

See these examples from 2019 for help as well. [https://github.com/biodataprogram/GEN220\\_2019\\_examples/tree/master/Bioinformatics\\_8](https://github.com/biodataprogram/GEN220_2019_examples/tree/master/Bioinformatics_8)

## DNA Sequence Read alignment

Align genomic DNA reads from three different experiments to the genome.

We will align SARS-CoV-19 genomes against the reference genome.

1. The genome is available at [NCBI RefSeq Accession NC\\_045512.2](#) - this is already downloaded for you in `/bigdata/gen220/shared/data/SARS-CoV-2`
2. I selected 3 random SARS-CoV [SRA](#) accessions `SRR11587604`, `SRR11140748`.
3. Align reads to the genome with `bwa` (or other tools you choose if you prefer). You'll need to index the genome - remember you want to make a symlink to this file.
4. Create the BAM files for this alignment using `samtools`.
5. Use `samtools` and the subcommand `flagstat` (or other tools if you want) to get a count for the number of reads which map to the genome.

I highlight recommend reading the [htslib tools](#) docs if you want to see more on `samtools` and later on SNP calling with `bcftools`.

Additional things you can explore:

See other options for `samtools` - such as try the option to [retrieve reads which are unmapped](#) `samtools view -f 4`

Try using the `samtools fastq` option to dump out reads which are unmapped. For example.

## SNP calling

Call SNPs from this same dataset to explore how variants are called to create a VCF file. Follow example from class.

Generate a table of SNP locations using `bcftools view` to reformat.

## RNAseq and comparisons

Reanalyze data in this published paper [Baker et al 2014](#) "Slow growth of *Mycobacterium tuberculosis* at acidic pH is regulated by *phoPR* and host associated carbon sources"

Data are downloaded to `/bigdata/gen220/shared/data/M_tuberculosis`

The Transcriptome file is also in the folder as `M_tuberculosis.cds.fasta`  
- I have already renamed the sequences to be the LOCUS names. It was downloaded from [https://www.ncbi.nlm.nih.gov/assembly/GCF\\_000008585.1/](https://www.ncbi.nlm.nih.gov/assembly/GCF_000008585.1/) and the specific file is [linked here](#)

There is a `sra_info.tab` file which lists the sample accessions and their metadata so you can see what are the data sets. This is from the BioProject [PRJNA226557](#) and the SRA Project [SRP032513](#)

Compare gene expression between two sets of conditions. - pH5.7 - pH7

And growth carbon source - Glycerol - Pyruvate

1. Run Kallisto to get the gene expression calculated from each sample - you will need the file `M_tuberculosis.cds.fasta` as the database and each of the 8 `.fastq.gz` files in the folder. You can make links to these files (`ln -s /bigdata/gen220/shared/data/M_tuberculosis/*.fastq.gz`). You do not need to uncompress the files, Kallisto can read gzip compressed files.

### extra credit / extra

2. Run pfam analysis to get the Protein domains found in each protein - you will need the file `M_tuberculosis.pep.fasta`
3. Construct a tab delimited file which lists on each line
  - The Gene (LOCUS) name
  - The Protein length
  - An average TPM across replicates for each condition (eg there will be 4 conditions, two replicates per condition)
  - The Pfam Protein domains, separated by comma found in each Protein

FYI - to process the file and move the locus\_tags as the sequence names I ran this regular expression (in Perl)

```
perl -p -e 's/>(\S+).+(\[locus_tag=(\[^\]]+\)\)/>$3 $1 $2/' GCF_000008585.1_ASM858v1_cds_from
```

I made the protein file of sequences using script from BioPerl. `bp_translate_seq.pl`  
`M_tuberculosis.cds.fasta > M_tuberculosis.pep.fasta`