

File Types in Bioinformatics

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Enabler for Life Science











HOW STANDARDS PROLIFERATE: (SEE: A/C CHARGERS, CHARACTER ENCODINGS, INSTANT MESSAGING, ETC.)

SITUATION: THERE ARE 14 COMPETING STANDARDS.

14?! RIDICULOUS! WE NEED TO DEVELOP ONE UNIVERSAL STANDARD THAT COVERS EVERYONE'S USE CASES. YEAH!

SOON: SITUATION: THERE ARE 15 COMPETING STANDARDS.

http://xkcd.com



- Overwhelming at first
- Overview
 - FASTA reference sequences
 - FASTQ reads in raw form
 - SAM aligned reads
 - BAM compressed SAM file
 - CRAM even more compressed SAM file
 - GTF/GFF/BED annotations



FASTA

- Used for: nucleotide or peptide sequences
- Simple structure

> header sequence



FASTA

- Used for: nucleotide or peptide sequences
- Simple structure

> H.Sapiens chr17:135135135-1313566

ACTCAGATCGGAATAGCATACGCATACTCAGATCGGAATAGCATACGCAT
GGATAGCTCACGACACATGACACTACAGCCAGACTACACGACTACACGAT
AAGGATATAGGACTACGACTAGCATCGACTAACTAGCTACATACG

>that random protein sequence i saw yesterday
ARGAEBAEUIRGHAERGIAEUAEILHGAEIGAHEGLAEJKRGNAERBIAE
AEGHAELGIHAEGOUIAENGAEBAERIOTYUGAEGHILAEHRGAEIRGYU
AEHAEHAEIOGAEGAERTBETHUETHIRTHJNRFS



- Just like FASTA, but with quality values
- Used for: raw data from sequencing (unaligned reads)

@ header

sequence

+

quality





- Just like FASTA, but with quality values
- Used for: raw data from sequencing (unaligned reads)

```
@SEQ_001
GATTTGGGGTTCAAAGCAGTATCGATCAAATAGTAAATCCATTTGTTCAACTCACAGTTT
+
!''*((((***+))%%++)(%%!''*((((***).1***-+*''))**55CC!''*(D
@SEQ_002
GATTTGGGGTTCAAAGCAGTATTTGGGGTTCATTGGGGTTCATTGTTCAACTCACAGTTT
+
!''*((((***+))%>>CCCCC%++((((**).1***-+*''))**55CCF>>>>>C5
@SEQ_003
AAGCAGTATCGAGATTTGGGGTTCAAAGCAGTATAAGCAGTATCGATAAATCCATTTGTT
+
!''*((((*!''*(((**)(%%%).1***-+*''))**55CCF>>>>>%%%).1B5
```



- Quality 0-40
 - 40 = best

(Illumina 1.8 + = 41)



- Quality 0-40
 - 40 = best
- ASCII encoded

| Dec | Hex | Char | Dec | Hex | Char | Dec | Hex | Char | Dec | Hex | Char |
|-----|-----|------------------|-----|------------|-------|-----|-----|------|-----|-----|------|
| 0 | 00 | Null | 32 | 20 | Space | 64 | 40 | 0 | 96 | 60 | , |
| 1 | 01 | Start of heading | 33 | 21 | ! | 65 | 41 | A | 97 | 61 | a |
| 2 | 02 | Start of text | 34 | 22 | ** | 66 | 42 | В | 98 | 62 | b |
| 3 | 03 | End of text | 35 | 23 | # | 67 | 43 | С | 99 | 63 | c |
| 4 | 04 | End of transmit | 36 | 24 | Ş | 68 | 44 | D | 100 | 64 | d |
| 5 | 05 | Enquiry | 37 | 25 | * | 69 | 45 | E | 101 | 65 | e |
| 6 | 06 | Acknowledge | 38 | 26 | ٤ | 70 | 46 | F | 102 | 66 | £ |
| 7 | 07 | Audible bell | 39 | 27 | 1 | 71 | 47 | G | 103 | 67 | g |
| 8 | 08 | Backspace | 40 | 28 | (| 72 | 48 | H | 104 | 68 | h |
| 9 | 09 | Horizontal tab | 41 | 29 |) | 73 | 49 | I | 105 | 69 | i |
| 10 | OA | Line feed | 42 | 2A | * | 74 | 4A | J | 106 | 6A | j |
| 11 | OB | Vertical tab | 43 | 2B | + | 75 | 4B | K | 107 | 6B | k |
| 12 | OC. | Form feed | 44 | 2 C | , | 76 | 4C | L | 108 | 6C | 1 |
| 13 | OD | Carriage return | 45 | 2 D | _ | 77 | 4D | M | 109 | 6D | m |
| 14 | OE | Shift out | 46 | 2 E | | 78 | 4E | N | 110 | 6E | n |
| 15 | OF | Shift in | 47 | 2 F | / | 79 | 4F | 0 | 111 | 6F | 0 |
| 16 | 10 | Data link escape | 48 | 30 | 0 | 80 | 50 | P | 112 | 70 | р |
| 17 | 11 | Device control 1 | 49 | 31 | 1 | 81 | 51 | Q | 113 | 71 | đ |
| 18 | 12 | Device control 2 | 50 | 32 | 2 | 82 | 52 | R | 114 | 72 | r |
| 19 | 13 | Device control 3 | 51 | 33 | 3 | 83 | 53 | ຮ | 115 | 73 | ន |
| 20 | 14 | Device control 4 | 52 | 34 | 4 | 84 | 54 | Т | 116 | 74 | t |
| 21 | 15 | Neg. acknowledge | 53 | 35 | 5 | 85 | 55 | U | 117 | 75 | u |
| 22 | 16 | Synchronous idle | 54 | 36 | 6 | 86 | 56 | V | 118 | 76 | v |
| 23 | 17 | End trans, block | 55 | 37 | 7 | 87 | 57 | ឃ | 119 | 77 | w |
| 24 | 18 | Cancel | 56 | 38 | 8 | 88 | 58 | X | 120 | 78 | x |
| 25 | 19 | End of medium | 57 | 39 | 9 | 89 | 59 | Y | 121 | 79 | У |
| 26 | 1A | Substitution | 58 | 3A | : | 90 | 5A | Z | 122 | 7A | z |
| 27 | 1B | Escape | 59 | 3 B | ; | 91 | 5B | [| 123 | 7B | { |
| 28 | 1C | File separator | 60 | 3 C | < | 92 | 5C | ١ | 124 | 7C | 1 |
| 29 | 1D | Group separator | 61 | 3 D | = | 93 | 5D |] | 125 | 7D | } |
| 30 | 1E | Record separator | 62 | 3 E | > | 94 | 5E | ^ | 126 | 7E | ~ |
| 3.1 | 1 F | Unit separator | 63 | 3 F | 2 | 95 | 5E | | 127 | 78 | П |



Quality 0-40

(Illumina 1.8 + = 41)

- 40 = best
- ASCII encoded

```
.....
   !"#$%&'()*+,-./0123456789:;<=>?@ABCDEFGHIJKLMNOPQRSTUVWXYZ[\]^ `abcdefghijklmnopqrstuvwxyz{|}~
33
                                   104
                                              126
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)
  (Note: See discussion above).
L - Illumina 1.8+ Phred+33, raw reads typically (0, 41)
```



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```
@SEQ_001
GATTTGGGGTTCAAAGCAGTATCGATCAAATAGTAAATCCATTTGTTCAACTCACAGTTT
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GATTTGGGGTTCAAAGCAGTATTTGGGGTTCATTGGGGTTCATTGTTCAACTCACAGTTT
+
!''*((((***+))%*>>CCCCC%++((((**).1***-+*''))**55CCF>>>>>C5
@SEQ_003
AAGCAGTATCGAGATTTGGGGTTCAAAGCAGTATAAGCAGTATCGATAAATCCATTTGTT
+
!''*((((*!''*(((**)(%%%).1***-+*''))**55CCF>>>>>%%%).1B5
```



SAM

- Used for: aligned reads
- Lots of columns...



SAM

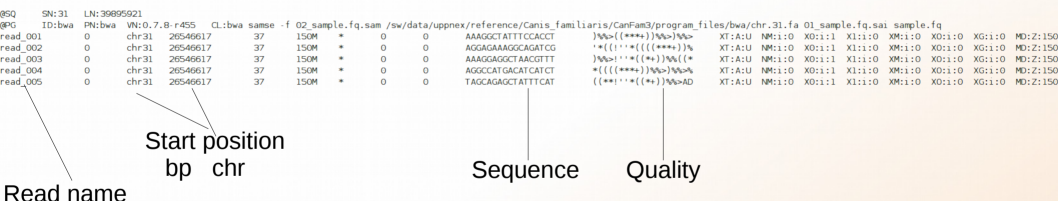
seguence_string.sam | <QNAME> <FLAG> <RNAME> <POS> <MAPQ> <CIGAR> <MRNM> <MPOS> <ISIZE> <SEQ> <QUAL> [<TAG>:<VTYPE>:<VALUE> [...]]

| Field | Regular expression | Range | Description | | |
|-------|-----------------------|--------------------------------------|---|--|--|
| QNAME | [^ \t\n\r]+ | | Query pair NAME if paired; or Query NAME if unpaired ² | | |
| FLAG | [0-9]+ | [0,216-1] | bitwise FLAG (Section 2.2.2) | | |
| RNAME | [^ \t\n\r@=]+ | | Reference sequence NAME 3 | | |
| POS | [0-9]+ | [0,2 ²⁹ -1] | 1-based leftmost POSition/coordinate of the clipped sequence | | |
| MAPQ | [0-9]+ | [0,28-1] | MAPping Quality (phred-scaled posterior probability that the mapping position of this read is incorrect) ⁴ | | |
| CIGAR | ([0-9]+[MIDNSHP])+ * | | extended CIGAR string | | |
| MRNM | [^ \t\n\r@]+ | | Mate Reference sequence NaMe; "=" if the same as <rname> 3</rname> | | |
| MPOS | [0-9]+ | [0,2 ²⁹ -1] | 1-based leftmost Mate POSition of the clipped sequence | | |
| ISIZE | -?[0-9]+ | [-2 ²⁹ ,2 ²⁹] | inferred Insert SIZE 5 | | |
| SEQ | [acgtnACGTN.=]+ * | | query SEQuence; "=" for a match to the reference; n/N/. for ambiguity; cases are not maintained 6,7 | | |
| QUAL | [!-~]+ * | [0,93] | query QUALity; ASCII-33 gives the Phred base quality 6,7 | | |
| TAG | [A-Z][A-Z0-9] | | TAG | | |
| VTYPE | [AifZH] | | Value TYPE | | |
| VALUE | [^\t\n\r]+ | | match <vtype> (space allowed)</vtype> | | |



SAM

- Used for: aligned reads
- Lots of columns..





- Binary SAM (compressed)
- 25% of the size
- SAMtools to convert
- .bai = BAM index

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| | | |





- Random order
- Have to sort before indexing





- Random order
- Have to sort before indexing



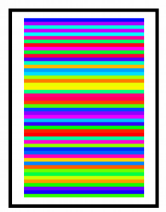


- Random order
- Have to sort before indexing

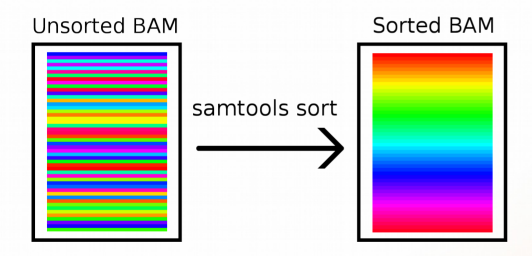




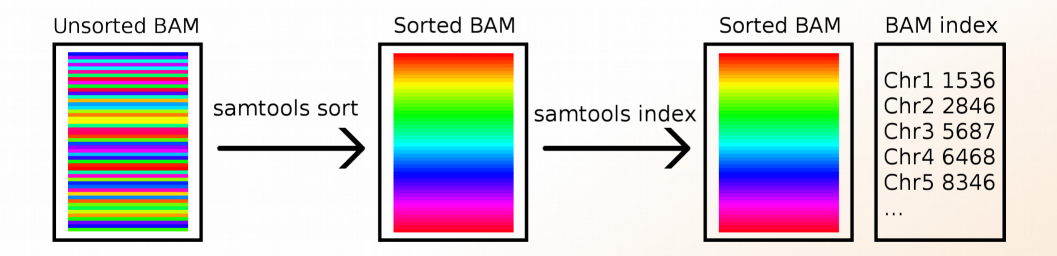
Unsorted BAM













CRAM

- Very complex format
- Used together with a reference genome

AGGCTGAGTCACGACGTGTTGAGA Reads

TAGATCGAGGCTGAGTCACGACG

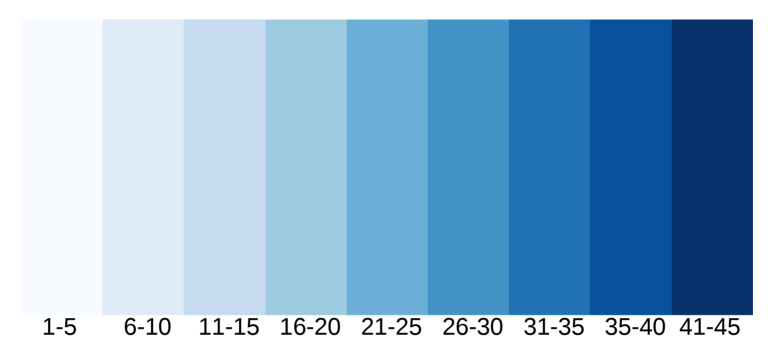
ATTCGGACGTAGATCGAGGCTGAG ACGTGTTGAGAGAGCCGTA

ATTCGGACGTAGATCGACGCTGAGTCACGACGTGTTGTGAGAGCCGTAGAC Ref:



CRAM

- Quality scores?
- 3 modes:
 - Lossless
 - Binned
 - No quality

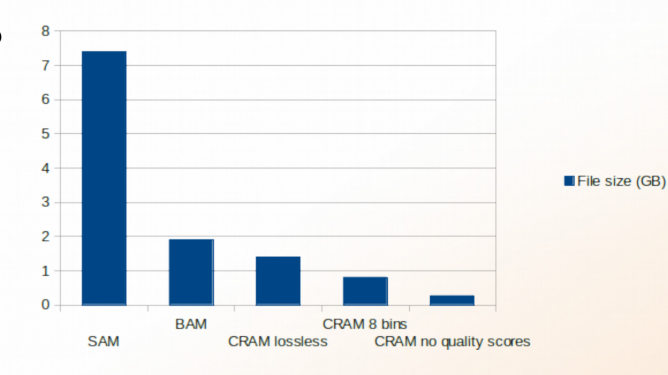


1 2 3 4 5 6 7 8 9 10 11 12 13 14 ... 32 33 34 35 36 37 38 39 40 41



CRAM

- Quality scores?
- 3 modes:
 - Lossless
 - Binned
 - No quality



Not widespread, yet



GTF/GFF/BED

- Used for: annotations
- Simple structure

- Usually:
 - chr start stop extra info



GTF/GFF/BED

- Used for: annotations
- Simple structure

Usually:chr start stop extra info

BED

```
chr22 1000 5000 cloneA 960 + 1000 5000 0 2 567,488, 0,3512 chr22 2000 6000 cloneB 900 - 2000 6000 0 2 433,399, 0,3601
```



GTF/GFF/BED

- Used for: annotations
- Simple structure

Usually:

chr start stop extra info

GFF

```
chr22 TeleGene enhancer 10000000 10001000 500 + . touch1 chr22 TeleGene promoter 10010000 10010100 900 + . touch1 chr22 TeleGene promoter 10020000 10025000 800 - . touch2
```



Laboratory time! (yet again)