Bulk RNAseq Data Pre-Processing

Snakemake Workflow

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Last modified: 24 Apr 2022

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

Install 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-withminiforge-bbc4e3924f2b

Create the virtual environment of named rna-seq and install all the required packages and download reference Genomes.

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 snakemake fastp bwa htseq samtools==1.11

 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 files

The work flow implementation via Snakemake here has following steps:

- Generate bwa index
- Perform QC (adapter trimming and filtering) on read fastq file using fastp package
- **bwa mem** to map fastq files to reference genome
- Use **samtools** to convert the mapped **sam** files to **bam** and sort and index the **bam** files
- Use *htseq-count* to generate read count to sample matrix data
- Use custom python script *merge_htseq_counts.py* to merge individual count matrix data

Let's begin:

- Make a folder for you working directory
 - mkdir rnaseq cd rnaseq
- Make other two folders reference and files
 - mkdir reference files
- move genome reference (fasta and gff3 or gtf) to reference and fastq reads to files
- In your working directly create a config.yaml file for Snakefile input
 - config.yaml:

```
core: 10
  ref:
     fa: reference/reference.fasta
     gff3: reference/reference.gff3
read dir: files/
read_ext: _R1_001.fastq.gz
```

• Feel free to edit this *config.yaml* file according to your data names and available cpu cores

```
Snakefile:
  configfile: "config.yaml"
  core = config["core"]-1
  ext = config["read_ext"]
  SAMPLES, = glob_wildcards(config['read_dir']+"{sample}" + ext)
  rule all:
         expand(config["read dir"] + "trimmed/{sample} trimmed R1 fastq.gz", sample = SAMPLES),
         expand("mapped_read/{sample}.bam", sample = SAMPLES),
         expand("mapped read/{sample} sorted.bam", sample = SAMPLES),
         expand("mapped_read/{sample}_sorted.bam.bai", sample = SAMPLES),
         expand("htseq_count/{sample}.txt", sample = SAMPLES),
         #expand("htseq_count/merged_counts.csv")
         #expand("htseq_count/metaData.csv")
  rule bwa_index:
      input:
          ref_fasta = config["ref"]["fa"]
          ref = touch("reference/bwa index")
      shell:
          "bwa index {input.ref_fasta} -p {output.ref}"
  rule fastp:
      input:
         R1 = config["read dir"]+ "{sample}" + config["read ext"]
         R2 = config["read_dir"]+ "{sample}" + config["read_ext"]
      output:
        R1= config["read dir"]+ "trimmed/{sample} trimmed R1 fastq.gz",
        R2= config["read dir"] + "trimmed/{sample} trimmed R2 fastq.gz",
        html= config["read_dir"]+ "trimmed/{sample}.fastp_report.html",
        json= config["read_dir"]+ "trimmed/{sample}.fastp_report.json",
      threads: core
      shell:
        "fastp -w {threads} -i {input.R1} -I {input.R2} --detect_adapter_for_pe -o {output.R1} -O {output
t.R2} -h {output.html} -j {output.json}"
  rule bwa_mem:
      input:
          ref = "reference/bwa_index",
          R1 = config["read_dir"]+ "trimmed/{sample}_trimmed_R1_fastq.gz",
          R2 = config["read_dir"]+ "trimmed/{sample}_trimmed_R2_fastq.gz"
      output:
         "mapped_read/{sample}.bam"
         "mapped_read/{sample}.bwa_map.log"
      threads: core
      shell:
         "bwa mem -t {threads} {input.ref} {input.R1} {input.R2} | samtools view -@ {core} -Sb - > {output}
2> {log}"
  rule samtools_sort:
      input:
          "mapped_read/{sample}.bam"
      output:
          "mapped_read/{sample}_sorted.bam"
      shell:
          "samtools sort -o {output} {input}"
  rule samtools_index:
      input:
          "mapped_read/{sample}_sorted.bam"
      output:
          "mapped_reads{sample}_sorted.bam.bai"
      shell:
          "samtools index {input}"
  rule htseq_count:
         gff3 = config["ref"]["gff3"],
         bam = "mapped_read/{sample}_sorted.bam",
         bai = "mapped_read/{sample}_sorted.bam.bai"
      output:
          "htseq_count/{sample}.txt"
      shell:
          "htseq-count -n {core} --format=bam --stranded=no "
          --type=gene --order=pos --idattr=ID {input.bam} {input.gff3} > {output}"
  rule merge_count:
      input:
          expand("htseq_count/{sample}.txt", sample = SAMPLES)
      output:
          "htseq_count/merged_counts.csv"
          "htseq_count/metaData.csv"
      shell:
        "cd htseq count"
        "wget https://raw.githubusercontent.com/githubrudramani/Bioinformatics/master/merge_htseq_counts.py"
        "python merge_htseq_counts.py"
```

Note: the indentation in config.yaml and Snakefile is not tap seperated.

Run command in your working directory *rnaseq:* snakemake --snakefile Snakefile --core 10

Note: While running this Snakefile, I found the clash of library between **snakemake** and **htseq**. In such case you can create individual environment for *htseq*.

following link: https://github.com/githubrudramani/Pipelines/tree/main/Bulk-RNAseq/DESeq2.

For that purpose modify the *rule htseq_count*:

```
rule htseq_count:
    input:
       gff3 = config["ref"]["gff3"],
      bam = "mapped_read/{sample}_sorted.bam",
      bai = "mapped_read/{sample}_sorted.bam.bai"
    output:
        "htseq count/{sample}.txt"
    conda:
        "htseq.yaml"
    shell:
        ".snakemake/conda/*/bin/htseq-count -n {core} --format=bam --stranded=no '
        --type=gene --order=pos --idattr=ID {input.bam} {input.gff3} > {output}"
```

In *htseq.yaml* file insert following:

```
channels:
  - bioconda
  - conda-forge
dependencies:
  - samtools =1.1
  - htseq
```

Then run command in your working directory *rnaseq:* snakemake --snakefile Snakefile --core 10 --use_conda

Output:

Sample Data file: metaData.csv

The output is inside the folder *htseq_count*. Count Matrix: *merged_counts.csv*

You can edit this metadata file accrording to your study design. After that you can use differential analysis statistical packages like **DESeq2**, **edgeR**, **Limma** from Bioconductor. For DESeq2 method use