RNAseq

Goal: Learn how to use various tools to extract information from RNAseq reads.

Input(s): magnaporthe_oryzae_70-15_8_supercontigs.fasta

Moryzae_70-15_liquid_culture{1-3}.fastq.gz

Moryzae_FR13_inPlanta{1-3}.fastq.gz

magnaporthe_oryzae-70-15_8_transcripts.gtf

Output(s): MoryzaeHS{1-8}.ht2

Mo_70-15_LC{1-3}_accepted_hits.bam Mo_FR13_IP{1-3}_accepted_hits.bam

6.1 Information to be gained from RNASeq analysis

For this exercise, we will analyze six RNASeq datasets. Three were generated from RNA isolated from fungal <u>mycelium</u> growing in a liquid culture broth (three biological replicates). Three were generated from excised lesions containing the fungus growing in rice leaves. For the first step of our analysis, we will use **HISAT2** to align the RNASeq reads to a genome assembly of one of the fungal strains from which the RNAseq reads were derived (strain 70-15).

Kim D, Langmead B and Salzberg SL. <u>HISAT: a fast spliced aligner with low memory requirements</u>. *Nature Methods* 2015

https://daehwankimlab.github.io/hisat2/

Mapping RNASeq reads to a reference genome can tell us several things:

- i) It will identify genomic regions that code for RNAs (i.e., regions that are genes).
- ii) The number of reads mapping to each gene will tell us how strongly they were expressed in the two conditions from which RNA was extracted (mycelium growing in culture, infectious hyphae growing *in planta*).
- iii) Gaps in the mappings will reveal the presence and locations of introns in the genes.
- iv) Differences in gap positions will identify variable splice junctions.
- v) Nucleotide differences can be used for variant calling (when the reads come from a genome that is different to the reference—which is true in this case).

6.2 Create an indexed reference genome

Ш	Mak	ke sure	you	are	in 1	the	rnaseq	direc	tory

☐ Start a screen called "rnaseq".

First, we need to create an index of the reference genome that will allow faster searching. We will accomplish this using **hisat2-build**.

Usage: hisat2-build <sequence-to-be-indexed> <prefix-for-index-name>

☐ Index the reference genome.

 hisat2-build ~/blast/magnaporthe_oryzae_70-15_8_supercontigs.fasta MoryzaeHS

6.3 Mapping RNASeq reads to a reference genome

HISAT2 is based on the **Bowtie2** alignment engine but uses a more efficient indexing system for faster searching of RNAseq read alignments to the genome assembly. HISAT2 outputs .sam format files that we will need to convert to .bam for downstream processing.

Usage:

hisat2 [options] -o <output_dir> -x <path-to-index> <input-file(s)>

- ☐ Create a new directory called *alignments* to receive the output files from **HISAT2**
- ☐ Use **HISAT2** to map the RNAseq reads for the *liquid_culture1* dataset to the HISAT index (note: the program won't give you any indication it is running unless you made an error during entry).
 - hisat2 -p 2 -x MoryzaeHS -U
 Moryzae_70-15_liquid_culture1_R1.fastq.gz --max-intronlen 2000
 --summary-file alignments/LC1_summary.txt --dta-cufflinks |
 samtools sort -@ 2 -0
 bam -o alignments/Mo_70-15_LC1_accepted_hits.bam
 - -p number of processors to use. Note: you only have two processors available to you for this exercise, but normally you would run as many as are available.
 - -x name of index to align against
 - **-U** comma-separated list of files containing unpaired reads to be aligned to reference. (we provide only one file here.)
 - --max-intronlen maximum length of intron (500000 if left blank)
 - --summary-file write alignment summary to this file
 - **--o** name of output file

Note that we pipe the output of **hisat2** straight into **samtools** so that we can sort the results by chromosome position and then convert the sorted alignments from *.sam* to *.bam* format.

samtools sort - -@ 2 -0 bam -o output.bam

By default, **HISAT2** generates a .sam alignment file, which needs to be converted to .bam format and then sorted according to genome coordinates. This can be conveniently accomplished using a single pipe into the **samtools sort** program.

_	•	use the output of the prior program as input				
_	·@	number of processors to use				
-0		output format (bam/sam)				
_	-0	output file path				
6.4 I	Examine t	he RNASeq mapping results				
		our command line to look at the <i>LC1_summary.txt</i> alignment summary file to get a nuch useable data we have.				
	How many tota	al reads were present in the input file?				
	How many read	ds mapped to the reference genome?				
	Use head to tal	ke a quick look at the first 10 lines of results in the alignments file.				
	• head al	ignments/Mo_70-15_LC1_accepted_hits.bam				
	☐ Does the output make any sense? No? That's because the file is in a binary format. Let's use samtools to convert the .bam file into the human-readable .sam format:					
	• samtool	s view alignments/Mo_70-15_LC1_accepted_hits.bam				
	Whoa! Did you	a catch all that? Quit the process (ctrl-c) and try piping the results through head.				
	The main fields	in sam/bam format are as follows:				
	1. QNAME 2. FLAG 3. RNAME 4. POS 5. MAPQ 6. CIGAR 7. RNEXT	Query name (read ID) Bitwise flag describing alignments (e.g., strand, paired info) Reference sequence name 1-based leftmost position of alignment on reference Mapping quality (Phred value, 0–255) Alignment representation Mate reference name (= on same chromosome, * mate unmapped, or single end read) Meta position				
	8. PNEXT 9. TLEN 10. SEQ 11. QUAL	Mate position Template length (inferred length of sequenced fragment based on paired read info) Sequence Base quality scores (ASCII)				
		ne sam/bam format can be found by searching on Google or ChatGPT. t alignment record:				

	What is the alignment length?
	What is the lowest quality value for the read represented by this alignment (in PHRED format)?
	Does this read align on the forward or the reverse strand of the reference? Use the SAM flag tool decoder to find out (https://broadinstitute.github.io/picard/explain-flags.html).
	Space bar through a few pages to view a few more alignments and then quit the program (ctrl-C) when you're done.
6.5	Align the remaining liquid_culture datasets
	Repeat the alignment commands process for the <i>liquid_culture2</i> and <i>liquid_culture3</i> datasets. You can use the up arrow to speed up the command line entry but remember to change both the <u>input</u> AND <u>output</u> file identifiers. (Remember, there are two output files.). For example,
•	hisat2 -p 2 -x MoryzaeHS -U Moryzae_70-15_liquid_culture2_R1.fastq.gz max-intronlen 2000summary-file alignments/LC2_summary.txt dta-cufflinks samtools sort@ 2 -0 bam -o alignments/Mo_70-15_LC2_accepted_hits.bam
	etc After each run is complete, list the <i>alignments</i> directory contents to make sure that the expected files were created.
6.6	Align the inPlanta datasets
	Now you have probably discovered that sitting and waiting for long alignment jobs to finish can make repetitive tasks very frustrating. Let's learn to appreciate the power of the command line for automation. We will perform the alignments for <u>all three</u> <i>inPlanta</i> datasets using a single command to loop through the three files, one at a time:
•	for f in {13}; do echo Working on dataset Moryzae_FR13_inPlanta\${f}_R1.fastq.gz; hisat2 -p 2 -x MoryzaeHS -U Moryzae_FR13_inPlanta\${f}_R1.fastq.gzmax-intronlen 2000dta-cufflinkssummary-file alignments/IP\${f}_summary.txt samtools sort@ 2 -0 bam -o alignments/Mo_FR13_IP\${f}_accepted_hits.bam; done
	Here the for loop sequentially assigns the values 1, 2, and 3 to the variable f. We then substitute the

Here the **for** loop sequentially assigns the values 1, 2, and 3 to the variable **f**. We then substitute the variable **\$**{**f**} for the numerical identifiers in our input and output files. The semicolons (";") allow us to run several commands in a single line sequentially without passing the input of one command to the next (unlike a pipe). For example, we first run **echo** and then **hisat2**.

You should see a run time message saying "Working on dataset...". If instead, you see a command prompt ">," you made a mistake in typing the for loop. In that case, quit the command (ctrl-c) and re-type the command.

☐ When the three runs are completed, list the *alignments* directory contents to make sure that all of the expected files were created.

6.7 Assembling transcripts from RNAseq data

We will use **StringTie** to build complete transcripts from the individual RNAseq read mappings.

Pertea et al. (2015) StringTie enables improved reconstruction of a transcriptome from RNA-seq reads. Nature Biotechnology 33(3):290-295.

http://ccb.jhu.edu/software/stringtie/index.shtml

The first step in differential gene expression analysis is to identify the gene from which each sequence read is derived. **StringTie** examines the RNA-seq read mapping results produced by a read aligner such as **TopHat** or **HISAT2** and attempts to reconstruct the complete transcripts, identify transcript isoforms, and estimate transcript abundance. **StringTie** does this by clustering reads and constructing a splice graph for each cluster from which it will identify transcripts. A flow network is then constructed for each identified transcript to estimate the abundance of the transcript via a maximum flow algorithm. StringTie can accept an existing gene annotation file as a guide for constructing transcripts.

Usage:

stringtie <path/to/accepted_hits.bam> [options] Take a quick peek at all the **StringTie** options that are available. stringtie --help ☐ Make sure you are in the *maseq* directory. ☐ First, take a look at the existing annotation file, magnaporthe_oryzae_70-15_8_transcripts.gtf, to be sure you understand its format. Note that the GTF format is very similar to the GFF3 format used in the .gff3 file that you produced in the gene finding exercise. Each feature entry has an MGG gene_id which identifies the gene to which the feature corresponds. Note also that each gene has one or more corresponding transcript_ids, with T0 denoting the first transcript isoform, T1, the second, etc. ☐ Create a directory named *transcripts*. Now we are ready to run **StringTie**, and we will provide the *.gtf* file as a reference transcriptome: stringtie alignments/Mo_70-15_LC1_accepted_hits.bam -p 2 -G magnaporthe_oryzae_70-15_8_transcripts.gtf -o transcripts/Mo_70-15_LC1_transcripts.gtf number of processors to use -р

tells **StringTie** to use the provided reference annotation to guide transcript

assembly but also to report novel transcripts/isoforms

name of output directory

-G

-0

Notes:

- A) Omitting the **-G** option (and accompanying *.gtf* file specification) from the above command would tell the program to generate a *de novo* transcript assembly. Alternatively, one can use **-G** with an additional **-e** flag, which will tell the program to assemble only those reads that correspond to previously identified genes/transcripts.
- B) It is recommended that you assemble your replicates individually, i) to speed computation; and ii) to simplify junction identification. Therefore, you will need to run **StringTie** separately for each of your .bam files.

Make sure that the <i>Mo_70-15_LC1_</i>	_transcripts.gtf output file	was created	indicating	that the
command completed successfully.				

 \square Look at the first few lines of the Mo_70 -15_LC1_transcripts.gtf in order to answer the questions below (use the information below to understand what the different fields mean).

The final column of a GTF file is a semicolon-separated list of tag-value pairs providing additional information about each feature. Depending on whether an instance is a transcript or an exon and on whether the transcript matches the reference annotation file provided by the user, the content of the attributes field will differ. The following list describes the possible attributes shown in this column.

- a. gene_id: A unique identifier for a single gene and its child transcript and exons based on the alignments' file name.
- b. transcript_id: A unique identifier for a single transcript and its child exons based on the alignments' file name.
- c. exon_number: A unique identifier for a single exon, starting from 1, within a given transcript.
- d. reference_id: The transcript_id in the reference annotation (optional) that the instance matched.
- e. ref_gene_id: The gene_id in the reference annotation (optional) that the instance matched.
- f. ref_gene_name: The gene_name in the reference annotation (optional) that the instance matched.
- g. cov: The average per-base coverage for the transcript or exon.
- h. FPKM: Fragments per kilobase of transcript per million read pairs. This is the number of pairs of reads aligning to this feature, normalized by the total number of fragments sequenced (in millions) and the length of the transcript (in kilobases).
- i. TPM: Transcripts per million. This is the number of transcripts from this particular gene normalized first by gene length, and then by sequencing depth (in millions) in the sample.

For the first gene reported for chromosome 8.1:					
a.	Does this gene correspond	to a previously annotated	Magnaporthe gene (does it	have an MGC	
	identifier), or is it novel?	previously annotated:	: novel:		

b. How long is the gene? _______ bp
c. Does it contain any introns? yes: _______; no: ______.
d. How abundantly is it expressed in fragments per kilobase per million reads? ______.
□ Now we will complete the rest of the transcript assemblies by using another kind of for loop.
for f in \$(ls alignments/*bam | grep -v LC1); do echo Working on \$f; g=\${f/*\//}; stringtie \$f -p 2 -G magnaporthe_oryzae_70-15_8_transcripts.gtf __o transcripts/\${g/accepted*/}transcripts.gtf; done

Here we are running the loop a little differently. We list all bam files in the alignments directory, and, after omitting the file that we've already processed (using grep -v LC1), we assign the results to the variable f. Then, for each value (i.e., each file) of f, we strip off everything before and including the first forward slash (\${f/*\//}}) and save the result to the variable g. Then, we run stringtie on each value of f (alignments/Mo_70-15_LC1_accepted_hits.bam, etc.) and output the results to the transcripts directory under a filename that we create by cutting off the accepted_hits.bam suffix from what is stored in the g variable (\${g/accepted*/}}). We then place what's left (e.g., "Mo_70-15_LC2_") in front of "transcripts.gtf".

6.8 Merging transcript assemblies

We will use **StringTie** with the **-merge** option to generate a "super-assembly" of transcripts based on the mapping information from all six RNAseq datasets. This will allow **StringTie** to identify overlaps between alignment data for different datasets. In this way, it can assemble complete transcripts for genes whose expression levels are too low to allow full transcript reconstruction from a single sequencing lane.

Usage:

stringtie --merge [options] <list_of_gtf_files>

- \square Make sure you are in the *maseq* directory.
- First, we need to create a text file that lists the paths to the various transcript assemblies that we want *stringtie* to merge. This document should have the format:

Here is an example of where a file created by a standard text editor such as **Word** will not be read properly by the **stringtie** program and would produce an error.

		ould open a test editor such as nano to create this document but a much more efficient watthis (while avoiding typing errors) is to use another for loop:	У
		<pre>for f in \$(ls transcripts/*gtf); do echo ./\$f >> assemblies_gtf_list.txt; done</pre>	
	>>	means redirect to a file and append new entries to the same file	
	Now	we can run stringtie with the merge option.	
		stringtiemerge -p 2 -o merged_asm/merged.gtf -G magnaporthe_oryzae_70-15_8_transcripts.gtf assemblies_gtf_list.txt	
	-р	number of processors to use	
	- o	output file name for the merged transcripts. Here, stringtie will create the <i>merged_asm</i> directory if it doesn't already exist (this is not the case for many programs)	
	- G	include the reference annotation in the merging operation	
	Exam	nine the merged.gtf file produced by stringtiemerge inside of the merged_asm directory.	
		ommand line tools to interrogate the <i>merged.gtf</i> file to identify novel transcripts that have no previously identified. Note: novel transcripts will lack MGG identifiers.	t
a bit of analysis	a prob s, this v und th	that the <u>gene-id</u> attribute for each transcript has a stringtie -specific prefix (<u>MSTRG</u>). This is blem because, if we use this file for downstream analyses such as differential gene expression will be the gene ID that is displayed, even if there is already an established MGG identifier. To problem, we will use a custom perl script to rename the gene_IDs and save to a new file.	n o
	Use the	he Inherit_IDs.pl script to convert the gene-id field to its corresponding MGG identifier (able).	if
		<pre>perl Inherit_IDs.pl merged_asm/merged.gtf > merged_asm/cufflinks.gtf</pre>	

6.9 Differential gene expression analysis

The **HISAT2/StringTie** pipeline normally feeds into the **Ballgown** program for differential expression analysis (Frazee AC et al. 2015. Nat Biotechnol. 33:243-246). However, this would require many of us to learn the R programming language and environment, which itself could take a whole week. Therefore, we will use the legacy **cuffdiff** program to determine which genes are differentially expressed in one of the RNAseq datasets.

What is the total number of genes that show at least one alternative transcription start site and/or alternative splice junction? (Recall that the first transcript isoform is given a T0 identifier.)

http://cole-trapnell-lab.github.io/cufflinks/cuffdiff/

For our purposes, we will have **cuffdiff** use the *cufflinks.gtf* file which combines the prior gene annotations with the new information (novel transcripts, isoforms, etc.) generated from our RNAseq data. **cuffdiff** then uses the alignment data (in the *.bam* files) to calculate and compare transcript abundances.

Usage:

Note: experimental replicates are separated with commas; datasets being compared are separated by a space (i.e.: Condition1_rep1,Condition21_rep2 (SPACE) Condition2_rep1,Condition2_rep2).

Since **bash** doesn't know about the way **cuffdiff** separates the experimental replicate files (and we have not provided any custom code to **bash** for tab-completing **cuffdiff**), tab completion will not work for an argument after you type the first comma. We can get around this limitation by initially separating with spaces and then replacing the appropriate spaces with commas.

For our experiment, we will compare transcript abundance in fungus grown in liquid culture (three replicates) versus fungus growing *in planta* (three replicates).

☐ Run cuffdiff as follows (DO NOT PUT A SPACE BETWEEN THE COMMAS AND THE FLANKING DATASETS—SEE ABOVE COMMENT IN RED BEFORE RUNNING):

```
    cuffdiff -o diff_out -p 2 -L culture,inPlanta
        -u merged_asm/cufflinks.gtf
        alignments/Mo_70-15_LC1_accepted_hits.bam,
        alignments/Mo_70-15_LC2_accepted_hits.bam,
        alignments/Mo_70-15_LC3_accepted_hits.bam (include a space here)
        alignments/Mo_FR13_IP1_accepted_hits.bam,
        alignments/Mo_FR13_IP2_accepted_hits.bam,
        alignments/Mo_FR13_IP3_accepted_hits.bam
```

- -0 output directory where results will be deposited
- -p number of processors to use
- -L Labels to use for the two conditions being compared. These labels will appear at the top of the relevant columns in the various output files.
- -u Tells **cufflinks** to do an initial estimation procedure to more accurately weight reads mapping to multiple locations in the genome
- This produces many output files in the *diff_out* output directory. Most of these are not really needed for us to identify the genes that have expression differences, so we will not go into them in detail. A brief explanation of files is provided in the appendix to this module.
- The main information about gene expression differences are written to the file named <code>gene_exp.diff</code> inside the defined output folder. View the header of this file and see if you can determine what information is contained in the various columns. If necessary, look at the description of the output columns in the appendix to this module, or you can find more information in the online <code>cufflinks</code> manual:

http://cole-trapnell-lab.github.io/cufflinks/manual/

UK/KY-INBRE BIOINFORMATICS WORKSHOP, 2025

	Practice using the command line to produce an output that shows only those genes that cuffdiff predicts to be differentially expressed between the two samples.
	Use awk to print the lines of results for genes that show a more than ten-fold higher expression <i>in planta</i> versus in liquid culture.
	Use the command line to produce a list that contains the identities of the genes that show significant differences in their expression levels (only the names of the genes and nothing else). Write this list to a file.
Hi	nt: You will need to use grep and/or awk .

Appendix

Explanation of cuffdiff output files:

The main types of files produced are:

bias_params.info The bias model learns how read coverage varies across transcript sequences,

and stores those parameters here.

model.info stores information about the empirical fragment length distribution

model that Cuffdiff builds when estimating transcript abundance.

count_tracking reports raw and normalized fragment counts for each feature

across individual replicates.

fpkm_tracking reports FPKM values for each feature (gene, isoform, etc.) across all samples

and replicates.

read_group_tracking contains per-replicate quantification data — including FPKM values,

fragment counts, and confidence intervals

Several differential expression output files are produced:

gene_exp.diff Gene-level differential expression. Tests differences in the summed FPKM

of all transcripts sharing the same gene_id

isoform_exp.diff Transcript-level differential expression. Tests differences in the summed

FPKM of each transcript isoform for a given gene

tss_group_exp.diff Primary transcript differential expression. Tests differences in the summed

FPKM of transcripts sharing the same transcription start site (tss)

cds_exp.diff Coding sequence differential expression. Tests differences in the summed

FPKM of transcripts that result in the same coding sequence, regardless of

tss.

Each of the differential expression output files has the following fields:

test_id identifier of the feature being tested

gene_id official gene/feature identifier (if it was previously annotated)

gene official name of gene (e.g. BRCA2)

locus genomic coordinates

sample_1 identifier for test group 1 (e.g. inCulture)
sample_2 identifier for test group 2 (e.g. inPlanta)
status whether a statistical test was performed
value_1 mean expression value for group 1
value_2 mean expression value for group 2

log2(fold_change) log2 fold change in expression for group 2 relative to group 1

test_stat test statistic value p_value raw p-value for the test

q_value adjusted p-value after Benjamini-Hochberg correction for false discovery

rate

significant whether the expression level is significantly different between group 1 and 2