### Introduction to Bioinformatics

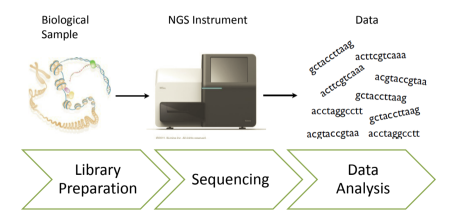
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### **NGS** Overview





# NGS "Hardware" I



**Roche GS-FLX** 



Illumina HiSeq



**Life Technologies SOLID** 



**Life Technologies Ion Torrent** 



**Life Technologies Ion Proton** 



Illumina MiSeq



# NGS "Hardware" II

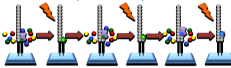


Pacbio Sequel

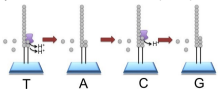


# NGS Technologies I

fluorescence-based (Illumina)



hydrogen ion /pH-mediated based (Life)





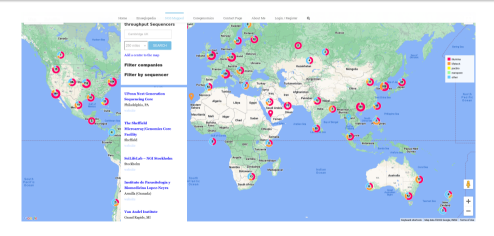
# NGS Technologies II

	Run Time	Read Length	Quality	Total nucleotides sequenced	Cost /MB	
454 Pyrosequencing	24h	700 bp	Q20-Q30	1 GB	\$10	
Illumina Miseq	27h	2x300bp	> Q30	15 GB	\$0.15	
Illumina Hiseq 2500	1 - 10days	2x250bp	>Q30	3000 GB	\$0.05	
Ion torrent	2h	400bp	>Q20	50MB-1GB	\$1	
Pacific Biosciences	30m - 4h	10kb - >40kb	>Q50 consensus >Q10 single	500 - 1000MB /SMRT cell	\$0.13 - \$0.60	

http://www.hindawi.com/journals/bmri/2012/251364



# Growing demand





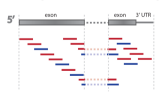
http://enseqlopedia.com/ngs-mapped/

# Sequencing options

#### Single-end

- Cheaper
- Suitable for more general purpose analyzes, e.g. DE

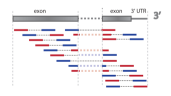
#### Single-end sequencing



#### Paired-end

- More information regarding the length and the position of a read
- Useful for spliced junctions, indels etc.

#### Paired-end sequencing



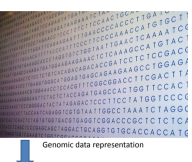


Zhernakova, Daria V., et al. "DeepSAGE reveals genetic variants associated with alternative polyadenylation and expression of coding and non-coding transcripts."



# Bioinformatics is a necessity I

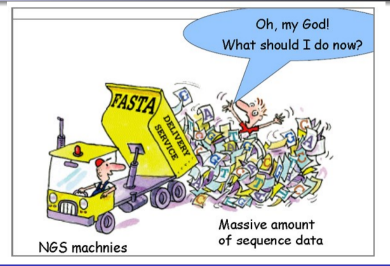








# Bioinformatics is a necessity II





# FASTQ file format

#### **FASTQ**

Useful file types

Text-based format for storing biological sequences

- Raw unaligned reads (nucleotides)
- Corresponding quality scores

@HWI-ST661:319:D28MYACXX:6:1101:1170:2180 1:N:0:GTGGCC NAGTGGTTTATGCCTGTAATCCCAGCATTTTGGGAGACGAAGTTGAGAN

+

#1:ADDFFHGHHHIJGHHIIJJJIIIEHIJJJIHEHIGHIJJHHJGHC#



# Phred quality score

Useful file types

$$Q = -10 \log_{10} P$$

P: base-calling error probability

Q: Phred quality score

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 10,000	99.99%
50	1 in 100,000	99.999%
60	1 in 1,000,000	99.9999%



# Phred quality score encoding

Useful file types

```
!"#$%&'()*+,-./0123456789:;<=>?@ABCDEFGHIJKLMNOPQRSTUVWXYZ[\]^ `abcdefghijklmnopqrstuvwxyz{|}~
33
                                       104
                                                    126
              26...31......40
                  0 9 40
                   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)
L - Illumina 1.8+ Phred+33, raw reads typically (0, 41)
```

https://en.wikipedia.org/wiki/FASTO format



# FASTA file format I

#### FASTA

Useful file types

Text-based format for representing either nucleotide sequences or peptides encoded as a character

- Starts with the character ">" followed by an (alpharithmetic) identification code
- One or more lines contain the sequence



# FASTA file format II

Useful file types

- During the alignment process the reads of a fastq file are mapped on a reference genome stored in fasta format
- There are various available genomes, e.g.:
  - Human: hg16 (2003), hg17 (2004), hg18 (2006), hg19 (NCBI)/GRCh37 (Ensembl) (2009), hg38/GRCh38 (2013)
  - Mouse: mm7 (2005), mm8 (2006), mm9 (2007), mm10 (2011)
  - D. melanogaster: dm1 (2003), dm2 (2004), dm3 (2006), dm6 (2014)
  - ...



# SAM/BAM format I

Useful file types

### SAM - Sequence Alignment Map

SAM format stores aligned reads and is independent of the sequencing technology used

- SAM: textbased
- BAM: binary



# SAM/BAM format II

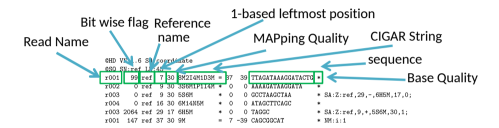
Useful file types

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



# SAM/BAM format III

Useful file types





# **BED** format

Useful file types

#### BED - Browser Extensible Data

#### Per line

- 3 mandatory fields
  - chrom The chromosome name, e.g. chr3, chrY, chr2\_random
  - chromStart The start position of a feature in the chromosome **0-based numbering**.
  - chromEnd The end position of a feature in the chromosome. The end position **in not** included in the feature. E.g. the first 100 bases of a chromosome are defined as chromStart=0, chromEnd=100, and are the bases 0-99
- 9 optional fields
  - name, score, strand, thickStart, thickEnd, itemRgb, blockCount, blockSizes, blockStarts



# VCF format I

Useful file types

#### VCF files - Variant Call Format

- For encoding structural variations
- Widely used by the 1000 Genomes Project
- Only the variants are stored along with the reference genome



# VCF format II

Useful file types

```
##fileformat=VCFv4.1
    ##fileDate=20090805
    ##source=mvImputationProgramV3.1
    ##reference=file:///seg/references/1000GenomesPilot-NCBI36.fasta
    ##contig=<TD=20.length=62435964.assembly=B36.md5=f126cdf8a6e0c7f379d618ff66beb2da.species="Homo_sapiens".taxonomy=x>
    ##phasing=partial
    ##INFO=<ID=NS.Number=1.Tvpe=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=g10, Description="Quality below 10">
14 ##FILTER=<ID=s50, Description="Less than 50% of samples have data">
    ##FORMAT=<ID=GT.Number=1.Type=String.Description="Genotype">
16 ##FORMAT=<ID=GO.Number=1.Type=Integer.Description="Genotype Quality">
    ##FORMAT=<ID=DP.Number=1.Type=Integer.Description="Read Depth">
    ##FORMAT=<ID=HO, Number=2, Type=Integer, Description="Haplotype Quality">
19 #CHROM POS
                   TD
                                            QUAL FILTER INFO
                                                                                          FORMAT
                                                                                                      NACCOCA
                                                                                                                     NACCOOCS
                                                                                                                                    NACCOCA
           14370
                   rs6054257 G
                                            29
                                                 PASS
                                                        NS=3:DP=14:AF=0.5:DB:H2
                                                                                          GT:GO:DP:HO 010:48:1:51.51 110:48:8:51.51 1/1:43:5:...
21 20
           17330
                                                 g10
                                                        NS=3:DP=11:AF=0.017
                                                                                          GT:GO:DP:HO 010:49:3:58.50 011:3:5:65.3 0/0:41:3
22 20
           1110696 rs6040355 A
                                    G.T
                                            67
                                                 PASS
                                                        NS=2:DP=10:AF=0.333.0.667:AA=T:DB GT:GO:DP:HO 1|2:21:6:23.27 2|1:2:0:18.2 2/2:35:4
           1230237
                                                 DASS
                                                        NS=3:DP=13:AA=T
                                                                                          GT:GO:DP:HO 010:54:7:56.60 010:48:4:51.51 0/0:61:2
           1234567 microsatl GTC
                                    G.GTCT 50
                                                 PASS
                                                        NS=3:DP=9:AA=G
                                                                                                      0/1:35:4
                                                                                                                     0/2:17:2
                                                                                                                                    1/1:40:3
                                                                                          GT:GQ:DP
```



# **VCF** format III

Useful file types

#CHROM	POS	ID	REF	ALT	QUAL	FILTER	INFO	FORMAT	NA00001	NA00002	NA00003
20	14370	rs6054257	G	A	29	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		T	A	3	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	67	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
20	1230237		T		47	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
20	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

- AF: allele frequency for each ALT allele
- DP: combined depth across samples
- DB: dbSNP membership
- H2: membership in hapmap2
- NS: Number of samples with data



# index files

Useful file types

Index creation, for faster data access since it enables non sequential search

- fasta  $\rightarrow$  fai
- bam  $\rightarrow$  bai
- $vcf \rightarrow idx$
- $vcf.gz \rightarrow tbi$



### samtools I

#### samtools

Samtools is a suite of programs for interacting with high-throughput sequencing data. It consists of three separate repositories:

Samtools Reading/writing/editing/indexing/viewing SAM/BAM/CRAM format **BCFtools** Reading/writing BCF2/VCF/gVCF files and calling/filtering/summarising SNP and short indel sequence variants

**HTSlib** A C library for reading/writing high-throughput sequencing data

http://www.htslib.org/



### samtools II

```
$ samtools
```

```
Program: samtools (Tools for alignments in the SAM format)
```

Version: 1.3 (using htslib 1.3)

Usage: samtools <command> [options]

#### Commands:

-- Indexing

dict create a sequence dictionary file

faidx index/extract FASTA

index index alignment



### samtools III

-- Editing

calmd recalculate MD/NM tags and '=' bases

fixmate fix mate information reheader replace BAM header

rmdup remove PCR duplicates

cut fosmid regions (for fosmid pool only) targetcut

addreplacerg adds or replaces RG tags

-- File operations

collate shuffle and group alignments by name

concatenate BAMs cat

merge sorted alignments merge

mpileup multi-way pileup



```
sort sort alignment file split splits a file by read group
```

quickcheck quickly check if SAM/BAM/CRAM file appears intact

fastq converts a BAM to a FASTQ fasta converts a BAM to a FASTA

-- Statistics

bedcov read depth per BED region

depth compute the depth

flagstat simple stats

idxstats BAM index stats

phase phase heterozygotes

stats generate stats (former bamcheck)



# samtools V

-- Viewing

flags explain BAM flags

tview text alignment viewer

view SAM<->BAM<->CRAM conversion

depad convert padded BAM to unpadded BAM



# samtools VI

• sort: after the alignment process, the final sam file contains the reads in a random order. For random access and also for the conversion to bam, the sam file needs to be sorted

```
samtools sort sample.bam sample.sorted -08
```

- index: for the faster data access of the bam file samtools index sample.sorted.bam
- view: for sam to bam conversion and vice versa. For filtering a sam/bam file based on conditions

```
samtools view sample.bam
samtools view -f 0x2 sample.sorted.bam
samtools view -F 0x2 sample.sorted.bam
```



### Bedtools I

Introduction

### bedtools: a powerful toolset for genome arithmetic

Collectively, the bedtools utilities are a swiss-army knife of tools for a wide-range of genomics analysis tasks. The most widely-used tools enable genome arithmetic: that is, set theory on the genome. For example, bedtools allows one to intersect, merge, count, complement, and shuffle genomic intervals from multiple files in widely-used genomic file formats such as BAM, BED, GFF/GTF, VCF. While each individual tool is designed to do a relatively simple task (e.g., intersect two interval files), quite sophisticated analyses can be conducted by combining multiple bedtools operations on the UNIX command line

http://bedtools.readthedocs.io/en/latest/



### **Bedtools II**

```
$ bedtools
```

bedtools: flexible tools for genome arithmetic and DNA sequence analysis.

bedtools <subcommand> [options] usage:

The bedtools sub-commands include:

```
[ Genome arithmetic ]
```

intersect Find overlapping intervals in various ways.

window Find overlapping intervals within a window around

an interval.

. . .



### **Bedtools III**

#### bedtools intersect

By far, the most common question asked of two sets of genomic features is whether or not any of the features in the two sets "overlap" with one another. This is known as feature intersection. bedtools intersect allows one to screen for overlaps between two sets of genomic features. Moreover, it allows one to have fine control as to how the intersections are reported. bedtools intersect works with both BED/GFF/VCF and BAM files as input.



### **Bedtools IV**

```
$ bedtools intersect
```

Tool: bedtools intersect (aka intersectBed)

Version: v2.21.0

Summary: Report overlaps between two feature files.

bedtools intersect [OPTIONS] -a <bed/gff/vcf> -b <bed/gff/vcf> Usage:

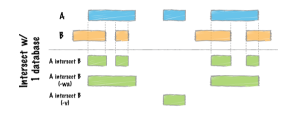
Note: -b may be followed with multiple databases and/or

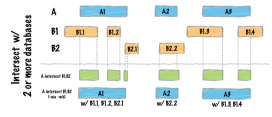
wildcard (\*) character(s).

. . .



# Bedtools V







# Bedtools VI

```
$ cat A.bed
chr1 10 20
chr1 30 40

$ cat B.bed
chr1 15 20

$ bedtools intersect -a A.bed -b B.bed
chr1 15 20
```



# **Bedtools VII**

#### bedtools merge

bedtools merge combines overlapping or "book-ended" features in an interval file into a single feature which spans all of the combined features.



### **Bedtools VIII**

```
$ bedtools merge
```

Tool: bedtools merge (aka mergeBed)

Version: v2.21.0

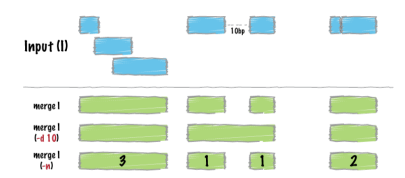
Summary: Merges overlapping BED/GFF/VCF entries into a single interval.

Usage: bedtools merge [OPTIONS] -i <bed/gff/vcf>

. . .



## Bedtools IX





## Bedtools X

```
$ cat A.bed
chr1
      100
           200
chr1
      180
           250
chr1
      250
           500
chr1
      501
           1000
$ bedtools merge -i A.bed
chr1
      100
           500
      501
           1000
chr1
```



### Bedtools XI

### bedtools genomecov

bedtools genomecov computes histograms (default), per-base reports (-d) and BEDGRAPH (-bg) summaries of feature coverage (e.g., aligned sequences) for a given genome.



### **Bedtools XII**

```
$ bedtools genomecov
```

Tool: bedtools genomecov (aka genomeCoverageBed)

Version: v2.21.0

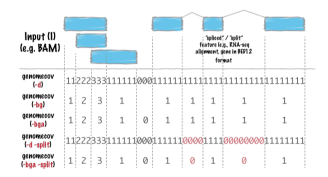
Summary: Compute the coverage of a feature file among a genome.

Usage: bedtools genomecov [OPTIONS] -i <bed/gff/vcf> -g <genome>

. . .



## **Bedtools XIII**





### **Bedtools XIV**

```
$ cat A.bed
chr1
      10
          20
chr1
      20
          30
chr2
          500
      0
$ cat my.genome
      1000
chr1
chr2
      500
$ bedtools genomecov -i A.bed -g my.genome
chr1
          980
               1000
                     0.98
chr1
          20
               1000
                     0.02
```



500

500

chr2

## **Bedtools XV**

genome 0 980 1500 0.653333 genome 1 520 1500 0.346667



### **VCFtools I**

#### Welcome to VCFtools

VCFtools is a program package designed for working with VCF files, such as those generated by the 1000 Genomes Project. The aim of VCFtools is to provide easily accessible methods for working with complex genetic variation data in the form of VCF files.

https://vcftools.github.io/index.html



### **VCFtools II**

\$ vcftools

VCFtools (0.1.15)

© Adam Auton and Anthony Marcketta 2009

Process Variant Call Format files

For a list of options, please go to: https://vcftools.github.io/man\_latest.html

Alternatively, a man page is available, type:

man vcftools

Questions, comments, and suggestions should be emailed to: vcftools-help@lists.sourceforge.net



### **VCFtools III**

#### vcf-isec

Creates intersections and complements of two or more VCF files. Given multiple VCF files, it can output the list of positions which are shared by at least N files, at most N files, exactly N files, etc. The first example below outputs positions shared by at least two files and the second outputs positions present in the files A but absent from files B and C.

```
vcf-isec -n +2 A.vcf.gz B.vcf.gz | bgzip -c > out.vcf.gz
vcf-isec -c A.vcf.gz B.vcf.gz C.vcf.gz | bgzip -c > out.vcf.gz
```



### VCFtools IV

#### vcf-merge

Merges two or more VCF files into one so that, for example, if two source files had one column each, on output will be printed a file with two columns. See also vcf-concat for concatenating VCFs split by chromosome.

\$ vcf-merge A.vcf.gz B.vcf.gz C.vcf.gz | bgzip -c > out.vcf.gz



### **VCFtools V**

### vcf-concat

Concatenates VCF files (for example split by chromosome). Note that the input and output VCFs will have the same number of columns, the script does not merge VCFs by position (see also vcf-merge).

\$ vcf-concat A.vcf.gz B.vcf.gz C.vcf.gz | gzip -c > out.vcf.gz



### VCFtools VI

### vcf-compare

Compares positions in two or more VCF files and outputs the numbers of positions contained in one but not the other files; two but not the other files, etc, which comes handy when generating Venn diagrams. The script also computes numbers such as nonreference discordance rates (including multiallelic sites), compares actual sequence (useful when comparing indels), etc.

\$ vcf-compare A.vcf.gz B.vcf.gz C.vcf.gz



### beftools

#### bcftools

BCFtools is a set of utilities that manipulate variant calls in the Variant Call Format (VCF) and its binary counterpart BCF. All commands work transparently with both VCFs and BCFs, both uncompressed and BGZF-compressed.

Most commands accept VCF, bgzipped VCF and BCF with filetype detected automatically even when streaming from a pipe. Indexed VCF and BCF will work in all situations.

Un-indexed VCF and BCF and streams will work in most, but not all situations. In general, whenever multiple VCFs are read simultaneously, they must be indexed and therefore also compressed.

BCFtools is designed to work on a stream. It regards an input file "-" as the standard input (stdin) and outputs to the standard output (stdout). Several commands can thus be combined with Unix pipes.



https://samtools.github.io/bcftools/bcftools.html

## Familiarizing with CLI tools

#### Lab Exercise 6 - Familiarizing with CLI tools

```
https://eclass.uoa.gr/modules/work/index.php?course=DI425&id=54428 or
```

https://eclass.uoa.gr/modules/document/file.php/DI425/2023-24/exercises/ITBI2023-2024-exercise5-ACD14112023.zip

**Due for 28/11/23** 



# Questions?



