# MG\_HW7: Taxonomic Classification Using Centrifuge Version 6

#### **James Thornton**

# **Abstract**

This protocol provides a procedure to generate taxonomic data from assembled contigs using centrifuge.

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#### **Guidelines**

Centrifuge documentation

## **Protocol**

#### Step 1.

Log in to the HPC cluster (ICE)

cmd COMMAND

\$ ssh hpc

\$ ice

## NOTES

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Option 3 for those with menu enabled.

#### Step 2.

Move into your class directory.

cmd COMMAND

\$ cd /rsgrps/bh\_class/username
Use YOUR username

#### Step 3.

Create a directory called "original" and move all of your current directories into this directory. We will use this directory to store your original work.

cmd COMMAND

```
mkdir original
mv * original
```

#### Bonnie Hurwitz 28 Oct 2016

Ignore the warning that: "

mv: cannot move `original' to a subdirectory of itself, `original/original'"

## Step 4.

Create directories for running the steps in this protocol. We will download the paired reads from the SRA, run quality control, pair up the reads follow QC, run bowtie2 to remove human contamination, and centrifuge for taxonomic analysis of the individual reads.

```
cmd COMMAND
mkdir fastq
mkdir unmapped
```

mkdir taxonomy

## Step 5.

Go into the fastq directory

```
cmd COMMAND cd fastq
```

## Step 6.

Create a file called "list" with all of your SRR file names.

```
cmd COMMAND nano list
```

#### NOTES

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My file has the following:

% cat list

SRR1647045

SRR1647046

SRR1647047

SRR1647048

SRR1647049

SRR1647141

SRR1647142

## Step 7.

Now we will download all of the SRR files again from the SRA, but this time we will use a few new tricks to download the files as separate paired end read files (1 and 2 files).

Create the following script called get-fastq.sh:

```
cmd COMMAND
#!/bin/bash
#PBS -W group_list=bh_class
#PBS -q windfall
#PBS -l jobtype=cluster_only
#PBS -l select=1:ncpus=2:mem=4gb
#PBS -l walltime=24:00:00
#PBS -l cput=24:00:00
#PBS -l place=pack:shared
#PBS -M netid@email.arizona.edu
#PBS -m bea
module load sratools
echo "my job_id is: ${PBS_JOBID}"
FASTQ_DIR="/rsgrps/bh_class/username/fastq"
cd $FASTQ_DIR
for file in `cat list`; do
   fastq-dump --outdir $FASTQ DIR --qzip --skip-technical --readids --dumpbase --split-
files --clip $file;
done
NOTES
Bonnie Hurwitz 28 Oct 2016
```

--clip

removes adapter sequences

## Bonnie Hurwitz 28 Oct 2016

--skip-technical

Only output the biological, not technical reads

#### Bonnie Hurwitz 28 Oct 2016

--readids

Gives a unique name for each read (for forward and reverse read ids), so they don't end up with the same read id in the R1 and R2 file and break downstream processes.

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```
--gzip
```

compresses your files with gzip

#### Bonnie Hurwitz 28 Oct 2016

--dumpbase ensures the bases are A, T, G, C not in color space (as in SOLiD sequencing)

#### Bonnie Hurwitz 28 Oct 2016

```
--split-files
```

outputs paired ends as \_1 and \_2 files for the forward and reverse read.

# Step 8.

Run the script on the cluster to get all of the fastq paired-end files

```
cmd COMMAND
mkdir std-err std-out
chmod 755 get-fastq.sh
qsub -e std-err/ -o std-out/ get-fastq.sh
Step 9.
```

You can look at the quality of the read files using fastqc as you did before.

But, because we already have an idea of the issues, we are going to run a streamlined QC process in the next step.

```
module load fastqc/0.11.2 fastqc ./*.fastq.gz

Step 10.
```

To speed things up, we are going to apply a general set of quality control parameters. Create a script called run-qualityctrl.sh.

```
cmd COMMAND
#!/bin/bash

#PBS -W group_list=bh_class
#PBS -q windfall
#PBS -l jobtype=cluster_only
#PBS -l select=1:ncpus=2:mem=4gb
#PBS -l walltime=24:00:00
#PBS -l cput=24:00:00
#PBS -l place=pack:shared
#PBS -M netid@email.arizona.edu
#PBS -m bea

module load fastx/0.0.14
echo "my job_id is: ${PBS_JOBID}"
```

```
FASTQ_DIR="/rsgrps/bh_class/userid/fastq"
export $FASTQ_DIR

for file in `cat list`; do
   R1=$file"_1.fastq.gz"
   R2=$file"_2.fastq.gz"
   R10UT=$file"_1.fastq"
   R20UT=$file"_2.fastq"
   gzip -dc $R1 | fastx_trimmer -f 12 | fastq_quality_filter -q 20 -p 80 | fastx_clipper -l 50 > $R10UT
   gzip -dc $R2 | fastx_trimmer -f 12 | fastq_quality_filter -q 20 -p 80 | fastx_clipper -l 50 > $R20UT
   done
```

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make sure you don't split the lines in the script below, or you will get an error.

## **Step 11.**

Run the quality control script on the cluster.

```
cmd COMMAND
chmod 755 run-qualityctrl.sh
qsub -e std-err/ -o std-err/ run-qualityctrl.sh

ANNOTATIONS
```

#### Emma Skidmore 01 Nov 2016

this is the email I've received after running both get-fasta.sh and run-qualityctrl.sh. Does this mean they are failing?

```
PBS Job Id: 831910.service0
Job Name: run-qualityctrl
Execution terminated
Exit_status=0
resources_used.cpupercent=0
resources_used.cput=00:00:00
resources_used.mem=0kb
resources_used.vmem=0kb
resources_used.walltime=00:00:00
```

#### **Step 12.**

After quality control, some of the paired-end sequences are lost and they become singletons. We need to create new files, where the pairs are in R1 (forward) and R2 (reverse), and singletons are in a separate file. We will create a perl script to do this. Please create a script called get-paired.pl and paste in the script from below.

```
cmd COMMAND
#! /uaopt/perl/5.14.2/bin/perl
use strict;
use Bio::SeqIO;
my $file1 = shift @ARGV;
my $file2 = shift @ARGV;
my $out = shift @ARGV;
my $seq_in1 = Bio::SeqIO->new( -format => 'fastq',
                                -file => $file1,
                              );
my $seq in2 = Bio::SeqIO->new( -format => 'fastq',
                                -file
                                        => $file2,
                              );
my $seq_out1 = Bio::SeqIO->new( -format => 'fastq',
                                 -file
                                         => ">$out.R1.fastq"
                              );
my $seq_out2 = Bio::SeqIO->new( -format => 'fastq',
                                 -file
                                         => ">$out.R2.fastq"
                              );
my $seq_out3 = Bio::SeqIO->new( -format => 'fastq',
                                        => ">$out.singletons.fastq"
                                 -file
                              );
my %read_to_count;
while ( my $seq1 = $seq_in1->next_seq() ) {
   my $id = \$seq1->id();
   my (\$srr, \$ct, \$end) = split(/\./, \$id);
   my $read = $srr . "." . $ct;
   $read_to_count{$read}++;
while ( my $seq2 = $seq_in2->next_seq() ) {
   my $id = \$seq2->id();
   my (\$srr, \$ct, \$end) = split(/\./, \$id);
   my $read = $srr . "." . $ct;
   $read_to_count{$read}++;
}
my $seq_in3 = Bio::SeqIO->new( -format => 'fastq',
                                -file
                                       => $file1,
                              );
my $seq in4 = Bio::SeqIO->new( -format => 'fastq',
                                        => $file2,
                                -file
                              );
while ( my $seq1 = $seq_in3->next_seq() ) {
   my $id = \$seq1->id();
   my (\$srr, \$ct, \$end) = split(/\./, \$id);
   my $read = $srr . "." . $ct;
   if ($read_to_count{$read} == 2) {
      $seq_out1->write_seq($seq1);
   }
   else {
      $seq_out3->write_seq($seq1);
   }
}
```

```
while ( my $seq2 = $seq_in4->next_seq() ) {
    my $id = $seq2->id();
    my ($srr, $ct, $end) = split(/\./, $id);
    my $read = $srr . "." . $ct;
    if ($read_to_count{$read} == 2) {
        $seq_out2->write_seq($seq2);
    }
    else {
        $seq_out3->write_seq($seq2);
    }
}
```

# Bonnie Hurwitz 28 Oct 2016

note that I am defining an explicit path here to the version of perl that has BioPerl installed. You can also "module load perl" and use the perl from your environment

#### **Step 13.**

Now we need to create a bash script to run the paired end script on the cluster.

create a script called:

run-pairedends.sh

```
cmd COMMAND
#!/bin/bash
#PBS -W group_list=bh_class
#PBS -q windfall
#PBS -l jobtype=cluster only
#PBS -l select=1:ncpus=2:mem=4gb
#PBS -l walltime=24:00:00
#PBS -l cput=24:00:00
#PBS -l place=pack:shared
#PBS -M netid@email.arizona.edu
#PBS -m bea
module load perl
echo "my job_id is: ${PBS_JOBID}"
FASTQ_DIR="/rsgrps/bh_class/username/fastq"
export $FASTQ DIR
cd $FASTQ DIR
for file in `cat list`; do
   R1=$file"_1.fastq"
   R2=$file"_2.fastq"
   OUT=$file
   ./get-paired.pl $R1 $R2 $OUT
done
```

#### NOTES

Bonnie Hurwitz 28 Oct 2016

Be sure to replace the "netid" and "username" in this script with your own.

## Step 14.

Run the paired ends script on the cluster.

```
cmd COMMAND
chmod 755 get-paired.pl
chmod 755 run-pairedends.sh
qsub -e std-err/ -o std-out/ run-pairedends.sh
NOTES
James Thornton Jr 29 Oct 2016
gsub -e std-err/ -o std-out/ run-pairedends.sh
```

#### **Step 15.**

Now we are ready to run the comparison against the human genome to remove contaminants, and then run the remaining sequences through centrifuge to look at the taxonomy.

change into the taxonomy directory.

```
cmd COMMAND
   cd ../taxonomy
Step 16.
```

Copy the following into a new script named centrifuge tax.sh:

```
cmd COMMAND
#!/bin/bash
#PBS -W group_list=bh_class
#PBS -q windfall
#PBS -l jobtype=cluster only
#PBS -l select=1:ncpus=12:mem=23gb
#PBS -l pvmem=22gb
#PBS -l walltime=24:00:00
#PBS -l cput=24:00:00
#PBS -M netid@email.arizona.edu
#PBS -m bea
#-----EDIT THESE-----
FASTQ_DIR="/rsgrps/bh_class/username/fastq"
OUT_DIR="/rsgrps/bh_class/username/taxonomy"
BT2_OUT_DIR="/rsgrps/bh_class/username/unmapped"
CENT DB="/rsgrps/bh class/b compressed+h+v/b compressed+h+v"
BT2 INDEX="/rsgrps/bh class/bowtie2 index/human index"
module load bowtie2/2.2.5
cd $FASTQ_DIR
for file in `cat list`; do
   # unfiltered
```

```
R1=$FASTQ_DIR/$file".R1.fastq"
R2=$FASTQ_DIR/$file".R2.fastq"
S=$FASTQ_DIR/$file".singletons.fastq"

# no human
NH_R1=$BT2_OUT_DIR/$file".paired.1.fastq"
NH_R2=$BT2_OUT_DIR/$file".paired.2.fastq"
NH_S=$BT2_OUT_DIR/$file".singletons.fastq"

bowtie2 -x $BT2_INDEX -1 $R1 -2 $R2 -U $S -q --very-sensitive-local -p 12 --
un $BT2_OUT_DIR/$file.singletons.fastq --un-conc $BT2_OUT_DIR/$file.paired.fastq
centrifuge -x $CENT_DB -1 $NH_R1 -2 $NH_R2 -U $NH_S -S $OUT_DIR/$file-classout --report-
file $OUT_DIR/$file-centrifuge_report.tsv -q
done
```

## James Thornton Jr 29 Oct 2016

Important: For this to work you Fasta files must end with the extension .fasta

## **Step 17.**

Submit centrifuge tax.sh using qsub:

```
cmd COMMAND
mkdir std-err std-out
qsub -o std-out/ -e std-err/ centrifuge_tax.sh
ten 18.
```

Once the job is running it will loop through all of your Fastq files, remove human reads from the Fastq files, then run Centrifuge on unmapped files to generate taxonomic data. This will take about 1 hour to generate reports for all of your fastq files. You can use qstat to check the status of your job.

```
sqstat -u username
Use YOUR username Under S (Status) 'Q' means queued, 'R' means running
Step 19.
```

Once the job is complete move into your taxonomy directory and ensure all output files are there. If the job was successful there should be a total of 6 "classout" files and 6 "centrifuge\_report.tsv" files.

```
cmd COMMAND
$ cd taxonomy
$ ls

∠ EXPECTED RESULTS
```

```
3. jamesthornton@service2:/rsgrps/bh_class/jetjr/taxonomy (ssh)
[jamesthornton@service2 jetjr]$ cd taxonomy/
[jamesthornton@service2 taxonomy]$ ls
                                  SRR1647238-centrifuge_report.tsv
SRR1647144-centrifuge_report.tsv SRR1647238-classout
SRR1647144-classout
                                  SRR1647239-centrifuge_report.tsv
SRR1647145-centrifuge_report.tsv SRR1647239-classout
SRR1647145-classout
                                  SRR1647240-centrifuge_report.tsv
SRR1647236-centrifuge_report.tsv SRR1647240-classout
SRR1647236-classout
                                  SRR1647260-centrifuge_report.tsv
SRR1647237-centrifuge_report.tsv SRR1647260-classout
SRR1647237-classout
[jamesthornton@service2 taxonomy]$
```

## Step 20.

In your taxonomy directory make a new directory called barplots

#### cmd COMMAND

\$ mkdir barplots

Make sure you are in /rsgrps/bh class/username/taxonomy for this to work correctly

#### **EXPECTED RESULTS**

```
3. jamesthornton@service2:/rsgrps/bh_class/jetjr/taxonomy (ssh)
[jamesthornton@service2 taxonomy]$ pwd
/rsgrps/bh_class/jetjr/taxonomy
[jamesthornton@service2 taxonomy]$ ls
                                  SRR1647237-classout
                                  SRR1647238-centrifuge_report.tsv
SRR1647144-centrifuge_report.tsv SRR1647238-classout
SRR1647144-classout
                                  SRR1647239-centrifuge_report.tsv
SRR1647145-centrifuge_report.tsv
                                  SRR1647239-classout
                                  SRR1647240-centrifuge_report.tsv
SRR1647145-classout
SRR1647236-centrifuge_report.tsv
                                  SRR1647240-classout
SRR1647236-classout
                                  SRR1647260-centrifuge_report.tsv
SRR1647237-centrifuge_report.tsv SRR1647260-classout
[jamesthornton@service2 taxonomy]$
```

#### Step 21.

Copy + Paste the following into a script called cent barplots.R

**Important:** Edit cent.dir and out.dir to include the correct paths

- Edit cent.dir to include the path to your taxonomy directory (/rsqrps/bh class/username/taxonomy/)
- Edit out.dir to include the path to your barplots diretory (/rsgrps/bh class/username/taxonomy/barplots/)

```
cmd COMMAND

#!/usr/bin/env Rscript

#------EDIT HERE------
cent.dir <- "/rsgrps/bh_class/username/taxonomy/"

✓ protocols.io

10
```

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```
out.dir <- "/rsgrps/bh class/username/taxonomy/barplots/"</pre>
file.names <- dir(cent.dir, pattern="-centrifuge report.tsv")
gen barplot <- function (data) {</pre>
  data_title <- gsub("-centrifuge_report.tsv", "", data)</pre>
  data <- read.delim(paste0(i, data))</pre>
  total_reads <- sum(data$numReads)</pre>
  proportion_classified <- data$numReads / total_reads</pre>
  data["proportion classified"] <- proportion classified</pre>
  read subset <-
 subset(data, proportion_classified > 0.005, select = c("name", "numReads", "proportion_cla
ssified"))
  read subset$numReads <- as.numeric(read subset$numReads)</pre>
  png(filename=paste0(out.dir,data_title,"_taxonomy.png"), width = 600, height = 600)
  op <- par(mar=c(15, 8, 4, 2) + 0.1, mgp = c(10, 1, 0))
  p1 <-
 barplot(read subset$proportion classified, main=paste0("Read Proportional Classification:
",data_title), names.arg = read_subset$name, las=2, cex.names = 1, cex.axis = 1, ylab="Prop
ortion Classified", ylim = c(0, 0.90)
  grid(nx=NA, ny=NULL)
  print(p1)
  dev.off()
}
for (i in cent.dir) {
  lapply(file.names, gen_barplot)
Make sure to edit username in cent.dir and out.dir to include YOUR path. Also ensure that both
```

#### James Thornton Jr 29 Oct 2016

cent.dir and out.dir end with the slash

This R script will calculate the total number of reads and then divide the classified reads by the total for each hit generating a proportion classified statistic. Only hits with a proportion of 0.5% of reads classified will be plotted.

#### **ANNOTATIONS**

## Dawson Fairbanks 26 Nov 2016

I'm getting this error message, is there an error in the R script?

directories should be okay-

[dawsonfairbanks@service0 taxonomy]\$ ./cent\_barplots.R

Error in plot.window(xlim, ylim, log = log, ...):

need finite 'xlim' values

Calls: lapply -> FUN -> barplot -> barplot.default -> plot.window

In addition: Warning messages:

1: In min(w.l): no non-missing arguments to min; returning Inf

2: In max(w.r): no non-missing arguments to max; returning -Inf

**Execution halted** 

## Step 22.

Once you have edited cent.dir and out.dir save and close the file. Make cent barplots.R executable.

```
cmd COMMAND
$ chmod +x cent_barplots.R
Step 23.
```

Load the module R:

```
cmd COMMAND
$ module load R
Step 24.
```

Execute cent barplots.R

```
cmd COMMAND
$ ./cent_barplots.R
```

#### **ANNOTATIONS**

Emma Skidmore 01 Nov 2016

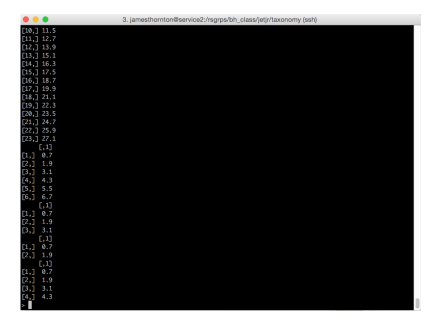
```
[service0@/rsgrps/bh_class/emmanoelskidmore/taxonomy]$ ./cent_barplots.R ./cent_barplots.R: line 6: -: No such file or directory ./cent_barplots.R: line 7: -: No such file or directory ./cent_barplots.R: line 10: syntax error near unexpected token `(' ./cent_barplots.R: line 10: `file.names <- dir(cent.dir, pattern="-centrifuge_report.tsv")'
```

I corrected the paths but I'm still getting this error.

## Step 25.

You should see something similar to what is shown below.

∠ EXPECTED RESULTS



# Step 26.

Move into your barplots directory and make sure you have 6 .png images.

```
cmd COMMAND
```

- \$ cd /rsgrps/bh\_class/username/taxonomy/barplots
- \$ ls

#### **► EXPECTED RESULTS**

```
3. jamesthornton@service2://rsgrps/bh_class/jetjr/taxonomy/barplots (ssh)

[jamesthornton@service2 taxonomy]$ cd barplots/
[jamesthornton@service2 barplots]$ is

SRR1647144_taxonomy.png SRR1647236_taxonomy.png

SRR1647145_taxonomy.png SRR1647237_taxonomy.png

SRR1647239_taxonomy.png

SRR1647260_taxonomy.png

SRR1647260_taxonomy.png

SRR1647260_taxonomy.png
```

# Step 27.

To view the images you must scp to your local machine. Open a new terminal (don't log into hpc). Determine where you want to store the files on your local machine and move into that directory.

# **P** NOTES

## James Thornton Jr 29 Oct 2016

Windows users using Cygwin, your file will be stored in C:/cygwin64/home/USER. Just open a new terminal window and proceed to next step (you can't move to a specific local directory).

## Step 28.

Execute the following command to scp the .png files to your local machine:

#### cmd COMMAND

\$ scp netid@hpc.arizona.edu:/rsgrps/bh\_class/username/taxonomy/barplots/\*.png .
Replace netid and username. (They may be different).

#### **ANNOTATIONS**

# Amy Hudson 31 Oct 2016

I keep getting

bash: /usr/bin/scp: Permission denied

any tips?

#### Farideh Farahnak 01 Nov 2016

scp does not copy files after duo two-factor authentication.

# Step 29.

You can now open the images on your local machine. Reminder that windows users will have their images in C:/cygwin64/home/USER.

## Step 30.

Report on what you've found for each sample. Make sure to state the method used to obtain these results.