

DATA FORMATS IN NGS INTRODUCTION TO GALAXY

Bioinformàtica per a la Recerca Biomèdica

Mireia Ferrer¹, Álex Sánchez^{1,2} Esther Camacho¹, Angel Blanco^{1,2}

1 Unitat d'Estadística i Bioinformàtica (UEB) VHIR 2 Departament de Genètica, Microbiologia i Estadística, UB















- 1. Data formats used in NGS
- 2. Introduction to Galaxy

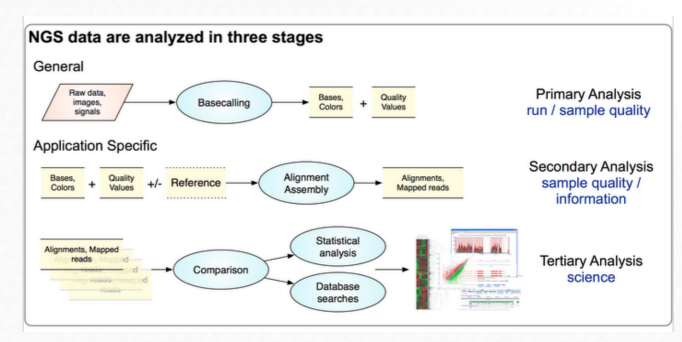


- 1. Data formats used in NGS
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There are many different types of file formats depending on:

- Type of information they contain
 - Raw Sequence files
 - Co-ordinate files
 - Parameter files
 - Annotation files
 - Metadata files
- Sequencing platform
- Analysis stage
- Data source





- Formats are designed to hold sequence data and other information about sequence
- All Sequence formats are ASCII text containing sequence ID, Quality Scores,
 Annotation details, comments, and other descriptions about sequence



FASTA format

- FASTA format is a text-based format for representing either nucleotide sequences or peptide sequences, in which base pairs or amino acids are represented using single-letter codes
- Header line starts with ">" followed by a sequence ID, and followed by lines of sequence data

>NG_016798.2 Homo sapiens DNA polymerase alpha 1, catalytic subunit (POLA1), RefSeqGene on chromosome X

>NP_001365232.1 DNA polymerase alpha catalytic subunit isoform 3 [Homo sapiens]
MAPVHGDDCEIGASALSDSGSFVSSRARREKKSKKGRQEALERLKKAKAGEKYKYEVEDFTGVYEEVDEE
QYSKLVQARQDDDWIVDDDGIGYVEDGREIFDDDLEDDALDADEKGKDGKARNKDKRNVKKLAVTKPNNI
KSMFIACAGKKTADKAVDLSKDGLLGDILQDLNTETPQITPPPVMILKKKRSIGASPNPFSVHTATAVPS
GKIASPVSRKEPPLTPVPLKRAEFAGDDVQVESTEEEQESGAMEFEDGDFDEPMEVEEVDLEPMAAKAWD
KESEPAEEVKQEADSGKGTVSYLGSFLPDVSCWDIDQEGDSSFSVQEVQVDSSHLPLVKGADEEQVFHFY
WLDAYEDQYNQPGVVFLFGKVWIESAETHVSCCVMVKNIERTLYFLPREMKIDLNTGKETGTPISMKDVY
EEFDEKIATKYKIMKFKSKAEMPQLPQDLKGETFSHVFGTNTSSLELFLMNRKIKGPCWLEVKSPQLLNQ
PVSWCKVEAMALKPDLVNVIKDVSPPPLVVMAFSMKTMQNAKNHQNEIIAMAALVHHSFALDKAAPKPPF
QSHFCVVSKPKDCIFPYAFKEVIEKKNVKVEVAATERTLLGFFLAKVHKIDPDIIVGHNIYGFELEVLLQ
RINVCKAPHWSKIGRLKRSNMPKLGGRSGFGERNATCGRMICDVEISAKELIRCKSYHLSELVOOILKTE



FASTQ format

- Output of most actual sequencing platforms for raw data
- A text-based format for storing both a nucleotide sequence and its corresponding quality scores
- Standard file extension for a FASTQ file are .fq and .fastq
- FASTQ files are uncompressed and quite large because they contain the following information for every single sequencing read.
- Compressed files are also possible: fastq.gz



FASTQ format

- File structure. 4 lines:
 - @ followed by the read ID and possibly information about the sequencing run
 - sequenced bases
 - + (perhaps followed by the read ID again, or some other description)
 - quality scores for each base of the sequence (ASCII-translated Phred scores)

```
@Seq description
GATTTGGGGTTCAAAGCAGTATCGATCAAATAGTAAATCCATTTGTTCAACTCACAGTTT
+
!''*((((***+))%%%++)(%%%%).1***-+*''))**55CCF>>>>>CCCCCCC65
```



FASTQ format

Phred Scores

- Sequencing systems assign quality scores to each peak, that represents the error probability that an individual base call is incorrect.
- Phred scores provide $\log(10)$ -transformed error probability values: If p is probability that the base call is wrong the Phred score is $Q = .10 \cdot log_{10}p$

PHRED Quality Score	Probability of incorrect base call	Base call accuracy
10	1 in 10	90 %
20	1 in 100	99 %
30	1 in 1000	99.9 %
40	1 in 10000	99.99 %
50	1 in 100000	99.999 %



- The base calling (A, T, G or C) is performed based on Phred scores.
- Ambiguous positions with Phred scores <= 20 are labeled with N.
- To assign each base a unique score identifier (instead of numbers of varying character length), Phred scores are typically represented as ASCII characters.
- Different sequencing platforms may use different ASCII ranges for Phred encoding

Description	ASCII characters		Quality score	
	Range	Offset	Type	Range
Solexa/early Illumina (1.0)	59 to 126 (; to ~)	64	Solexa	-5 to 62
Illumina $1.3+$	64 to 126 (@ to $\tilde{\ }$	64	Phred	0 to 62
Sanger standard/Illumina $1.8+$	33 to 126 (! to $\tilde{\ }$)	33	Phred	0 to 93

Base call quality scores are represented with the Phred range. Different Illumina (formerly Solexa) versions used different scores and ASCII offsets. Starting with Illumina format 1.8, the score now represents the standard Sanger/Phred format that is also used by other sequencing platforms and the sequencing archives.



SAM / BAM formats

- The **Sequence Alignment/Map (SAM)** format is a generic nucleotide alignment format that describes the alignment of sequencing reads to a reference.
- SAM files typically contain:
 - a short header section with information about the genomic loci of each read
 - a very long alignment section where each row represents a single read alignment.
 - Each alignment line has 11 mandatory fields for essential alignment information such as mapping position, and variable number of optional fields for flexible or aligner specific information



Mandatory Alignment Section Fields

Position	Field	Description	
1	QNAME	Query template (or read) name	
2	FLAG	Information about read mapping (see next section)	
3	RNAME	Reference sequence name. This should match a @SQ	
		line in the header.	
4	POS	1-based leftmost mapping position of the first	
		matching base. Set as 0 for an unmapped read without	
		coordinate.	
5	MAPQ	Mapping quality of the alignment. Based on base	
		qualities of the mapped read.	
6	CIGAR	Detailed information about the alignment (see relevant	
		section).	
7	RNEXT	Used for paired end reads. Reference sequence name of	
		the next read. Set to "=" if the next segment has the	
		same name.	
8	PNEXT	Used for paired end reads. Position of the next read.	
9	TLEN	Observed template length. Used for paired end reads	
		and is defined by the length of the reference aligned to.	
10	SEQ	The sequence of the aligned read.	
11	QUAL	ASCII of base quality plus 33 (same as the quality	
		string in the Sanger FASTQ format).	
12	OPT	Optional fields (see relevant section).	



```
QHD VN:1.6 SO:coordinate
@SQ SN:ref LN:45
r001
     99 ref 7 30 8M2I4M1D3M = 37 39 TTAGATAAAGGATACTG *
r002 0 ref 9 30 3S6M1P1I4M *
                                     O AAAAGATAAGGATA
r003 0 ref 9 30 5S6M
                                     O GCCTAAGCTAA
                                                         * SA:Z:ref,29,-,6H5M,17,0;
r004
     0 ref 16 30 6M14N5M
                              * O O ATAGCTTCAGC
r003 2064 ref 29 17 6H5M
                                     O TAGGC
                                                         * SA:Z:ref,9,+,5S6M,30,1;
r001 	 147 	 ref 	 37 	 30 	 9M 	 = 	 7 	 -39 	 CAGCGGCAT
                                                         * NM:i:1
```



SAM / BAM formats

- A BAM file is a binary version of a SAM file.
- Both contain identical information about reads and their mapping.
- A BAM file requires a header but a SAM file may not have one.
- Many operations (such as sorting and indexing) work only on BAM files.
- For almost any application that requires SAM input, this can be created on the fly from a BAM,
- BAM files take up much less space than SAM files.
- For archiving purposes, keep only the BAM file. The SAM file can easily be regenerated (if ever needed).



BED / GFF / GTF formats

- Formats for genome annotations
- One line per genomic feature
- The BED format is the simplest way to store annotation tracks. It has three required fields (chromosome, start, end) and up to 9 optional fields (name, score, strand, thickStart, thickEnd, itemRgb, blockCount, blockSizes, blockStarts).

```
# 6-column BED file defining transcript loci
chr1
     66999824
               67210768
                         NM_032291 0 +
     33546713
               33586132
                         NM_052998 0 +
chr1
     25071759
               25170815
                         NM_013943 0 +
chr1
     48998526
               50489626
                         NM_032785 0 -
chr1
```



BED / GFF / GTF formats

 The General Feature Format (GFF) and General Transfer Format (GTF) has nine required fields; the first three fields form the basic name, start, end tuple that allows for the identification of the location in respect to the reference genome.

```
Sample GTF output from Ensembl data dump:
 1 transcribed unprocessed pseudogene gene
                                                   11869 14409 . + . gene id "ENSG00000223972"; gene name "DDX11L1"; gene source "havana"; gene biotype "transcribed unprocessed pseudogene";
                                       transcript 11869 14409 . + . gene id "ENSG00000223972"; transcript id "ENST00000456328"; gene name "DDX11L1"; gene sourc e "havana"; gene biotype "transcript
 1 processed transcript
Sample GFF output from Ensembl export:
         Ensembl Repeat 2419108 2419128 42
                                                                 hid=trf; hstart=1; hend=21
         Ensembl Repeat 2419108 2419410 2502
                                                                 hid=AluSx; hstart=1; hend=303
         Ensembl Repeat 2419108 2419128 0
                                                                 hid=dust; hstart=2419108; hend=2419128
         Ensembl Pred.trans. 2416676 2418760 450.19 -
                                                                         genscan=GENSCAN00000019335
         Ensembl Variation 2413425 2413425 .
         Ensembl Variation
                                2413805 2413805 .
```



- 1. reference sequence: coordinate system of the annotation (e.g., "Chr1")
- 2. **source**: describes how the annotation was derived (e.g., the name of the annotation software)
- 3. **method**: annotation type (e.g., gene)
- 4. start position: 1-based integer, always less than or equal to the stop position
- stop position: for zero-length features, such as insertion sites, start equals end and the implied site is to the right of the indicated base
- score: e.g., sequence identity
- strand: "+" for the forward strand, "-" for the reverse strand, or "." for annotations that are not stranded
- 8. phase: codon phase for annotations linked to proteins; 0, 1, or 2, indicating the frame, or the number of bases that should be removed from the beginning of this feature to reach the first base of the next codon
- group: contains the class and ID of an annotation which is the logical parent of the current one ("feature is composed of")



VCF format

- Variant Call Format (VCF) is a text file format.
- It contains meta-information lines, a header line, and then data lines each containing information about a position in the genome.
- Also has the ability to contain genotype information on samples for each position.
- The header line names the 8 fixed, mandatory columns.
 - 1.
 - #CHROM
 - 2. POS
 - 3. ID
 - 4. REF
 - 5. ALT
 - 6. QUAL
 - 7. FILTER
 - 8. INFO



```
##fileformat=VCFv4.2
##fileDate=20090805
##source=myImputationProgramV3.1
##reference=file:///seq/references/1000GenomesPilot-NCBI36.fasta
##contig=<ID=20,length=62435964,assembly=B36,md5=f126cdf8a6e0c7f379d618ff66beb2da,species="Homo sapiens",taxonomy=x>
##phasing=partial
##INFO=<ID=NS.Number=1.Type=Integer,Description="Number of Samples With Data">
##INFO=<ID=DP, Number=1, Type=Integer, Description="Total Depth">
##INFO=<ID=AF, Number=A, Type=Float, Description="Allele Frequency">
##INFO=<ID=AA, Number=1, Type=String, Description="Ancestral Allele">
##INFO=<ID=DB, Number=0, Type=Flag, Description="dbSNP membership, build 129">
##INFO=<ID=H2, Number=0, Type=Flag, Description="HapMap2 membership">
##FILTER=<ID=q10, Description="Quality below 10">
##FILTER=<ID=s50,Description="Less than 50% of samples have data">
##FORMAT=<ID=GT, Number=1, Type=String, Description="Genotype">
##FORMAT=<ID=GQ, Number=1, Type=Integer, Description="Genotype Quality">
##FORMAT=<ID=DP, Number=1, Type=Integer, Description="Read Depth">
##FORMAT=<ID=HQ, Number=2, Type=Integer, Description="Haplotype Quality">
#CHROM POS
                                ALT
                                        QUAL FILTER INFO
                                                                                       FORMAT
                                                                                                    NA00001
                                                                                                                   NA00002
                                                                                                                                  NA00003
20
       14370 rs6054257 G
                                             PASS
                                                    NS=3;DP=14;AF=0.5;DB;H2
                                                                                       GT:GQ:DP:HQ 0|0:48:1:51,51 1|0:48:8:51,51 1/1:43:5:...
       17330
                                                     NS=3:DP=11:AF=0.017
                                                                                       GT:GQ:DP:HQ 0|0:49:3:58,50 0|1:3:5:65,3
                                                                                                                                  0/0:41:3
                                             q10
                                                    NS=2;DP=10;AF=0.333,0.667;AA=T;DB GT:GQ:DP:HQ 1|2:21:6:23,27 2|1:2:0:18,2
       1110696 rs6040355 A
                                G,T
                                             PASS
                                                                                                                                  2/2:35:4
      1230237 .
                                             PASS
                                                    NS=3:DP=13:AA=T
20
                                                                                       GT:GQ:DP:HQ 0|0:54:7:56,60 0|0:48:4:51,51 0/0:61:2
                                                    NS=3; DP=9; AA=G
       1234567 microsat1 GTC
                                G.GTCT 50
                                             PASS
                                                                                       GT:GQ:DP
                                                                                                    0/1:35:4
                                                                                                                   0/2:17:2
                                                                                                                                  1/1:40:3
```

https://samtools.github.io/hts-specs/VCFv4.2.pdf



To sum up

- Raw data: .fastq (.fastq.gz)
- Aligned data: .sam / .bam
- Annotation data: .gtf / .gff / .bed
- Results data: .vcf