Aligners

J Fass | 20 June 2018

Definitions

Assembly:

I've found the shredded remains of an important document; put it back together!

Definitions

Alignment:

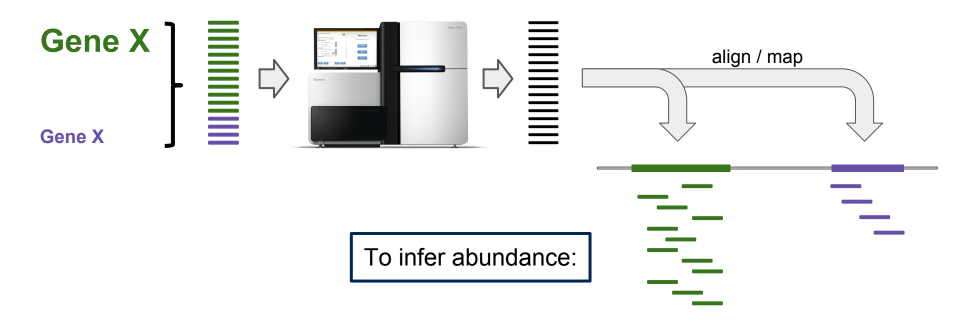
Somebody plagiarized parts of my document; where did they copy paragraphs from and where were each of the words and letters copied (perhaps with mistakes or changes) from?

Definitions

Mapping:

Somebody plagiarized parts of my document; where did they copy paragraphs from and where were each of the words and letters copied (perhaps with mistakes or changes) from?

Why align (or map)?



Why align (or map)?



<u>ATGATAGCATCGTCGGGTGTCTCAATAATAGTGCCGTATCATGCTGGTGTTATAATCGCCGCATGACATGATCAATGG</u>

CAATAA**A**AGTGCCGTATCATGCTGGTGTTA**C**AATCGCCGCA

CGTATCATGCTGGTGTTACAATCGCCGCATGACATGATCAATGG

TGTCTGCTCAATAA**A**AGTGCCGTATCATGCTGGTGTTACAATC

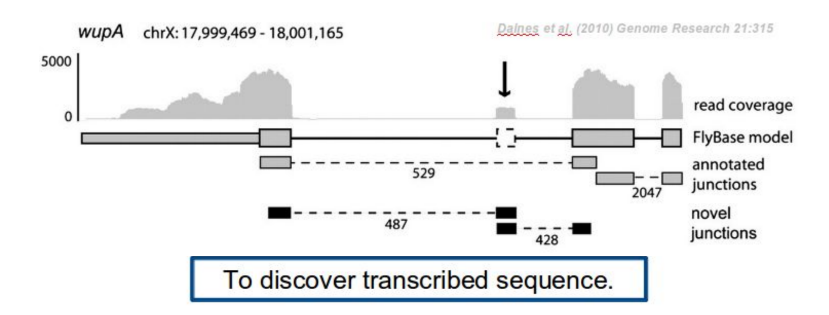
ATCGTCGGGTGTCTCAATAAAAGTGCCGTATCATG--GGTGTTATAA

CTCAATAAGAGTGCCGTATCATG--GGTGTTATAATCGCCGCA

GTTATAATCGCCGCATGACATGATCAATGG

To measure variation

Why align (or map)?



More Definitions: "Global" and "local"

Global aligners try to align all provided sequence, end to end, both "query" and "subject / target" ...

E.g.

- Aligning two Salmonella genomes
- Aligning human and gorilla orthologous coding regions

"Global" and "local"

Local aligners try to find "hits" or chains of hits within each provided sequence ...

E.g.

- Finding mitochondrial "splinters" in nuclear chromosomes
- Finding genes that share a domain with a gene of interest

"Glocal ... ?"

Short read aligners generally assume that the *whole read* came from somewhere within the target (reference) sequence ...

... so, *global* with respect to the read, and *local* with respect to the reference.

Short Read (Non-splicing) Aligners

Li, H and Homer, N (2010) *Briefings in Bioinformatics* 11:473 "A survey of sequence alignment algorithms for next-generation sequencing"

Table 1:Popular short-read alignment software

Program	Algorithm	SOLID	Long	Gapped	PE	Q°
Bfast	hashing ref.	Yes	No	Yes	Yes	No
Bowtie	FM-index	Yes	No	No	Yes	Yes
BWA	FM-index	Yes ^d	Yes ^e	Yes	Yes	No
MAQ	hashing reads	Yes	No	Yes ^f	Yes	Yes
Mosaik	hashing ref.	Yes	Yes	Yes	Yes	No
Novoalign ^g	hashing ref.	No	No	Yes	Yes	Yes

These two were fastest, at ~7 Gbp (vs human) per CPU day ... HiSeq 2500 generated 50-100 Gbp per day (at the time)

(Fall '12-'13) ... 150-180 Gbp per day

(Summer '16) ... 600 Gbp per day

(Summer '17) ... 1-3 Tbp per day

https://www.illumina.com/systems/sequencing-platforms.html

Burrows-**W**heeler Aligners

Burrows-Wheeler Transform used in bzip2 file compression tool; FM-index (Ferragina & Manzini) allow efficient finding of substring matches within compressed text – algorithm is *sub-linear* with respect to time and storage space required for a certain set of input data (reference 'ome, essentially).

Reduced memory footprint, faster execution.

BWA

BWA is a fast gapped aligner. Long read aligners (bwasw and mem) also fast, and can perform well for 454, Ion Torrent, Sanger, and PacBio reads. BWA is actively maintained and has a strong user community.

bio-bwa.sourceforge.net

'bwa aln' (BWA "backtrack") for reads < 70 bp

'bwa bwasw'

'bwa mem' (seeds with *maximal exact matches*, extends via *Smith-Waterman*)

Bowtie

(now Bowtie 2) ... comparable to BWA. Bowtie is part of a suite of tools (Bowtie, Tophat, Cufflinks, CummeRbund) that address RNAseq experiments.

http://bowtie-bio.sourceforge.net

Written by same folks as Tophat ... so, full compatibility.

Tophat2

Aligns full reads to genome, to determine "coverage islands." Creates simulated spliced exons based on these islands, then aligns remaining short reads to the simulated cDNA (to find reads crossing splice junctions).

STAR

Spliced Transcripts Aligned to a Reference

Aligns short reads and full length cDNA

Similar to BWA MEM algorithm, but searches uncompressed version (suffix array) of genome (faster, but more RAM required!). Claims better sensitivity and specificity than previous short read aligners, and 50x speed vs TopHat2. Requires ~27 GB RAM for human genome!

HISAT2

Improved replacement for Tophat2, multiple high level and low level indexes of compressed sequence (similar to graph-based assemblers using De Bruijn graphs to represent overlapping k-mers). Graph structure can represent multiple sequence paths, i.e. a population of references, allowing rapid genotyping of samples.

HISAT2 is currently probably the best combination, *in a full aligner*, of speed (faster than STAR) and memory (~4 GB for human genome).

Kim, Langmead, Salzberg (2015) Nature Methods 12:357

Kallisto

A pseudoaligner (mapper? see Pachter <u>blog</u> <u>post</u>) that compares read k-mers (overlapping subsequences) to a *transcriptome de Bruijn* graph (*T-DBG*) to find transcripts compatible with the read. Also uses expectation maximization (EM) and bootstrapping to determine most likely transcript and abundance uncertainty.

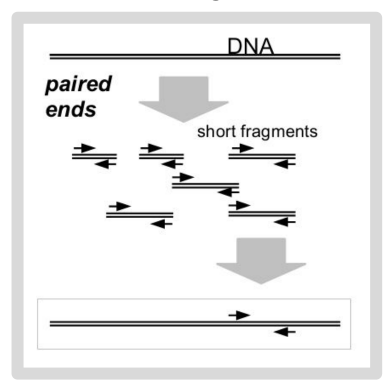
Bray ... Pachter (2016) *Nat Biotech* 34:525 See *sleuth* for DGE analysis. b

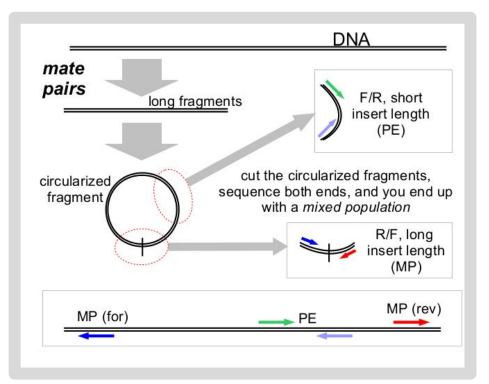
http://pachterlab.github.io/kallisto/ http://pachterlab.github.io/sleuth/

Salmon

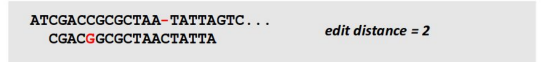
Supersedes Sailfish, an older k-mer based transcript analysis tool. Performs *quasi-mapping* to a set of transcripts (not genome), similar in methods to kallisto. Also performs bias correction for multiple modes of bias (sequence, position) to more accurately determine abundance.

Patro, Duggard, Kingsford (2015) bioRxiv http://dx.doi.org/10.1101/021592





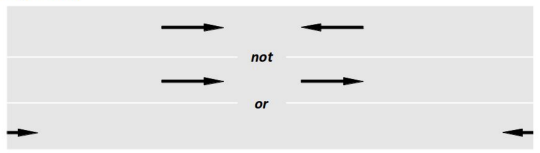
Edit Distance:

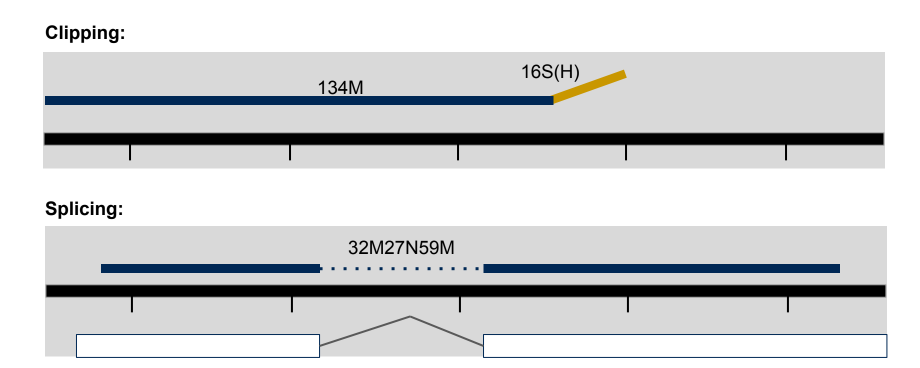


Mapping Quality:

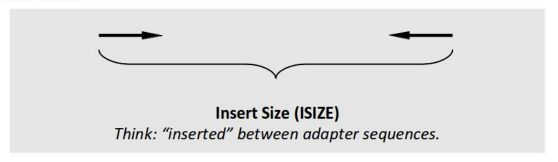
```
prob. of incorrect position = 10^{-MQ/10} ... (BWA)
```

Proper Pairs:

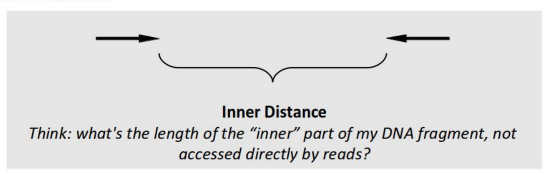




Insert Size:



Inner Distance:



Multimappers:

Reads that align equally well to more than one reference location.

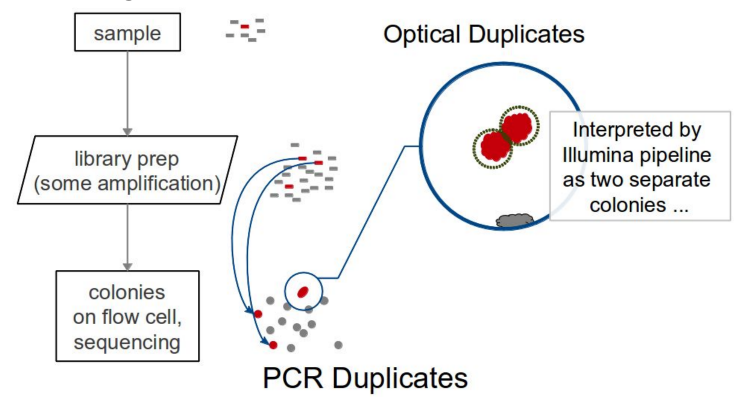
Generally, multimappers are discounted in variant detection, and are often discounted in counting applications (like RNA-Seq ... would "cancel" out anyway).

Note: *multimapper "rescue"* in some algorithms (RSEM, Express?).

Duplicates:

Reads or read pairs arising from the same original library fragment, either during library preparation (PCR duplicates) or colony formation (optical duplicates; not an issue anymore).

Generally, duplicates can only be detected reliably with paired-end sequencing. If PE, they're discounted in variant detection, and discounted in counting applications (like RNA-Seq).





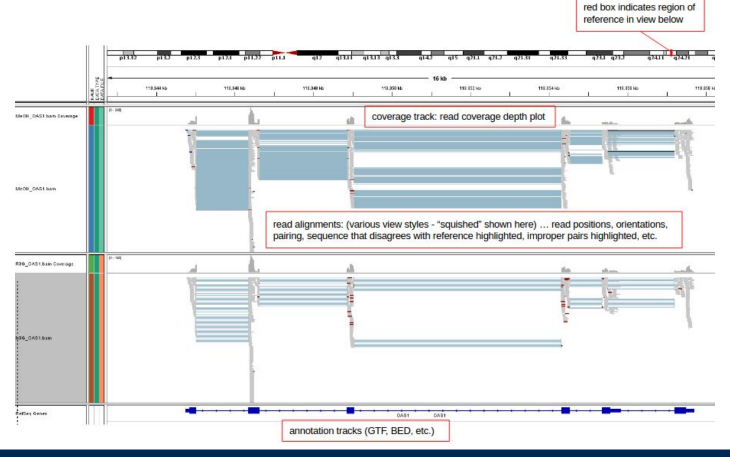
Alignment Viewers

- IGV (Integrated Genomics Viewer)
 - www.broadinstitute.org/igv/
- BAMview, tview (in SAMtools),
 IGB, GenomeView, SAMscope

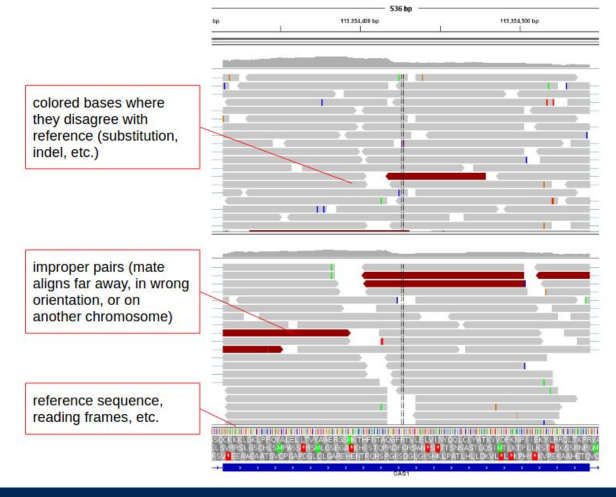
 UCSC Genome Browser, GBrowse



IGV



IGV



IGV

More on IGV's interface, file formats, and display can be found here:

http://www.broadinstitute.org/igv/AlignmentData

More on interpreting and customizing IGV's display can be found here:

http://www.broadinstitute.org/software/igv/interpreting insert size