# Interoperability: File formats

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Presented: April 21, 2022

#### About file formats

What is a file format?

A standardized layout/structure of information storage

Why is file formatting important?

File formats are created with specific goals in mind

#### File formats and FAIR data

- Interoperability
  - Most software relies on correctly formatted data
  - Many researchers are not able to reformat data themselves
- Reusability
  - Datasets are easier to use the more similar they are
  - Using established, sustainable file formats ensures datasets remain similar

#### Common file formats in omics research

- Fasta
- Fastq
- BED
- SAM/BAM
- VCF
- GFF
- JSON
- XML

### Examples: Fasta unaligned

Used to store nucleotide and amino acid sequences with headers

>tr|A0A679IRD3|A0A679IRD3\_9ENTE Multifunctional fusion protein OS=Enterococcus saigonensis OX=1805431 GN=rphA PE=3 SV=1
MRHDGRQVQQIRPVIIKTNVFKHPEGSVVISFGDTQVVCSATIEERVPHFLRDTGKGWVN
AEYSMLPRATQTRNRRESAKGKLSGRTMEIQRLIARSLRAVVDLEKLGERSIIVDCDVLQ
ADGGTRTASITGAFVALRLAINKLLQANVLTEDPIKEHLAAISVGILSDGTCVTDLDYVE
DFEASVDMNVVMTESGQFVELQGTGEESTFNGEELNEMLVYAKHAINDLIAFQKEALLGN
VALKEVIINENPKTIVIATKNPGKAKEFDALFSAAGYQVKTLLDYPEIPEVEETGHTFEE
NARLKAETIAHLLNRPVLADDSGLSVDALNGMPGVYSARFAGEMKSDAANNAKLLHELTH
VPDEDRTAHFHCTLVFAAPEKNSLSVSADWPGRIGRIPRGDDGFGYDPLFIPQGMEKTAA
ELSRTEKNAISHRGOAMKKLOKEWRTWLEA

> header
NUCLEOTIDEORAMINOACIDSEQUENCE

### Examples: Fasta aligned

>protein1
MRHDGRQVQQIRPVIIKTNVFKHPEGSVVISFGDTQVVCSATIEERVPHFLRDTGKGWVN
AEYSMLPRATQTRNRRESAKGKLSGRTMEIQRLIARSLRAVVDLEKLGERSIIVDCDVL\*
>protein2
MRHDGRQVQ-----TNVFKHPEGSVVISFGDTQVVCSATIEERVPHFLRDTGKGWVN
AEYSMLPRATQTRNRRESAKGKLSGRTMEIQRL-ARSLRAVVDLEKLGERSIIVDCDVL\*
>protein3
MRHDGRQVQQIRPVIIKTNVFKHPEGSVVISFGDTGIICSATIEERVPHFLRDTGKGWVN
AEYSMLPRATQTRNRRESAKGKLSGRTMEIQRLIARSLRAVVDLEK--ERSIIVDCDVL\*

### Examples: Fastq

Used for DNA sequences and their quality scores

```
@information:about:sequencing
DNASEQUENCE
+ (optional comment)
QUALITYSCORE
```

# Quality (Phred) scores

| Q  | P_error | ASCI | I        | Q  | P_error | ASCI | I | Q  | P_error | ASC | II | Q  | P_error | ASCII |
|----|---------|------|----------|----|---------|------|---|----|---------|-----|----|----|---------|-------|
| 0  | 1.00000 | 33   | !        | 11 | 0.07943 | 44   | , | 22 | 0.00631 | 55  | 7  | 33 | 0.00050 | 66 B  |
| 1  | 0.79433 | 34   | •        | 12 | 0.06310 | 45   | - | 23 | 0.00501 | 56  | 8  | 34 | 0.00040 | 67 C  |
| 2  | 0.63096 | 35   | ŧ        | 13 | 0.05012 | 46   |   | 24 | 0.00398 | 57  | 9  | 35 | 0.00032 | 68 I  |
| 3  | 0.50119 | 36   | \$       | 14 | 0.03981 | 47   | / | 25 | 0.00316 | 58  |    | 36 | 0.00025 | 69 E  |
| 4  | 0.39811 | 37   | ŧ        | 15 | 0.03162 | 48   | 0 | 26 | 0.00251 | 59  |    | 37 | 0.00020 | 70 F  |
| 5  | 0.31623 | 38   | <u>.</u> | 16 | 0.02512 | 49   | 1 | 27 | 0.00200 | 60  | <  | 38 | 0.00016 | 71 0  |
| 6  | 0.25119 | 39   |          | 17 | 0.01995 | 50   | 2 | 28 | 0.00158 | 61  | =  | 39 | 0.00013 | 72 H  |
| 7  | 0.19953 | 40   | (        | 18 | 0.01585 | 51   | 3 | 29 | 0.00126 | 62  | >  | 40 | 0.00010 | 73 I  |
| 8  | 0.15849 | 41   | )        | 19 | 0.01259 | 52   | 4 | 30 | 0.00100 | 63  | ?  | 41 | 0.00008 | 74 J  |
| 9  | 0.12589 | 42   | *        | 20 | 0.01000 | 53   | 5 | 31 | 0.00079 | 64  | 0  | 42 | 0.00006 | 75 K  |
| 10 | 0.10000 | 43   | +        | 21 | 0.00794 | 54   | 6 | 32 | 0.00063 | 65  | A  |    |         |       |

| Q | P_error | ASCII | Q  | P_error | ASCII | Q  | P_error | ASCII | Q  | P_error | ASCII |
|---|---------|-------|----|---------|-------|----|---------|-------|----|---------|-------|
| 0 | 1.00000 | 64 @  | 11 | 0.07943 | 75 K  | 22 | 0.00631 | 86 V  | 33 | 0.00050 | 97 a  |
| 1 | 0.79433 | 65 A  | 12 | 0.06310 | 76 L  | 23 | 0.00501 | 87 W  | 34 | 0.00040 | 98 b  |
| 2 | 0.63096 | 66 B  | 13 | 0.05012 | 77 M  | 24 | 0.00398 | 88 X  | 35 | 0.00032 | 99 c  |
| 3 | 0.50119 | 67 C  | 14 | 0.03981 | 78 N  | 25 | 0.00316 | 89 Y  | 36 | 0.00025 | 100 d |
| 4 | 0.39811 | 68 D  | 15 | 0.03162 | 79 0  | 26 | 0.00251 | 90 Z  | 37 | 0.00020 | 101 e |
| 5 | 0.31623 | 69 E  | 16 | 0.02512 | 80 P  | 27 | 0.00200 | 91 [  | 38 | 0.00016 | 102 f |
| 6 | 0.25119 | 70 F  | 17 | 0.01995 | 81 Q  | 28 | 0.00158 | 92 \  | 39 | 0.00013 | 103 g |
| 7 | 0.19953 | 71 G  | 18 | 0.01585 | 82 R  | 29 | 0.00126 | 93 ]  | 40 | 0.00010 | 104 h |
| 8 | 0.15849 | 72 H  | 19 | 0.01259 | 83 S  | 30 | 0.00100 | 94 ^  | 41 | 0.00008 | 105 i |
| 9 | 0.12589 | 73 I  | 20 | 0.01000 | 84 T  | 31 | 0.00079 | 95    | 42 | 0.00006 | 106 j |
| 0 | 0.10000 | 74 J  | 21 | 0.00794 | 85 U  | 32 | 0.00063 | 96 -  |    |         |       |

### Examples: BED (Browser Extensible Data)

- Tab-separated file used for alignment to a reference
- First three columns are required
- Example: http://useast.ensembl.org/info/website/upload/bed.html

```
Chrom chromStart chromEnd
                           name score strand thickStart thickEnd otherOptionalCols
     127471196
                127472363
                                       127471196
                                                  127472363
chr7
                           Pos1 0
chr7
     127472363
                127473530
                           Pos2 0 +
                                       127472363
                                                  127473530
     127473530
                127474697
                           Pos3 0 +
                                       127473530
                                                  127474697
chr7
chr7
     127474697
                127475864
                           Pos4 0 +
                                       127474697 127475864
                                       127475864
chr7
     127475864
                127477031
                           Neg1
                                 0 -
                                                  127477031
chr7
     127477031
                127478198
                           Neg2
                                 0
                                       127477031
                                                  127478198
chr7
     127478198
                127479365
                           Neg3 0
                                       127478198
                                                  127479365
chr7
     127479365
                127480532
                           Pos5 0 +
                                       127479365
                                                  127480532
     127480532
                127481699
                                       127480532
                                                  127481699
chr7
                           Neq4
```

### Examples: BAM/SAM

- SAM: Sequence Alignment Map
- BAM: Binary SAM
- Tab-separated table with a header
- Describes the full alignment of a sequence against a reference
- This tool can help you to decode flags

#### Annotated SAM example

```
Coor
              12345678901234 5678901234567890123456789012345
    ref -
             AGCATGTTAGATAA**GATAGCTGTGCTAGTAGGCAGTCAGCGCCAT
    +r001/1
                   TTAGATAAAGGATA*CTG
    +r002
                  aaaAGATAA*GGATA
    +r003
                gcctaAGCTAA
    +r004
                              ATAGCT.....TCAGC
    -r003
                                     ttagctTAGGC
    -r001/2
                                                   CAGCGGCAT
      The corresponding SAM format is:1
Version
    QHD VN:1.5 SO:coordinate
                                Length of reference
    QSQ SN:ref LN:45
    r001
                                          39 TTAGATAAAGGATACTG *
    r002
                        3S6M1P1I4M *
                                          O AAAAGATAAGGATA
    r003
                   9 30 5S6M
                                          O GCCTAAGCTAA
                                                               * SA:Z:ref,29,-,6H5M,17,0;
    r004
            0 ref 16 30 6M14N5M
                                          O ATAGCTTCAGC
    r003 2064 ref 29 17 6H5M
                                          O TAGGC
                                                               * SA:Z:ref,9,+,5S6M,30,1;
          147 ref 37 30 9M
                                                               * NM:i:1
    r001
                                        -39 CAGCGGCAT
                                         TEN
```

## Examples: VCF (Variant Call Format)

- Tab-separated table with a header
- After headers lines, contains one line per variant found
- Required columns:
  - o #CHROM
  - o POS

  - > REF
  - ALT
  - QUAL
  - FILTER
  - INFO
- A very thorough breakdown of VCF: https://samtools.github.io/hts-specs/VCFv4.2.pdf

## Examples: GFF3 (Generic Feature Format)

- Tab-separated table with 9 columns
- Official specifications: https://github.com/The-Sequence-Ontology/Specifications/blob/master/g ff3.md

```
##gff-version 3
ctg123 . exon 1300 1500 . + . ID=exon00001
ctg123 . exon 1050 1500 . + . ID=exon00002
ctg123 . exon 3000 3902 . + . ID=exon00003
ctg123 . exon 5000 5500 . + . ID=exon00004
ctg123 . exon 7000 9000 . + . ID=exon00005
```

### Examples: JSON (JavaScript Object Notation)

- Not specific to bioinformatics
- Used to store objects and can indicate structure

## Examples: XML (Extensible Markup Language)

- Not specific to bioinformatics
- Use case is similar to JSON, but XML is more complex

```
<menu id="file" value="File">
    <popup>
        <menuitem value="New" onclick="CreateNewDoc()" />
        <menuitem value="Open" onclick="OpenDoc()" />
        <menuitem value="Close" onclick="CloseDoc()" />
        </popup>
</menu>
```

#### Additional resources

https://genome.ucsc.edu/FAQ/FAQformat.html#format1

https://bioinformatics.uconn.edu/resources-and-events/tutorials-2/file-format s-tutorial/

http://useast.ensembl.org/info/website/upload/bed.html

https://www.toptal.com/web/json-vs-xml-part-1#:~:text=JSON%20is%20a%20data%20interchange,of%20any%20XML%20sub%2Dlanguage.