

## Bulk RNA-seq Workflow

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#View reference and FASTQ files.
#FASTQ files are from the Griffith lab.
#https://github.com/griffithlab/rnaseq_tutorial/wiki/RNAseq-Data
seqkit stats hg38.fasta
cat hg38.fasta | head -5
seqkit stats *.fastq.gz

#Make file of root names.
cat > rootNames.txt
brain1
brain2
brain3
cancer1
cancer2
cancer3

#Align sequencing reads to genome using hisat2.
mkdir -p bam

cat rootNames.txt | parallel "hisat2 -x ../Genomics/OptimizedPipeline/hg38 -1 {}.read1.fastq.gz -2
{}.read2.fastq.gz | samtools sort > bam/{}.bam"

#Index BAM files.

cat rootNames.txt | parallel "samtools index bam/{}.bam"

#Load BAM files into IGV.

#Count features, remembering that data is stranded.
cat rootNames.txt | parallel -j 1 echo 'bam/{}.bam' | xargs featureCounts -p -s 1 -a
../Genomics/OptimizedPipeline/hg38.ensGene.gtf -o counts.csv

#Perform transcript classification analysis.
#Activate salmon environment.
conda activate salmon_env

#Build salmon index.
salmon index -t GRCh38_latest_rna.fna -i salmon.idx

#Run salmon quantification.
cat rootNames.txt | parallel -j 4 "salmon quant -i salmon.idx -l A --validateMappings -1
{}.read1.fastq.gz -2 {}.read2.fastq.gz -o salmon/{}"

#Save quant.sf files as TSV files. Then open as spreadsheets and copy and paste each NumReads
#column into a new spreadsheet to make one file of counts (salmonCounts.csv).
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#Run edgeR on counts files to perform differential expression analysis. I will use code from the
#“Biostars Handbook” to run edgeR.
#First for hisat2 featureCounts data.
Rscript code/edger.r
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#Next, for salmon data.
Rscript code/edger_salmon.r
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```
#Make heatmap of hisat2 featureCounts data with FDR < .05.
Rscript code/create_heatmap.r
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#Brain and cancer samples segregate very well.
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#Make heatmap of salmon data with FDR < .05.
Rscript code/create_salmon_heatmap.r
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#The salmon heatmap has more genes with FDR < .05. It segregates well by group, although there are
#two or three genes that do not segregate that well.
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```
#Calculate transcript integrity number (TIN) as a QC step. Use tin.py to calculate TIN for each
#transcript using hisat2 BAM files.
python tin.py -i bam -r hg38_RefSeq.bed
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#This outputs a .tin.xls file for each BAM file, as well as a .summary.txt file giving the average TIN for
#that BAM file. I will only use genes that have TIN > 60.
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#The next step is to perform gene set enrichment analysis (GSEA) on genes with FDR < .05 and TIN >
#60 using agriGO, DAVID, Panther, goatools, and g:Profiler. These tools will return gene ontology
#(GO) annotations that are statistically over-represented compared to background.
```