## EpiSyStem MS & Single Cell Workshop: Hands-On Tutorial

#### Objectives:

- Inspect the read quality
- Map reads on a reference genome
- Compare coverage files
- Call enriched regions or peaks
- Infer cell-type specific histone modification levels in the bone marrow

Estimated time required: approximately 3 hours.

#### Take home messages:

- Sequence affinity of PA-MNAse can be seen in the fastq files.
- Although scChIC-seq data is sparse at the single cell level, there is enough
  information to cluster cells into pseudobulk samples that reflect cell-type
  specific chromatin structures.
- We can define cell-type specific genomic regions by finding enriched signal along genomic regions.

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The document has been adopted from the galaxy training material

Software needed (hopefully already installed):

- fastqc
- samtools
- deeptools
- bwa
- hiddenDomains
- macs2
- R
- perl
- python

Software and computing environments needed:

- CoCalc computing environment (please see cocalc\_manual.pdf for instructions found on https://github.com/avolab/episystem-workshop)
- IGV genome browser (http://software.broadinstitute.org/software/igv/download)

## Introduction

Within a cell nucleus, the DNA is tightly-packed and the chromatin is spatially distributed with different levels and scales of organizations. At the smallest scale, DNA is packaged into units called nucleosomes, made of eight histone proteins.

Transcription factors (TFs) in concert with histone modifications shape the chromatin landscape of the genome, and thus regulate cell types and cell states. Histone modifications form an adaptable epigenetic regulatory layer that mediate dynamic transcriptional programs. Functional genomics assays, the most popular involving chromatin immunoprecipitation (ChIP), have revealed active and repressive chromatin structures in bulk tissues. However, inefficiencies of ChIP hinder its application in single cells, preventing genome-wide analysis of histone modifications along the continuum of cellular phenotypes. Therefore, how chromatin landscapes change between repressed, poised, and active states during development and homeostasis is relatively unexplored at the single-cell level.

Binding certain proteins to each of the eight histone proteins may modify the chromatin structure and may result in changes in transcription level. For example, the H3K4me3 is adding 3 methyl-group of the 4th Lysine in the histone 3 amino-acid. This modification is known to activate the transcription on nearby genes by opening the chromatin.

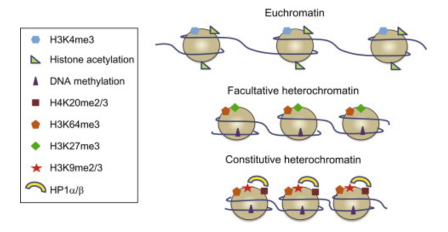


Figure 1: Fadloun et al, 2013

In the upcoming tutorial, we will look at the activator marks H3K4me1 and H3K4me3 scChIC-seq data from mouse bone marrow. We have already performed a dimensionality reduction (using a method called Latent Dirichlet Allocation) on the scChIC-seq in order to cluster cells (using a method called Louvain community detection) with similar histone modification profiles. The cell-cell relationships calculated from this analysis can be visualized in a 2-dimensional plot (see Figure 2, Figure 3)

## H3K4me1

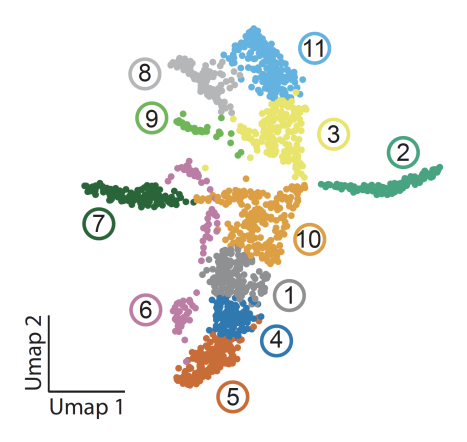


Figure 2: H3K4me1 UMAP (Uniform Manifold Approximation and Projection) plot of cell-cell relationships. Colors and labels show different clusters of cells, inferred from Louvain algorithm.

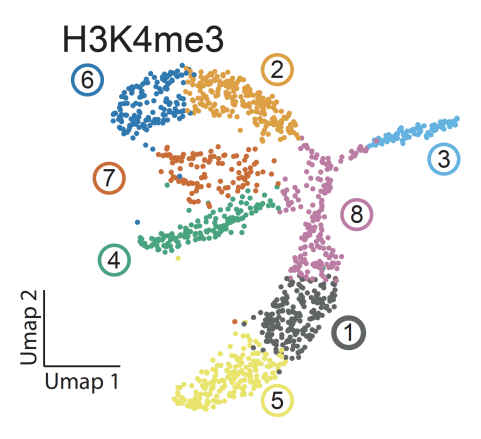


Figure 3: H3K4me3 UMAP plot of cell-cell relationships. Colors and labels show different clusters of cells, inferred from Louvain algorithm.

We will use this pre-defined clustering to explore scChIC-seq data. We suspect that the differences across cells could be coming from distinct cell types. In this exercise, we will focus on two clusters for each histone mark: for H3K4me1 clusters 2 and 5; for H3K4me3 clusters 3 and 5. We have already prepared the scChIC-seq data for you such that the individual cells are grouped into three clusters. Your job is to infer which cluster corresponds to which cell type.

## Step 0: Checking the files

#### File formats

Below are the file-formats you will have to deal with in a typical "Seq" data analysis.

#### sequenced DNA (reads) GGTTCAAAGCAGTAT **FASTQ** GATCAAATAGTAAATCCATTTGTTCAACTCA read alignment DNA sequence + r1 163 chr1 7 30 8M2I4M1D3M = 37 39 TTAGATGAT r2 0 chr1 9 30 3S6M1P1I4M \* 0 0 AAAAGGATA \* SAM <--> BAM genomic counting reads, localization normalizing for sequencing read numbers per depth etc. bedGraph <-> bigWig genomic region Peak calling Specific genomic Chr1 10 20 region1 1.5 + Chr1 30 50 region2 1.7 + Chr1 60 70 region3 2.0 -Chr1 80 95 region4 1.8 coordinates with Bed strand/score/ other information DOWNSTREAM ANALYSES

## File formats during typical analysis of sequencing data

Figure 4:

## Overview of files provided in this tutorial:

All file paths are relative to the data directory: \$HOME/Handouts/EpiSyStem\_Workshop\_Files Remember you can navigate around the directories using cd and explore the files present in each direction using 1s.

NOTE: \${variable} represents a bash variable

- fastq\_raw/raw\_fastq\_R\${read}.fastq: raw fastq files to see what raw data looks like before demultiplexing. You already had a look at these in the previous tutorial. \${read} denotes a bash variable that in this case can be equal to either 1 or 2.
- fastq\_full/demultiplexedR\${read}\_10000rows.fastq.gz : demultiplexed fastq files for quality control checks before mapping. Again, \${read} is a bash variable equal to 1 or 2. Note that this fastq files are zipped.
- references/Mus\_musculus.GRCm38.dna\_rm.primary\_assembly.fa: reference genome fasta file.
- sorted\_bams\_filtered/\${hmark}\_cluster\_\${clstrID}.filtered.bam : single cell scChIC-seq profiles grouped by clusters. Here, \${hmark} is either H3K4me1 or H3K4me3; and \${clstrID} will be either 2, 5 or 11 for H3K4me1, and 3, 5 or 6 for H3K4me3. We have already assigned cells to clusters for you, you just have to infer the biological meaning of these clusters (i.e., infer the cell type). We will use these to visualize bam files with IGV, explore how to calculate number of reads by MAPQ quality, and do peak calling. bam files are subset to include only four main regions in order to reduce file size (defined in regions/regions\_to\_filter.txt).
- regions/regions\_to\_filter.txt: File containing the four genomic regions that contain signal in the bam files.
- sorted\_bigwigs/\${hmark}\_cluster\_\${clstrID}.bw: bigwig files of scChIC-seq profiles, providing genome-wide coverage of scChIC-seq grouped by their clusters (pseudobulk). We will use bigwig files to correlate across pseudobulk samples and visualize the on the IGV.
- chromsizes/chromsizes.\${genome}.filt.txt: size of genomes which are used as input in hiddenDomains.

# Step 1: Quality control and treatment of the sequences

Let's first assess the quality of the demultiplexed fastq files. Demultiplexed means UMI and cell barcodes have been clipped from the sequences in the original fastq file and placed in the header.

#### Hands-on: First look at the fastq files

- 1. Using the terminal, go to the directory Handouts/EpiSyStem\_Workshop\_Files/fastq\_full
- 2. Inspect the fastq files by using the less command. What differences do you detect when comparing the reads in these files to the reads in the fastq files available in the fastq\_raw directory? Hint you can add a -N flag (i.e., less -N) in order to show the line number explicitly.
- 3. Run the command head on any of the two files. What is the output? Why do you think this does not work?

4. In order to explore zipped files, the head/tail/cat commands are not good enough. For this reason, there is the command zcat which has the same function as cat but only works with zipped files. Try it out by running the command zcat demultiplexedR1\_10000rows.fastq.gz.

During sequencing, errors are introduced, such as incorrect nucleotides being called. These are due to the technical limitations of each sequencing platform. Sequencing errors might bias the analysis and can lead to a misinterpretation of the data.

#### Hands-on: Quality control

- 0. Rename the fastq files to H3K4me3\_demultiplexedR1\_10000rows.fastq.gz and H3K4me3\_demultiplexedR1\_10000rows.fastq.gz.
- 1. Run FastQC on the fastq files
- fastqc H3K4me3\_demultiplexedR1\_10000rows.fastq.gz and fastqc H3K4me3\_demultiplexedR2\_10000rows.fastq.gz
- 2. Inspect the generated HTML files. To do so, run the command open demultiplexedR1\_10000rows\_fastqc.html and Enable the view.
- 3. Go back to the terminal by selecting with your mouse the corresponding tab in the top of your screen, and run the command open demultiplexedR2\_10000rows\_fastqc.html. Again by clicking with the mouse in the corresponding tab, you will be available to explore the contents of each HTML file.

#### Questions

- 1. How is the quality of the read 1 and read 2? Why do the lengths of the reads differ? Hint: Remember that the fastq files have been demultiplexed.
- 2. What are the most common start sequences for the fastq files? How does it differ for R1 and R2?
- 3. Have the adapters been trimmed in the demultiplexed fastq files?

## Step 2: Mapping of the reads

We obtain sequences corresponding to a portion of DNA linked to the histone mark of interest (e.g., H3K4me1, H3K4me3). The output file contains a list of sequences that need to be mapped to the genome for further analysis. After mapping, we can then ask whether there are cell-type specific differences in different genomic regions.

## Running BWA

Mapping requires a database of the genome to which you are mapping. These files often are downloaded from publicly available genome browsers such as Ensembl or UCSC Genome Browser. We have already downloaded the mouse genome (genome assembly mm10) and created an index used for mapping. The index file is used often in mapping programs to allow fast and efficient access to the large genome.

bwa is a widely used software that allows to map reads in a fastq file to their most likely position in a reference genome or reference transcriptome. This software can be used as a command in the terminal. If you type bwa alone in the terminal you will see a list of options that you can use to run it and perform the mapping. In this tutorial we will use the option mem. Now, if you type bwa mem in the terminal, a new list of options will be displayed on screen.

#### Hands-on: Mapping

 Run bwa on the fastq files, using an mm10 reference genome (the forward slash \ is only used to split a long single line into multiple lines, so the long command can be printed on a PDF page).

bwa mem ../references/Mus\_musculus.GRCm38.dna\_rm.primary\_assembly.fa  $\$  demultiplexedR1\_10000rows.fastq.gz demultiplexedR2\_10000rows.fastq.gz  $\$  -o demux\_map.sam

The output file containing all the mapping information is a sam file (in the example above, we gave it the name demux\_map.sam, but you can change it an give it another name). An explanation of the SAM format can be found on wikipedia.

2. Inspect the reference file, which can be found in:

Handouts/EpiSyStem\_Workshop\_Files/references with the name Mus\_musculus.GRCm38.dna\_rm.primary\_assembly.fa.

There is another command line, grep, which is very useful to find string patterns in your files. Using the cd command, go to the Handouts/EpiSyStem\_Workshop\_Files/references directory, and type there:

grep '>' Mus\_musculus.GRCm38.dna\_rm.primary\_assembly.fa

This command line prints on your screen only the lines of the file Mus\_musculus.GRCm38.dna\_rm.primary\_assembly.fa that contain the character >. What are these lines?

3. Inspect the mapping stats. Go back to the folder where you created your sam file (Handouts/EpiSyStem\_Workshop\_Files/fastq\_full)

and explore your sam file with the commands that we learned so far (less, head, tail, cat,...).

#### Questions

- How many lines does the sam file have?
- Have a look at the first 67 lines of your sam file. What do you see?
- How do we see the length of each chromosome used in the mapping?

Find all the lines in the sam file containing mapping information for the read with name (long line is split into multiple lines with  $\$ ):

```
Is:NS500414;RN:518;Fc:H2GV2BGX9;La:1;Ti:11101;CX:23815;CY:1073;\
Fi:N;CN:0;aa:CACTCA;aA:CACTCA;aI:32;\
LY:PZ-BM-m1-H3K4me1-2_H2GV2BGX9_S11;RX:CCT;RQ:GGG;BI:175;bc:\
TGCTAATG;BC:TGCTAATG;QT:GGKKKKKK;MX:NLAIII384C8U3
```

- What is each line? Help: visit the wikipedia page https://en.wikipedia.org/wiki/SAM\_%28file\_format%29
- Where has this read been mapped?
- Which is the quality of the mapping?

Now have a look at the mapping information for read:

```
Is:NS500414;RN:518;Fc:H2GV2BGX9;La:1;Ti:11101;CX:23241;\
CY:4823;Fi:N;CN:0;aa:CACTCA;aA:CACTCA;aI:32;\
LY:PZ-BM-m1-H3K4me1-2_H2GV2BGX9_S11;RX:ACA;RQ:GGG;BI:175;\
bc:TGCTAATG;BC:TGCTAATG;QT:GGKKKKKK;MX:NLAIII384C8U3
```

• Is it mapped? Where? What is the quality of the mapping?

#### **SAMTOOLS**

The sam file is so important that is has a command line on its own to speed up its inspection. This is samtools. First, let's type samtools in the terminal with no arguments: a summary of all the options will be displayed.

We will first inspect the flagstat option. Type in the terminal:

```
samtools flagstat demux_map.sam
```

#### Questions

- What is the output telling us?
- How many pair reads are mapped to the same chromosome in demux\_map.sam?

- How many reads are not mapped at all?
- In how many pair reads only one of the reads is mapped?

We will now inspect the view option. What happens when you type the following in the terminal?

```
samtools view demux_map.sam
```

Importantly, commands in the terminal can be concatenated (or pipped). For that, the character | is used. For instance, type to following examples in the terminal. What are we getting in each case?

```
samtools view demux_map.sam | wc
samtools view demux_map.sam | head
samtools view demux_map.sam | grep 'BC:CGTAGTGC'
```

If you now type:

```
samtools view
```

a full list of new options is displayed. Answer the following questions using samtools view with the different options. You might need to pipe some commands to get the answer.

#### Questions

- How many reads do have the flag 99?
- How many reads do not have the flag 99?
- How many reads do have a mapping quality equal or higher than 60?
- List the first 5 reads that have a mapping quality equal or higher than 60 with a flag equal to 163.
- Print the SAM header only.
- How many reads with a mapping quality equal or higher than 60 do have the string BC:CGTAGTGC in their name (BC stands for cell barcode)?

#### From SAM to BAM format

Once you get used, the sam format is very convenient to explore how your sequencing data maps to the reference. However, the files become very big very fast. For this reason, the bam file was invented, which contains the same information as the sam file but in a compress manner. To convert your sam file into a bam file, type in the terminal:

```
samtools view -Sb demux_map.sam > demux_max.bam
```

, and you will see that the new bam file has been created (use 1s to check). Commands such as cat, head, tail will not work on the bam file, but they do

in combination with samtools. For instance, compare the outputs of the two following command lines:

```
head demux_max.bam
samtools view demux_max.bam | head
```

## Correlation between samples

We will compare genome-wide correlation of H3K4me3 and H3K4me1 for different cell clusters.

To compute the correlation between the samples we are going to to use the QC modules of <code>deepTools</code> (http://deeptools.readthedocs.io/), a software package for the QC (quality check), processing and analysis of NGS data. Before computing the correlation a time consuming step is required, which is to compute the read coverage (number of unique reads mapped at a given nucleotide) over a large number of regions from each of the inputed BAM files. For this we will use the tool <code>multiBamSummary</code> . Then, we use <code>plotCorrelation</code> from <code>deepTools</code> to compute and visualize the sample correlation. This is a fast process that allows to quickly try different correlation methods and visual outputs.

In this tutorial we are interested in assessing H3K4me3 and H3K4me1 scChIC-seq samples. To save time, we have already converted the bam to bigwig files, which contains the read coverage.

#### Hands-on: Correlation between samples

- 1. Find the bigwig files (wit extension .bw) in the EpiSyStem\_Workshop\_Files directory. You should find two files for each chromatin modification (H3K4me1 clusters 2 and 5; for H3K4me3 clusters 3 and 5).
- 2. Compare all bigwigs using multiBigwigSummary. You can type multiBigwigSummary --help in the terminal to see all the options. We will use the following options:
- (Choose computation mode) bins
- (Bin size in bp) -bs 100000
- (Input bigwig files): the four imported -b <bigwig> files
- (Output file): -o results.npz

Using these parameters, the tool will take bins of 100000 bp. For each bin the overlapping reads in each sample will be computed and stored into a matrix.

3. Check results using plotCorrelation. Remember that you can type plotCorrelation --help in the terminal to see the list of options. Use to following parameters to explore the file results.npz:

- "Correlation method": Pearson or Spearman (which do you think is more appropriate here? Check by inspecting the scatterplots)
- Plot heatmap or scatterplot.
- In the scatter plot, you can plot output in log scale (-log1p) for visualization. What happens if you do not use this option?

You can visualize the output by using the command open in the terminal. To go back to the terminal, select the corresponding tab in the top of your screen with the mouse.

#### Questions

 From the correlation plot, can you infer which clusters correspond to the same cell type in H3K4me1 and H3K4me3?

## Step 4: Exploring bam files on the IGV browser

Now, go to the directory EpiSyStem\_Workshop\_Files/sorted\_bams\_filtered, which contains a 6 different bam files. These files have been prepared by the instructors and only contain reads falling in specific genomic regions, in order to reduce the file size. For each modification, we have clustered single cells into three separate bam files, associated with one of three cell types: erythroblast, granulocytes, and B-cells. Your job is to explore which bam file is associated with which cell type by looking at cell-type specific regions in the genome browser.

Download the .bam and .bam.bai files onto your computer (.bam.bai files are indexes used by the IGV browser to quickly read the contents of the bam file). In order to download the files, select with your mouse the Files tab, go to the EpiSyStem\_Workshop\_Files/sorted\_bams\_filtered folder, select all the files present and create a zip folder. Next, download the zip folder onto your computer and unzip it.

Open first the 3 bam files belonging to the modification H3K4me1 using IGV, which should already be installed on your computer. Explore the following cell-type specific regions:

chr7	114972165	116898716
chr7	103325741	104425479
chr3	90523036	90870443
chr11	44114099	45269522

#### Questions:

1. Can you infer which clusters correspond to which cell types based on the coverage around the four regions?

Hint: Hbb is a standard marker for erythroblast, S100a8 is a standard marker for granulocytes, and Ebf1 is a standard marker for B cells.

Repeat the same procedure with the modification H3K4me3.

## Step 5: Detecting enriched regions (peak calling)

We could see in the scChIC-seq data some enriched regions that differ across samples. We now would like to call these regions to obtain their coordinates, using hiddenDomains. Perhaps the number of peaks predicted in each region across each sample may be clues to what cell type each sample is.

Go back to CoCalc, and using the terminal go to the directory (in case you are not there):

EpiSyStem\_Workshop\_Files/sorted\_bams\_filtered

#### Hands-on: Peak calling with hiddenDomains

- Calling peaks with hiddenDomains. Remember you can write hiddenDomains by itself in the terminal and a list of options will be shown. You can also find more information on their website (http://hiddendomains.sourceforge.net). For now, we need the following required inputs:
- -g Size of chromosomes for the mouse genome. Can you find this file?
- -q Minimum MAPQ score. A low MAPQ score may include reads that are poor quality, while high MAPQ score keeps only high quality reads. For now, we will keep reads with a quality threshold equal to 60.
- -p A threshold to remove domains called with probabilities less than p. Set to a value such as 0.5. You can play around with this value to see how it changes the output.
- -b minimum length. Use the default 1000 bp.

Run this for all the bam files.

#### Questions

- 1. Which type of files were generated? What do they include?
- 2. How do the peaks called differ between samples? Can you infer cell types based on this analysis?
- 3. Download all the BED files that have been produced by hiddenDomains in your computer and open them

using IGV. For that, you need to go the tab Files in the top of your screen with your mouse, and navigate throught the directories until you find the BED files. Once there, select them, create a compressed folder, and download that folder onto your computer. In your computer, you can unzip the folder and load the files in IGV.

#### Questions

- 1. How do the cell-type specific regions look like?
- 2. How do the peaks called differ between samples? Can you infer cell types based on this analysis?

#### Conclusion

We learned to explore fastq and bam files as well as do calculations on them. We visualized bam files and bigwig files to see how reads are mapped on the genome. Finally, we inferred genomic regions of interest by defining cell-type specific levels of histone marks.