## RNAseq Walkthrough part 3/3

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### Overview

- tophat2 paths
- htseq-count
- 3 DESeq2
  - cluster
  - RStudio
- Running DESeq2

## Today's Goals

- Check tophat2 paths
- Discuss htseq-count flags
- Format data for DESeq
- Run DESeq

## tophat

- tophat2 alignment software
- In: Sequence data
- Out: .bam file where that data aligns to/fits in the genome

### **SLURM Interactive Node**

```
srun -mem-per-cpu=4000MB -pty bash -i
```

- When you're done with an interactive node type exit
- Also, check your SLURM queue squeue -u <netID>
- If you're still running a "bash" job, use scancel <job ID number> to cancel it

## tophat

- tophat2 is the command/software that aligns the reads to the genome
- It is already installed on the cluster
- So put the following location in your path, or in your bash\_profile

```
export PATH=/opt/apps/tophat-bowtie/:$PATH
tophat2
export PATH=/opt/apps/tophat/bin/samtools/bin/:$PATH
samtools
```

## tophat

### tophat2 example format (all one line):

```
tophat2 -p <number of threads> -o <output dir> -G <gff file,
annotations> <bowtie2 index> <R1 fastq> <R2 fastq>
```

- Help can be found by running "tophat2"
- Or in the tophat2 manual online
- http://ccb.jhu.edu/software/tophat/manual.shtml

## **HTSeq**

- python-based program to count reads
- Input:
- .bam file <u>and</u> .gtf/.gff
- Output:
- A table of counts by gene

### htseq-count

- We'll be using htseq-count
- This will count the number of reads mapped to each gene
- That data will be taken into DESeq2

export PATH=/opt/apps/rhel7/Python-2.7.11/bin/:\$PATH

### htseq-count

• What are its flags and options?

htseq-count <options> <alignment bam> <gff file> > <count
output>

### Files to Use

- I have some example files to use for the DESeq tutorial
- They're human RNAseq files from a hypoxia experiment:

ls -lthr /work/cc216/490S/cc216/RNAseq\_pt3

• There are four samples of htseq-count output:

```
s01.norm.counts, s02.norm.counts, s03.hypo.counts, s04.hypo.counts
> head s01.norm.counts
3.8-1.4 0
3.8-1.5 0
5-HT3C2 0
A1BG 252
A1BG-AS1 47
```

 Check that they're all the same length (have the same rows):

```
> wc -l s*counts
45381 s01.counts
45381 s01.norm.counts
45381 s02.counts
45381 s02.norm.counts
```

### • Use paste to combine these files:

```
> paste s01.norm.counts s02.norm.counts s03.hypo.counts
s04.hypo.counts
3.8-1.4 0 3.8-1.4 0 3.8-1.4 0 3.8-1.4 0
3.8-1.5 0 3.8-1.5 0 3.8-1.5 0
5-HT3C2 0 5-HT3C2 0 5-HT3C2 0 5-HT3C2 0
A1BG 252 A1BG 192 A1BG 175 A1BG 153
A1BG-AS1 47 A1BG-AS1 28 A1BG-AS1 31 A1BG-AS1 35
```

### • And cut to eliminate redundant columns:

```
> paste s01.norm.counts s02.norm.counts s03.hypo.counts
s04.hypo.counts | cut -f1,2,4,6,8
3.8-1.4 0 0 0 0
3.8-1.5 0 0 0 0
5-HT3C2 0 0 0 0
A1BG 252 192 175 153
A1BG-AS1 47 28 31 35
```

### Finally write to a file:

```
> paste s01.norm.counts s02.norm.counts s03.hypo.counts
s04.hypo.counts | cut -f1,2,4,6,8 > hypoxia_hsap.counts
```

# Move the count table down to your laptop

### On your laptop's terminal (not slogin):

```
scp -v <yournetID>@dcc-slogin-02.oit.duke.edu:
/work/cc216/490S/cc216/RNAseq_pt3/*.counts
/<your laptop location>/
```

(pay attention to the spaces! the spacing above isn't correct because the slide is too narrow)

- See the website for installation instructions
- Needs two things:
  - A matrix of counts = "cts"
  - A matrix of sample conditions = "coldata"

### In R:

```
> head(cts, n = 4)
s01norm s02norm s03hypox s04hypox
3.8-1.4 0 0 0 0
3.8-1.5 0 0 0 0
5-HT3C2 0 0 0 0
A1BG 252 192 175 153
> coldata
condition type
s01norm "untreated" "paired-end"
s02norm "untreated" "paired-end"
s03hypox "treated" "paired-end"
s04hypox "treated" "paired-end"
```

- Order must match (01, 02, 03...)
- Name the columns and rows appropriately

```
> rownames(cts)
[1] "s01norm" "s02norm" "s03hypox" "s04hypox"
> colnames(cts)
[1] "condition" "type"
```

You use the same function to call (see above) define (see below) column and row names:

```
> colnames(coldata) <- c("condition","type")
```

- Get your data in this format
- Keep track of your work in a .Rmd file!!
- Then (and only then) proceed to running DESeq (see further slides)

## Today's Goals

- Check tophat2 paths
- Discuss htseq-count flags
- Format data for DESeq

## DESeq2 guides

- Here are the DESeq guides that I have summarized in this walkthrough:
- Walkthrough Link
- Focus on "Quick Start" and more specifically:
- Setting the R objects cts and coldata correctly
- Using paste (a unix command) to format your data into cts

http://bioconductor.org/packages/devel/bioc/vignettes/ DESeq2/inst/doc/DESeq2.html

### To run DESeq, first create a "DESeq dataset"

```
> dds <- DESeqDataSetFromMatrix(countData = cts,
colData = coldata,
design = ~ condition)
> dds
```

Where cts and coldata are the files described earlier Output:

```
class: DESeqDataSet
dim: 45381 4
metadata(1): version
```

 Let's throw out genes with low expression levels (here, below 10)

```
> keep <- rowSums(counts(dds)) >= 10
> dds <- dds[keep,]
> dds$condition <- relevel(dds$condition, ref = "untreated")
> dds
```

### Output:

```
class: DESeqDataSet
dim: 17380 4
metadata(1): version
```

We've shrunk our gene list by roughly 60 percent

### Now... Run DESeq!!!

```
> dds <- DESeq(dds)
> res <- results(dds)
> res
```

### Output:

```
log2 fold change (MAP): condition treated vs untreated
Wald test p-value: condition treated vs untreated
DataFrame with 17360 rows and 6 columns
baseMean log2FoldChange lfcSE stat pvalue

A1BG 189.927175 -0.20541335 0.16773404 -1.22463727 0.2207119
```

A1BG-AS1 34.902293 0.02668120 0.26939459 0.09904134 0.9211054

- A few final things to do and explore:
- Sort the results by p-value
- > resOrdered <- res[order(res\$pvalue),]</pre>
  - Summarize the results
- > summary(res)
  - How many genes are significant (at 0.10 and 0.05)?
- > sum(res\$padj < 0.1, na.rm=TRUE)</pre>
- > sum(res\$padj < 0.05, na.rm=TRUE)</pre>

## The End