

# Day 4 Worksheet – Read mapping and visualization

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1. Make sure you have the following files in your `.../day4/trimmomatic/` directory:

```
26M Jul 13 14:31 chr21Eric_repA.RNA.end1.trimmed.fastq
0 Jul 13 14:31 chr21Eric_repA.RNA.end1.unpaired.fastq
26M Jul 13 14:31 chr21Eric_repA.RNA.end2.trimmed.fastq
0 Jul 13 14:31 chr21Eric_repA.RNA.end2.unpaired.fastq
```

2. Create new directory (`mkdir`) named **hisat2**, under `.../day4/`, for the output directory for mapped reads.

```
[arer2562@ip-172-31-29-36 day4]$ mkdir hisat2
[arer2562@ip-172-31-29-36 day4]$ ls -lsh
total 16K
4.0K drwxrwxr-x 2 arer2562 arer2562 6.0K Jul 20 13:24 eofiles
4.0K drwxrwxr-x 2 arer2562 arer2562 6.0K Jul 20 16:11 hisat2
4.0K drwxrwxr-x 2 arer2562 arer2562 6.0K Jul 20 14:54 scripts
4.0K drwxrwxr-x 2 arer2562 arer2562 6.0K Jul 20 13:24 trimmomatic
[arer2562@ip-172-31-29-36 day4]$
```

3. Wget the `d4_mapping` script from the `sread2023` github to your scripts directory.

Start by navigating to github day4 directory at: <https://github.com/Dowell-Lab/sr2023/tree/main/day04/scripts>

Once there, copy link address for desired script

The screenshot shows a GitHub repository page for 'sr2023 / day04 / scripts'. A table lists files and their commit messages. A context menu is open over the 'd4\_mapping.sbatch' file, showing options like 'Open link in new tab', 'Open link in new window', 'Open link in incognito window', 'Save link as...', and 'Copy link address'.

Name	Last commit message	Last commit date
..		
d4_fastqc.sbatch	Updated day4 scripts	3 days ago
d4_mapping.sbatch	Added Day4 scripts	3 days ago
d4_trim_qc.sbatch	Updated d4 script	3 days ago
ext...	Added resources	3 days ago

If successful screen should look like this:

```
[arer2562@ip-172-31-29-36 scripts]$ wget https://github.com/Dowell-Lab/sr2023/tree/main/day04/scripts
--2023-07-20 16:19:00-- https://github.com/Dowell-Lab/sr2023/tree/main/day04/scripts
Resolving github.com (github.com)... 140.82.114.3
Connecting to github.com (github.com)[140.82.114.3]:443... connected.
HTTP request sent, awaiting response... 200 OK
Length: 2864 (2.8K) [text/plain]
Saving to: 'scripts'

100%[=====] 2,864 --.-K/s in 0s

2023-07-20 16:19:00 (66.5 MB/s) - 'scripts' saved [2864/2864]

[arer2562@ip-172-31-29-36 scripts]$ ls
ALE_d4_trim_qc.sbatch d4_mapping.sbatch d4_trim_qc.sbatch scripts
```

4. Edit the new “d4\_mapping.sbatch” using the text editor **vim**. First, edit the SBATCH configuration to meet the needs of read mapping:
- Change the name of the job to something more useful, such as “hisat2\_mapping”.
  - Replace <EMAIL> with your own email address to which you want to receive any notifications.
  - Replace <USERNAME> with your own username to complete the path directory to where to store the error and output files.
  - Complete the following fields: nnodes, ntasks, mem and time. Hisat2 can use multiple processors per input file. So, 1 node, 8 tasks/processors/CPU, 2 Gb for memory and 90 minutes for wall-time should be enough.

```
#!/bin/bash
#SBATCH --job-name=d4_mapping           # Job name
#SBATCH --mail-type=ALL                 # Mail events (NONE, BEGIN, END)
#SBATCH --mail-user=<YOUR_EMAIL_HERE>    # Where to send mail
#SBATCH --nodes=1                       # Number of nodes requested
#SBATCH --ntasks=8                     # Number of CPUs (processors)
#SBATCH --mem=2gb                       # Memory limit
#SBATCH --time=01:30:00                 # Time limit hrs:min:sec
#SBATCH --partition=short                # Partition/queue requested
#SBATCH --output=/scratch/Users/<USERNAME>/day4/eofiles/%x.%j.out
#                                       # Output file (replace %x with job_name and %j by the job id)
#SBATCH --error=/scratch/Users/<USERNAME>/day4/eofiles/%x.%j.err
```

5. Next, assign path variables. In this case, we will specify two directories, both under **DATADIR**. **TRIM** stores the directory path to trimmed reads. **HISAT2** stores the directory path to output mapped reads.

```
### Assigns path variables

DATADIR=/scratch/Users/<USERNAME>/day4/
HISAT2=${DATADIR}/hisat2
TRIM=${DATADIR}/trimmomatic
```

6. Next, load the modules/software needed for mapping reads and file conversion:

```
### Loads modules
module load hisat2/2.1.0
module load samtools/1.8
```

7. And finally, specify the read mapping and file conversion commands. Note that you could instead break up the command onto many lines using the character “\” at the end of every line. These \ characters are ignored by the computer, but will help you identify each part of the command more easily:

**NOTE:** The genome index is located at  
**/scratch/Shares/public/genomes/hisatfiles/hg38/HISAT2/genome**

```
##### Software Specifics #####
## Map trimmed reads to reference genome
hisat2 --very-fast -x /scratch/Shares/public/genomes/hisatfiles/hg38/HISAT2/genome \
-1 ${TRIM}/chr21Eric_repA.RNA.end1.trimmed.fastq \
-2 ${TRIM}/chr21Eric_repA.RNA.edn2.trimmed.fastq \
> ${HISAT2}/chr21Eric_repA.RNA.sam \
2> ${HISAT2}/chr21Eric_repA.hisat2_mapstats.txt

## Convert mapped reads to sorted bam file
### Convert SAM to BAM
samtools view -@ 8 -bS -o ${HISAT2}/chr21Eric_repA.RNA.bam \
> ${HISAT2}/chr21Eric_repA.RNA.sorted.sam

### sort bam file
samtools sort -@ 8 ${HISAT2}/chr21Eric_repA.RNA.bam \
> ${HISAT2}/chr21Eric_repA.RNA.sorted.bam

### index sorted bam file
samtools index ${HISAT2}/chr21Eric_repA.RNA.bam \
${HISAT2}/chr21Eric_repA.RNA.sorted.bam.bai
```

8. Before you close vim, make sure to save your edits by press Esc button to exit insertion mode, then type in **:wq** to save and quit vim.

9. Now that the job script is complete, submit the job by type in **sbatch** command. While waiting for the job to execute, you can check the job status using the command **squeue -u <USERNAME>**:

```
-bash-4.2$ sbatch mapping.sbatch
Submitted batch job 7730124
-bash-4.2$ squeue -u qiya9811
```

JOBID	PARTITION	NAME	USER	ST	TIME	NODES	NODELIST(REASON)
7730124	short	hisat2_m	qiya9811	R	0:07	1	fjinode-12

10. Finally, check the output directory **.../day4/hisat2/** - there should be 5 different files:

```
-bash-4.2$ ls -lsh
total 172M
1.0K -rw-rw-r-- 1 qiya9811 dowelldegrp 613 Jul 13 16:33 chr21Eric_repA.hisat2_maptstats.txt
24M -rw-rw-r-- 1 qiya9811 dowelldegrp 24M Jul 13 16:33 chr21Eric_repA.RNA.bam
127M -rw-rw-r-- 1 qiya9811 dowelldegrp 127M Jul 13 16:33 chr21Eric_repA.RNA.sam
19M -rw-rw-r-- 1 qiya9811 dowelldegrp 19M Jul 13 16:33 chr21Eric_repA.RNA.sorted.bam
1.7M -rw-rw-r-- 1 qiya9811 dowelldegrp 1.7M Jul 13 16:33 chr21Eric_repA.RNA.sorted.bam.bai
```

11. To visualize the mapped reads using IGV, you will need to transfer the sorted.bam and sorted.bam.bai files to your local machine. **rsync** the files from the AWS using a terminal on your local machine. Note that here, I've navigated to the directory for my desktop before rsyncing (Windows machine).

```
lsanford@DESKTOP-3GP5MRN:/mnt/c/Users/lsanford/Desktop$ rsync lynn-sanford@3.136.149.251:
/scratch/Users/lynn-sanford/day4/hisat2/chr21Eric_repA.RNA.sorted* ./
```

