### 1. Running Bowtie2

- Reminder to take notes, share info here: Etherpad
- · For this part of the tutorial, we're each going to work with a single long scaffold from the Red Siskin assembly.
  - They are here: /data/genomics/dikowr/SMSC/assembly scaffolds
  - Copy one of these to a new directory (PopGen) in your /pool/genomics/ space.
- First we'll need to index the fasta file before we can map raw data to it. Create a job file for this:
  - o module: bioinformatics/bowtie2
  - command: bowtie2-build --threads \$NSLOTS <SCAFFOLD.fasta> NAME
  - · Hint: it doesn't need much RAM or time.
- Then we can start mapping the reads to our scaffolds.
  - There are 9 individuals plus the one from which we are assembling the genome.
  - $\circ~$  The read data are here:  $\begin{tabular}{ll} /data/genomics/dikowr/SMSC/resequence\_data \end{tabular}$
  - Don't copy it to your space.
  - Create job files to map each individual's reads to your scaffold.
  - o module: bioinformatics/bowtie2
  - Here is an example **command** (this will need editing to make it work for your data):

bowtie2 --very-sensitive -N 1 -I 100 -X 600 -x siskin -p \$NSLOTS --phred33 --rg-id "MB-12866" --rg SM: "MB-12866" --rg PL: "ILLUMINA" -

· Check out the output file using head

### 2. Manipulating the Bowtie2 output

- In order to get the Bowtie2 outputs ready to go for variant calling, we'll have to do some file manipulation.
- · First, we'll have to convert our sam file output to bam format.
- · Create a job file to do this.
  - module: bioinformatics/samtools/1.6
  - o command: samtools view -b <YOUR\_OUTPUT.sam> > <YOUR\_OUTPUT.bam>
  - o Try head on the output bam file to see what happens.
- · Next we'll need to sort the bam file.
  - **module**: bioinformatics/samtools/1.6
  - o command: samtools sort <YOUR\_BAM.bam> -o <YOUR\_BAM\_sorted.bam>

## 3. Mark Duplicates with picard-tools

- For future steps, we will need an sequence dictionary and fasta index.
- Create a job file to create a fasta index on your scaffold:
  - module: module load bioinformatics/samtools/1.6
  - o command: samtools faidx <YOUR\_SCAFFOLD.fasta>
- Create a job file to create a sequence dictionary:
  - module: module load bioinformatics/picard-tools/2.5.0
  - command: runpicard CreateSequenceDictionary R=<YOUR\_SCAFFOLD.fasta> O=<YOUR\_SCAFFOLD.dict>
- Now we can create a job file to mark duplicates:
  - module: module load bioinformatics/picard-tools/2.5.0
  - command: runpicard MarkDuplicates I=<YOUR\_SORTED\_BAM.bam> O=<YOUR\_SORTED\_BAM\_marked.bam> M=marked\_dup\_metrics\_SAMPLE\_NAME.txt

#### 4. Realign indels before calling variants

- Now we'll look for regions with indels in our bam files and realign them.
- Create a job file to use GATK RealignerTargetCreator:
  - **module**: module load bioinformatics/gatk/3.7
  - command:
    - rungatk -T RealignerTargetCreator -R <YOUR\_SCAFFOLD.fasta> -I <YOUR\_SORTED\_BAM\_marked.bam> -o <YOUR\_SORTED\_BAM\_marked.bam>.list
- Create a job file to use GATK IndelRealigner:
  - module: module load bioinformatics/gatk/3.7
  - o command

rungatk -T IndelRealigner -R <YOUR\_SCAFFOLD.fasta> -I <YOUR\_SORTED\_BAM\_marked.bam> -o indels-<YOUR\_SORTED\_BAM\_marked.bam> -targetInte

# 5. Call variants with GATK

- First, we will run GATK HaplotypeCaller on each bam file individually:
  - module: module load bioinformatics/gatk/3.7
  - o command:

rungatk -T HaplotypeCaller --emitRefConfidence GVCF --variant\_index\_type LINEAR --variant\_index\_parameter 128000 -R <YOUR\_SCAFFOLD.fa

- This job might need to be run on himem.
- Now we'll run GATK GenotypeGVCFs across all vcf files:
  - module: module load bioinformatics/gatk/3.7
  - o command

rungatk -T GenotypeGVCFs -variant\_index\_type LINEAR -variant\_index\_parameter 128000 -R <YOUR\_SCAFFOLD.fasta> -V HERE\_LIST\_ALL\_YOUR\_VC

- $\circ~$  This job might need to be run on himem.
- Use head to look at the VCF file.