

Introduction to RNA-Seq – Mapping & Aligning

Wandrille Duchemin







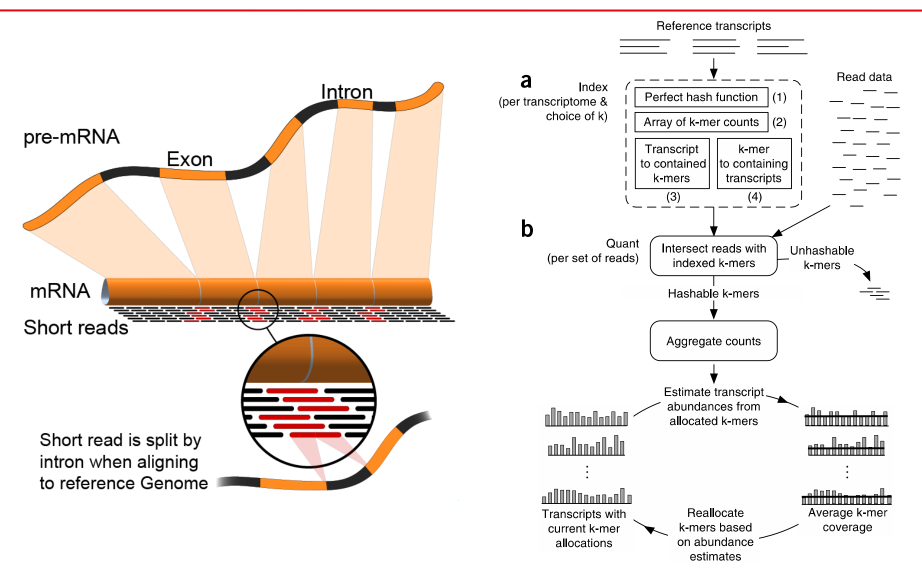


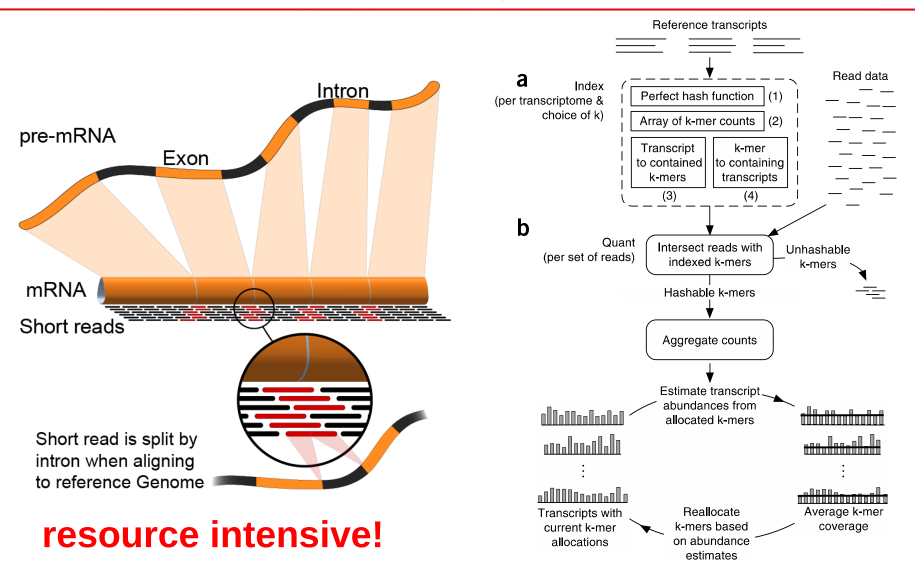




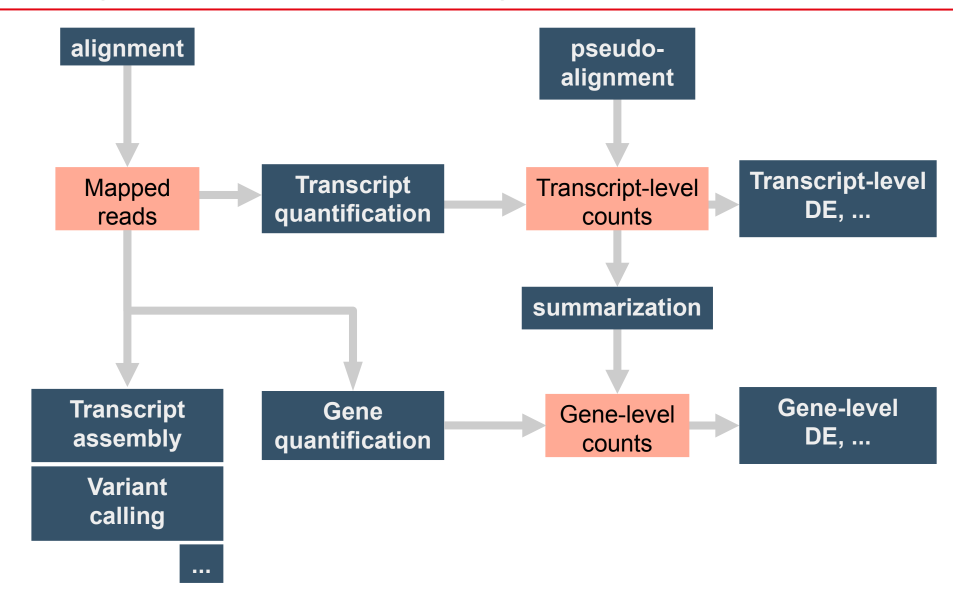


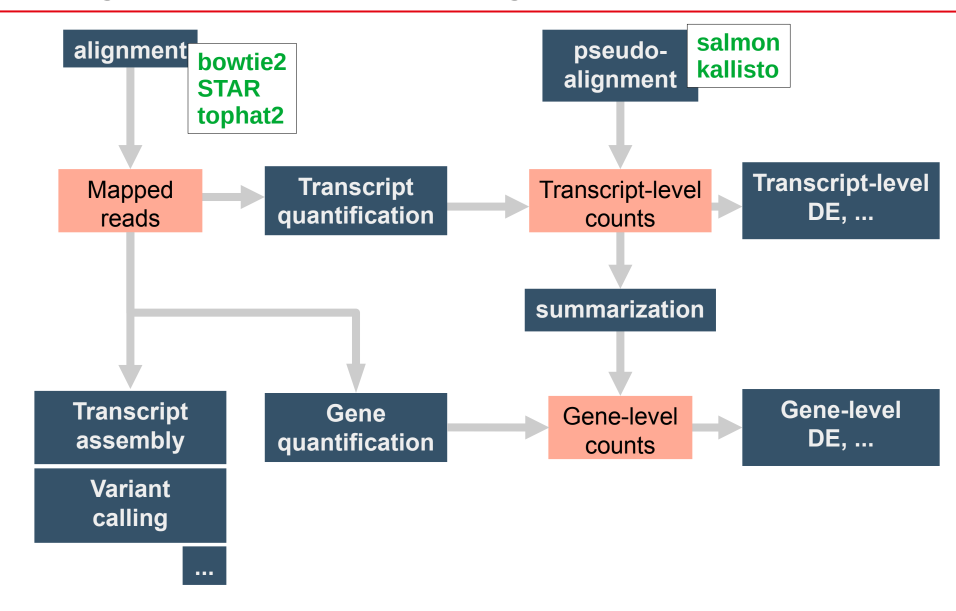


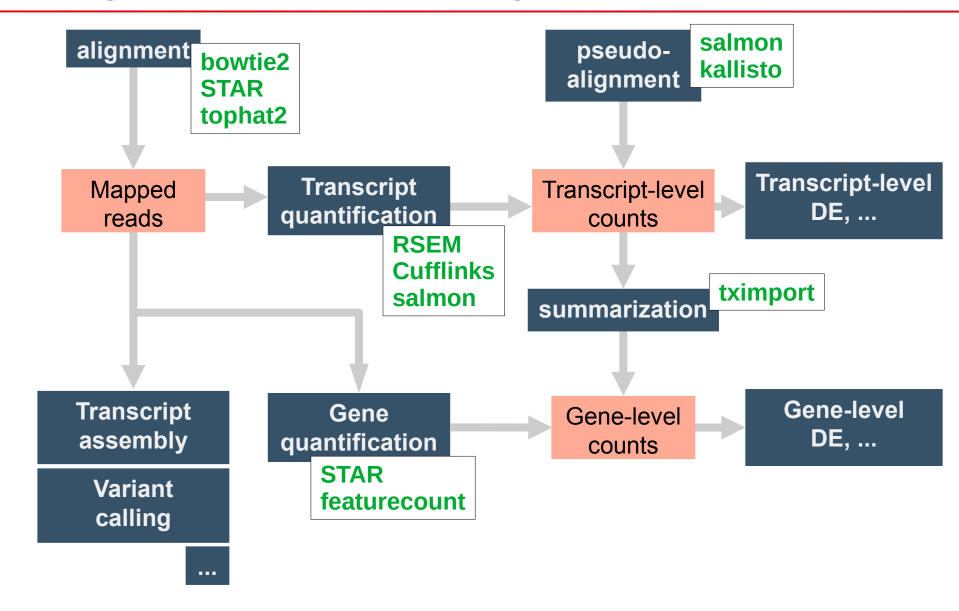


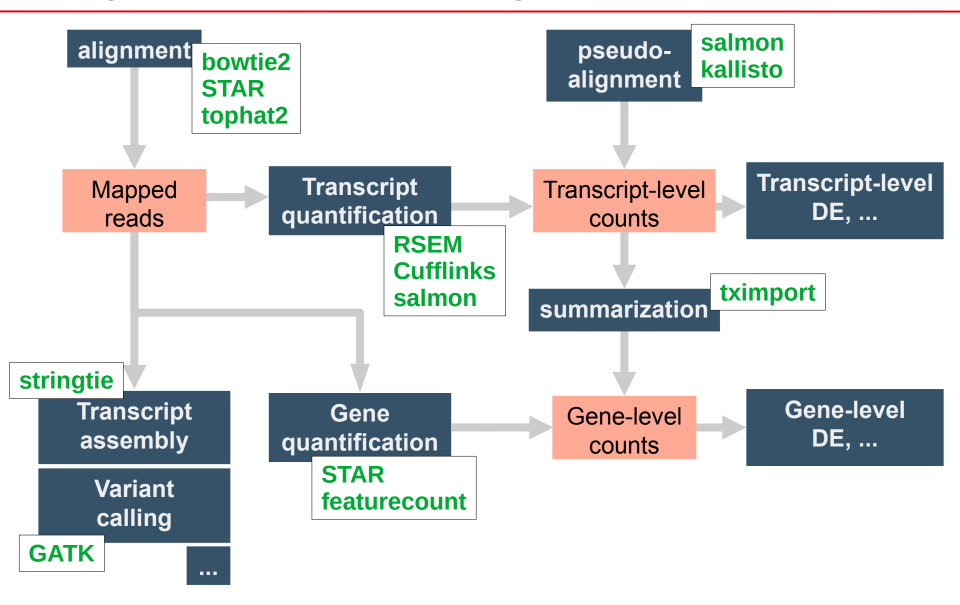




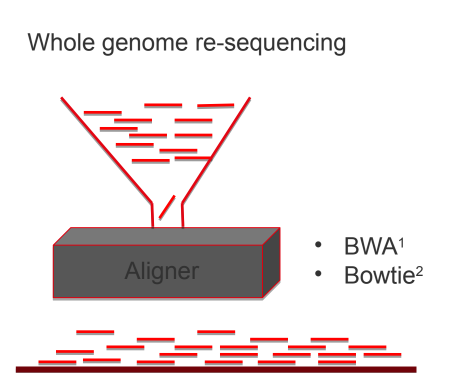


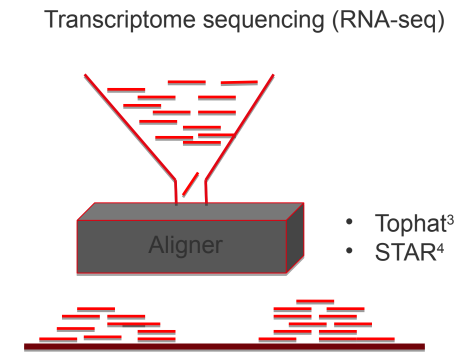






"Aligning" & "Mapping" Sequencing Reads





- 1. Li and Durbin 2009
- 2. Langemead et al. 2009
- 3. Trapenell et al. 2009; Kim et al. 2013
- 4. Dobin et al. 2013

Alignment using STAR

Phase 1 – Mapping using "Maximum Mappable Prefix

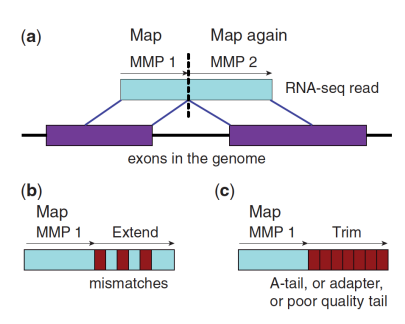
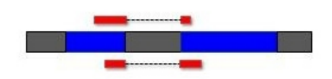


Fig. 1. Schematic representation of the Maximum Mappable Prefix search in the STAR algorithm for detecting (a) splice junctions, (b) mismatches and (c) tails

Phase 2 – "Stitching"



Benchmarking the Aligners (simulated dataset)

	Correctly mapped 200	>=150 bases correctly mapped	Unmapped	True positive junctions		False positive junctions	
	bases			Number	Sensitivity	Number	FDR
Aligner	1	2	4	5	6	7	8
STAR	81.3%	95.0%	4.82%	148,487	92.7%	409	0.3%
TopHat2	82.6%	83.7%	6.70%	135,006	84.3%	1,228	0.9%

• Star : x20 faster than Tophat2

Tophat2 : x6 more memory efficient (can be run on recent laptops)

Dobin & Gingeras 2013

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Essentially, if you have access to a cluster you should be using STAR

Reference Index Preparation

Different for each software!

- Need a suitable reference genome
 - sequence
 - annotation

https://www.ensembl.org/info/data/ftp/index.html

https://hgdownload.soe.ucsc.edu/downloads.html

Genome Annotation Files

- text file describing genomic features
 - Gene, CDS, exon, intron, miRNA, etc
 - Chromosome, start, end, strand, attributes, etc

most common format: gtf / gff3

http://www.ensembl.org/info/website/upload/gff.html

Example GTF

```
MT
    RefSeq
                            2751 3707 .
                                                     gene id "ENSMUSG00000064341";
                                                                                    gene ver
              gene
                                                     gene id "ENSMUSG00000064341";
ΜT
    RefSeq
              transcript
                            2751 3707 .
                                                                                    gene ver
    RefSea
                            2751 3707 .
                                                     gene id "ENSMUSG00000064341";
MT
              exon
                                                                                    gene ver
    RefSea
              CDS
MT
                            2751 3704 .
                                                     gene id "ENSMUSG00000064341";
                                                                                    gene ver
MT
    RefSeq
              start codon
                            2751 2753 .
                                                     gene id "ENSMUSG00000064341";
                                                                                    gene ver
                            3705 3707 .
                                                     gene id "ENSMUSG0000064341";
MT
    RefSeq
              stop codon
                                                                                    gene ver
```

Practical

Go to the website and do the mapping practical

REFERENCES

<u>Li H & Durbin R (2009) "Fast and accurate short read alignment with Burrows-Wheeler transform" Bioinformatics 25(14):1754-60</u>

<u>Langmead et al (2009) "Ultrafast</u> <u>and memory-efficient alignment of short DNA sequences to the human</u> <u>genome" Genome Biology 10(3):R25.</u>

<u>Trapnell *et al* (2009) "TopHat: discovering splice junctions with RNA-Seq" Bioinformatics 25(9):1105-11.</u>

Kim et al (2013) "TopHat2

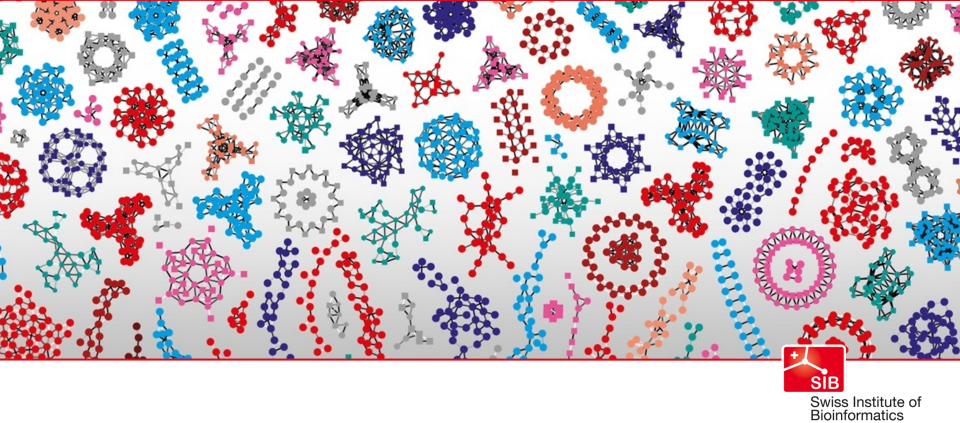
: accurate alignment of transcriptomes in the presence of insertions, deletions and gene fusions" Genome Biology 14(4):R36.

Dobin et al (2013) "STAR: ultrafast universal RNA-seq aligner" Bioinformatics 29(1):15-21.

Patro et al (2014) "Sailfish enables alignment-free isoform quantification from RNA-seq reads using lightweight algorithms" Nature Biotechnology 32(5):462-4.

Patro et al (2017) "Salmon provides fast and bias-aware quantification of transcript expression" Nature Methods 14(4):417-419.

Bray et al (2016) "Near-optimal probabilistic RNA-seq quantification (Kallisto)" Nature Biotechnology 34(5):525-7.



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Annex: Pseudoalignment

transcript-level quantification →

pseudo alignment generally better than alignment

https://www.nature.com/articles/s41598-017-01617-3

https://bmcbioinformatics.biomedcentral.com/articles/10.1186/s12859-021-04198-1

kallisto & salmon : very close, maybe small advantage for salmon

See: https://github.com/mikelove/salmon_kallisto_diffs + publications above

Annex: GTF (GFF2) Annotation Format

- http://www.ensembl.org/info/website/upload/gff.html
- Tab-delimited, empty columns denoted with "."
- Column order:
 - seqname chromosome, scaffold, etc
 - source origin of the annotation, db/project
 - feature gene, transcript, exon, etc
 - start feature start coordinate (1-based)
 - end feature end coordinate (1-based)
 - score floating point, eg quality score
 - **strand** + (forward) or (reverse)
 - **frame** reading frame, 0, 1, or 2
 - attribute semicolon-delimited feature descriptions

GTF vs GFF3

GTF2	GFF3		
same	same		
same	same		
CDS, start_codon, end_codon are required. feature requirements depend on software.	can be anything		
same	same		
same	same		
not used	optional		
same	same		
same	same		
 tag/value delimited by a space each attribute must end with a semi-colon must begin with gene_id and transcript_id attributes Text values must be in quotes ex. gene_id "gene01"; transcript_id "transcript01"; created_by "Damian"; 	 tag/value delimited by '=' each attribute delimited by semi-colon there are a list of pre-defined attributes here must have a unique ID attribute ex. ID=geneA;Parent=geneAP;Name=geneA 		
	same CDS, start_codon, end_codon are required. feature requirements depend on software. same not used same same • tag/value delimited by a space • each attribute must end with a semi-colon • must begin with gene_id and transcript_id attributes • Text values must be in quotes • ex. gene_id "gene01"; transcript_01";		

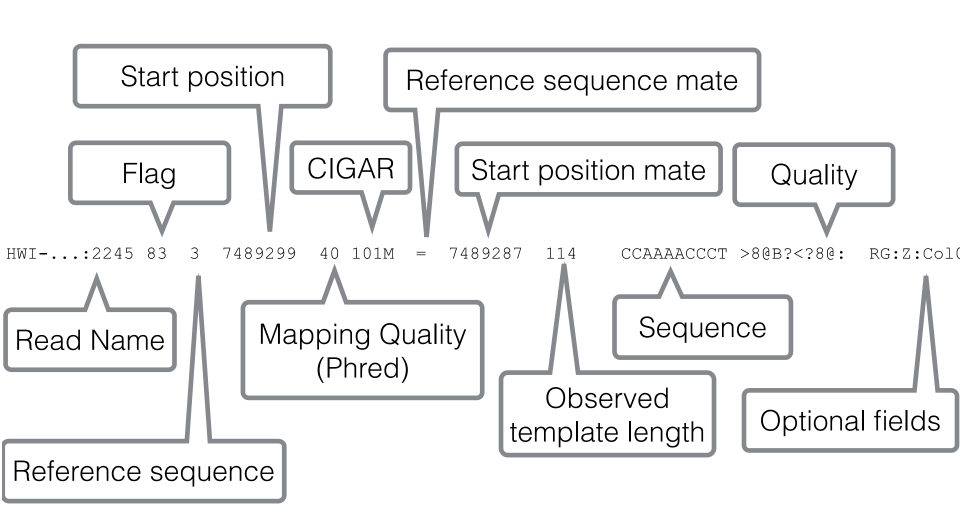
http://blog.nextgenetics.net/?e=27

Annex: SAM Alignment Format

```
@SQ
       SN:1 LN:30427671
@SQ
       SN:2 LN:19698289
@SQ
       SN:3 LN:23459830
@SQ
       SN:4 LN:18585056
@SQ
       SN:5 LN:26975502
@SQ
       SN:M LN:366924
@SQ
       SN:C LN:154478
@RG
       ID:Col0 R1 PL:Illumina LB:1342 SM:Col0 R1
```

@SQ Reference Sequence: SN name, LN length
@RG Read Group: e.g. grouping samples

SAM alignments



SAM Alignment Format: Flags

Ε	m Bit	Description
1	0x1	template having multiple segments in sequencing
2	0x2	each segment properly aligned according to the aligner
4	0x4	segment unmapped
8	0x8	next segment in the template unmapped
16	0x10	SEQ being reverse complemented
32	0x20	SEQ of the next segment in the template being reverse complemented
64	0x40	the first segment in the template
128	0x80	the last segment in the template
256	0x100	secondary alignment
512	0x200	not passing quality controls
1024	0x400	PCR or optical duplicate
2048	0x800	supplementary alignment

Example, flag 83 = 64+16+2+1 means it's first read (0x40) of pair-end reads (0x1) and it's mapped on minus strand (0x10) and both reads mapped (0x2).

https://broadinstitute.github.io/picard/explain-flags.html

SAM format: CIGAR string

- Summary of alignment to the reference
- **■** *eg*, 101M, 1S92M, 15M87N70M90N16M

Code	Description	
M	Alignment match	Base-level match + mismatch
I	Insertion	
D	Deletion	
N	Skipped	eg intron
S	Soft clipping	Kept in SAM
Н	Hard clipping	Not in SAM

SAM format: optional fields

- Used by some aligners to encode additional information for downstream analyses
- Can cause incompatibilities among workflows

Code	Description
RG	Read Group e.g. sample or lane
MD	String for mismatching positions
NH	Number of reported alignments that contains the query in the current record
HI	Query hit index, indicating the alignment record is the i-th one stored in SAM

BAM format

- Binary SAM format
- Lossless compression of SAM format
- ~4-fold smaller file size
- Genome viewers and many downstream applications require the BAM file to be sorted and have an index (typically .bai extension)

Annex: Assessing read coverage for biases

- The RSeQC package includes a function for evaluating "gene body coverage"
- This can be used to assess 5' or 3' bias, which might happen if your RNA is degraded or otherwise biased
- Requirements:
 - Genome annotations in the 12-column BED format
 - Index (.bai) for sorted BAM file, which can be generated using the SAMtools package

Annex - CRAM format

- Binary SAM format, significantly improved over BAM lossless compression
- Compatible with BAM files
- Both lossless and lossy compression possible
- https://samtools.github.io/hts-specs/CRAMv3.pdf

Annex - Other relevant formats: BED

- Tab-delimited text file used to describe intervals
- Minimally:
 - Sequence ID
 - Start
 - End
- Optional:
 - Name
 - Score
 - Strand
- For large files, use binary index format bigBED
- BEDtools (<u>http://code.google.com/p/bedtools</u>)

Annex - Other relevant formats: VCF (Variant Call Format)

- Tab-delimited text to describe SNPs, structural variants, indels etc
- Contains:
 - Chromosome
 - Position
 - Reference allele, alternative allele(s)
 - Various statistical metrics
- BCF: indexed binary format
- https://samtools.github.io/hts-specs/VCFv4.2.pdf