Analysis of ATAC-seq data

Introduction

ATAC-seq (Assay for Transposase-Accessible Chromatin using sequencing) is a technique used to assess chromatin accessibility in a genome-wide manner. Compared with alternatives, such as MNaseq-seq or DNase-seq, ATAC-seq has multiple advantages: it requires significantly less input materials and has much shorter preparation times. In addition, it allows assessing regions of open chromatin and nucleosome positions from the same dataset, whereas the other techniques only allow for one or the other.

The advantages of ATAC-seq are based on the usage of an hyperactive mutant Tn5 transposase. In a process called *tagmentation* (tag + fragmentation), the Tn5 transposases cleaves the DNA double-strand and tags the ends of the cut DNA with sequencing adaptors. This leads to the production of short, double-stranded DNA fragments that have sequencing adaptors on each end. These fragments are then purified, PCR amplified and finally sequenced.

In this module you will learn how prepare ATAC-seq data, going from raw sequencing reads to calling regions of open chromatin. Along the way you will learn how to control the quality of your data and which special considerations are necessary when working with ATAC-seq data.

Practical 1: Pre-processing of reads

The first step in every analysis is controlling the quality of your sequencing data. This will allow you to spot any peculiarities of your data that might give you problems during later analysis steps and allow you to deal with them effectively. In addition, we will trim the sequencing data, removing adapter sequences, low-quality bases and other artefacts that complicate mapping the reads

 Create a new directory for your raw sequencing data and copy/symlink the files there

```
mkdir raw_data
cp <path-to-sequencing-data> raw_data
```

2) To avoid accidentally altering or deleting the raw data, remove write permissions from the files

```
chmod a-w raw data/*
```

For quality control, we will be using FastQC. This simple tool will run a number of checks to give you a quick impression of your data and alert you to any problems your data might have.

1) Make sure that fastqc is accessible from the command line by running:

```
fastqc --version
```

This step is a simple way to test whether a tool is installed and accessible and important for making your analysis replicable: results can often change slightly between different versions of the same tool. Therefore, keeping track of which version you used for your analysis will allow others to replicate your results!

2) To run fastqc on your samples:

```
mkdir -p data/fastqc/untrimmed/
fastqc -t 4 -o data/fastqc/untrimmed/ raw_data/*
```

Using 4 threads in parallel (one file per thread) fastqc will produce quality reports for each file and store them in the data/fastqc/untrimmed/ directory. One file will be an .html file, containing some basic information and plots of the quality metrics. The other will be a .zip archive containing the data underlying the report.

- Open a report in your browser and familiarise yourself with the quality checks.
 - What is the sequence length for your data?
 - Is there a specific trend for the per base sequence quality?
 - What about the per base sequence content?
 - Do you reads have high levels of PCR duplicates?
 - Are the read contaminated with adapter sequences?

To remove any problematic basecalls and other sequencing artefacts, we will run fastp a very-fast trimming tool. This will perform many trimming steps: - Remove low quality reads - Trim low quality base calls from the 3' end of reads - Remove adapter sequences from the 3' end of reads - Trim low-complexity and poly-X basecalls - Correct basecalls of overlapping reads (paired-end only)

4) Run fastp for your reads:

```
mkdir -p data/{trimmed,fastp}
fastp --thread 4 --in1 <raw_data/${SAMPLE}_R1.fastq.gz> --in2 <raw_data/${SAMPLE}_R2.fastq.g
    --out1 <data/trimmed/x_R1.fastq.gz> --out2 <data/trimmed/x_R2.fastq.gz> \
    -3 --cut_tail_window_size 1 --cut_tail_mean_quality 20 \
    --low_complexity_filter --trim_poly_x --correction --detect_adapter_for_pe \
    --json <data/fastp/${SAMPLE}.json> --html <data/fastp/${SAMPLE}.html>
```

5) Control quality after trimming with fastqc:

```
mkdir -p data/fastqc/trimmed/
fastqc -t 4 -o data/fastqc/trimmed/ data/trimmed/*
```

Practical 2: Alignment of trimmed reads to the genome

After trimming, we use Bowtie2 to align the short reads to the genome. While Bowtie2 offers many settings to fine-tune the alignment process, the default options work very well in almost all situations. We only set a few additional options to restrict the alignments that Bowtie2 will output: - Limit the fragment size to 2,000bp - Ignore reads that do not align to the genome - Ignore reads that are not properly paired (only 1 read aligns, or reads that align too far from each other)

1) Align reads with Bowtie2:

The output of Bowtie2 are SAM files, but we prefer to work with BAM files, as they are compressed and therefore take up less space. Converting from SAM to BAM is a straightfoward process during which we can already remove low-quality alignments (this includes multi-mapping reads).

2) Convert from SAM to BAM:

```
samtools\ view\ -@\ 4\ -q\ 10\ -b\ data/alignment/\$\{SAMPLE\},\$\{SAMPLE\}.bam\ data/alignment/\$\{SAMPLE\}, and alignment/\$\{SAMPLE\}, and alignment/\$\{SA
```

Practical 3: Filtering to remove uniformative and artefactual alignments

After aligning the reads and converting them into BAM files we want to remove reads that are either uninformative or are artefacts of the library preparation and sequencing. Uninformative reads are those coming from the mitochondrial genome which is histone-free and therefore always open and usually not interesting to your research. Artefacts that arise during library preparation are PCR duplicates, as these do not give any additional information and actually bias your results towards regions that PCR-amplify more easily. Another class of duplicates are so-called optical duplicates that arise during sequencing, caused by a single cluster of fragments being interpreted as two or more separate clusters. Both types of duplicates appear the same, as alignments with exactly the same sequence, aligning to exactly the same position.

1) Remove mitochondrial alignments by filtering based on the name of the reference sequence:

```
sambamba view --nthreads 4 --filter "ref_name != 'chrM'" --format bam \
     --output-filename data/alignment/${SAMPLE}/${SAMPLE}.noM.bam data/alignment/${SAMPLE}/$-
```

2) Mark PCR and optical duplicates:

```
sambamba markdup --nthreads 4 --show-progress \
   data/alignment/${SAMPLE}/${SAMPLE}.noM.bam data/alignment/${SAMPLE}.dupmarked
```

Lastly, we will remove alignment from so-called "blacklisted regions". These are regions that show extremly high signal independent of the experimental setup. For the most used model organisms (human, mouse, worm, fly) these can easily be downloaded.

3) Remove alignments coming from blacklisted regions

One final step that is sometimes done is to shift the alignments. Due to the mechanisms by which the Tn5 transpose insert the sequencing adaptors the ends of the sequencing fragments are shifted by 9bp. To center the ends of the alignments on the center of the Tn5 binding site, reads need to be shifted by 5/-4bp. But for most analysis this nucleotide resolution is not necessary.

Practical 4: Calling regions of open chromatin (and nucleosomes)

One very common question for ATAC-seq data is to identify regions of open chromatin. This translates to finding regions where our reads pile up. In theory, this process is very similar to calling peaks, for example when analysing CUT&RUN data. Because of this, many people use peak callers originally developed for ChIP-seq data, with minor adaptions to make them work on ATAC-seq data. One often used peak caller is MACS2.

For this practical we will go a different route and use a tool developed specifically for ATAC-seq data, called HMMRATAC. It will take our alignments as input and call not only regions of open chromatin but is also able to detect the flanking nucleosomal regions.

1) Create a list of chromosome sizes (needed for HMMRATAC to work):

There are multiple ways to get this information, e.g. downloading it, or extracting it from Fasta indexes, but this approach does not require any additional data.

2) Run HMMRATAC on your alignments:

```
java -jar HMMRATAC_V1.2.4_exe.jar -b ${SAMPLE}.bam -i ${SAMPLE}.bam.bai -g genome.info
```

Practical 5: Quality control of alignments

With our alignments being filtered, we can now assess some additional quality metrics. This includes things like: - Number of non-duplicate, non-mitochondrial aligned fragments - Alignment rate - Library complexity - Correlation of replicates - Fragment size distribution - TSS enrichment

Whereas the first 4 metrics are useful to check for most sequencing experiments, the last 2 metrics (fragment size distribution and TSS enrichment) are especially important to check.

1) Count the number of sequenced fragments and alignments:

```
# The number of sequenced fragments is equal to the number of lines in the fastq divided by
zcat raw_data/*_1.fastq.gz | wc -1
zcat data/trimmed/*_1.fastq.gz | wc -l
# We can easily count the number of alignments with `sambamba`
# The count directly after alignment
sambamba view -c data/alignment/${SAMPLE}/${SAMPLE}.bam
# The count after removing mitochondrial reads
sambamba view -c data/alignment/${SAMPLE}/${SAMPLE}.noM.bam
# The count after removing duplicates
sambamba view -c -F "not duplicate" data/alignment/${SAMPLE}.dupmarked.bam
# The count after removing alignments from blacklisted regions
sambamba view -c -F "not duplicate" data/alignment/${SAMPLE}/${SAMPLE}.filtered.bam
From the number of alignments we can calculate the library complexity as
the fraction of distinct (non-duplicated) alignments / total alignments. The
alignment rate will be part of the output from Bowtie2.
  2) Estimate the correlation between replicates by calculating the pairwise
```

correlation between samples:

```
multiBamSummary -p 4 bins --samFlagExclude 1024 <bamfiles> -out data/bamsummary.npz mkdir results
```

 $\verb|plotCorrelation -in data/bamsummary.npz --corMethod pearson --skipZeros --whatToPlot scattered from the correlation of the$

There are a range of tools that allow you to easily assess the fragment size distribution and TSS enrichment. For this practical we will use ataqv, which can produce interactive HTML report you can view in your browser.

3) Create JSON reports for each sample with ataqv:

```
mkdir -p data/ataqv

ataqv --peak-file ${PEAKFILE} \
    --name ${SAMPLE} \
     --metrics-file data/ataqv/${SAMPLE}.ataqv.json.gz \
     --tss-file ${TSSFILE} \
     mouse \
     ${SAMPLE}.bam

4) Combine all .json reports into the .html report:
mkarv results/ataqv_report data/ataqv/*
     5) Open the interactive report in your browser:
xdg-open results/ataqv report/index.html
```

Practical 6: Converting alignments into coverage tracks for visualization

With all basic analysis steps done and after having checked the quality of our data we can finally have a look at our data in a genome browser. For this we could use the final BAM files, but this has 2 big drawbacks: - BAM files tend to contain a lot of information not needed for visualising the signal, making the files quite big - Signal strength directly depends on sequencing depth, making comparisons between samples difficult

To solve these problems we can convert our alignments into simpler coverage tracks that only contain information on how strong the signal is at a specific point in the genome. In addition, this allows us to scale the signal based on the sequencing depth, making samples more comparable.

For this step we will use one of Deeptools many useful tools: bamCoverage.

1) Convert your BAM files into bigWig files:

```
mkdir -p data/coverage

bamCoverage -p 4 --binSize 50 \
    --normalizeUsing CPM \
    --ignoreForNormalization chrX chrY \
    --bam ${SAMPLE}.bam -o data/coverage/${SAMPLE}.all.bw
```

This will create coverage tracks of all alignments, summarising the signal into bins of 50bp and normalising it using Counts Per Million (CPM). With this normalisation strategy, a signal of 1 means you would expect to find 1 alignment at this position for every 1 Million alignments in your sample.

But as you remember, depending on the size of the sequencing fragment it contains different information, e.g. whether we have a nucleosome free region (NFR)

or mono-nucleosomes. Luckily bamCoverage has options to filter alignments by size before creating the coverage tracks. With those, we can create coverage tracks for just the NFR or mono-nucleosomal alignments.

2) Create coverage tracks for the NFR (fragment length < 147bp):

```
bamCoverage -p 4 --binSize 50 \
    --normalizeUsing CPM \
    --ignoreForNormalization chrX chrY \
    --extendReads \
    --maxFragmentLength 135 \
    --bam ${SAMPLE}.bam -o data/coverage/${SAMPLE}.nfr.bw
```

3) Create coverage tracks for mono-nucleosomal regions (147bp < fragment length < 294 bp):

```
bamCoverage -p 4 --binSize 50 \
    --normalizeUsing CPM \
    --ignoreForNormalization chrX chrY \
    --extendReads \
    --minFragmentLength 150 \
    --maxFragmentLength 290 \
    --bam ${SAMPLE}.bam -o data/coverage/${SAMPLE}.mono.bw
```

4) Visualize your coverage tracks in the genome browser, together with the regions of open chromatin you called before.