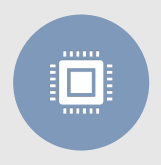
INTRODUCTION TO BIOINFORMATICS & TOOLS

Steve Kemp 24th October

What is Bioinformatics?



An interdisciplinary field that uses software to understand biological data.



Combination of biology, computer science, information engineering, mathematics and statistics.



Bioinformatics tools can be used individually to answer a specific question or chained together into a "pipeline" for rapid analyses.



For genetic analyses, bioinformatics analyses sequence data and allows us to annotate genomes, determine mutations and reconstruct predicted ancestral phylogenies.

Sequence Data

- A "sequence" is the order of nucleotides or amino acids that make up a strand of DNA.
- Many common sequencing methods produce short fragments of DNA called "reads".
- Raw reads must be processed and assembled to obtain a complete genome.
- Raw reads are comprised of forward and reverse reads.

TATCAGATCGCATACTGATCGTTGTACGCGATGCAACGCTGCATTGATG TACCCTCATCCTACCCCCCCATATCCCCCATCTCCCCTCATATCCCATCCCT <u>ACAATCGATCGATCGACGTGACTAGCTAGCTAGCATGACGCGCTAGCGCATGCGCATGCCGATI</u> ATATAC@C@C@ATCA@TCTACTACTCCCACTA@CTACAAAC@ATCACTC@C@CC@C

Basic **Bioinformatics Pipeline**



Download raw forward + reverse reads (.fastq)



Pre-alignment quality control (FastQC)



Mapping reads to a reference (bwa)



Sort & Index BAM files (samtools)



Eyeballing the alignment (IGV)



Post-alignment quality control (FastQC)



Downstream analysis

Bioinformatic Tools

- Analysis of whole genomes can be **computationally expensive**.
- Many tools are run on Linux cluster computers which have faster CPUs with more threads.
- Most common (free) bioinformatic tools are used by typing code into a terminal it is best to use
 Linux or a Mac for Bioinformatics (both are based on UNIX). If using a Windows PC/Laptop, consider installing Ubuntu 18.04.
- Installing various software on a Mac or Linux system can be managed using **Homebrew** or **Conda.**
 - If you already installed Ubuntu according to the instructions sent out last week, then feel free to click along with me.

Mac vs Linux

- This webinar is primarily geared towards **Linux users**.
- Installation of all software is tailored to **Linux machines** installation inputs (anything with sudo in front of it) **will not work** on a mac.
- Once you have installed the software, then the usage commands are identical on Linux or Macs.
- If you need help installing any software, please email me or have a google.
- If using a mac, almost all of the discussed software can be installed via homebrew or conda, or by downloading the .dmg files from the software websites on a mac ©

Terminals



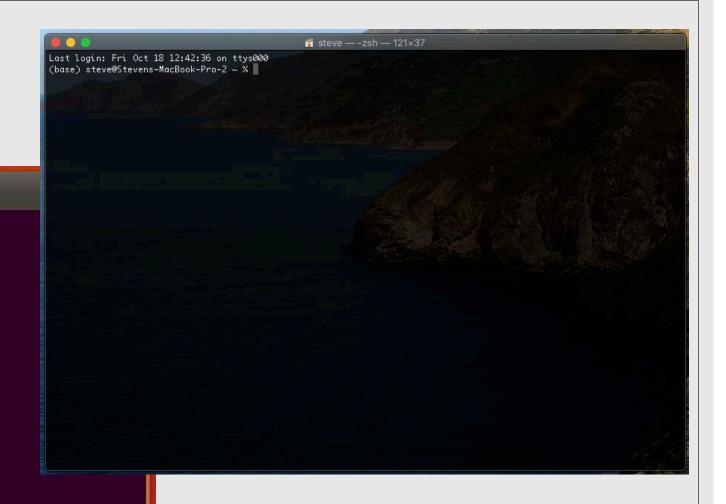


Terminals

parallels@parallels-Parallels-Virtual-Platform: ~

File Edit View Search Terminal Help

(base) parallels@parallels-Parallels-Virtual-Platform:~\$



Basic Linux/Unix commands

- To navigate the terminal you type code, rather than point and click.
- The directory that you are in when you first open the terminal is the HOME directory.
- To view the contents of the HOME directory, type **Is** and hit enter.
- Most of this tutorial will take place in the DOWNLOADS directory. To change directory, type cd
 Downloads/
- Is = list directory contents
- cd = change directory // cd ../ go up one directory
- o mkdir {name} = make directory
- mv = change filename or move from one folder to another
- ∘ rm = remove file // rm -r = remove directory

Installing Homebrew

- Homebrew is a large repository of tools which can be searched and will install software with a very simple command.
- To install
 - Open Terminal
 - Type sudo apt install linuxbrew-wrapper
 - Follow instructions (usually type Y a few times)
 - Add Linuxbrew to your PATH:

```
export PATH="/home/linuxbrew/.linuxbrew/bin:$PATH"" >>~/.bash_profile

export MANPATH="/home/linuxbrew/.linuxbrew/share/man:$MANPATH"" >>~/.bash_profile

echo 'export PATH="/home/linuxbrew/.linuxbrew/bin:$PATH"" >>~/.bash_profile
```

• When you install software, adding to your PATH means you can call the software anywhere in the terminal.



Installing "Conda"



- Miniconda2 and Minoconda3 are alternative managers similar to Homebrew.
- ∘ To Install:
 - Open terminal:
 - wget https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86_64.sh
 - Type bash Miniconda3-latest-Linux-x86_64.sh
 - Follow prompts on screen (usually accept user agreement by typing yes and then choose installation path)
 - **IMPORTANT:** After installing, **close and re-open** the terminal.

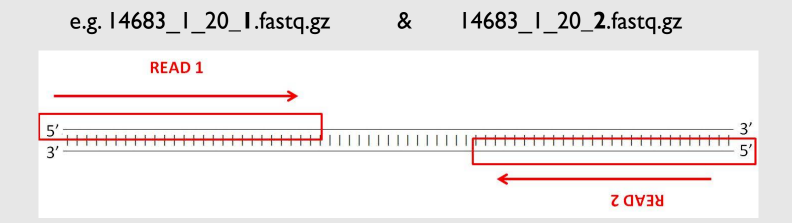
Installing a Java Environment

- Java is a popular programming language and several common bioinformatics tools are written in this language. You need to install a java environment.
- Very easy to do, type:
 - sudo apt install default-jre
- You can check to see if this installed correctly by typing:
 - java -version



1. Download Raw reads

- A **forward** and **reverse** read is created as part of the **amplification** process during sequencing.
- At the end of the sequencing, each DNA sample will have a **raw forward** and **raw reverse** read in a file.



Fastq files

• Text file containing a sequence and quality scores. Quality score is coded in ASCII characters.

@MS8_14683:1:1101:1856:12754#20/1
CCCTTTCTAAGGTATTATTCCAGACCCTTCTAGTAATGTTACAATGTGCTCGTCTTATGTCTCCTAT
TATGTTCTGTGCATAGAATACCTGTCCTGGTCCTATCCTTACACTTTTTCTTGTATTATTGTTGGGT
CTTGCACACGTGATTTCTACAGATTCATTAAGATGTACTATTATTGTCTTAGCATTGTTTGATATATT
TTCAGATCTAATTATTATATCTTTTTTCTGCTAGGCTACCATTTAACAGTAGTTGAGTTGACACCACT
GGCTTAATCCCATGTGTACATTGTACTATGC

@@@CCCEF9-

"C,CE9ECCE9<,C@@CC<E,,C,,C,CC<C<,C,C,6CC7@@ED9C,C@CC@F,CC,C<CECE,C,<,E9,,,<,CC C,CECF<,:CCF9CCEEFCE9CEEFEFDEEEE9EE9FF9EE8,+B=EE,,,:?A7A8,?ED?E9A<<,CAA9EF9,;,C<A;AF;A E,EE9EAFF99,@=@EEEFF9>ADADEGGC;D,@=F9=DDADD:FCDEGFFGGGGF8EA,66A??DDD?FCD==+C?9=F7;*BF77;=:5??FF*69AFCFCEA@A6E6A@CACECAEAEEE?*

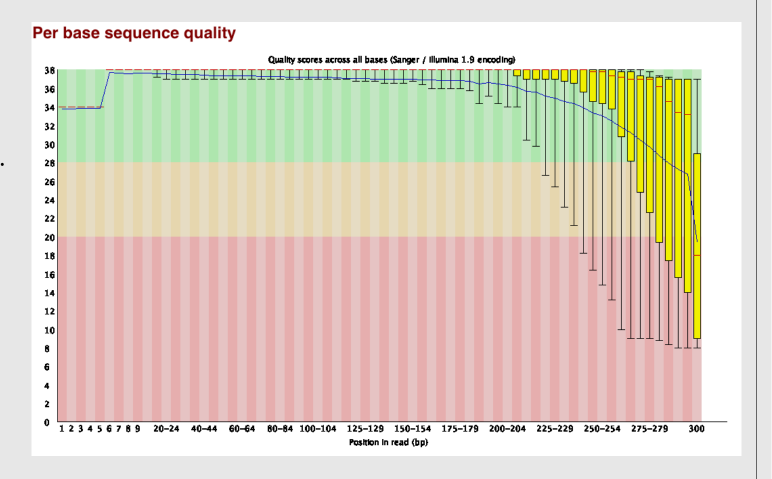
2. Quality Control of raw reads

- Modern sequencing technology can generate huge number of sequences, but no method is perfect.
- We need to assess the quality of reads and then determine if we need to trim or filter reads.
- FastQC is a good program to indicate if raw reads are 'good' or 'bad'.
- Install by typing brew install fastqc into terminal
- FastQC is used by typing fastqc {reads_I} {reads_2} etc.
- e.g. fastqc 14683_I_201.I.fastq.gz

FastQC outputs a html file and a .zip file. The .html file is the user-friendly report you can read.

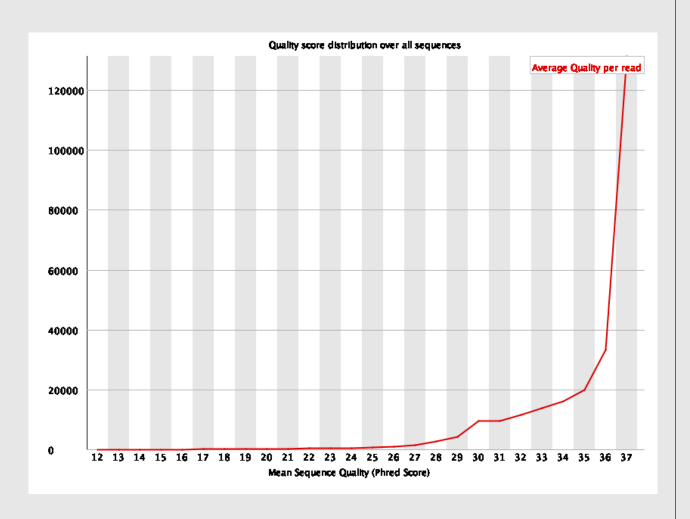
Assessing the Read Quality

- FastQC looks at quality scores collectively across all reads in a sample.
- X-axis shows the position of the sample.
- Y-axis shows the quality scores.
- Higher score = better base call.
- The quality of the 1st 5-7 bases is often lower due to signal decay during a run.
- Where **Phred scores** (quality scores) are <20, we consider trimming them.



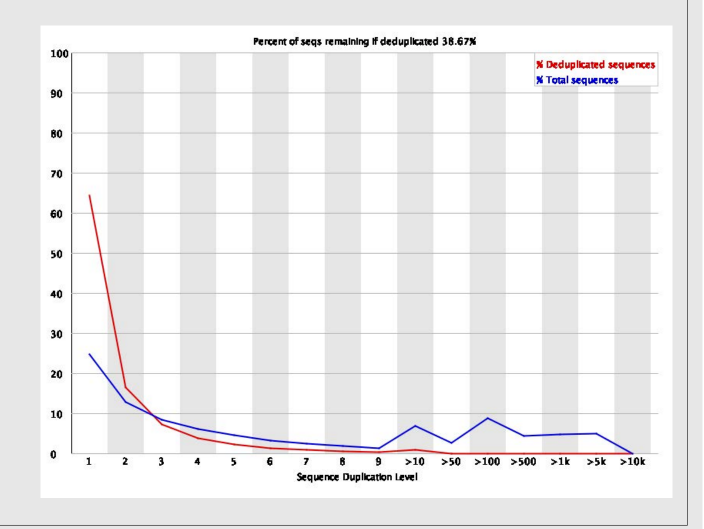
Per sequence quality scores

- A plot of the average quality score of all reads. This example shows a good quality score as the peak rise exponentially.
- Where he mean quality score is <27 this means there has been an error rate of 0.2% you will see a warning.
- If mean quality score is <20, this is a 1% error rate and is cindered by FastQC as a failure.



Sequence Duplication Levels

- In diverse sequences, many reads only occur once.
- A high level of duplication may be due to PCR enrichment – PCR duplicates misrepresent the actual proportion of sequences.
- Here, the blue line is what we actually have
- The red line is what the distribution would look like when deduplicated



Over-represented sequences

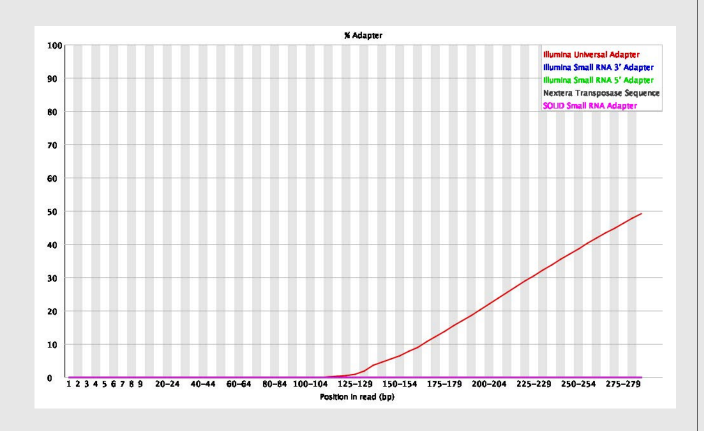
- No one sequence should make up a large proportion of the sequences.
- FastQC shows all sequences which make up >0.1%
 of the total.
- FastQC also looks at these sequences and matches it to a known contamination database.
- These sequences may be:
 - highly biologically significant
 - indicate a **contaminated library**
 - suggest that sample is not very diverse

Overrepresented sequences

Sequence	Count	Percentage	Possible Source
$\tt AGCCCGGGAGCTCTCTGGCTGCTTAGGGAACCCACTGCTTAAGCCTCAAT$	7271	2.779774284316124	No Hit
$\tt CCTCCAATTCCTCCTATCATTTTTCGCCTTTTAATACTGACGCTCTCGCA$	5827	2.227718987032053	No Hit
${\tt AGCCTGGGAGCTCTCTGGCTGCTTAGGGAACCCACTGCTTAAGCCTCAAT}$	3169	1.211539637876193	No Hit
$\tt CCTCCAATTCCCCCTATCATTTTTCGCCTTTTAATACTGACGCTCTCGCA$	1820	0.6958037680450208	No Hit
$\tt CCTATGGCAGGAAGAAGCTCCGGATGGAGGCCAGCTAAGCCACTCGGGGC$	1647	0.6296641791044776	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGAGGGGGGGGGGGGGG$	1326	0.5069427452899438	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGGGGTACAGTGCAGGAGAAAGAA$	1297	0.49585576217274285	No Hit
${\tt AGCCCGGGAGCTCTCTGTGGCCTCTTTTATTAAGGCGTGAATCGCATCCC}$	1169	0.44692011255199415	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGCAGCAGAAACCTGTCCCAGTGAGGCC}$	1146	0.4381269880107659	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGGGGTACAGTGCAGGAGAGAATAATA}$	1079	0.4125122339124052	No Hit
${\tt AGCCCGGGAGCTCTCTGGGACTTCACTGGTAATGTCACTAAGCCAAGTCA}$	993	0.3796335943234646	No Hit
${\tt CCTCCAATTCCTCCTATCATTTTTGGTTTCCATTTCCCTGGCAAATTTAT}$	894	0.3417849278199168	No Hit
${\tt CCTATGGCAGGAAGAAGCGGGCCTCCTGGTGAACGCAGGCCTGGCCTTA}$	852	0.3257279177881086	No Hit
${\tt TGGCTTGTACCGTCAGCGCTGCGGAGATCTACAATGAGGTCCTCAGGTGG}$	825	0.31540555419623195	No Hit
$\tt CTTTTATGCAGCTTCTGAGGGCTCTACCAACGAACTGAAAGGAGGATGAG$	806	0.30814166870565207	No Hit
${\tt CCTCCAATTCCTCCTATCATTTTATGTACACAATAGAGAGTTGCTACTGT}$	736	0.2813799853193051	No Hit
${\tt AGCCCGGGAGCTCTCTGACCTCTAAATTCTTATAAGTCAGAGAGTGACTG}$	735	0.280997675556643	No Hit
${\tt CCTCCAATTCCTCCTATCATTTTTGGTTTCCATTTCCCTGGCAAGTTTAT}$	650	0.2485013457303646	No Hit
$\tt CTTATATGCAGCTTCTGAGGGCTCTACCAACGAACTGAAAGGAGGATGAG$	648	0.2477367262050404	No Hit
${\tt TGGCCTGTACCGTCAGCGTATCCCCCAGGTTTTGGGTTATCTGCTATAGG}$	620	0.23703205285050158	No Hit
$\tt CCTCCAATTCCTCCTATCATTTTTCACCACACACTGCCAGTGTGGTCTT$	598	0.2286212380719354	No Hit
$\tt TGGCCTGTACCGTCAGCGTCAATGACGCCGCGCCCATAGTGCTCCCGGCT$	591	0.2259450697333007	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGGACTGAGAGGGTATGGACTCAGGTC}$	545	0.20835882065084413	No Hit
${\tt TGGCTTGTACCGTCAGCGTTAATGACGCCGCGCCCATAGTGCTCCCGGCT}$	543	0.20759420112551993	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGCTGGCCCTGGTAACAAAAGCCATCCGG}$	539	0.20606496207487154	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGGGGAAGACTGGAGGTGACCATCTTGGG}$	516	0.19727183753364325	No Hit
$\tt CTTTTATGCAGCATCTGAGGGGCTCTACCAACGAACTGAAAGGAGGATGAG$	514	0.19650721800831905	No Hit
${\tt AGCCTGGGAGCTCTCTGGGACTTCACTGGTAATGTCACTAAGCCAAGTCA}$	497	0.19000795204306337	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGGGTTGGGAGATGCAGAAGGAGAGTGAG}$	477	0.1823617567898214	No Hit
${\tt TTAAAAGAAAAGGGGGGATTGGGAGGAAAGTCAGTGCATTTCAGACAAGC}$	476	0.1819794470271593	No Hit

Adapters

- Adapters are oligonucleotides which can be attached to the ends of DNA or RNA recognised during sequencing.
- The plot shows the cumulative percentage of reads with adapter sequences at each position.
- Only adapters specific to the library type are searched.
- Ideally Illumina sequence data should not have any adapter sequence present.
- These should be removed as they can interfere with downstream analyses.



Filtering and Trimming Reads



- Any issues such as high duplication, low-quality sequences, inclusion of adapter sequences etc. can introduce bias in downstream analyses. Sequences must be treated to reduce that bias. Both steps can be completed using Trimmomatic.
- Cutting/**Trimming**/masking sequences
 - from low quality score regions
 - beginning/end of sequence
 - removing adapters
- Filtering of sequences
 - with low mean quality score
 - too short
 - with too many ambiguous (N) bases

Trimmomatic

- A program which can remove Illumina adapters and remove low-quality sequences all in one go.
- To install:
 - cd Downloads/
 - wget http://www.usadellab.org/cms/uploads/supplementary/Trimmomatic/Trimmomatic-0.39.zip
 - unzip Trimmomatic-0.39.zip
 - cd Trimomatic-0.39.zip
 - ∘ mv /adapters/*.
- To run:
 - java -jar trimmomatic-0.39.jar PE -threads {number_threads} -phred33 {forward_read.fq.gz}
 {reverse_read.fq.gz} -baseout {examplename} ILLUMINACLIP:TruSeq3-PE.fa:2:30:10 LEADING:3 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:36
- This outputs 4 files: two unpaired reads and two paired reads the ones you want are called {example} IP and {example} 2P.

3. Mapping reads to a reference genome

- Once we have good quality reads, we can assemble these into a genome:
 - de novo
 - map to reference
- Map to reference using **BWA** (Burrows-Wheeler Aligner)
- Open terminal
- Type sudo apt install bwa
- o Download **reference genome** fasta you wish to assemble to e.g. HXB2 or subtype-specific reference
- Type bwa index {reference genome}
- Type bwa mem {reference genome} {forward read} {reverse read} -o eg.sam

```
SN:K03455|HIVHXB2CG
                             LN:9719
       ID:bwa PN:bwa VN:0.7.17-r1188 CL:bwa mem K03455.fasta 14683 1 20 1.fastq.qz 14683 1 20 2.fastq.qz -o 14483.sam
MS8 14683:1:1101:1856:12754#20 77
{\sf GTTACAATGTGCTCGTCTTATGTCTCCTATTATGTTCTGTGCATAGAATACCTGTCCTGGTCCTATCCTTACACTTTTTCTTGTATTATTGTTGGGTCTTGCACACGTGATTTCTACAGATTCATTAAGATGT
ACTATTATTGTCTTAGCATTGTTTGATATATTTTCAGATCTAATTATTATATCTTTTTCTGCTAGGCTACCATTTAACAGTAGTTGACACCACTGGCTTAATCCCATGTGTACATTGTACTATGC
+B=EE,,,:?A7A8,?ED?E9A<<,CAA9EF9,;,C<A;AF;AE,EE9EAFF99,@=@EEEFF9>ADADEGGC;D,@=F9=DDADD:FCDEGFFGGGGF8EA,66A??DDD?FCD==+C?9=F7;*BF77;=:
5??FF*69AFCFCEA@A6E6A@CACECAEAEEE?*
MS8 14683:1:1101:1856:12754#20 141
                                                                                          ATTCAATGGGACAGGACCATGCAGTAACGTCAGCATA
{f GTACAATGTACACATGGGATTAAGCCAGTGGTG}TCAACTCAACTACTGTTAAATGGTAGCCTAGCAGAAAAAGATATAATAATTAGATCTGAAAAATATATCAAACAATGCTAAGACAATAGTACATAGTACATCTTA
ATGAATCTGTAGAAATCACGTGTGCAAGACCCAACAATAATACAAGAAAAAGTGTAAGGATAGGACCAGGACAGGTATTCTATGCACAGAACATAATAGGAGACATAAGACGAGCACATTGTAACATTAC
-8A@--<--,;,,,,:9,;,6,,=,,,,;,,,,C,,<,,<,,,,C,,,6,C,,,,;C,,,6;,,;,,,;C,,;@,,;CC,,:C,99,<6,,,C,,<,,55C,,95,,,,,,,;,A,,C,,9A,,,@EFA9,,,
A,C,AE,,,@,,=>@D9,4,4,4@9,CD9E;C9DBAEFCE8=8=ADDDFC88,@;ED+=1=8==,++===++=++@+6=+?+6+***0*59<C779*+;****895))08)0)37;77;*@76*6*0)31:<4
*:**.*0)-):6964>22(59B3;C4>4C?9EEE) AS:i:0 XS:i:0
MS8 14683:1:1101:1980:10573#20 77
CCTGATGACCTCATTTGACCCAAATTACCTCCAAAGAAGGTCATACCCTGAAGTCCTGGGGGGCGAGGACTTCAACACCCCTGCTCTGGGGAATGGGGTTGAGCCACTGCCTGAGGCACAGGACGGGGCTCCC
TGGATCACGGGGGTCAAAGCTGAAGCCATCACATCTGTCCCAGGGAGACCCTGAGCTGGCTTTGGGTAATGGGACACGGGGCCGTACGGCCCATGTGCGGGGACTGAGCCAGGGTTCGGCAGGGGTGTTG
@CCCCCEGGGFEE@FGGGGGGGCFDEGEFFGGGDECF@FGCFFEFCGGFFGGFGGFC@FEFGGDEFFGFCCEFGGGGFEGGCCEE<EEGGGDGGGFG>FGEEGEGGGGEFBEEFEEGGGCFEGGGGE7BFF
GGGGGGCFCCCFGGGFGGCFFE@FCGGFGGDGGGGGECEGGGG:FBFGCFEGEFFCCFEGGFGGCFGGGGGCCGGGFF9BFFGFGE==FFF6FGGGGDGCFGGDGGGGGGGGDDFFGGCC>DGD5DFD375<FF
F=?EDEFFFF:4@(54>F:<((302-70(((-3:(
                                    AS:i:0 XS:i:0
```

- The output file we have requested is a **SAM file** (Sequence Alignment Map)
- Tab-delimited file of alignments of short reads mapped against a reference sequence.
- Read Name → SAM flag → contig name → mapped position of base I of a read on the reference sequence → mapping quality → CIGAR string → name of mate → position of mate → template length → read sequence → read quality → Other information in TAG:TYPE:VALUE format.

Converting SAM to BAM file

- SAM files are extremely large (but readable by humans). We store alignments as BAM files (Binary Alignment Map) which is compressed version of a SAM.
- Install samtools:
- sudo apt install samtools
- To change a SAM to a BAM type: samtools view -h -b -S {example.sam} > {example.bam}
- When aligning to a reference genome, many reads may not be mapped to the reference.
- We can remove these to further reduce to BAM size.
- Type samtools view -b -F 4 {example.bam} > {mapped.bam}

4. Sorting and Indexing a BAM File

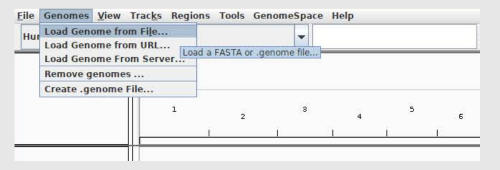
- BAM files can be sorted by read name or by coordinates (chromosomal position of an alignment).
- Some downstream analyses require that BAM files be sorted and indexed.
- We can do both of these things using samtools.
- Samtools sort sorts alignments by leftmost coordinates
- Samtools index indexes a co-ordinate sorted BAM for fast random access
- To sort, type: samtools sort {input.bam} -o {sorted_output.bam}
- To index, type: samtools index {sorted_output.bam}

5. Viewing your sorted BAM file

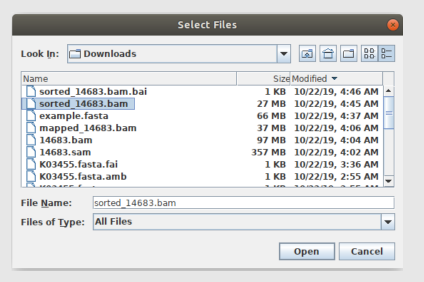
- After creating the BAM and then sorting and indexing it, we can use a viewer to identify any areas of poor coverage.
- A well-known BAM viewer is the Integrative Genomics Browser (IGV).
- To install:
 - Download the Linux archive: go to Downloads/ in terminal and type
 - wget https://data.broadinstitute.org/igv/projects/downloads/2.7/IGV_Linux_2.7.2.zip
 - unzip IGV_Linux_2.7.2.zip
 - cd IGV 2.7.2
 - ∘ sh igv.sh
- This will open the IGV viewer.

How to use IGV to view alignment

You must first load the reference genome that the BAM file
 was made from. Click Genomes → Load genome from File...
 and select the fasta file that we used earlier.



You can then load the sorted BAM file, and index file. Click
 File → Load from File... and select the sorted BAM file.





6. Post-alignment Quality Control

- Once you have eyeballed the alignment and are happy with it, you can perform some additional QC to see if there are any issues.
- We can do this by using FastQC again.
- Type:
 - fastqc {input.bam}
- After this has completed, you can open the .html file again to see the report.

7. Downstream Analysis



GENERATE PHYLOGENETIC TREES



ANALYSE SEQUENCES FOR DRUG-RESISTANCE MUTATIONS

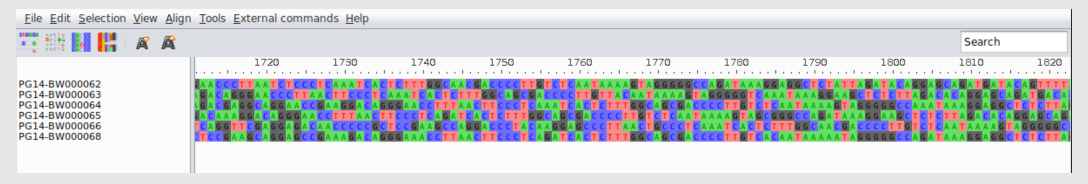


ANCESTRAL RECONSTRUCTION

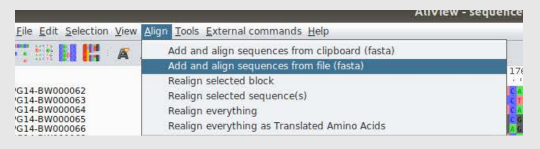
Phylogenetic Trees

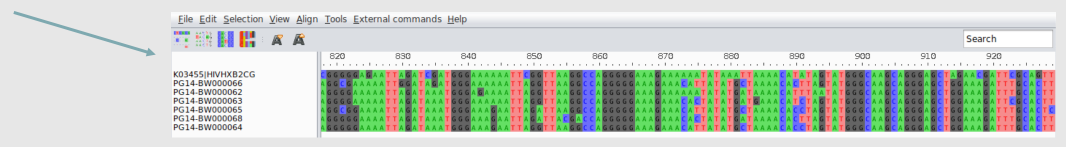
- If using Phyloscanner input is the BAM files you have made.
- If creating a maximum-likelihood or neighbourhood-joining tree, these use the **consensus sequence** of the BAM files. PANGEA create a consensus sequence using a tool called **SHIVER**.
- Researchers who are **accredited** by PANGEA can request these sequences which saves you from having to make them yourselves.
- You can then use these to create a multiple sequence alignment (MSA):
 - Place all files into a folder
 - concatenate all sequences into a single file; type: cat * > sequences.fasta
- You can then install a sequence viewer I recommend Aliview. To install:
 - wget https://ormbunkar.se/aliview/downloads/linux/linux-version-1.26/aliview.install.run
 - chmod +x aliview.install.run
 - sudo ./aliview.install.run
 - aliview

- After installing and typing 'aliview' into the terminal, AliView will open!
- Open your concatenated consensus sequences by clicking File → Open File and then selecting your sequences.
- It will look like this:



- We need to add a reference to align everything to now
- Click on Align → Add and align sequences from file (fasta) and choose your reference sequence.
- The programme will now insert the reference genome and perform a multiple sequence alignment.





- Alternatively, you can use online tools such as HIValign or Gene Cutter from LANL:
 - https://www.hiv.lanl.gov/content/sequence/HIV/HIVTools.html
- HIVAlign takes your unaligned fasta file and aligns to HXB2 using MAFFT.
- Gene Cutter aligns nucleotide sequences and codon aligns all coding regions. This can output a nucleotide alignment and protein alignments.
- Whichever tool you use, you'll get an aligned fasta file at the end. Use this to build trees.

Alignment and sequence manipulation

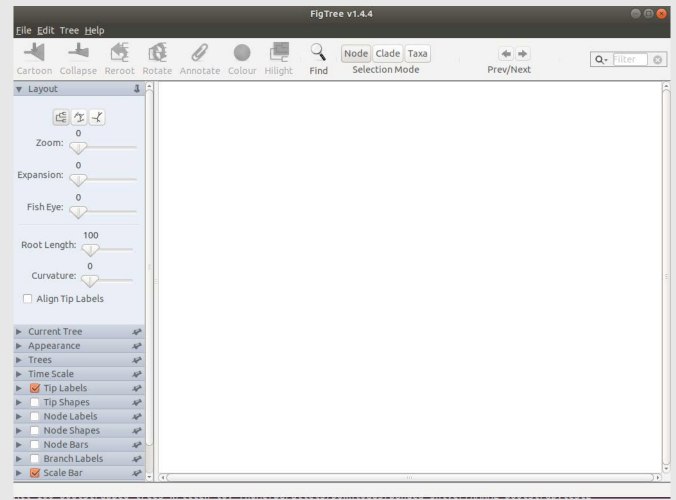
- Align Multi-tool manipulates sequence alignments, including sorting, pruning, and renaming
- Alignment Slicer cuts vertical slices from sequence alignments
- Analyze Align shows weblogos, calculates frequency by position, and finds variants in an alignment
- Codon Alignment takes a nucleotide alignment and returns a codon alignment and translation
- Consensus Maker computes a customizable consensus
- <u>ElimDupes</u> compares the sequences within an alignment and eliminates any duplicates
- Gap Strip/Squeeze removes columns with more than a given % of gaps
- Gene Cutter clips genes from a nucleotide alignment, codonaligns, and translates
- HIValign uses our HMM alignment models to align your sequences
- PepMap can be used to map epitopes, functional domains, or any protein region of interest
- <u>Pixel</u> generates a PNG image of an alignment using 1 or more colored pixel(s) for each residue
- QuickAlign (formerly Epilign and Primalign) aligns short nucleotide or protein sequences (e.g., primers, epitopes) to our prebuilt genome or protein alignments, or to a user alignment
- Sequence Locator finds the standard numbering of your HIV or SIV nucleotide or protein sequence
- SynchAlign aligns overlapping alignments to one another
- Translate nucleotide sequences to 1-letter amino acids

Tree-Building

- Three major types of tree: **Neighbourhood-Joining** (NJ), **Maximum-Likelihood** (ML) & **Bayesian** (BEAST).
- Argument for both, NJ is faster but considered to be less accurate. ML is slower and uses more complex evolutionary models. BEAST uses markov-chain monte-carlo (MCMC) to generate rooted, time-measured phylogenies.
- Recommend using software called **raxml** for building ML trees with bootstraps.
- Install by
 - sudo apt install raxml
- Build trees and perform rapid bootstrapping on one go:
 - raxmlHPC –f a -m GTRGAMMA -p \${RANDOM} -x \${RANDOM} -# {number_bootstraps} -s {alignment.fasta} -n {name}
- This will do x bootstrap searches, 20 ML searches and return the best ML tree with bootstrap values.
 - 100 bootstraps = fast tree
 - 500-1000 bootstraps publication quality.

Viewing Trees

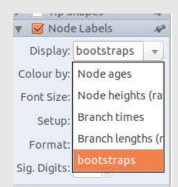
- Use a tree-viewer such as Figtree. Can also do some manipulation of trees in this programme.
- To install:
 - sudo apt install figtree
 - figtree
- Open the correct file to view your tree. Click File → Open, then select the file called RAxML.bipartitions.example_name
- An input window will open type bootstraps and click OK.



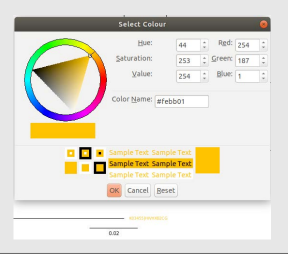
Manipulating Trees

Eroot

- Can do various things in Figtree:
- Re-root trees
 - Click on nodes you wish to reroot at (outgroup). Click Reroot
- Show bootstrap values
 - Tick Node Labels and change Display: to bootsraps
- Sort nodes by increasing or decreasing evolution.
 - Select Trees, then Order Nodes (by increasing)
- Colour branches or Taxa or Nodes
 - Choose Node/Clade/Taxa as appropriate
 - Select the Node/Clade/Taxa you want to colour
 - Select Colour and Choose a colour!

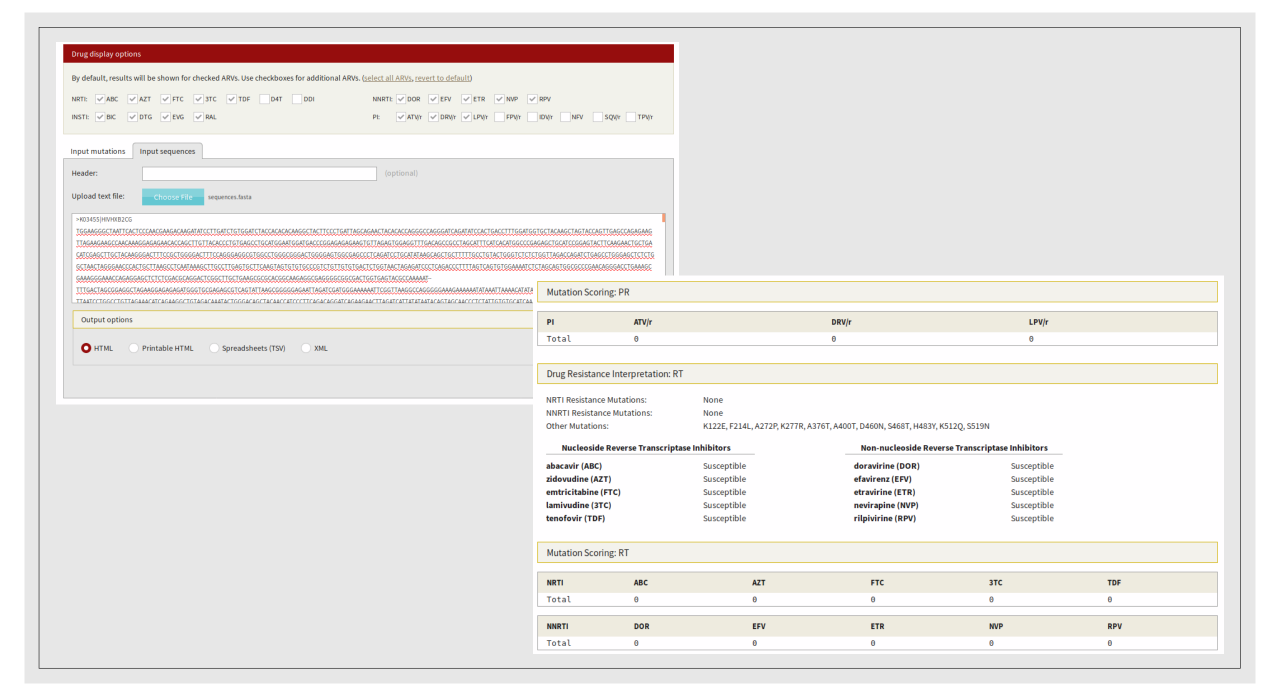






Looking for Drug Resistance Mutations

- Easiest way is to use the **Stanford HIV database** https://hivdb.stanford.edu/hivdb/by-sequences/
- Input the sequences you wish to analyse, and it gives a report which you can view on-screen or a spreadsheet that you can download, for many sequences. Limited to 1000 sequences of up to 3000 nucleotides.
- This tells you the resistance mutations and which drugs would be susceptible or resistant.



Tools for Help

- Many of these processes can be automated using an online system called Galaxy: https://galaxyproject.org/tutorials/g101/
- This is a user-interface where you upload all of your sequences and choose the tools you want to use, and this will run on their servers.
 - Can be a little slow and steep initial learning curve
 - Means you don't have to leave computer running or worry about power interruption etc.
- A paid tool exists which is useful for many analyses including sequence building, multiple alignments, SNP calling, tree building etc: https://www.geneious.com/
 - User-friendly, no coding needed
 - Expensive \$450 per user, per year.
 - I4-day unrestricted free trial

Getting Accredited by PANGEA



To become a **PANGEA-accredited researcher** and get access to sequences, you must fill in a **PANGEA** accreditation form and attached a certificate of a course on **Human Subjects Research**.



If you are a member of PANGEA and are employed by MRC Uganda, Rakai, Partners/UW, Botswana/Harvard, and PopART, there is funding for you to take these courses.



Please email Lucie if unsure, some institutions have free institutional access



Once you are accredited, you can formally request data from PANGEA and do some analyses.

Any other information?



- Are there any tools or specific analyse which you want to perform?
- Send me a list of tools you may wish to use, or questions with specific analyses and PANGEA will try and help you with this ☺
- Any tools that you may wish to use can be explored in next month's webinar (14th November).