Read Recruitment Using Bowtie2 Version 2

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Abstract

This protocol details how to perform read recruitement to the contigs generated from the previous protocol using Bowtie2.

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Protocol

Step 1.

Log into the HPC.

```
cmd COMMAND
```

- \$ ssh hpc
- \$ ocelote

PAIRED END CLEAN UP

Step 2.

For paired end only:

Move into your fasta directory. Ensure only 1 pair from each file is in the fasta directory. Keep the pair that is "1.fasta". Move all "2.fasta" files into the fastq directory.

```
cmd COMMAND
```

- \$ cd /rsgrps/bh_class/username/assembly/fasta
- \$ mv *_2.fasta ../../fastq

NOTES

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In the previous protocol we ended up only using 1 pair for assembly to simplify the process. In this protocol we only want to map the one pair that we used during assembly. Make sure the fasta directory only contains the 1.fasta of the pair.

Step 3.

Move into the assembly/megahit-out directory created from the previous protocol.

cmd COMMAND

\$ cd /rsgrps/bh_class/username/assembly/megahit-out

Step 4.

Simpifly the fasta headers of your final.contigs.fa file using fasta renamer from the Fastx toolkit.

```
$ module load fastx
$ fastx_renamer -n COUNT -i final.contigs.fa -o fixed-contigs.fa
Step 5.
```

Append your fixed-contigs.fa file with your partners fixed-contigs.fa file.

```
cmd COMMAND
```

```
$ cat fixed-contigs.fa /rsgrps/bh_class/partnerusrname/assembly/megahit-out/fixed-
contigs.fa > final-contigs.fa
```

The first argument to cat is YOUR fixed-contigs.fa file while the second is the path to your partners fixed-contigs.fa file. The combined files will be named final-contigs.fa

NOTES

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IMPORTANT: This step requires that your partner has also done the previous step. May require some coordination...

Step 6.

Move into your project directory. Then create a 'read recruit' directory. Move into that directory.

```
cmd COMMAND
$ cd /rsgrps/bh_class/username
$ mkdir read_recruit
$ cd !$

Step 7.
```

Create a bam and bowtie2 index directory.

Move into the contig indexing directory. And create the contig index.

```
cmd COMMAND
$ cd bt2_index
$ module load bowtie2
$ bowtie2-build -f /rsgrps/bh_class/username/assembly/megahit-out/final-contigs.fa contig_index
• NOTES
```

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This step could take awhile depending on the size of your contig file.

Step 9.

Move into your bam directory found at /rsgrps/bh class/username/read recruit/bam

```
cmd COMMAND
$ cd ../bam
Step 10.
```

Make directories for standard error and standard out.

```
cmd COMMAND
$ mkdir std-err std-out
Step 11.
```

Copy the following script named bt2_align.sh. Edit the username variables found in the script.

```
cmd COMMAND
#!/bin/bash
#PBS -W group_list=bh_class
#PBS -q windfall
#PBS -l select=1:ncpus=4:mem=15qb
#PBS -l pvmem=14gb
#PBS -l walltime=24:00:00
#PBS -l cput=24:00:00
#PBS -M netid@email.arizona.edu
#PBS -m bea
echo "my job id is: ${PBS JOBID}"
#####change here ######
FASTA_DIR="/rsgrps/bh_class/username/fasta"
BT2_INDEX="/rsgrps/bh_class/username/read_recruit/bt2_index/contig_index"
OUT_DIR="/rsgrps/bh_class/username/read_recruit/bam"
CONTIGS="/rsgrps/bh_class/username/assembly/megahit-out/fixed-contigs.fa"
###########################
cd $FASTA DIR
export FASTA_LIST="$FASTA_DIR/fasta-list"
ls *fasta > $FASTA LIST
echo "Samples to be processed:" $(cat $FASTA_LIST)
module load bowtie2
module load samtools
while read FASTA; do
    FASTA_N=$(basename $FASTA | cut -d '.' -f 1)
    bowtie2 -x $BT2_INDEX -U $FASTA -f --maxins 800 --fr --very-sensitive-local -p 4 -
S $OUT_DIR/$FASTA_N.sam
    cd $0UT DIR
    echo "Converting $FASTA_N.sam using reference $CONTIGS"
    samtools view -@ 16 -bT $CONTIGS $FASTA_N.sam > $FASTA_N.temp
    echo "Sorting $FASTA N"
    samtools sort -@ 16 $FASTA_N.temp > $FASTA_N.bam
```

```
echo "Removing $FASTA_N.temp"
rm $FASTA_N.temp
cd $FASTA_DIR
```

done < \$FASTA_LIST</pre>

NOTES

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Many of you actually had your fasta files located in:

/rsgrps/bh class/username/assembly/fasta

Which is fine, just make sure the script points to the correct location.

ANNOTATIONS

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Important: CONTIGS variable should be the final-contigs.fa file that you combined with your partner.

Step 12.

Submit the job.

```
sub -e std-err/ -o std-out/ bt2_align.sh
$ sub -e std-err/ -o std-out/ bt2_align.sh
```

Upon job completion navigate to your bam std-err directory.

```
cmd COMMAND
$ cd /rsgrps/bh_class/username/read_recruit/bam/std-err
Step 14.
```

Cat the standard error file to view the alignment rate for each file.

```
cmd COMMAND
$ cat 881767.head1.cm.cluster.ER
Your file name will differ
```