# NGS - quality control, alignment, visualisation

File types

### File types

fasta	sequences
fastq	reads
sam/bam	alignments
bed	regions
gff	annotations
vcf	variants

#### fasta

- Plain sequence: \*.fasta or \*.fa
- Nucleotides or amino acids (proteins)
- Useful command:grep -c "^>" sequence.fasta

#### sequence.fasta

>sequence title1

**ATCGTATCT** 

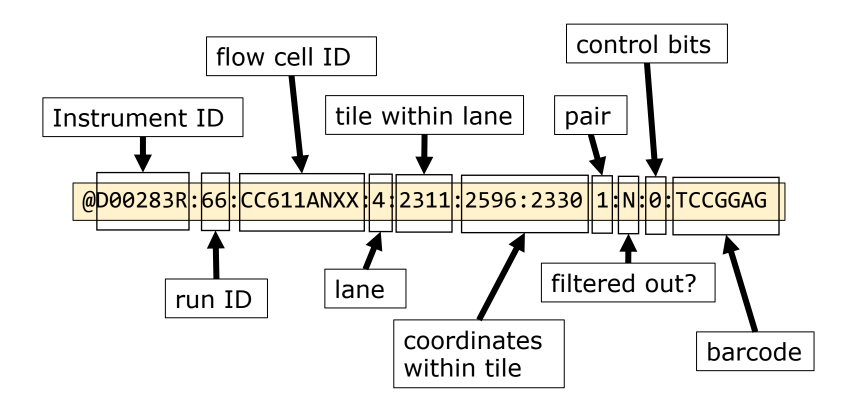
>sequence title2

**ATGATGACGT** 

#### fastq



### fastq header (Illumina)



### Base quality (phred)

```
.....
!"#$%&'()*+,-./0123456789:;<=>?@ABCDEFGHIJKLMNOPQRSTUVWXYZ[\]^_`abcdefghijklmnopqrstuvwxyz{|}~
33
                                       104
                                                   126
0.2......41
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, 41)
  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)
```

### Quiz Question 5

#### sam

sequence alignment format Aim: alignments

#### sam header

```
@HD VN:1.0 S0:coordinate
@SQ SN:U00096.3 LN:4641652
@PG ID:bowtie2 PN:bowtie2 VN:2.4.1
CL: "/opt/miniconda3/envs/ngs/bin/bowtie2-align-s \
--wrapper basic-0 \
-x /home/ubuntu/ecoli/ref_genome//ecoli-strK12-MG1655.fasta \
-1 /home/ubuntu/ecoli/trimmed_data/paired_trimmed_SRR519926_1.fastq \
-2 / home/ubuntu/ecoli/trimmed_data/paired_trimmed_SRR519926_2.fastq"
```

SAM column	example
read name	SRR519926.5
flag	89
reference	U00096.3
start position	61
mapping quality	42
CIGAR string	214M
reference name mate is mapped	=
start position mate	476
fragment length	515
sequence	CATCACCATTCCCAC
base quality	@>4:4C@89+&9CC@
optional	AS:i:-2
optional	XN:i:0

### Quiz Question 6

### sam flags

I	Bit	Description
1	0x1	template having multiple segments in sequencing
$^2$	0x2	each segment properly aligned according to the aligner
4	0x4	segment unmapped
8	0x8	next segment in the template unmapped
16	0x10	SEQ being reverse complemented
32	0x20	SEQ of the next segment in the template being reverse complemented
64	0x40	the first segment in the template
128	0x80	the last segment in the template
256	0x100	secondary alignment
512	0x200	not passing filters, such as platform/vendor quality controls
1024	0x400	PCR or optical duplicate
2048	0x800	supplementary alignment

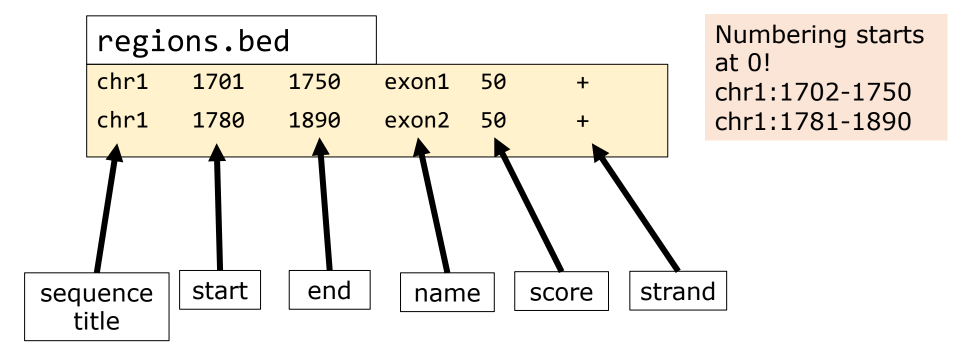
	read paired?	properly aligned?	unmapped?	next unmapped?	flag
read1	0	1	0	0	2
read2	1	0	1	1	13
read3	1	1	0	1	11
	1	2	4	8	

### Quiz Question 7

#### bed

#### Browser Extensible Data

Aim: specify regions



### gff

#### General Feature Format

Aim: annotation

seq name	source	feature	start	end	score	strand	frame	attributes
1	ensembl	mRNA	339070	346959		-		ID=;
1	ensembl	exon	339070	339312	•	-		Parent=;
1	ensembl	CDS	339070	339312		ı	0	ID=;

#### vcf

##fileformat=VCFv4.2
##fileDate=20090805

##source=myImputationProgramV3.1

## Variant Call Format Aim: variants

##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
                                                                                         FORMAT
                                 ALT
                                         QUAL FILTER INFO
                                                                                                      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:.,.
20
       17330
                                              q10
                                                      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
20
       1110696 rs6040355 A
                                 G, T
                                              PASS
                                                      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
                                                                                                                                     2/2:35:4
       1230237 .
                                              PASS
                                                      NS=3; DP=13; AA=T
                                                                                         GT:GQ:DP:HQ 0|0:54:7:56,60 0|0:48:4:51,51 0/0:61:2
       1234567 microsat1 GTC
                                 G,GTCT 50
                                              PASS
                                                      NS=3; DP=9; AA=G
                                                                                         GT:GQ:DP
                                                                                                      0/1:35:4
                                                                                                                     0/2:17:2
                                                                                                                                     1/1:40:3
```