1. Running FastQC on raw data

- Reminder to take notes, share info here: <u>Etherpad</u>
- FastQC is a program that can quickly scan your raw data to help figure out if there are adapters or low quality reads present. Create a job file to run FastQC on one of the fastq files here:

```
/data/genomics/dikowr/SMSC/fastq for fastqc
```

- **module**: bioinformatics/fastqc/0.11.5
- o command: fastqc <FILE.fastq>
- after your job finishes, find the results and download some of the images, e.g.

 per base quality.png to your local machine using scp, CyberDuck, or Filezilla.

2. Trimming adapters with TrimGalore!

- PacBio data will be error-corrected solely by the assembler, but Illumina data trimming and thinning are common.
- Most assemblers these days don't want you to trim/thin for quality before assembling, but trimming is
 important for downstream applications. TrimGalore will auto-detect what adapters are present and remove
 very low quality reads (quality score <20) by default.
- Create a job file to trim adapters and very low quality reads for the Illumina data here:

```
/data/genomics/dikowr/SMSC/fastq for fastqc
```

command:

```
trim_galore --paired --retain_unpaired <FILE_1.fastq> <FILE_2.fastq>
```

- **module**: bioinformatics/trimgalore/0.4.0
- You can then run FastQC again to see if anything has changed.

3. Run Genomescope

- Genomescope can be used to estimate genome size from short read data:
 - Genomescope
- To run Genomescope, first you need to generate a Jellyfish histogram.
- You'll need two job files for Jellyfish, one to count the kmers and the second to generate a histogram to give to Genomescope:
- Here is a copy of the Red Siskin Illumina data: /data/genomics/dikowr/SMSC/Illumina all

- Hint: don't copy these data to your own space.
- First job file: kmer count:

```
    Module: bioinformatics/jellyfish/2.2.3
```

Commands:

```
jellyfish count -C -m 21 -t $NSLOTS -s 800000000 *.fastq -o reads.jf
```

- ∘ -m = kmer length
- ∘ -s = RAM
- Hint: this job needs to run on the high memory queue.
- This will take a while, so we can move on and then come back to it when it finishes.
- Second job file: histogram:

```
• Module: bioinformatics/jellyfish/2.2.3
```

- Commands: jellyfish histo -t \$NSLOTS reads.jf > reads.histo
- Download the histogram to your computer, and put it in the Genomescope webservice: Genomescope

4. Run the fasta metadata parser to get statistics about the PacBio data

We use a python script to grab some statistics from assembly files, but we can also use it to look at our PacBio data. These are the stats it produces:

Total number of base pairs: Total number of contigs:

N90:

N80:

N70:

N60:

N50:

L90:

L80:

L70:

L60:

L50:

GC content:

Median contig size:

Mean contig size:

Longest contig is: Shortest contig is:

- Create job files to run this script for each assembly file. There are 7 assembly files.
 - + Module: bioinformatics/bioinformatics/fastametadata/1.0
 - Commands: fasta meta data parser.py <PACBIO.fasta> > pacbio stats.out
 - The PacBio data are here: /data/genomics/dikowr/SMSC/PacBio/all pacbio.fasta
- How long is the longest read?
- What is the read N50?

5. Setting up MaSuRCA Illumina + PacBio Hybrid Assembly

- We are going to split up into three groups of 5 people each to submit whole genome assembly jobs. These
 will take a while and create a lot of large files.
- MaSuRCA runs with 2 job files. The first uses a configuration file to generate an sh script called assemble.sh. Then you just execute the sh script to complete the actual assembly.
- Here is a sample MaSuRCA config file that you will need to copy to your space and modify:

/data/genomics/dikowr/SMSC/masurca_config.txt

- * Edit the file so that it points to your files and familiarize yourself with the parts.
 - Reminder, the raw data are here: /data/genomics/dikowr/SMSC/Illumina_all and /data/genomics/dikowr/SMSC/PacBio
 - Do not copy the data to your own space (look how big the files are!)
- To keep things tidy, create a directory for the MaSuRCA assembly in your space.
 - Hint: use mkdir
- Create a job file for this first part of MaSuRCA.
 - Queue: Short, high CPU
 - Threads & RAM: single thread, 2GB RAM
 - **Module:** module load bioinformatics/masurca/3.2.8
 - o Commands: masurca <CONFIG FILE>
- This job should complete in a few seconds and result in a file called assemble.sh
- Create a second job file for the second part of MaSuRCA.
 - Queue: Long, himem
 - Threads & RAM: 16 threads, 30GB RAM each
 - Module: module load gcc/4.9.2

- Command: ./assemble.sh
- Submit this second job.

6. Run the fasta metadata parser to get statistics about the assembly

- I have put a finished assembly here: /data/genomics/dikowr/SMSC/finished_assembly
- Create job files to run this script for each assembly file. There are 7 assembly files.
 - + Module: bioinformatics/bioinformatics/fastametadata/1.0
 - Commands: fasta_meta_data_parser.py <ASSEMBLY> > assembly_stats.out
- How long is the longest read?
- What is the read N50?