

## **Full reproducible codes**

### **1) Create directory 'data', 'src', 'results' in a CA2 folder**

```
(base) weiqi@LSLab5-22:~$ cd Desktop
(base) weiqi@LSLab5-22:~/Desktop$ cd CA2
(base) weiqi@LSLab5-22:~/Desktop/CA2$ mkdir data src results
```

### **2) Download relevant packages**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install curl
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install fastqc
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install trimmomatic
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install picard-tools
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install bowtie bowtie2
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install samtools
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install libbz2-dev
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install zlib1g-dev
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install libncurses5-dev
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install libncursesw5-dev
(base) weiqi@LSLab5-22:~/Desktop/CA2$ sudo apt-get install liblzma-dev
(base) weiqi@LSLab5-22:~/Desktop/CA2$ wget
https://github.com/samtools/samtools/releases/download/1.9/(base) weiqi@LSLab5-
22:~/Desktop/CA2$ samtools-1.9.tar.bz2
(base) weiqi@LSLab5-22:~/Desktop/CA2$ tar -xjf samtools-1.9.tar.bz2
(base) weiqi@LSLab5-22:~/Desktop/CA2$ cd samtools-1.9
(base) weiqi@LSLab5-22:~/Desktop/CA2/samtools-1.9$ make
(base) weiqi@LSLab5-22:~/Desktop/CA2/samtools-1.9$ sudo make install
(base) weiqi@LSLab5-22:~/Desktop/CA2/samtools-1.9$ sudo apt-get install bedtools
```

### **3) Download mutated yeast clone (A0158131M.tgz) from nBox and put it in data folder**

### **4) Prepare data by extracting out all the files from the "tarball"**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2/samtools-1.9$ cd ../data
(base) weiqi@LSLab5-22:~/Desktop/CA2/data$ tar -zxvf A0158131M.tgz
```

### **5) Quality control - Assessing quality using FastQC**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2/data$ cd ../results
(base) weiqi@LSLab5-22:~/Desktop/CA2/results$ mkdir fastqc
(base) weiqi@LSLab5-22:~/Desktop/CA2/results$ cd ../
(base) weiqi@LSLab5-22:~/Desktop/CA2$ fastqc data/*.fq -o results/fastqc
(base) weiqi@LSLab5-22:~/Desktop/CA2$ cd results/fastqc
(base) weiqi@LSLab5-22:~/Desktop/CA2/results/fastqc$ for filename in *.zip; do unzip $filename;
done
(base) weiqi@LSLab5-22:~/Desktop/CA2/results/fastqc$ cd ../..
(base) weiqi@LSLab5-22:~/Desktop/CA2$ mkdir -p docs
(base) weiqi@LSLab5-22:~/Desktop/CA2$ cat results/fastqc/*/summary.txt >
docs/fastq_summaries.txt
```

#### **6) check for any failed category: No failed results**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ grep FAIL docs/fastq_summaries.txt
```

#### **7) Assess HTML files and decided not to trim as all categories passed.**

#### **8) Align reads to reference genome - prepare by downloading yeast genome (sacCer.fa) and transposon (ty5\_6p.fa) from IVLE, and put it in data folder**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ mkdir combined
(base) weiqi@LSLab5-22:~/Desktop/CA2$ mkdir -p results/sam results/bam
(base) weiqi@LSLab5-22:~/Desktop/CA2$ bowtie2-build 'data/sacCer3.fa,data/ty5_6p.fa'
combine.indexing
```

#### **9) Put combine.indexing into a new folder called "combine"**

#### **10) Next, prepare 3 bash(.sh) scripts and put them in "src" folder. The scripts can be found in the github repository too.**

#### **11) Align paired-ends to reference genome**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ bash src/run-bowtie2.sh
```

#### **12) Convert SAM file format to BAM file format**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ bash src/convert-to-bam.sh
```

#### **13) Sort BAM file by coordinates**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ bash src/sort-bam-files.sh
```

#### **14) Learn more about the BAM file**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ samtools flagstat results/bam/data-sorted.bam
```

**15) Using SAM flags to filter to get improperly paired, and read and mates mapped**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ samtools view -h -f 1 -F 14 results/bam/data-sorted.bam | grep -e 'TY5' -e '^@' > data-sorted.sam
```

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ head data-sorted.sam
```

**16) Make it into a .txt file so as to open on excel sheet to further filter and narrow down the regions to find the paired-ends and insertions on IGV later on**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ samtools view -h -f 1 -F 14 results/bam/data-sorted.bam | grep -e 'TY5' -e '^@' > data-sorted.txt
```

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ samtools view -S -b data-sorted.sam > data-sorted.bam
```

**17) Index BAM file to allow visualisation of alignment on IGV**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ samtools index results/bam/data-sorted.bam
```

**18) Check the number of reads; 150**

```
(base) weiqi@LSLab5-22:~/Desktop/CA2$ samtools view data-sorted.bam | wc -l
```