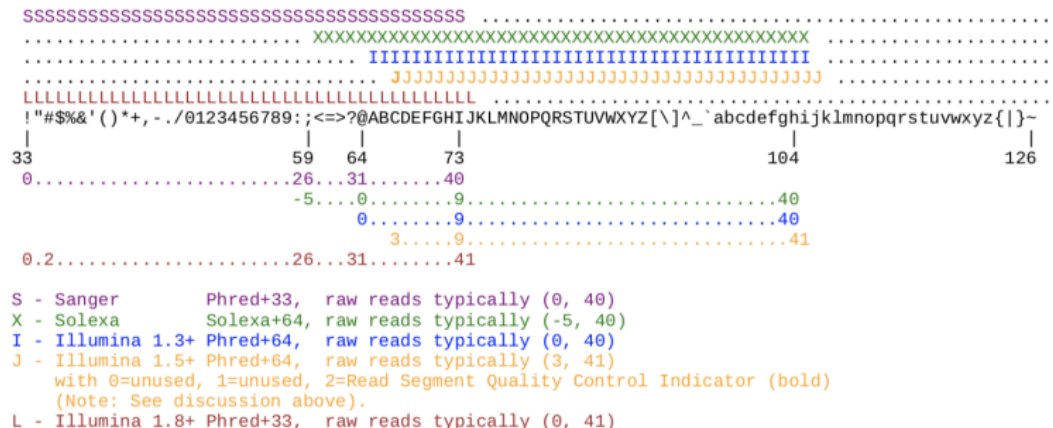
- The Phred score, Q , associates each base with the probability that the base is called incorrectly, P .
- Phred score: $Q = -10\log_{10}P$.
- A Phred score of 30 or more is considered to be good enough: the error probability for a single base $< 0.1\%$.

Phred Quality Score	Probability of incorrect base call	Base call accuracy
10	1 in 10	90%
20	1 in 100	99%
30	1 in 1000	99.9%
40	1 in 10,000	99.99%
50	1 in 100,000	99.999%
60	1 in 1,000,000	99.9999%

Wikipedia

Phred score

- The association between the Phred score and the characters of the FASQT file changes with the technology.
- For recent Illumina technologies, capital letters have good quality: error probability for a single base < 0.1%.



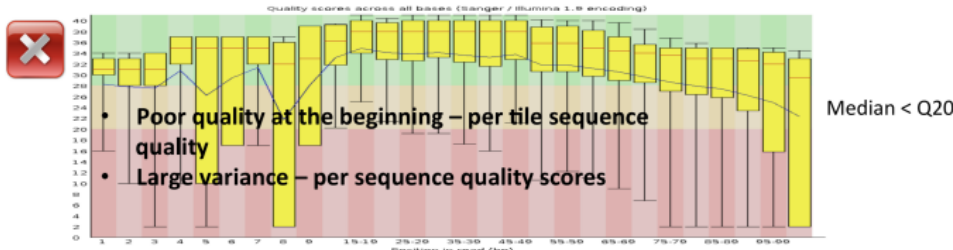
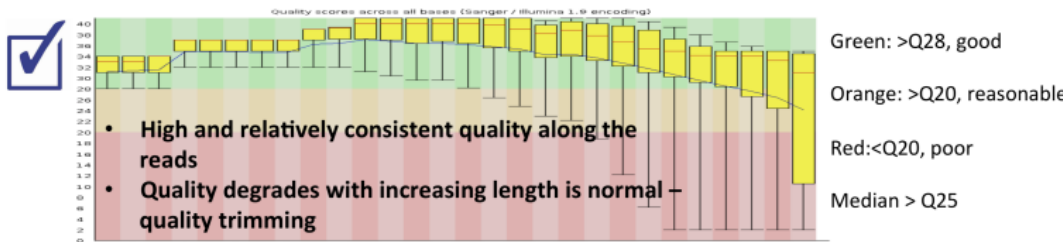
Assessing the quality of reads

- We can assess the quality of the raw reads, before aligning them to the genome/transcriptome, via **FastQC**.
- FastQC outputs one file per sample for single-end reads and two files for paired-end reads.
- To aggregate and jointly visualize the output of FastQC for multiple samples, you can use **MultiQC**.
- In next slides we will inspect the output from FastQC in individual samples.
- Nice interpretation of FastQC output in:
https://www.youtube.com/watch?v=GnWSXwQeJ_U
How to Check the Quality of Illumina Sequencing Reads with FastQC (Part 2).
- Interpretation of MultiQC output in:
[https://www.youtube.com/watch?v=qPbI1O_KWNO.](https://www.youtube.com/watch?v=qPbI1O_KWNO)
Using MultiQC Reports.

FastQC

Per base sequence quality - FastQC

- Range of quality values across all bases at each position

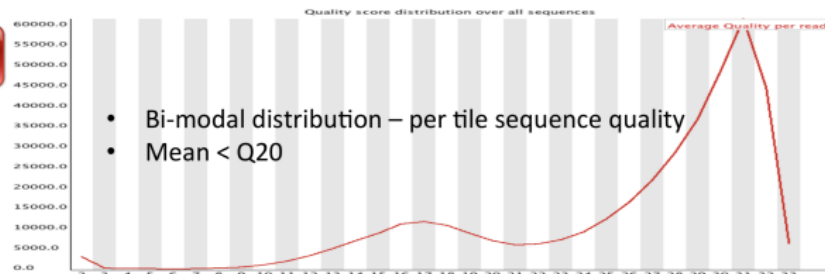
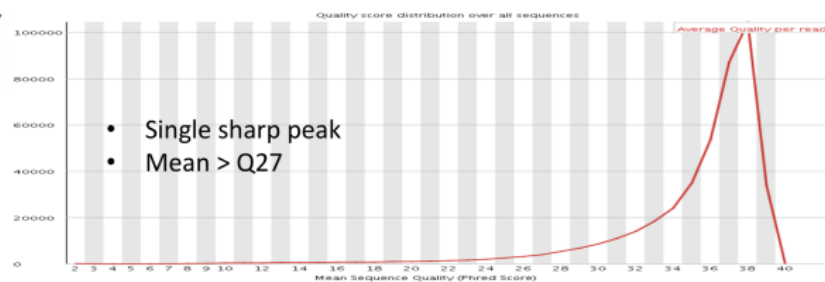


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FastQC

Per sequence quality scores - FastQC

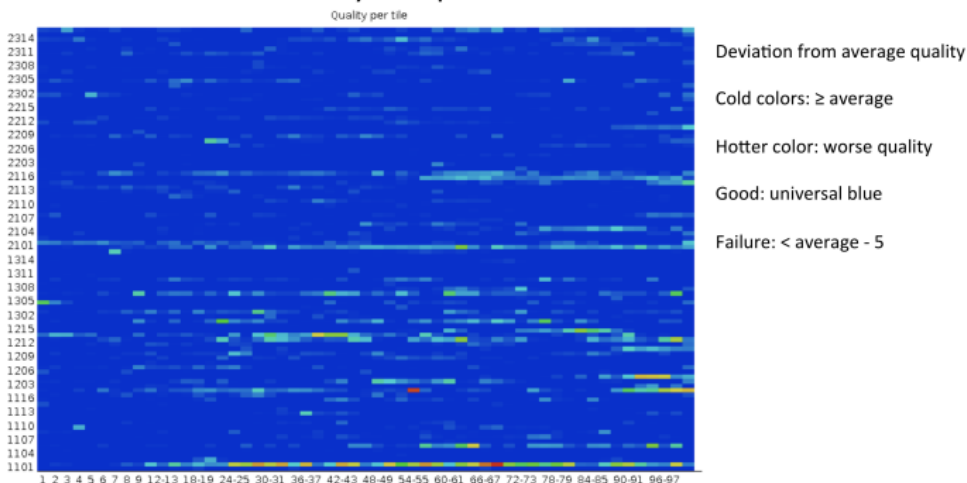
- Subset of sequences with universally low quality values



FastQC

Per tile sequence quality - FastQC

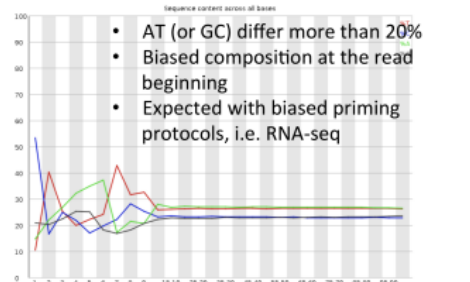
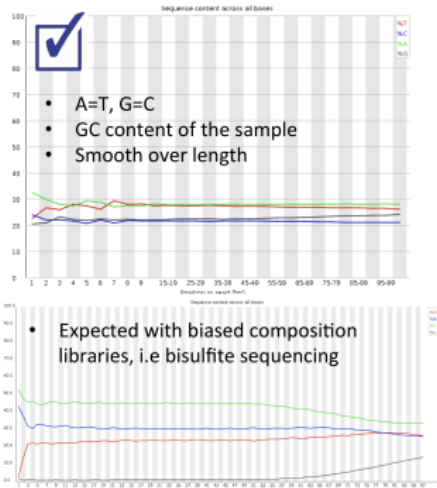
- Quality scores from each tile across all bases - loss in quality associated with only one part of the flowcell



FastQC

Per base sequence content - FastQC

- The portion of A, T, G, and C at each position



Biases in Illumina transcriptome sequencing caused by random hexamer priming

Kasper D. Hansen^{1,*}, Steven E. Brenner² and Sandrine Dudoit^{1,3}

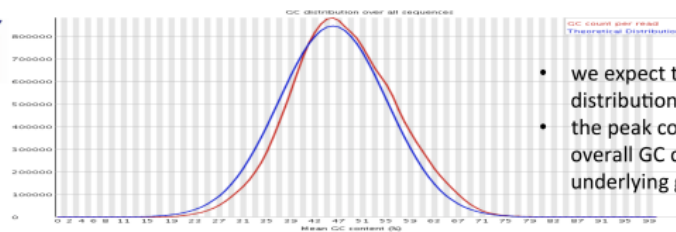
Treatment of DNA with bisulfite converts cytosine to uracil, but leaves methylated cytosine unaffected. Therefore, DNA that has been treated with bisulfite retains only methylated cytosines.

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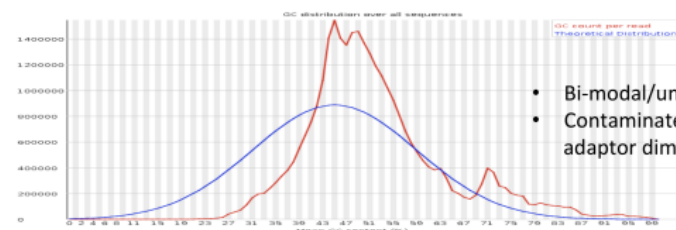
FastQC

Per sequence GC content - FastQC

- Distribution of average GC in all reads



- we expect to see a roughly normal distribution of GC content
- the peak corresponds to the overall GC content of the underlying genome



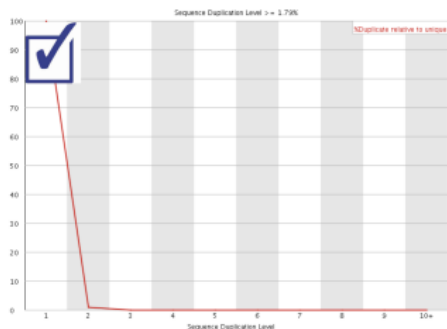
- Bi-modal/unusual distribution
- Contaminated/biased subset, i.e. adaptor dimmers, rRNA etc

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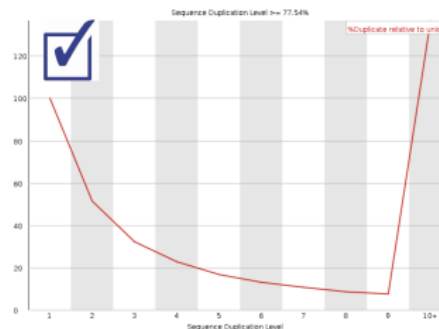
FastQC

Sequence duplication - FastQC

- Relative number of sequences with different degrees of duplication



- Essentially no duplication



High duplication levels:

- DNA-seq: PCR over amplification, too little input material
- Normal in RNA-seq: high expression

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FastQC

Overrepresented sequences - FastQC

- Sequences make up >0.1 % of the total
- Compare those with a contamination database for finding contamination (i.e. adaptor dimmers)



Overrepresented sequences

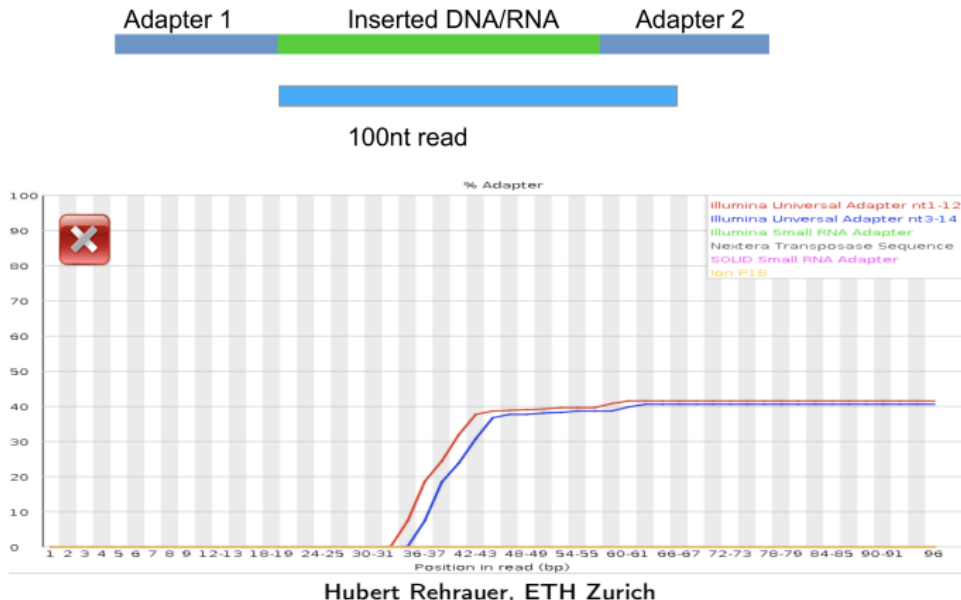
Sequence	Count	Percentage	Possible Source
GGAAGAGCACACGTCTGAACTCCAGTCACCGATCATCTGTATGCCGTC	75874	1.5613887498682963	TruSeq Adapter, Index 7 (100% over 50bp)
GGAAGAGCACACGTCTGAACTCCAGTCACCGATGTATCTGTATGCCGTC	7636	0.15713900010536297	TruSeq Adapter, Index 2 (100% over 50bp)
GGAAGAGCACACGTCTGAACTCCAGTCACACAGTGATCTGTATGCCGTC	7539	0.1551428656095248	TruSeq Adapter, Index 5 (100% over 50bp)
GGAAGAGCACACGTCTGAACTCCAGTCACGCCAATATCTGTATGCCGTC	5117	0.10530123933199874	TruSeq Adapter, Index 6 (100% over 50bp)

- Can be normal and biologically meaningful
 - highly expressed transcripts
 - high copy number repeats
 - Less diverse library (amplicons)

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FastQC

Adapter Content - FastQC



FastQC



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- If the fragment of DNA/RNA is too short, our read could include part of the adapter.

FastQC

Data preprocessing common tasks

1. Trimming: remove bad bases from (end(s) of) reads
 - Adaptor sequence
 - Low quality bases
2. Filtering: remove bad reads
 - Low quality reads
 - Contaminating sequences
 - Low complexity reads (repeats)
 - Short (<20bp) reads – they slow down mapping software

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Trimming

- Trimming is always recommended.
- Trimming removes “low” quality bases from the end (and eventually the start) of reads and adaptor sequences; it also removes entire reads if, after trimming, they are too short (e.g., < 20 bps).
- Two popular tools for trimming reads are **trimmomatic** and **Trim Galore**.
- Trim Galore user guide: https://github.com/FelixKrueger/TrimGalore/blob/master/Docs/Trim_Galore_User_Guide.md
- We can either remove the end of the reads when they fall below a minimum quality (preferred choice) or we can trim after a fixed number of bases.
- After trimming, we need to check again for the quality of trimmed reads to make sure quality is now adequate.

QC examples

- Visualize FastQC and MultiQC outputs before and after trimming reads.

Considerations

- In DNA studies there is a lot of focus on the accuracy of the individual bases, because a base calling error (when we identify the wrong nucleotide) might indicate a variation in the genome, e.g. a single nucleotide polymorphism (SNP).
- In RNA-seq studies, single base calling errors are less relevant: the main aim here is to align reads correctly to a reference (genome or transcriptome) and quantify the relative abundance; i.e., how many reads map to a gene/transcript/exon/locus.
- In this regard the alignment of a fragment is unlikely to be compromised by a single base calling error.

QC & trimming ex. code

FastQC:

```
fastqc -o out_directory sample_1_R1.fastq.gz
```

```
fastqc -o out_directory sample_1_R2.fastq.gz
```

MultiQC:

```
cd out_directory
```

```
multiqc . --interactive
```

% multiqc will merge all fastQC files present in the folder.

Trim Galore:

```
trim_galore -q 20 -phred33 -length 20 -illumina -o out_directory \
-paired sample_1_R1.fastq.gz sample_1_R2.fastq.gz -fastqc -gzip
```

% -q 20, bps with Phred scores < 20 will be trimmed

% -length 20, remove reads if, after trimming, they become shorter than
20 bps

% -illumina, use illumina standard adapters

% -o, directory where the output will be stored

% -paired, use paired end reads (if one read is removed, also remove the
other one)

% -fastqc, run fastqc on the outputted fastq files

% -gzip, compress the output file

3. Alignment

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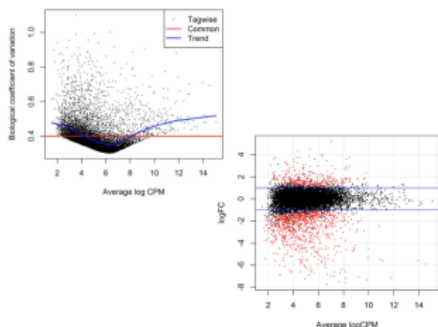
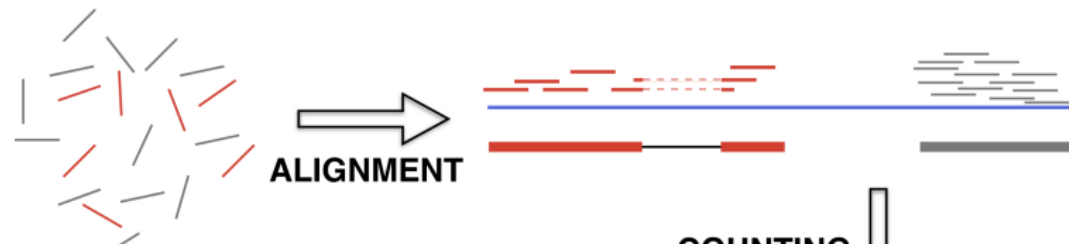
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Alignment



COUNTING

ANALYSIS

Gene A 7 . . .
Gene B 13 . . .
.
. . .
Gene X . . .

A large bracket on the right side of the table groups all the rows together.

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Genome alignment

- In most analyses, after checking the quality of our reads, we align them to a reference.
- We can use a full aligner to align reads to a reference genome (**STAR**, **TopHat**, ...).
- Full aligners attempt to align the entire read to a reference genome (splice-aware alignment).
- To align our reads to the genome we need:
 - ▶ a reference genome (DNA), usually in a fasta format;
 - ▶ a gene transfer format (GTF) file which contains the location of genes on the reference genome.
- Both can be downloaded from
<http://www.ensembl.org/info/data/ftp/index.html> under the DNA and Gene sets columns.

Considerations on genome alignment

- Genome aligners take into account mismatches with respect to the reference, due to sequencing error (approx. 1/500) or mutations (approx. 1/10,000 in humans).
- A read could align in multiple positions of the reference genome/transcriptome, with the same or similar alignment scores. The output is represented by both unique aligning reads and multi mapping reads. Some reads (a minority) remain unmapped because they don't align well enough to any location of the genome.
- Alignment is an optimization problem: for every read, it looks for the alignment with the highest score.
- Aligners do not run a complete search of all possible alignments of all read, the optimization is heuristic, not optimal (yet very sophisticated).

Transcriptome alignment

- Alternatively, we can align our reads directly to a reference transcriptome.
- Transcript aligners are typically pseudo/quasi aligners (**Salmon**, **kallisto**, ...).
- Pseudo/quasi aligners don't align the full reads, instead they use a low cost pseudo alignment:
 - ▶ from each read they create many k-mers (substrings of the read of k base pairs);
 - ▶ they map the k-mers to the reference transcriptome and check their compatibility with the transcripts.
- To align our reads to the transcriptome we need a reference transcriptome (cDNA) alone, which can be downloaded from <http://www.ensembl.org/info/data/ftp/index.html> under the cDNA column.

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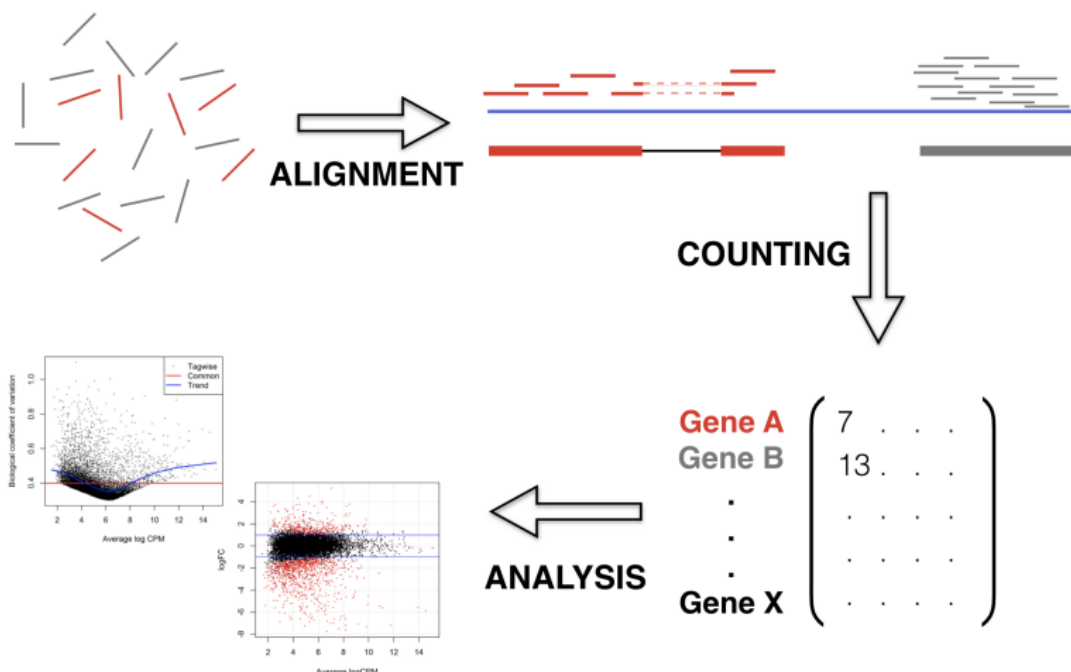
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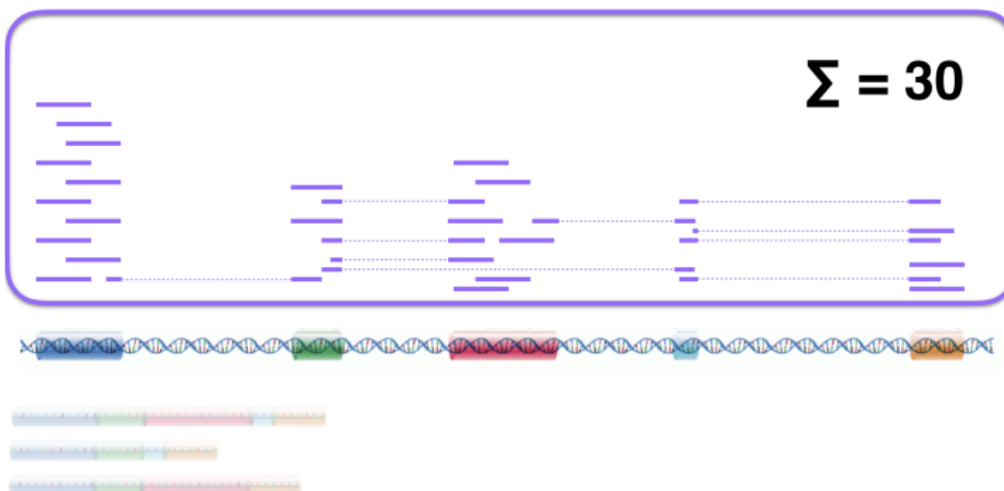
Counting



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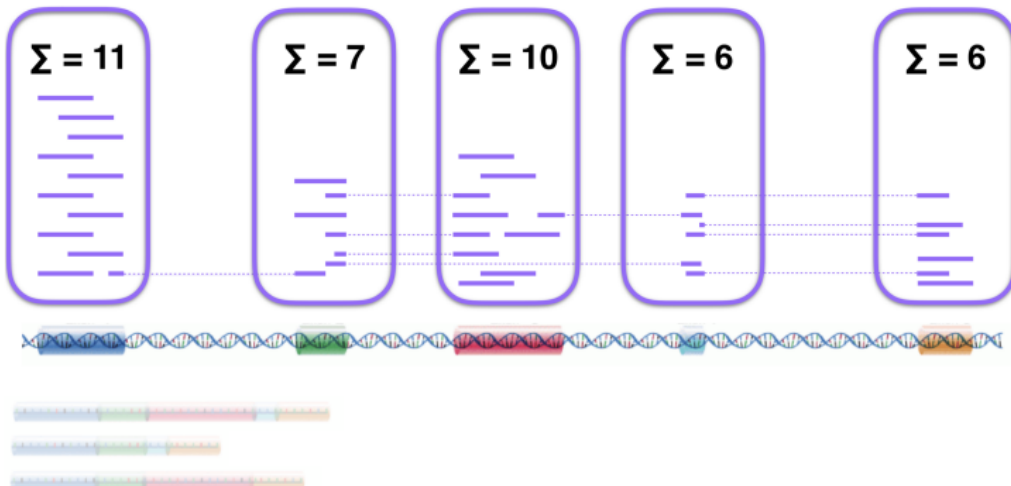
Quantification with genome aligners: gene counts

- Once the reads are aligned to the genome, we need to sort them (for STAR use the *SortedByCoordinate* option).
- We can then count how many reads overlap each gene.
- STAR** and **TopHat** are the most popular genome aligners (we'll see STAR in the tutorial).
- Remember that RNA-seq reads only map to exons!



Quantification with genome aligners: exon counts

- The counts of each individual transcript cannot be obtained due to the overlapping regions between transcripts.
- To study the transcript, the attention sometimes shifts towards the exons, for which we can observe the counts.



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Quantification with genome aligners: junction counts

- An alternative is to only consider junction counts: counts of reads that span over two exons (the ones with the dotted lines in the previous image).
- Pros:
 - ▶ we know what exons reads connect;
 - ▶ useful for discovering new transcripts with non annotated exon junctions (STAR also outputs non-annotated junction reads).
- Cons:
 - ▶ we still miss what transcript each read maps to (there could be > 1 transcripts associated to the two exons the reads spans over);
 - ▶ we only use a sub-set of the data.
- STAR outputs junction counts in the SJ.out.tab file.

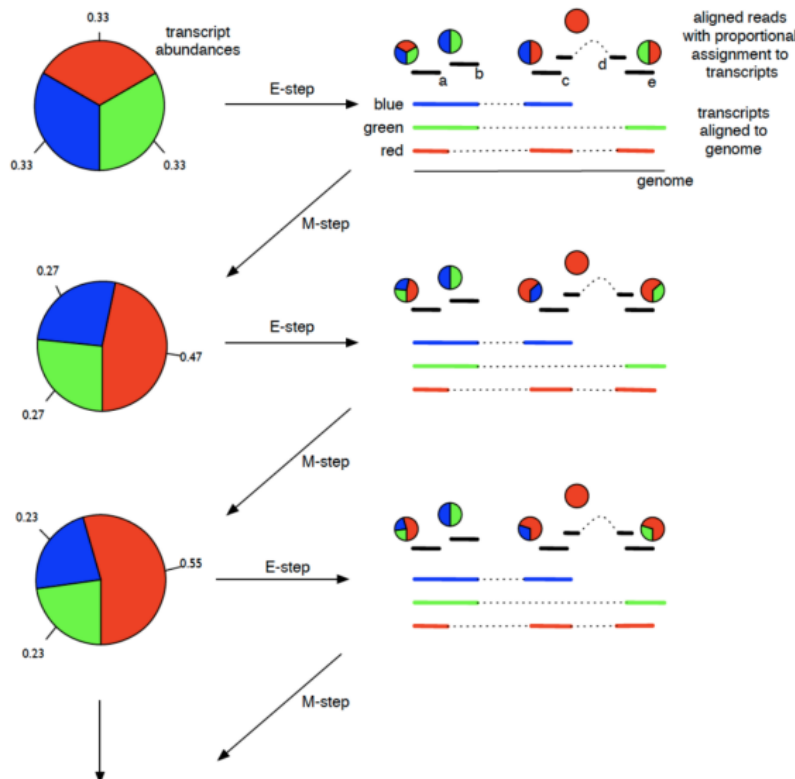
Quantification with transcriptome aligners

- Reads can be mapped directly to the transcriptome, skipping genome mapping.
- A well annotated transcriptome is required: if a transcript is missing in the reference, the reads coming from that transcript will either be mapped to other transcripts (if compatible), or remain unmapped.
- Multi-mapping reads are a lot more frequent when mapping to the transcriptome: for every exon, there can be multiple transcripts that contain it.
- Pseudo aligners don't actually map the full reads, but k-mers built from the reads.
- **Salmon** and **kallisto** are the most popular transcript aligners.

Quantification with transcriptome aligners

- After mapping reads to transcripts, an expectation-maximization (EM) algorithm is used to estimate the expected number of reads mapping each transcript.
- The algorithm outputs estimated transcript level counts, from which we can easily obtain gene level counts.
- Note that these numbers are estimates, not real counts.
- Salmon and kallisto provide bootstrap replicates to measure the uncertainty in the estimated counts.

Quantification with transcriptome aligners



Reference-free assembly

- An alternative to genome and transcript mapping is to use a reference-free approach and assemble the reads into transcripts, build a reference transcriptome and then map them back to the reference.
- Very challenging to reconstruct full transcripts from Illumina short reads, even if paired-end reads.
- **Trinity** is the most popular tool for reference-free assembly: Grabherr et al. (2011). Trinity: reconstructing a full-length transcriptome without a genome from RNA-Seq data, Nature Biotechnology.
- In this context it is highly suggested to have:
 - ▶ paired-ended,
 - ▶ stranded,
 - ▶ “long” RNA-seq reads (e.g., 150 bps).
- Trinity will use all reads from all samples (across all experimental conditions) when building the transcriptome.
- Rarely used in well studied organisms.

Trinity wiki

- Trinity wiki:
<https://github.com/trinityrnaseq/trinityrnaseq/wiki>
[https://github.com/trinityrnaseq/trinityrnaseq/wiki/
Running-Trinity](https://github.com/trinityrnaseq/trinityrnaseq/wiki/Running-Trinity)
- Trinity tutorial: [https://biohpc.cornell.edu/lab/doc/
trinity_workshop_part1.pdf](https://biohpc.cornell.edu/lab/doc/trinity_workshop_part1.pdf).

Importing the counts

- For each sample, the main output of the quantification will be a matrix of genes/transcripts and respective counts, i.e. the number of reads mapping to a gene/transcript.
- Remember that counts are discrete, not continuous.
- The estimated counts from Salmon or kallisto are continuous though because they are expected counts.
- We can import the counts from STAR/TopHat output (stored in a .bam or .sam file) in R via featureCounts (**Rsubread**), see tutorial.
- We can import the counts from Salmon/kallisto output (stored in a .bam or .sam file) in R via tximport (**tximport**), see tutorial.

Genome alignment ex. code I

% STAR requires a genome annotation (in a fasta format) and the location of the annotated transcripts on the genome (in a gtf file).

STAR:

% Generate genome index:

```
star --runMode genomeGenerate --runThreadN 20 --genomeDir
```

```
genome_dir \
```

```
--genomeFastaFiles ensembl_Homo_sapiens.GRCh37.71.fa \
```

```
--sjdbGTFfile Homo_sapiens.GRCh37.71.gtf --sjdbOverhang 100
```

% --runMode genomeGenerate, to generate the genome index.

% --runThreadN 20, runs on 20 cores.

% --genomeDir genome_dir, specifies the output directory for the genome index.

% --sjdbGTFfile specifies the path to the file with annotated transcripts in the standard GTF format: STAR will extract splice junctions from this file and use them to greatly improve accuracy of the mapping. % While this is optional, and STAR can be run without annotations, using annotations is highly recommended whenever they are available. % sjdbOverhang should be equal to the (average) read length - 1 (i.e., 100 for 101 bp long reads).

Genome alignment ex. code II

STAR:

% Align (paired-end) reads; for each sample:

```
star --runMode alignReads --runThreadN 20 --genomeDir genome_dir \
--readFilesIn sample_1_R1_trim.fastq.gz sample_1_R2_trim.fastq.gz \
--outFileNamePrefix sample1 --outSAMtype BAM SortedByCoordinate
```

% STAR download: <https://github.com/alexdobin/STAR>

% STAR manual: <https://github.com/alexdobin/STAR/blob/master/doc/STARmanual.pdf>

Transcriptome alignment ex. code

% salmon documentation:

<https://salmon.readthedocs.io/en/latest/>

% salmon release:

<https://github.com/COMBINE-lab/salmon/releases>

% salmon only requires a transcript annotation (transcriptome.fasta).

Salmon:

% Build the index:

```
salmon index -i index_dir -t transcriptome.fasta -p 30 -type quasi -k 31
```

% For every sample:

```
salmon quant -i index_dir -l A \
-1 sample_1_R1_trim.fastq.gz -2 sample_1_R2_trim.fastq.gz \
-p 20 -o sample1_out_dir -dumpEq -numBootstraps 100 \
-seqBias -gcBias
```

% -p 20 specifies that salmon will use 20 cores

% -numBootstraps 100, computes 100 bootstrap replicates (used by some differential tools)

% -l A specifies the library type (A = automatic detection, if unknown).

% -seqBias and -gcBias are options to correct for sequence-specific and GC content biases.

De novo assembly ex. code

% Trinity relies on salmon and bowtie2: both need to be installed.

Trinity:

```
Trinity -seqType fq -max_memory 400G \
-left sample_1_R1_trim.fastq.gz,sample_2_R1_trim.fastq.gz \
-right sample_1_R2_trim.fastq.gz,sample_2_R2_trim.fastq.gz \
-CPU 40 -output out_dir -SS_lib_type RF
% By default, reads are treated as not strand-specific.
% -SS_lib_type, specifies the strand orientation (RF = reverse forward).
% -genome_guided_bam: genome guided mode, in case a (reliable)
genome is available.
```

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Comparability

- When comparing the expression of different genes/transcripts, keep in mind that genes/transcripts have different lengths: if two genes/transcripts are equally expressed, but one is twice as long, it will have about twice as many reads mapping to it.
Keep in mind that, given the same expression levels, we have more power* to detect differential behaviour in longer genes/transcripts than in shorter ones. *Statistical power = $\Pr(\text{detecting an effect if the effect is present})$.
- When comparing one gene/transcript across different samples, beware of:
 - possible batch effects;
 - sequencing biases (Salmon tries to correct for it via the option *seqBias*);
 - library size, i.e. the total number of reads in the sample, across all genes.
- We can calculate normalization factors to make counts comparable between different experiments. Remember that, except a few cases, the counts are not scaled themselves: we compute normalization factors to include in the model.

Normalization

How to calculate normalization factors?

- Attempt 1: **total count** (library size)
 - Define a reference sample (one of the observed samples or a “pseudo-sample”) - gives a “target library size”
 - Normalization factor for sample j is defined by

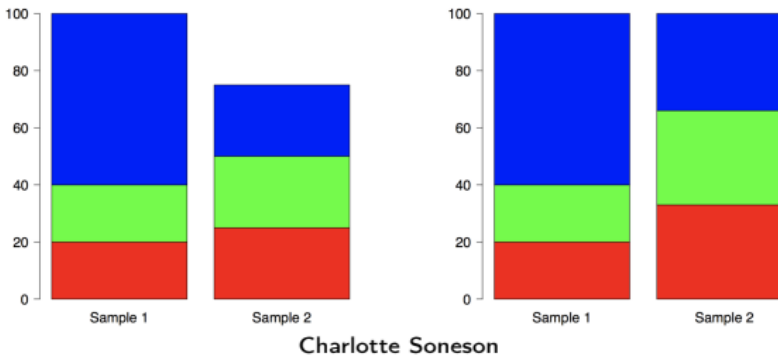
$$\frac{\text{total count in sample } j}{\text{total count in reference sample}}$$

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Normalization

The influence of RNA composition

- Observed counts are relative
- High counts for some genes are “compensated” by low counts for other genes



Normalization

How to calculate normalization factors?

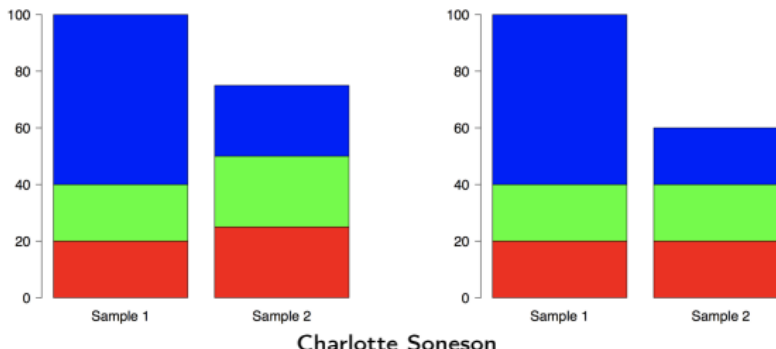
- Attempt 2: total count (library size) * compensation for differences in composition
- Idea: use only non-differentially expressed genes to compute the normalization factor
- Implemented by both edgeR (TMM) and DESeq2 (median count ratio)
- Both these methods assume that most genes are not differentially expressed

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Normalization

How to calculate normalization factors?

- Attempt 2: total count (library size) * compensation for differences in composition



Definition of expression levels

- We introduce the following elements:
 - ▶ X_t = number of reads arising from transcript t ;
 - ▶ N = total number of reads sequenced from all genes, i.e. the library size;
 - ▶ l_t = effective length of the transcript t .
We do not consider the actual length of transcripts, but the effective transcript length: the part of the transcript I can map my read on: $l_t = \text{length_transcript} - \text{length_read} + 1$.
 - ▶ $Z = \sum_t \pi_t l_t$, where π_t represents the relative abundance of transcript t . Z is a normalization factor representing the mean effective length of transcripts, weighted by transcript relative abundance.
- CPM = $\frac{X_t}{N} \times 10^6$, counts per million; how many reads map to transcript t every 10^6 reads. CPM can also be expressed at the gene level by replacing the counts mapping to gene g , X_g , instead of X_t . CPM normalizes for the library size.

Definition of expression levels

- $\text{RPKM} = \frac{X_t}{I_t N} \times 10^9$, reads per kilobase per million of mapped reads. It normalizes counts for transcript length (or gene length) and for sequencing depth, i.e. library size.
It is the per base rate of reads/fragments mapping to transcript t , every 10^9 reads. It is also called fragment per kilobase per million of mapped reads, FPKM: RPKM and FPKM are the same, one name refers to reads and one to fragments (paired-end reads).
- $\text{TPM} = \frac{X_t}{I_t N} \times Z \times 10^6$, transcripts per million. TPM is the number of transcripts of type t you expect when sequencing 10^6 transcripts. TPM is usually preferred over RPKM/TPKM. It normalizes for transcript length and for library size.
- Remember, these normalized counts are NOT used for differential expression analyses; however, they are useful for plotting the data.

1. Introduction
2. Quality control & trimming
3. Alignment
4. Quantification
5. Normalizations

References

References |

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Questions?