

High Performance Computing for Genomics

Part II: for Genomics

Project description

Assume this population study:

We have 2 populations, population 1 has 14 individuals, population 2 only 5.
There is no reference genome available.

Are there 'diagnostic' variations between the 2 populations.

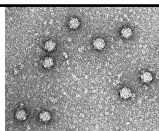
Project parts:

- Assembly
- Mapping
- Variant calling

Actual data

Population 1: Sequenced PhiX in the Genomics Core

Population 2: Simulated data from the ncbi reference genome



Phi X 174:

- Bacteriophage
- First sequenced DNA virus (1977)
- Circular, single stranded DNA genome of 5386bp, GCcontent 44%
- 95% are coding genes, total of 11 genes
- Used as positive control in Illumina sequencing



Project description

Project parts:

- **Assembly**
Population 1 has most individuals, so best to choose one of these individuals.
- Mapping
- Variant calling

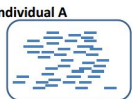
Assembly

Simple description: put **overlapping reads** together in order to get a **contig**

Used software: ABySS

Sequencing Reads

Individual A



De novo assembly



Contigs

Contigs



Which partition would you use for the assembly?

Exercise 4

- Load Cerebro
- Load ABySS (in the 2015a toolchain)
- Check the abyss.pbs file, change the header/variables if needed
- Launch the abyss.pbs script
- Check the output of the assembly

Project description

- Project parts:
- Assembly
 - **Mapping**
19 individuals need to be mapped to the newly created reference genome
 - Variant calling

Mapping

Compare your sequences against a reference in order to get the location

Used software: Bowtie2

Sequencing Reads



Which partition would you use for the mapping?

Go to ThinKing

- List all loaded modules
- Purge
- Load thinking

Job efficiency

Run every task **separately**, using all available cores in a node

- Multithreading
Using 20 cores (ivybridge)
- Needs 1 pbs per task
- Is applicable for all tasks,
but each pbs file needs adjustments

You can do this on your own now :)

Run all tasks in **parallel**

- Single threads
19 processes + master process (worker)
- Needs 1 pbs in total
- Tasks must be exact the same,
except for some parameters

This requires some more theory :(

Parallel Jobs

Assume following pbs script

```
#!/bin/bash -l
#PBS -l nodes=1:ppn=1
#PBS -l walltime=00:15:00
cd $VSC_SCRATCH
map -s sample1 -r homo_sapiens -l 100
```

Now you want to use 'map' for 100 different samples => 100 tasks

Parallel Jobs: parameters

The script will be changed to:

```
#!/bin/bash -l
#PBS -l nodes=1:ppn=8
#PBS -l walltime=04:00:00
cd $VSC_SCRATCH
map -s $sample -r $reference -l $length
```

The 'map' parameters are now variables, these will be listed in a separate file:

```
sample,reference,length
sample1,homo_sapiens,100
sample2,homo_sapiens,50
```

Parallel Jobs: Parameter file

The parameter file is a simple comma separated value file (CSV)
Which can be created in any text editor (NOT word)
or in a spreadsheet program (as excel)

```
sample,reference,length
sample1,homo_sapiens,100
sample2,homo_sapiens,50
sample2,mus_musculus,50
sample3,homo_sapiens,100
sample3,homo_sapiens,50
```

Parallel Jobs: PBS header

Original file

```
#!/bin/bash -l
#PBS -l nodes=1:ppn=1
#PBS -l walltime=00:15:00
```

- Uses 1 core
- Walltime is 15 minutes
100 tasks, 15 minutes each
=> 1500 minutes or 25 hours

Parallel file

```
#!/bin/bash -l
#PBS -l nodes=1:ppn=8
#PBS -l walltime=04:00:00
```

- 8 cores in use: 7 for the job, 1 for delegating the work
- Walltime is 4 hours
15 minutes per job, 7 jobs at the time
=> 215 minutes (1500/7) or 3.56 hours
So 4 hours is a safe time

Parallel Jobs: start a job

Load the worker module:

```
$ source switch_to_2015a
$ module load worker
$ wsub -batch run.pbs -data data.csv
```

Use wsub instead of qsub

-batch	The pbs script (.pbs)
-data	The parameter file (.csv)

Parallel Jobs: Map Reduce

Parallel computations can be abstracted into 3 steps:

1. Preparation
2. The work items
3. Aggregating results, clean-up, ...

The worker framework also supports this:

-prolog	Preparation step	e.g.: copying the reference genome to scratch
-batch	The parallel jobs	e.g.: mapping the reads to the genome
-epilog	Clean-up step	e.g.: cleaning the scratch storage

```
$ wsub -prolog split-data.sh -batch run.pbs -epilog distr.sh -data data.csv
```

Exercise 5a

- Open bowtie_batch.pbs
- Change script
- Variable name?
- input/output path?
- Reference genome?

Exercise 5b

- Open prolog script
- What happens?
- Check genome path
- Where is the genome stored?

Exercise 5c

- Open epilog script
- What happens?

Exercise 5d

- Start job on thinking
- Check mapping statistics (especially the data for the assembly)

Project description

Project parts:

- Assembly
 - Mapping
 - **Variant calling**
- The 2 populations have the be called together in order to find common variants (if no variant found, we want to know if it was reference, or not seen)

Homework: do population variant calling

```
source switch_to_2015a
module load freebayes/1.0.2-33-foss-2015a

#set some variables (see other scripts)
GENOME="$GIT_DIR/results/denovo/genome_k31-unlftgs.fa";
BAM_DIR="$GIT_DIR/results/mapping";
FREEBAYES_OPTIONS="-m 20 -q 15 -p plotdy 2 ";
FREEBAYES_OUTPUT_FILE_NAME="freebayes-no2-q15-plotdy2.vcf";

#copy the GENOME to SVSC_SCRATCH_NODE/genome/genome.fa
#copy the bam files to the SVSC_SCRATCH_NODE/bams directory
banfiles="";
for i in $(ls -l -d SVSC_SCRATCH_NODE/bams/*bam);
do
    banfiles="$banfiles $i";
done

#do the variant calling
freebayes -f $GENOME -r $BAM_DIR/genome/genome.fa $FREEBAYES_OPTIONS $banfiles > $SCRATCH_DIR/FREEBAYES_OUTPUT_FILE_NAME;

#copy the data
```
