

Day 2 PM, Working with NGS data on Hydra-3

How to begin

Copy the Day2 directory in /pool/cluster0/workshop/Hydra_workshop_2015/Day2/ to your working directory in /pool/cluster0/workshop/username/.

Running jobs on Hydra

1. Quality Assessment/Quality check (QA/QC) with FastQC - module bioinformatics/fastqc/0.10.1

FastQC (www.bioinformatics.babraham.ac.uk/projects/fastqc/) aims to provide a simple way to do some quality control checks on raw sequence data coming from high throughput sequencing pipelines.

A. Generate a gsub file to run your data through FastQC - Qsub generator (https://hydra-3.si.edu/tools/QSubGen/) commands:

fastqc sequences.fq

- -- Please use all minimum values for memory and CPUs for this job.
 - Example Qsub Generator output:

```
# /bin/bash
# ----- #
#$ -S /bin/bash
#$ -q sThC.q
#$ -cwd
#$ -j y
#$ -N FastQC
#$ -o FastQC.log
#$ -m abe
#$ -M gonzalez@si.edu
# ----- #
module load bioinformatics/fastqc/0.10.1
# ----- #
echo + `date` job $JOB_NAME started in $QUEUE with jobID=$JOB_ID on $HOSTNAME
fastqc sequences.fq
echo = `date` job $JOB_NAME done
```

- B. Transfer Qsub generator file to hydra from your local computer. *Hint: you can use* scp or copy into a new file in vi.
- C. Execute qsub script to run FastQC on Hydra-3
 - qsub <filename>.qsub
- D. Download output (zip file) to your local computer
 - HINT: use can use the scp command
- E. Open HTML file and assess results.
- 2. Adaptor/Quality Trimming with TrimGalore module bioinformatics/trimgalore/0.4.0

Trim Galore! (www.bioinformatics.babraham.ac.uk/projects/trim_galore/) is a wrapper script to automate quality and adapter trimming as well as quality control.

A. Generate a qsub file to run your data through FastQC - Qsub generator commands:

trim_galore -q 30 --fastqc --clip_R1 5 --three_prime_clip_R1 5 sequences.fq

-- *Note: Default quality encoding is Phred 33 (Sanger/Illumina 1.9). Please verify that the encoding is correct

from the results of FastQC.

Please use all minimum values for memory and CPUs for this job.

Example Qsub Generator output:

```
# /bin/bash
# ----- Parameters ---- #
#$ -S /bin/bash
#$ -q sThC.q
#$ -cwd
#$ -j y
#$ -N Trimgalore
#$ -o Trimgalore.log
#$ -m abe
#$ -M gonzalez@si.edu
# ------ #
module load bioinformatics/trimgalore/0.4.0
# ------ Your Commands----- #
echo + `date` job $JOB NAME started in $QUEUE with jobID=$JOB ID on $HOSTNAME
trim galore -q 30 --fastgc --clip R1 5 --three prime clip R1 5 sequences.fq
echo = `date` job $JOB NAME done
```

B. Run TrimGalore on Hydra-3

- \$ qsub <filename>.qsub
- C. Download output of FastQC run (zip file) to your local computer HINT: use can use the scp command.
- D. Open HTML file and assess results.

3. Assembly with Velvet - module bioinformatics/velvet/1.2.10

Velvet (https://www.ebi.ac.uk/~zerbino/velvet/) is a de novo genomic assembler specially designed for short read sequencing technologies. Velvet runs in parallel (OPENMP). The velvet assembler runs in a two set process; by first running velveth and then by running velvetg. velveth helps you construct the dataset for the following program, velvetg, and indicate to the system what each sequence file represents. velvetg is the core of Velvet where the de Bruijn graph is built then manipulated.

A. Generate a qsub file to assemble your data (sequences.fq) with velveth velveth takes in a number of sequence files, produces a hashtable, then outputs two files in an output directory (creating it if necessary), Sequences and Roadmaps, which are necessary to velvetg.

Qsub generator commands for velveth :

```
export OMP_NUM_THREADS=$NSLOTS
export MAXKMERLENGTH='51'
velveth ecoli.33 33 -short -fastq sequences.fq
```

- -- Please use all minimum values for memory and CPUs for this job.
 - Example Qsub Generator output:

```
# /bin/bash
# ----- #
#$ -S /bin/bash
#$ -pe orte 2
#$ -q sThC.q
#$ -1 mres=1G
#$ -cwd
#$ -j y
#$ -N velveth_short
#$ -o velveth short.log
#$ -m abe
#$ -M gonzalez@si.edu
#
# ------ #
module load bioinformatics/velvet/1.2.10
# ------ #
echo + `date` job $JOB_NAME started in $QUEUE with jobID=$JOB_ID on $HOSTNAME
echo + NSLOTS = $NSLOTS distributed over:cat $PE HOSTFILE
export OMP_NUM_THREADS=$NSLOTS
export MAXKMERLENGTH='51'
velveth ecoli.33 33 -short -fastq sequences.fq
echo = `date` job $JOB_NAME done
```

B. Run velveth on Hydra-3

• \$ qsub velveth.qsub

- C. Generate a qsub file for velvetg to assemble your data in the directory generated from velveth.
 - Qsub generator commands for velvetg:

```
export OMP_NUM_THREADS=$NSLOTS
export MAXKMERLENGTH='51'
velvetg ecoli.33
```

- -- Please use all minimum values for memory and CPUs for this job.
 - Example Qsub Generator output:

```
# /bin/bash
# ------ #
#$ -S /bin/bash
#$ -pe orte 2
#$ -q sThC.q
#$ -1 mres=1G
#$ -cwd
#$ -j y
#$ -N velvetg short
#$ -o velvetg short.log
#$ -m abe
#$ -M gonzalez@si.edu
# ----- #
module load bioinformatics/velvet/1.2.10
# ------ #
echo + `date` job $JOB NAME started in $QUEUE with jobID=$JOB ID on $HOSTNAME
echo + NSLOTS = $NSLOTS distributed over:cat $PE_HOSTFILE
export OMP_NUM_THREADS=$NSLOTS
export MAXKMERLENGTH='51'
velvetg ecoli.33
echo = `date` job $JOB_NAME done
```

- B. Transfer Qsub generator file to hydra from your local computer. *Hint: you can use scp or copy into a new file in vi*.
- C. Run velvetg on Hydra-3

- \$ qsub velvetg.qsub
- D. Count the number of sequences in your output assembly.
 - Navigate to the output directory with the output: cd ecoli.33
 - Assembly output is called contigs.fa
 - Count number of sequences; Hint use skills from Day 1 PM Practical