# RnaSeq Data Pre-Processing

Dr. Rudramani Pokhrel,

Last modified: 14 Apr 2022

Immunobiology Department, The university of Arizona, Tucson Arizona

email: rpokhrel@email.arizona.edu

The aim of this tutorial is to generate gene vs sample count matrix from raw fastq Bulk RNAseq read files. It starts with preparing the virtual environment and step by step process to generate count data. At the end I have provided the necessary script files to process the method in batches.

### Download the miniconda package from conda repo for linux platform.

Type following commands in the terminal:

- Download: wget https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86\_64.sh
- Install: bash Miniconda3-latest-Linux-x86 64.sh

Initialize the conda environment: conda init bash

Note: You can download your version according to the platforms (linux, macOS, Windows) from this link: https://docs.conda.io/en/latest/miniconda.html. For Apple M1 please follow this link: https://naolin.medium.com/conda-on-m1-mac-with-

miniforge-bbc4e3924f2b

#### Create the virtual environment of named rna-seq and installing all the required packages.

Run following commands in the terminal:

- Create virtual environment named rna\_seq: conda create -y -n rna\_seq python=3
- Activate the environment: conda activate rna\_seq
- Install required packages: conda install -y -c bioconda -c conda-forge -c anaconda fastp bwa htseq glob2 pandas samtools==1.11

#### Mappping read files to the reference genome and get transcriptomic counts

- Use Ensembl (https://uswest.ensembl.org/info/data/ftp/index.html) or from USCS genome browser (https://genome.ucsc.edu) or from NCBI (https://www.ncbi.nlm.nih.gov) or from GENECODE (https://www.gencodegenes.org) to download reference genome and annotation
- Make a folder in which you will put all of you genome data and fastq read files
  - mkdir rnaseq
  - cd rnaseq
- copy all the files into this directory using cp command or simply copy/paste
- Next process would be doing quality control of your fastg reads. I would prefer **fastp** package which can be used to inspect the read data and at the same time perform QC (adapter trimming and filtering) on read fastq file:
  - First create a folder for filtered reads
    - mkdir filtered read
  - Run fastp for single end read
    - fastp -i read1 -o filtered\_read/read1 -h filtered\_read/filtered.html
  - For paired end reads:
    - fastp -i read1 -l read2 -detect\_adapter\_for\_pe -o filtered\_read/read1 -O filtered\_read/read2 -h filtered\_read/filtered.html

The filtered.html file gives the summary of read before and after filtering.

#### Mapping to genome

I am using bwa aligner package to map the filtered read to the reference genome.

- First prepare the index file:
  - bwa index reference.fasta -p bwa\_index
  - Note: -p is for prefix for all index files
- Now run following command for mapping to genome:
  - For single end read:
  - bwa mem -t 8 bwa\_index filtered\_read/read1 > mapped\_read.sam For paired end read:
  - Note: -t option is for number of cores/cpus you want to use. I recommend using minimum of 8GB of memory
- Use samtools to convert sam to bam file, sort the bam file and index it samtools view -Sb mapped\_read.sam > mapped\_read.unsorted.bam
  - samtools sort -o mapped\_read.sorted.bam mapped\_read.unsorted.bam
  - samtools index mapped\_read.sorted.bam
- Use htseq-count to obtain gene/transcriptomic count matrix
  - htseq-count -format=bam -stranded=no -type=gene -order=pos -idattr=ID mapped\_read.sorted.bam reference\_annnotation.gff3 > count.txt

bwa mem -t 8 bwa\_index filtered\_read/read1 filtered\_read/read2 > mapped\_read.sam

Note: you can change **-type** and **-idattr** options according to your study need. Look into the **Annotation.gff3** file for available type and ID.

## Employ above methods in multiple samples

Generally, we have multiple samples (eg. control vs disease) read files. Here I present a bash scripting to loop over all the samples for paired end reads.

# Create directory for filtered read and count matrix

mkdir -p filtered\_read mkdir -p htseq\_count

# Looping over read1 for R1 in \*\_R1\_\*.gz do

# replacing R1 with R2 for reverse read. please look at your pattern

 $R2=($\{R1//_R1_/_R2_\})$ 

# you can print with echo command to see whether it is working or not #echo \$R1 \$R2

# create a variable B for base name for reading convenience which only captures the sample information

 $B=\$\{R1::-16\}$ 

# Here B trims all the last 16 character from read file. You can choose your own number fastp -i \$R1 -I \$R2 --detect\_adapter\_for\_pe -o filtered\_read/\$R1 -O filtered\_read/\$R2 -h filtered\_read/\$B.html

bwa mem -t 8 bwa\_index filtered\_read/\$R1 filtered\_read/\$R2 > \${B}.sam

samtools view -Sb \${B}.sam > \${B}.unsorted.bam

samtools sort -o \${B}.bam \${B}.unsorted.bam samtools index \${B}.bam

htseq-count --format=bam --stranded=no --type=gene --order=pos --idattr=ID \${B}.bam \$gff3 > htseq\_count/\${B}.txt done

## **Alternatively**

You can download the bash script from my github ripo and use it directly. Download script in current rnaseq folder from:

wget https://raw.githubusercontent.com/githubrudramani/Bioinformatics/master/rna\_seq\_preprocess.sh Note: change the path/file name and location of reference in rna\_seq\_preprocess.sh script.

bwaIndex=/path/to/bwa\_index

gff3=/path/to /annotation.gff3

Run command:

chmod +x rna\_seq\_preprocess.sh ./rna\_seq\_preprocess.sh

Now you have count matix for all the samples in *htseq\_count* directory. You can use python or R to merge the count matrix and make sample vs transcriptomic data matrix. I have written a python script to merge the data. To download the script run following command in the inside htseq\_count folder:

cd htseq\_count

wget https://raw.githubusercontent.com/githubrudramani/Bioinformatics/master/merge\_htseq\_counts.py

Then run: **python merge\_htseq\_counts.py** The name of output file is merged\_counts.csv in csv format.

-----Code Summary-----

Following lines of code is all needed for pre-procession bulk RNAseq raw fastq read files. Make sure you have changed the respective file names and paths in the scripts.

wget https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86\_64.sh bash Miniconda3-latest-Linux-x86\_64.sh

conda init bash

conda create -y -n rna\_seq python=3

conda activate rna\_seq

conda install -y -c bioconda -c conda-forge -c anaconda fastp bwa htseq glob2 pandas samtools==1.11

bwa index reference.fasta -p bwa\_index

wget https://raw.githubusercontent.com/githubrudramani/Bioinformatics/master/rna\_seq\_preprocess.sh

chmod +x rna\_seq\_preprocess.sh

./rna\_seq\_preprocess.sh

cd htseq\_count

wget https://raw.githubusercontent.com/githubrudramani/Bioinformatics/master/merge\_htseq\_counts.py

python merge\_htseq\_counts.py

## Output:

merged\_counts.csv

After that you can use differential analysis statistical packages like **DEseq2**, **edgeR**, **Limma** from Bioconductor.