High-throughput sequencing: Alignment and related topic

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HTS Platforms

- Established platforms
 - Illumina HiSeq, ABI SOLiD, Roche 454
- Newcomers:
 - Benchtop machines: Illumina MiSeq, etc
 - Long reads: Pacific Bioscience SMRT



The enzyme incorporates nucleotides to build double-stranded bridges on the solidphase substrate. Denaturation leaves single-stranded templates andhored to the substrate.

Several million dense dusters of doublestranded DNA are generated in each channel of the flow cell.

Bridge PCR



ML Metzker, Nat Rev Genetics (2010)



After laser excitation, collect the image data as before. Record the identity of the second base for each duster.

Repeat cycles of sequencing to determine the sequence of bases in a given fragment a single base at time. Align data, compare to a reference, and identify sequence differences.

Applications of HTS

- Sequencing of (genomic) DNA
 - de-novo sequencing
 - resequencing (variant finding)
 - enrichment sequencing (ChIP-Seq, MeDIP-Seq, ...)
 - targeted sequencing (exome sequencing, ...)
 - CCC-like (4C, HiC)
 - Metagenomics
 - Barcodes (tagged cells, yeast deletion collection, ...)
- Sequencing of RNA (actually: cDNA)

Applications of HTS

- Sequencing of (genomic) DNA
- Sequencing of RNA (actually: cDNA)
 - whole transcriptome*: RNA-Seq, Tag-Seq, ...
 - enriched fraction: HITS-CLIP, ...
 - labeled material: DTA, ...

* or: polyadenylated fraction

HTS: Bioinformatics challenges

Solutions specific to HTS are required for

- assembly
- alignment
- statistical tests (counting statistics)
- visualization
- segmentation
- ...

Two types of experiments

- Discovery experiments
 - finding all possible variant
 - getting an inventory of all transcripts
 - finding all binding sites of a transcription factor
- Comparative experiments
 - comparing tumour and normal samples
 - finding expression changes due to a treatment
 - finding changes in binding affinity

Assembly and Alignment

• First step in most analyses is the *alignment* of reads to a genome

• Except the point is to get the genome: *de-novo assembly*

• Special cases: Transcriptome assembly, metagenomics

The data funnel: ChIP-Seq, non-comparative

- Images
- Base calls
- Alignments
- Enrichment scores
- Location and scores of peaks (or of enriched regions)
- Summary statistics
- Biological conclusions

The data funnel: Comparative RNA-Seq

- Images
- Base calls
- Alignments
- Expression strengths of genes
- Differences between these
- Gene-set enrichment analyses

Where does Bioconductor come in?

- Processing of the images and determining of the read sequencest
 - typically done by core facility with software from the manufacturer of the sequencing machine
- Aligning the reads to a reference genome (or assembling the reads into a new genome)
 - Done with community-developed stand-alone tools.
- Downstream statistical analysis.
 - Write your own scripts with the help of Bioconductor infrastructure.

Alignment

Alignment

• Many different aligners:

Eland, Maq, Bowtie, BWA, SOAP, SSAHA, TopHat, SpliceMap, GSNAP, Novoalign, ...

- Main differences:
 - Publication year, maturity, development after publication, popularity
 - usage of base-call qualities, calculation of mapping qualities
 - Burrows-Wheeler index or not
 - speed-vs-sensitivity trade-off
 - suitability for RNA-Seq ("spliced alignment")
 - suitability for special tasks (e.g., color-space reads, bisulfite reads, variant injection, local re-alignment, ...)

Alignment: Workflow

• Preparation: Generate an *index* from FASTA file with the genome.

- Input data: FASTQ files with raw reads (demultiplexed)
- Alignment
- Output file: SAM file with alignments

Raw reads: FASTQ format

"FASTA with Qualities"

Example:

@HWI-EAS225:3:1:2:854#0/1
GGGGGGAAGTCGGCAAAAATAGATCCGTAACTTCGGG
+HWI-EAS225:3:1:2:854#0/1
a`abbbbabaabbababb^`[aaa`_N]b^ab^``a
@HWI-EAS225:3:1:2:1595#0/1
GGGAAGATCTCAAAAACAGAAGTAAAACATCGAACG
+HWI-EAS225:3:1:2:1595#0/1

FASTQ format

Each read is represented by four lines:

- '@', followed by read ID
- sequence
- '+', optionally followed by repeated read ID
- quality string:
 - same length as sequence
 - each character encodes the base-call quality of one base

FASTQ format: quality string

• If *p* is the probability that the base call is wrong, the Phred score is:

$$Q = -10 \log_{10} p$$

• The score is written with the character whose ASCII code is Q+33 (Sanger Institute standard).

- Before SolexaPipeline version 1.8, Solexa used instead the character with ASCII code Q+64.
- Before SolexaPipeline version 1.3, Solexa also used a different formula, namely $Q = -10 \log_{10} (p/(1-p))$

FASTQ: Phred base-call qualities

quality score Q _{phred}	error prob. <i>p</i>	characters
09	1 0.13	! "# \$%&'()*
1019	0.1 0.013	+,/01234
2029	0.01 0.0013	56789:;<=>
3039	0.001 0.00013	?@ABCDEFGH
40	0.0001	I

Quality scales

		JJJJJJJJJJJJJJJJJJJJJJJJJJJJJJJ			
!"#\$%&'()*+,/0123456789:;<=>?@ABCDEFGHIJKLMNOPQRSTUVWXYZ[\]^_`abcdefghijklmnopqrstuvwxyz{ }~					
1			I		
33	59 64 73	104	126		
<pre>S - Sanger Phred+33, raw reads typically (0, 40) X - Solexa Solexa+64, raw reads typically (-5, 40) I - Illumina 1.3+ Phred+64, raw reads typically (0, 40) J - Illumina 1.5+ Phred+64, raw reads typically (3, 40) with 0=unused, 1=unused, 2=Read Segment Quality Control Indicator (bold) L - Illumina 1.8+ Phred+33, raw reads typically (0, 41)</pre>					

[Wikipedia article "FASTQ format"]

FASTQ and paired-end reads

Convention for paired-end runs:

The reads are reported two FASTQ files, such that the nth read in the first file is mate-paired to the nth read in the second file. The read IDs must match.

Alignment output: SAM files

A SAM file consists of two parts:

- Header
 - contains meta data (source of the reads, reference genome, aligner, etc.)
 - Most current tools omit and/or ignore the header.
 - All header lines start with "@".
 - Header fields have standardized two-letter codes for easy parsing of the information
- Alignment section
 - A tab-separated table with at least 11 columns
 - Each line describes one alignment

A SAM file

[...]

HWI-EAS225_309MTAAXX:5:1:689:1485 16 XIII 863766 25 36M * 0 0 CTACAATTTTGCACATCAAAAAAGACCTCCAACTAC =8A=AA784A9AA5AAAAAAAAAAAAAAAAAAAAAA NM:i:0 X0:i:1 MD:Z:36

[...]

SAM format: Alignment section

The columns are:

- QNAME: ID of the read ("query")
- FLAG: alignment flags
- RNAME: ID of the reference (typically: chromosome name)
- POS: Position in reference (1-based, left side)
- MAPQ: Mapping quality (as Phred score)
- CIGAR: Alignment description (gaps etc.) in CIGAR format
- MRNM: Mate reference sequence name [for paired end data]
- MPOS: Mate position [for paired end data]
- ISIZE: inferred insert size [for paired end data]
- SEQ: sequence of the read
- QUAL: quality string of the read
- extra fields

Reads and fragments



SAM format: Flag field

FLAG field: A number, to be read in binary

bit	hex	decimal	
0	00 01	1	read is a paired-end read
1	00 02	2	read pair is properly matched
2	00 04	4	read has not been mapped
3	80 00	8	mate has not been mapped
4	00 10	16	read has been mapped to "-" strand
5	00 20	32	mate has been mapped to "-" strand
6	00 40	64	read is the first read in a pair
7	00 80	128	read is the second read in a pair
8	01 00	256	alignment is secondary
9	02 00	512	read did had not passed quality check
10	04 00	1024	read is a PCR or optical duplicate

SAM format: Optional fields

last column

- Always triples of the format TAG : VTYPE : VALUE
- some important tag types:
 - NH: number of reported aligments
 - NM: number of mismatches
 - MD: positions of mismatches

SAM format: CIGAR strings

Alignments contain gaps (e.g., in case of an indel, or, in RNA-Seq, when a read straddles an intron).

Then, the CIGAR string gives details.

Example: "M10 I4 M4 D3 M12" means

- the first 10 bases of the read map ("M10") normally (not necessarily perfectly)
- then, 4 bases are inserted ("I4"), i.e., missing in the reference
- then, after another 4 mapped bases ("M4"), 3 bases are deleted ("D4"), i.e., skipped in the query.
- Finally, the last 12 bases match normally.

There are further codes (N, S, H, P), which are rarely used.

SAM format: paired-end and multiple alignments

- Each line represents one *alignments*.
- Multiple alternative alignments for the same read take multiple lines. Only the read ID allows to group them.
- Paired-end alignments take two lines.

• All these reads are not necessarily in adjacent lines.

sorted SAM/BAM files

- Text SAM files (.sam): standard form
- BAM files (.bam): binary representation of SAM
 - more compact, faster to process, random access and indexing possible
- BAM index files (.bai) allow random access in a BAM file that is sorted by position.

SAMtools

- The SAMtools are a set of simple tools to
 - convert between SAM and BAM
 - sort and merge SAM files
 - index SAM and FASTA files for fast access
 - calculate tallies ("flagstat")
 - view alignments ("tview")
 - produce a "pile-up", i.e., a file showing
 - local coverage
 - mismatches and consensus calls
 - indels
- The SAMtools C API facilitates the development of new tools for processing SAM files.

Visualization of SAM files



Integrative Genomics Viewer (IGV): Robinson et al., Broad Institute

Special considerations for RNA-Seq

RNA alignment

- Only some aligners (e.g., TopHat, GSNAP, STAR) deal with spliced read.
- Use these for RNA-Seq data.



Strand-specific protocols

- Standard RNA-Seq loses strand information.
- If you want to distinguish sense from anti-sense transcripts, you need a strand-specific one.

• Make sure you know whether the library you analyse is strand-specific.

Solexa standard protocol for RNA-Seq



Strand-specific RNA-Seq with random hexamer priming



Coverage in RNA-Seq

• When sequencing genomic DNA, the coverage seems reasonably even.

• In RNA-Seq, this quite different



Levin et al., Nature Methods, 2010

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