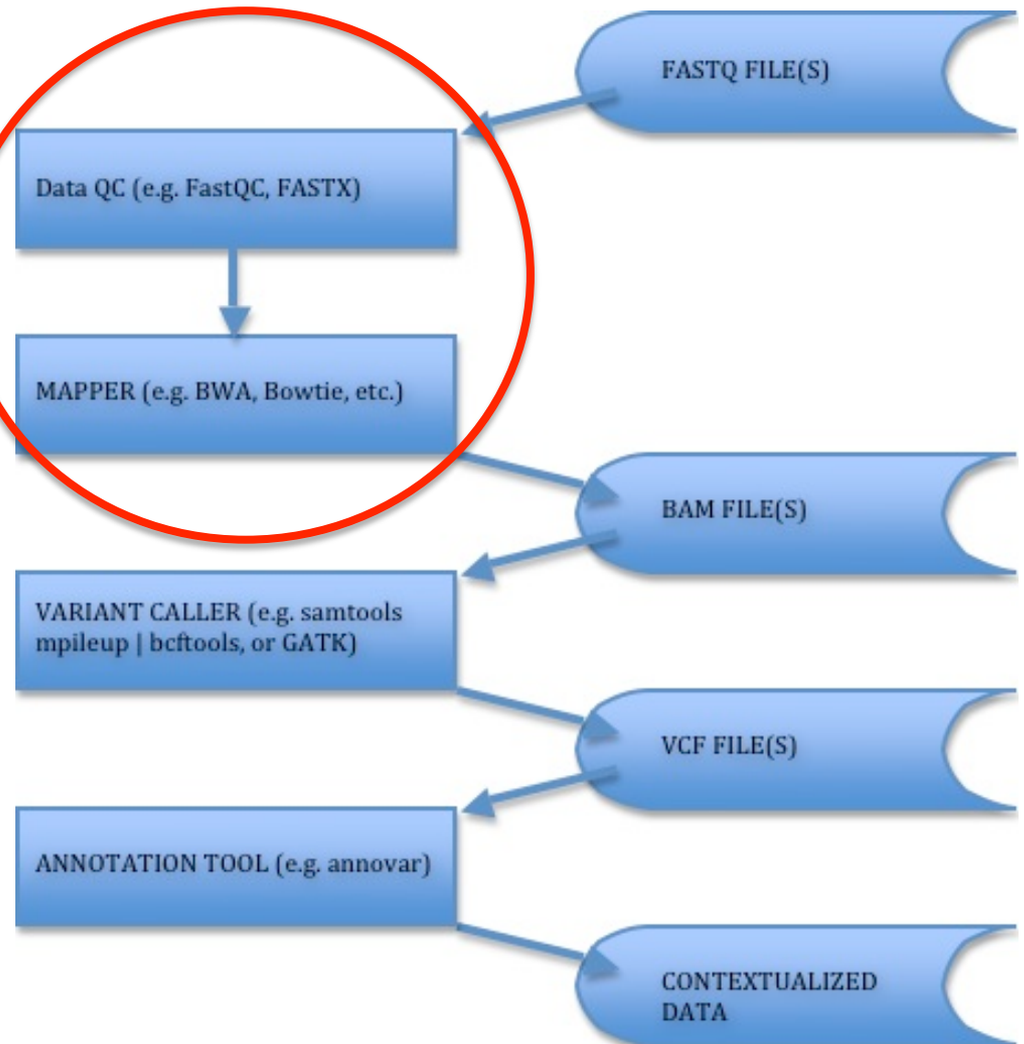


# Introduction to Mapping

**You are here!**



# Basic steps of mapping reads

1. Read file quality control
2. Build reference sequence index
3. Map DNA sequencing reads
  - Exact tool/approach depends on sequencing technology and DNA fragment library type
4. Convert result to SAM/BAM database
5. Application specific analysis...
  - These steps are common to any reference-based (opposed to *de novo*) data analysis.
  - We will look at variant calling first.

# Read sequences

## FASTQ Format

```
@HWI-EAS216_91209:1:2:454:192#0/1
GTTGATGAATTTCTCCAGCGCGAATTTGTGGGCT
+HWI-EAS216_91209:1:2:454:192#0/1
B@BBBBBB@BBBBAAAA>@AABA?BBBAAB??>A?
```

Line 1: @read name

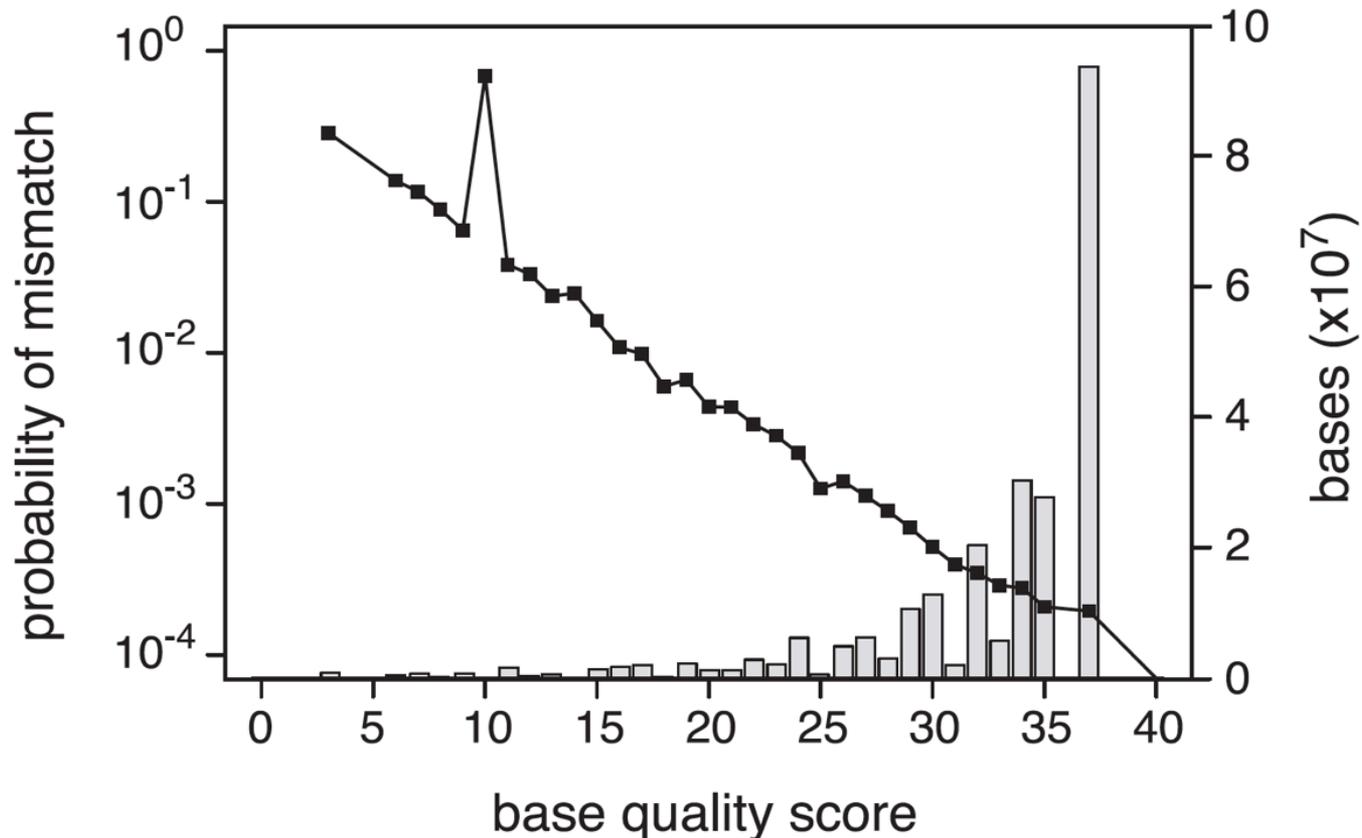
Line 2: called base sequence

Line 3: +read name (optional after +)

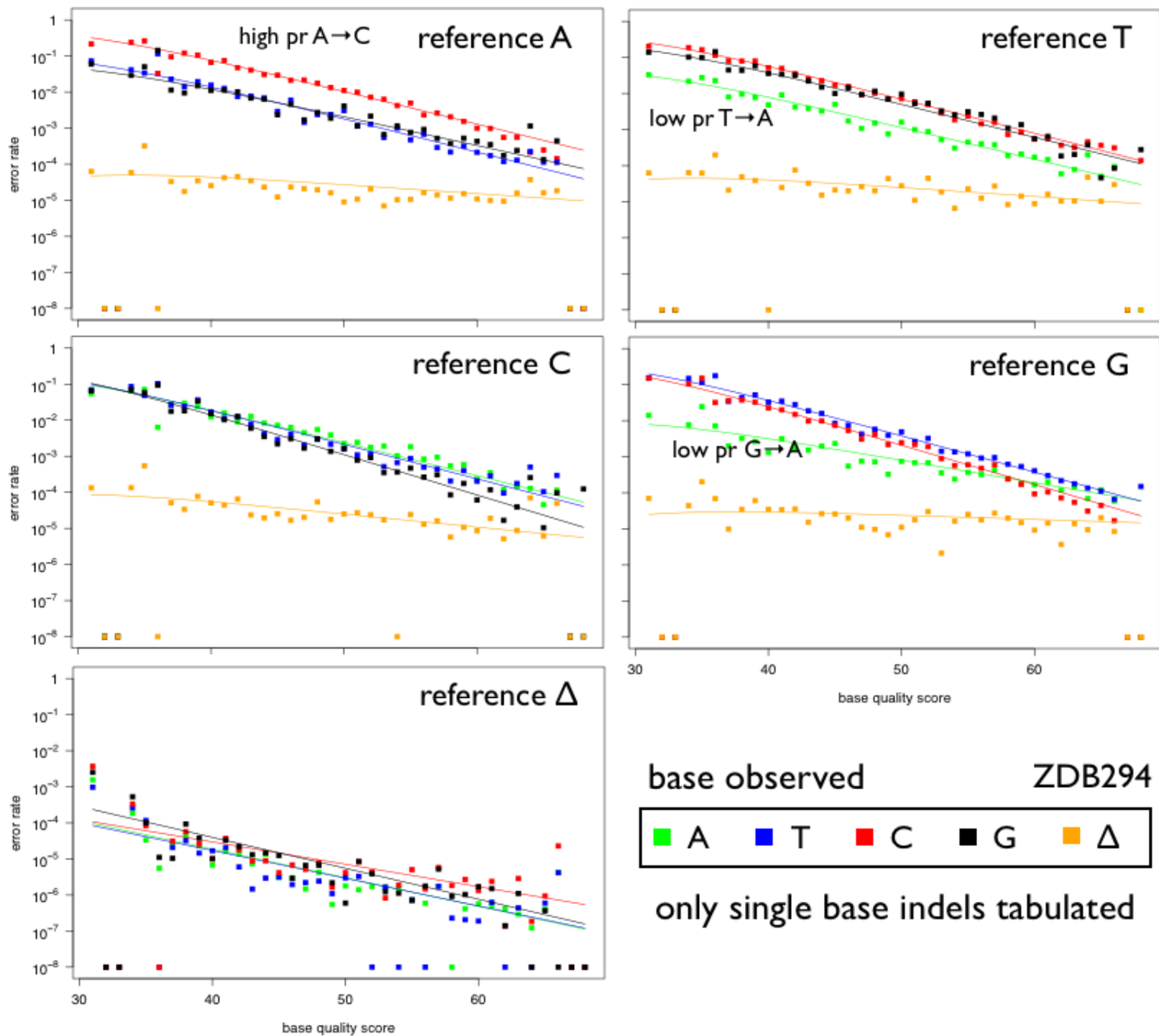
Line 4: base quality scores



# Example of Illumina data



- Most bases have high qualities ( $Q > 30$ ).
- Overall qualities are well calibrated\*.



# Read sequences

## Garbage in = garbage out

- Contaminated with other sequences?
- Sample barcodes removed?
- Adaptor sequences trimmed?
  - RNAseq
- Trim ends of reads with poor quality?
- Know your data
  - Paired reads? Relative orientations?
  - Technology specific concerns?
    - Indels with 454



# Types of Illumina fragment libraries

## single-end



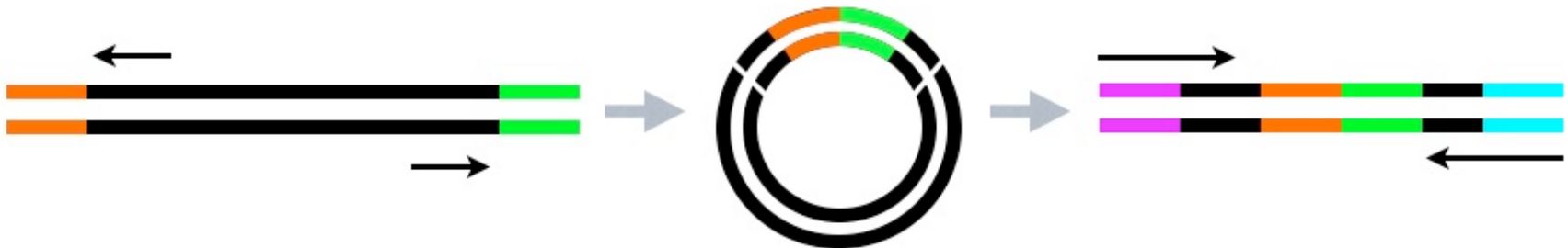
independent reads

## paired-end



two inwardly oriented reads separated by ~200 nt

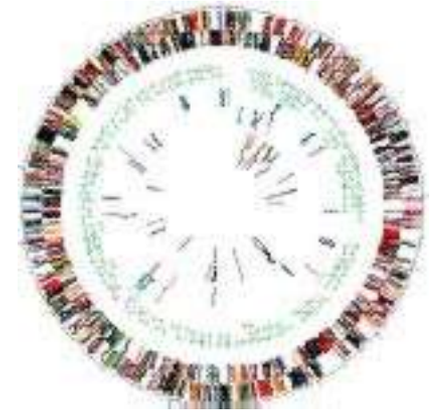
## mate-paired



two outwardly oriented reads separated by ~3000 nt

# Reference considerations

- Is it appropriate to your study?
  - Close enough to your species?
  - Complete?
- Which version?
  - Make sure you use an agreed-on standard
- Does it contain repeats?
  - Know this up front or you will be confused
- What annotations exist?
  - References lacking feature annotations are much more challenging to use



<http://microbialgenomics.energy.gov>

# Reference sequences

## FASTA Format

```
>gi|254160123|ref|NC_012967.1| Escherichia coli B str. REL606
agcttttcattctgactgcaacgggcaatatgtctctgtgtggattaaaaaaagagtgtc
tgatagcagcttctgaactggttacctgccgtgagtaaattaaattttattgacttagg
tcactaaataactttaaccaatataggcatagcgcacagacagataaaaattacagagtac
acaacatccatgaaacgcattagcaccaccattaccaccaccatcaccattaccacaggt
aacggtgcgggctgacgcgtacaggaaacacagaaaaaagcccgcacctgacagtgcggg
cttttttttcgaccaaggtaacgaggtaacaacatgcgagtggttgaagttcggcggtg
....
```

Using complex reference sequence names is a common problem during analysis. Might rename:

```
>REL606
agcttttcattctgactgcaacgggcaatatgtctctgtgtggattaaaaaaagagtgtc
tgatagcagcttctgaactggttacctgccgtgagtaaattaaattttattgacttagg
....
```

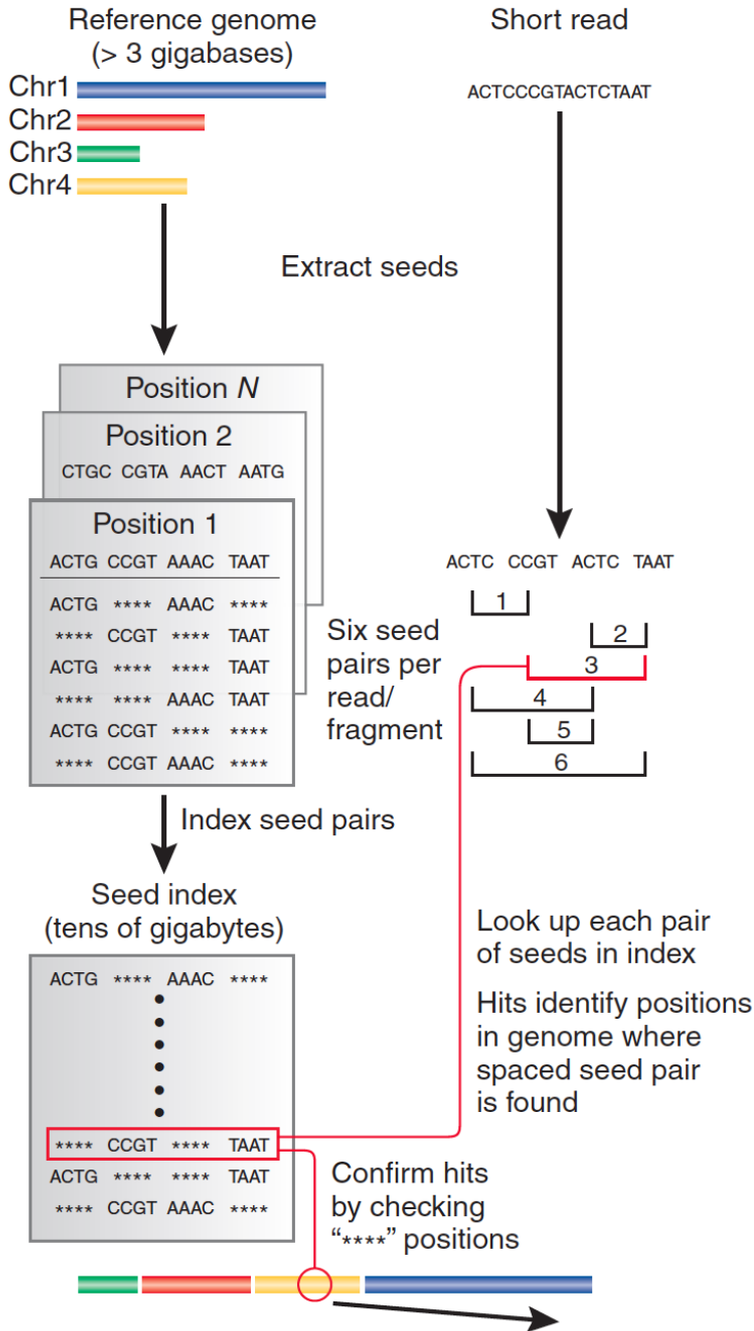
# Finding a Reference

- **Microbes (<10Mb)**: download FASTA containing in sequence and/or GenBank/EMBL/GFF flatfiles encapsulating both sequence and features.
- **Macrobies (>100Mb)**: download specific consortium "build" of reference (Ex: hg19), consisting of FASTA, and various files used to construct a database of features.
- **Non-model organisms**: build your own?  
*See de novo assembly topic.*

# Mappers/Aligners

- Algorithms
  - Spaced-seed indexing
  - Burrows-Wheeler transform (BWT)
- Differences
  - Input data (read length, colorspace aware/useful)
  - Speed and scalability (multithreading, GPUs)
  - Memory requirements (RAM, fat nodes)
  - Sensitivity: esp. indels (gaps)
  - Ease of installation and use. Development phase.
  - Uses base qualities? Outputs mapping scores?
  - Handles of multiple matches, paired end matches
  - Configurability and transparency of options

# a Spaced seeds



**Hash table** enables lookup of exact matches.

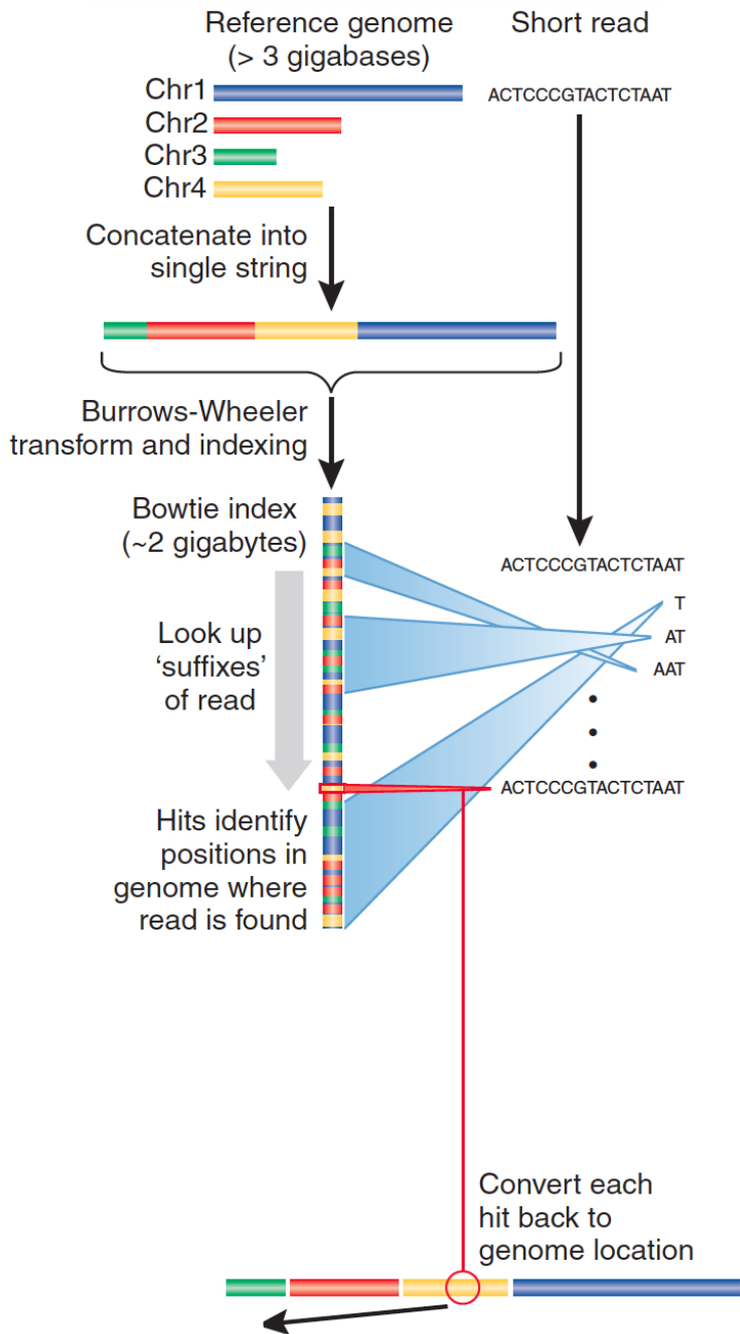
| Subsequence    | Reference Positions |
|----------------|---------------------|
| ATAGCTAATCCAAA | 2341, 2617264       |
| ATAGCTAATCCAAT |                     |
| ATAGCTAATCCAAC | 134, 13311, 732661, |
| ATAGCTATCCAAAG |                     |
| ATAGCTAATCCATA |                     |
| ATAGCTAATCCATT | 3452                |
| ATAGCTAATCCATC |                     |
| ATAGCTATCCAATG | 234456673           |

Table is sorted and complete so you can jump immediately to matches. (But this can take a lot of memory.)

May include N bases, skip positions, etc.

Trapnell, C. & Salzberg, S. L. How to map billions of short reads onto genomes. *Nature Biotech.* **27**, 455–457 (2009).

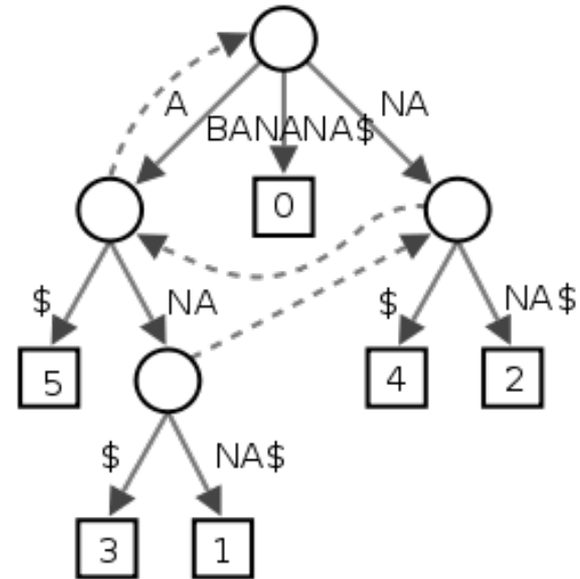
## b Burrows-Wheeler



**Burrows-Wheeler transform compresses sequence.**

|               |  |
|---------------|--|
| <b>Input</b>  | SIX.MIXED.PIXIES.SIFT.SIXTY.PIXIE.DUST.BOXES |
| <b>Output</b> | TEXYDST.E.IXIXIXSSMPPS.B..E.S.EUSFXDIIIOIIIT |

**Suffix tree** enables fast lookup of subsequences.



[http://en.wikipedia.org/wiki/Suffix\\_tree](http://en.wikipedia.org/wiki/Suffix_tree)

Exact matches at all positions below a node.

Trapnell, C. & Salzberg, S. L. How to map billions of short reads onto genomes. *Nature Biotech.* **27**, 455–457 (2009).

# From Mapped Read to Alignment

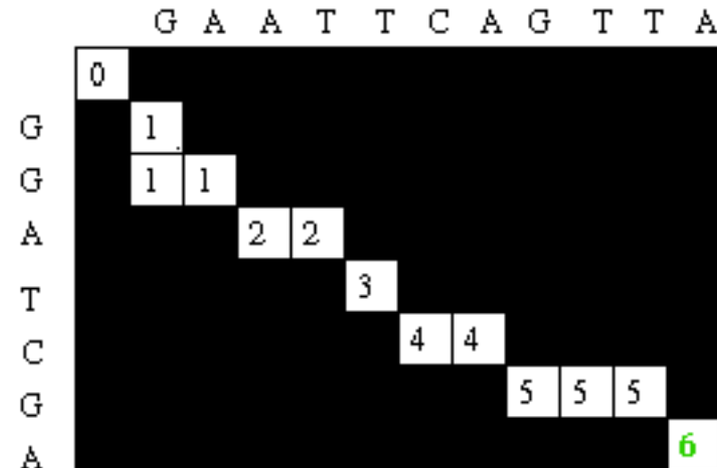
- **Mapping** determines a position where the read shares a subsequence with the reference. But, is this the best match?
- **Alignment** starts with the seed and determines how the read is best aligned on a base-by-base basis.

**Seed**→**Alignment score**→**Mapping quality**

# Alignment

- Dynamic programming algorithm (Smith-Waterman | Needleman-Wunsch)

|   | G | A | A | T | T | C | A | G | T | T | A |
|---|---|---|---|---|---|---|---|---|---|---|---|
| G | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| G | 0 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| A | 0 | 1 | 1 | 2 | 2 | 2 | 2 | 2 | 2 | 2 | 3 |
| T | 0 | 1 | 2 | 2 | 3 | 3 | 3 | 3 | 3 | 3 | 3 |
| C | 0 | 1 | 2 | 2 | 3 | 3 | 4 | 4 | 4 | 4 | 4 |
| G | 0 | 1 | 2 | 2 | 3 | 3 | 4 | 4 | 5 | 5 | 5 |
| A | 0 | 1 | 2 | 3 | 3 | 3 | 4 | 5 | 5 | 5 | 6 |



```

G _ A A T T C A G T T A
| | | | | | | | | |
G G _ A _ T C _ G _ _ A
  
```

- Various scoring schemes possible...

# Alignment Score

- Dynamic programming algorithm (Smith-Waterman | Needleman-Wunsch)
- **Alignment score** =  $\Sigma$ 
  - match reward
  - base mismatch penalty
  - gap open penalty
  - gap extension penalty
  - rewards and penalties may be adjusted for quality scores of bases involved
- Local versus global alignment of read

# Mapping Quality

**Mapping quality**– what is the probability that the read is correctly mapped to this location in the reference genome?

**Read 1**

**Read 2**

ATCGGGAGATCC

or

ATCGGGAGATCC

GCGTAGTCTGCC

|||||

|||||

|| ||| |||

...TAATCGGGAGATCCGC...TTATCGGGAGATCCGC... ..TAGCCTAGTGTGCCGC...

**Reference Sequence**

High **alignment** score  $\neq$  high **mapping** quality.

Phred score:  $P(\text{mismapped}) = 10^{-MQ/10}$

# Types of DNA fragment libraries

## single-end



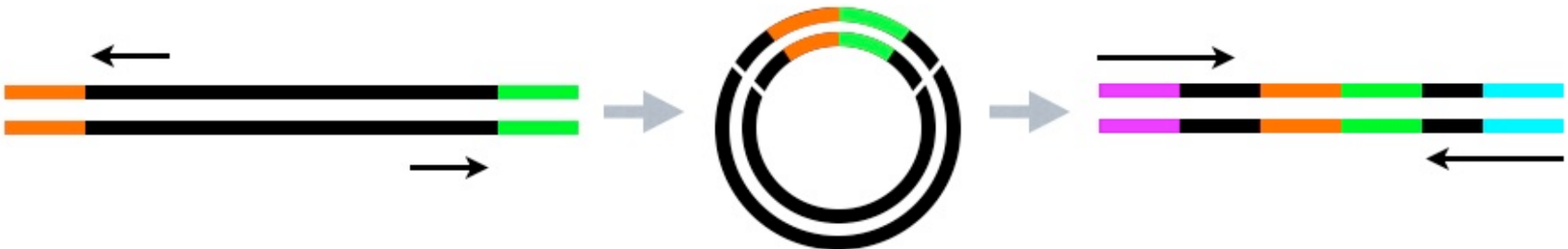
independent reads

## paired-end



two inwardly oriented reads separated by ~200 nt

## mate-paired



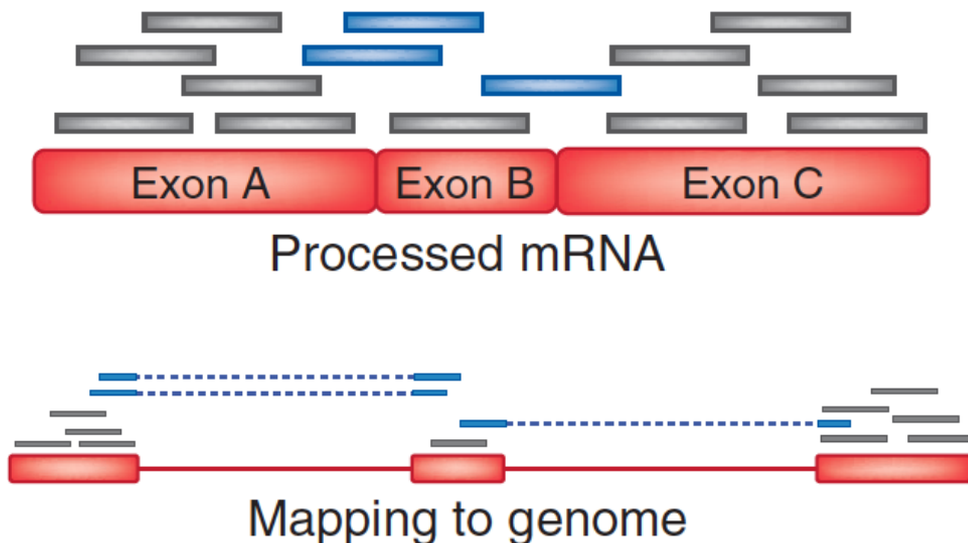
two outwardly oriented reads separated by ~3000 nt

# Paired-end mapping (PEM)

- There is an expected insert size distribution based on the DNA fragment library.
- Mapping one read anchor the paired read to a specific location, even if the second read alone maps multiple places in the reference.
- Only one read in a pair might be mappable. (**singleton/orphan**)
- Both reads can map with an unexpected insert size or orientation (**discordant pair**)

# Split-read alignment (SRA)

- Useful for predicting splice variants or structural variants.
- Not many mappers do this directly, usually happens in a post-processing step.



Trapnell, C. & Salzberg, S. L. How to map billions of short reads onto genomes. *Nature Biotech.* **27**, 455–457 (2009).

# List of Mappers/Aligners

|            | Algorithm                  | Gapped | Quality-aware | Colorspace aware |
|------------|----------------------------|--------|---------------|------------------|
| BLAST      | Hash table                 | Y      | N             | N                |
| BLAT/SSHA2 | Hash table                 | N      | N             | N                |
| MAQ        | Spaced seed                | N      | N             | N                |
| RMAP       | Spaced seed                | N      | Y             | N                |
| ZOOM       | Spaced seed                | N      | --            | N                |
| SOAP       | Spaced seed                | N      | N             | N                |
| Eland      | Spaced seed                | N      | N             | N                |
| SHRIMP     | Q-gram/multi-seed          | Y      | Y             | Y                |
| BFAST      | Q-gram/multi-seed          | Y      | Y             | Y                |
| Novoalign  | Multi-seed + Vectorized SW | Y      | Y             | Y                |
| clcBio     | Multi-seed + Vectorized SW | Y      | Y             | Y                |
| MUMmer     | Tries                      | Y      | N             | N                |
| OASIS      | Tries                      | Y      | --            | --               |
| VMATCH     | Tries                      | Y      | --            | --               |
| BWA/BWA-SW | Tries                      | Y      | Y             | Y                |
| BOWTIE     | Tries                      | Y      | Y             | Y                |
| SOAP2      | Tries                      | Y      | N             | N                |
| Saruman    | Exact (GPU)                | Y      | --            | N                |

**trie** = tree structure for fast text **retrieval**.

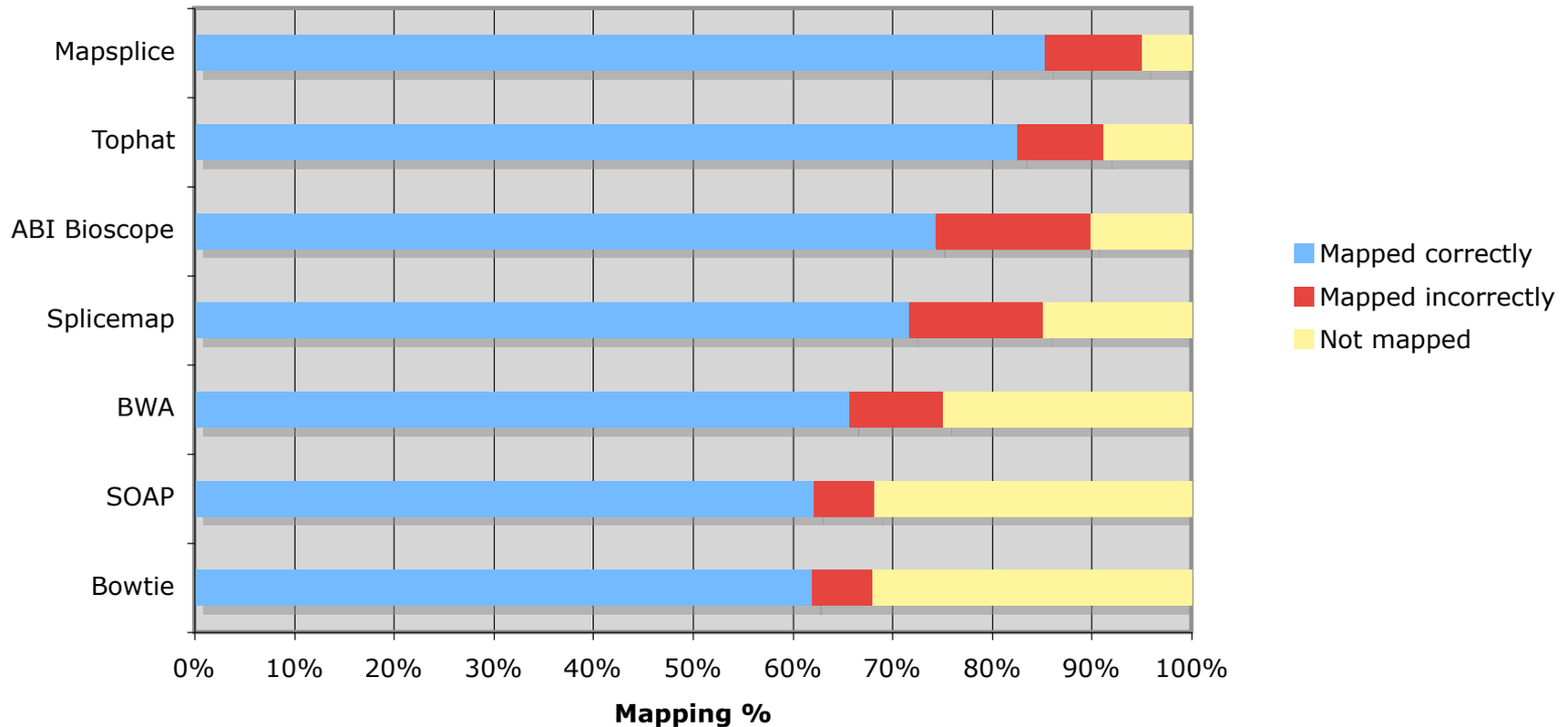
# Indexing time

| Aligner | Time (mins) to Index 3GB genome |
|---------|---------------------------------|
| SOAP2   | 98.06                           |
| BWA     | 110.73                          |
| Bowtie  | 220.82                          |
| Bfast   | 941.10*                         |

Data courtesy Dhivya Arasappan, GSAF Bioinformatician

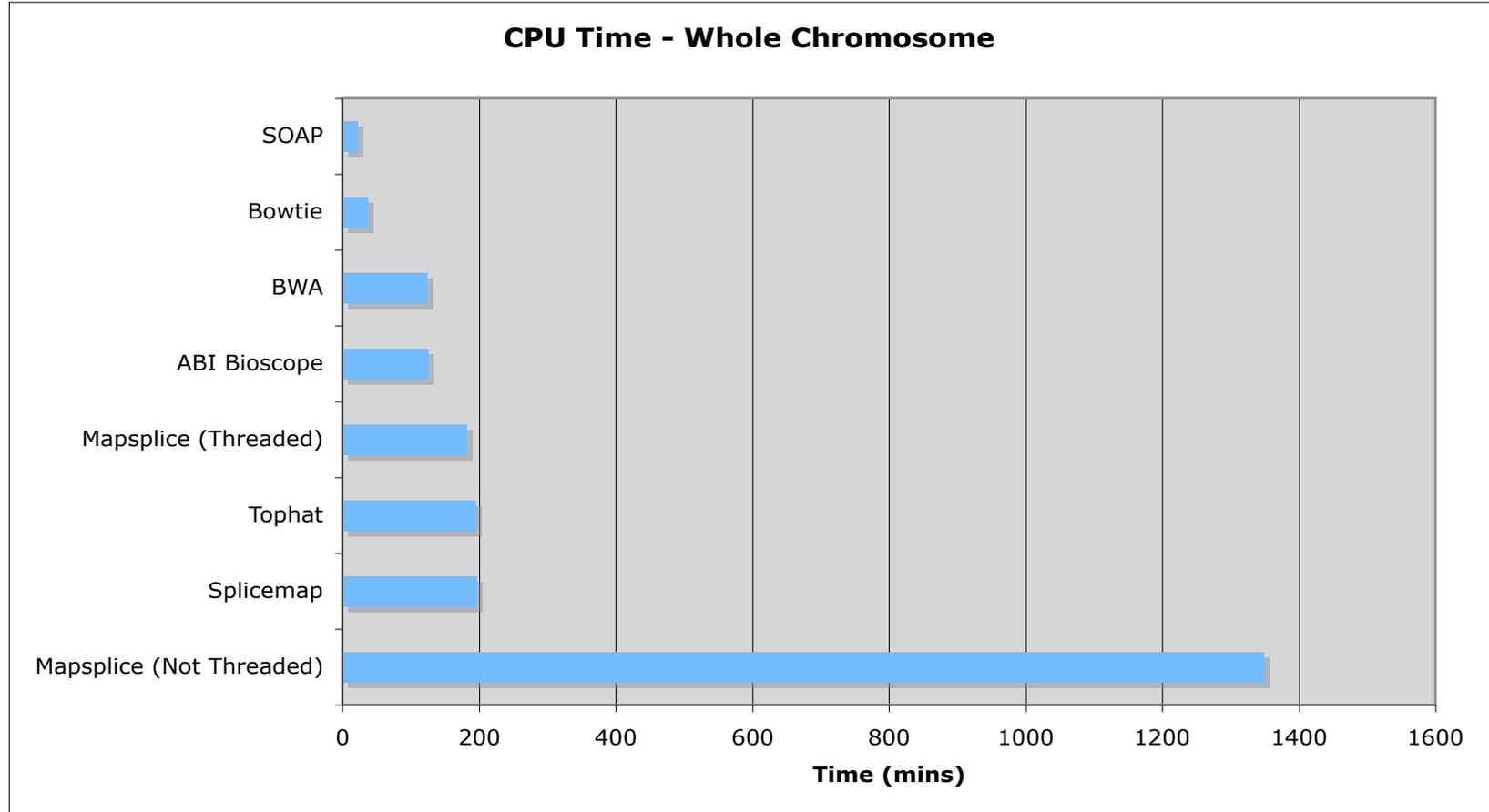
# Some Comparisons

**Mapping Accuracy - Spliced Data to Whole Genome**



Data courtesy Dhivya Arasappan, GSAF Bioinformatician

# Some Comparisons



Data courtesy Dhivya Arasappan, GSAF Bioinformatician

# Final Words

- My personal favorites of the moment...
  - Bowtie2
  - BWA
- Pay close attention to the details...
  - Methylation (bisulfite) analysis
  - Hypervariable region analysis  
(Ex: contingency loci / microsatellites)
  - To understand what you might be missing!

# SAM File Format

- Community flat file/database format that describes how reads align to a reference (and can also include unmapped reads).
- Can tag reads as being from different instrument runs / technologies / samples.
- Going forward you need the reference file and the SAM, no longer need the FASTQ.
- Tab delimited with fixed columns followed by arbitrary user-extendable key:data values.

# SAM File Format

Two example SAM lines:

```
SRR030257.264529    99  NC_012967   1521    29  34M2S   =   1564    79
CTGGCCATTATCTCGGTGGTAGGACATGGCATGCC
AAAAAA;AA;AAAAAA??A%.;?&'3735',()0*,
XT:A:M NM:i:3 SM:i:29 AM:i:29 XM:i:3 XO:i:0 XG:i:0 MD:Z:23T0G4T4

SRR030257.2669090  147 NC_012967   1521    60  36M     =   1458   -99
CTGGCCATTATCTCGGTGGTAGGTGATGGTATGCGC
<<9:<<AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
XT:A:U NM:i:0 SM:i:37 AM:i:37 X0:i:1 X1:i:0 XM:i:0 XO:i:0 XG:i:0 MD:Z:36
```

# SAM File Format

SAM fixed fields:

<http://samtools.sourceforge.net/>

| Col | Field | Type   | Regexp/Range                             | Brief description                     |
|-----|-------|--------|--|---------------------------------------|
| 1   | QNAME | String | [!-?A-~]{1,255}                          | Query template NAME                   |
| 2   | FLAG  | Int    | [0,2 <sup>16</sup> -1]                   | bitwise FLAG                          |
| 3   | RNAME | String | \*  [!-( )+-<>-~] [!-~]*                 | Reference sequence NAME               |
| 4   | POS   | Int    | [0,2 <sup>29</sup> -1]                   | 1-based leftmost mapping POSition     |
| 5   | MAPQ  | Int    | [0,2 <sup>8</sup> -1]                    | MAPping Quality                       |
| 6   | CIGAR | String | \*  ([0-9]+[MIDNSHPX=])+                 | CIGAR string                          |
| 7   | RNEXT | String | \* =  [!-( )+-<>-~] [!-~]*               | Ref. name of the mate/next segment    |
| 8   | PNEXT | Int    | [0,2 <sup>29</sup> -1]                   | Position of the mate/next segment     |
| 9   | TLEN  | Int    | [-2 <sup>29</sup> +1,2 <sup>29</sup> -1] | observed Template LENgth              |
| 10  | SEQ   | String | \*  [A-Za-z=.]+                          | segment SEQuence                      |
| 11  | QUAL  | String | [!-~]+                                   | ASCII of Phred-scaled base QUALity+33 |

```
SRR030257.264529    99  NC_012967    1521    29  34M2S    =    1564
79  CTGGCCATTATCTCGGTGGTAGGACATGGCATGCCC
AAAAAA;AA;AAAAAA??A%.;?&'3735',()0*,
XT:A:M NM:i:3 SM:i:29 AM:i:29 XM:i:3 XO:i:0 XG:i:0 MD:Z:23T0G4T4
```

# Sometimes a CIGAR is a just way of describing how a read is aligned...

Ref CTGGCCATTATCTC--GGTGGTAGGACATGGCATGCCC  
Read aaATGTCGCGGTG.TAGGAaggatcc



2S5M2I4M1D4M6S

| Op  | BAM | Description   |
|-----|-----|---|
| M   | 0   | alignment match (can be a sequence match or mismatch) |
| I   | 1   | insertion to the reference                            |
| D   | 2   | deletion from the reference                           |
| * N | 3   | skipped region from the reference                     |
| S   | 4   | soft clipping (clipped sequences present in SEQ)      |
| * H | 5   | hard clipping (clipped sequences NOT present in SEQ)  |
| * P | 6   | padding (silent deletion from padded reference)       |
| * = | 7   | sequence match  |
| * X | 8   | sequence mismatch                                     |

\*Rarer / newer

CIGAR = "Concise Idiosyncratic Gapped Alignment Report"

# BAM format

- "Human readable" text (SAM) and GZIP compressed binary (BAM) versions.
- BAM files can be **sorted** and **indexed**, so that all reads mapped to a given window of the reference genome can be retrieved rapidly (for display or processing).