# Predicting Chromatin Conformation From ChIPseq Transcription Factor

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## 1 Introduction

#### The Idea:

ChIPseq could identify not only the direct binding sites of the chromatin protein of interest, but also the indirect ones. This owes to the existence of chromatin conformations or complexes. The formation of such complexes will cluster co-factors in close proximity.

Here, we are interested in the identification of not only the *direct* binding sites but also the *indirect* ones. With such knowledge, we could predict the chromatin conformation of the protein of interest.

#### **Question:**

• Could we cluster the direct and indirect bindings of chromatin proteins from ChIP-seq experiment using Mixture Models (MMs) ?

Using M-Component MM, each cluster is called into each component. Therefore each component corresponds to a cluster of binding sites of the chromatin protein.

## 2 Implementation and Test

```
# export PATH for scripts required
PATH=$PATH:/home/ricky/Rlim/ChromatinConformation/Script;export PATH;
```

For more details information concerning the datasets please contact Nicolas Bertin for NF- $\kappa$ B Zhao dataset and Cebp $\alpha$  with Samuel Collombet

## 2.1 Read Aligns

## NF-κB ChIPseq Transcription factor from Zhao, et al 2014

The read dataset was obtained and aligned using bowtie by Nicolas Bertin (Fullwood Lab). The dataset was in BAM format and sorted accordingly, as follows:

```
ls *.bam | parallel -j 3 'samtools sort \{\} \{.\}.sorted' &
```

Only the sorted BAM files were stored.

## Cebpa ChIPseq in different cell lines from Samuel

The read dataset was obtained and aligned by Samuel. The dataset was downloaded from tblab-server at the following directory:

```
|/DAS/TBlab/Samuel/raw/new/|
```

The dataset was in BAM format and sorted accordingly, as follows:

```
ls *.bam | parallel -j 3 'samtools sort {} {.}.sorted' &
```

Only the sorted BAM files were stored.

## Cebp ChIPseq in from Porse

The read dataset was obtained from NCBI with GEO GSE42321. The download link was stored in:

/home/ricky/Rlim/ChromatinConformation/Input/SRA/Porse/sampleDownloadLinks.txt

The dataset was stored in the following directory:

/home/ricky/Rlim/ChromatinConformation/Input/ReadAligns/Porse

Script/getBamFromSRA -c 2 -g /home/ricky/Rlim/Biotools/Genomes/mm10\_bowtie2\_index/bowtie2\

- -a /home/ricky/Rlim/ChromatinConformation/Input/ReadAligns/Porse/annot.txt
- -i /home/ricky/Rlim/ChromatinConformation/Input/ReadAligns/Porse\
- -o /home/ricky/Rlim/ChromatinConformation/Output/ReadAligns/Porse 2> Log/getBamFromSRA\_Porse.txt

## Cebp ChIPseq in from Benner

The read dataset was obtained from NCBI with GEO GSM537984. The download link was stored in:

/home/ricky/Rlim/ChromatinConformation/Input/FA/Benner/sampleDownloadLinks.txt

The dataset was stored in the following directory:

/home/ricky/Rlim/ChromatinConformation/Input/ReadAligns/Benner

Script/getBamFromFA -c 2 -g /home/ricky/Rlim/Biotools/Genomes/mm10\_bowtie2\_index/bowtie2\

- -a Input/ReadAligns/Benner/annot.txt -i Input/ReadAligns/Benner/
- -o Output/ReadAligns/Benner/ 2> Log/getBamFromFA\_Benner.txt

## 2.2 Quality Check

## NF-κB ChIPseq Transcription factor from Zhao, et al 2014

#### Cebpα ChIPseq in different cell lines from Samuel

```
ls Input/Bam/Samuel/*.bam |\
parallel -j 2 'fastqc -o Output/QC/Samuel {}' &\
```

## Cebp ChIPseq in from Porse

```
ls Input/QC/Porse/*.bam | parallel -j 2 'fastqc -o Output/QC/Porse/ {}'
```

## Cebp ChIPseq in from Benner

```
ls Input/QC/Benner/*.bam | parallel -j 2 'fastqc -o Output/QC/Benner/ {}'
```

#### 2.3 Peak Calls

We used jaHMM for peak callings and macs2 peak caller. jahmm library was downloaded and stored locally at /home/ricky/Rlim/ChromatinConformation/Library/jahmm

## NF-kB ChIPseq Transcription factor from Zhao, et al 2014

```
Script/jahmmPeakCalling.sh -c 4 -g hg19 -r 300\
-i Input/PeakCalls/ZhaoB_etal.CellRep2014/\
-o Output/PeakCalls/Jahmm/ZhaoB_etal.CellRep2014/\
2> Log/jahmmPeakCall_ZhaoB.txt
```

Number of peaks called by jahmm from Zhao datasets are in millions.

```
8599068 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_cREL_Rep1.sorted_peaks.bed 9382822 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_cREL_Rep2.sorted_peaks.bed 6144560 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p50_Rep1.sorted_peaks.bed 832975 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p50_Rep2.sorted_peaks.bed 8622964 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p52_Rep1.sorted_peaks.bed 8627622 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p52_Rep2.sorted_peaks.bed 9384884 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p55_Rep1.sorted_peaks.bed 9363618 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p65_Rep2.sorted_peaks.bed 9398329 300bin-ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_RELB_Rep0.sorted_peaks.bed
```

```
Script/macsPeakCalling.sh -c 4 -g hs\
-i Input/PeakCalls/ZhaoB_etal.CellRep2014/\
-o Output/PeakCalls/ZhaoB_etal.CellRep2014\
2> Log/macsPeakCall_Zhao.txt
```

Number of peaks called by macs2 from Zhao datasets in majority are in millions.

```
1138227 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_cREL_Rep1.sorted_summits_peaks.bed 1045436 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_cREL_Rep2.sorted_summits_peaks.bed 445594 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p50_Rep1.sorted_summits_peaks.bed 465586 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p50_Rep2.sorted_summits_peaks.bed 984604 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p52_Rep1.sorted_summits_peaks.bed 385493 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p52_Rep2.sorted_summits_peaks.bed 3591979 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p65_Rep1.sorted_summits_peaks.bed 3406538 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_p65_Rep2.sorted_summits_peaks.bed 3706260 ZhaoB_etal.CellRep2014.ChIP-Seq_GM12878_RELB_Rep0.sorted_summits_peaks.bed
```

#### Cebp from Samuel

Note that we used macs2 as the dataset from Samuel has not input and jahmm could call peaks without an input sample.

```
Script/macsPeakCalling.sh -c 4 -g hs\
-i Input/PeakCalls/Samuel/\
-o Output/PeakCalls/Macs/Samuel/\
2> Log/macsPeakCall_Samuel.txt
```

## Cebp ChIPseq in from Porse

```
Script/jahmmPeakCalling.sh -c 2 -g mm10 -r 300 -i Input/PeakCalls/Porse/\
-o Output/PeakCalls/Jahmm/Porse/ 2> Log/jahmmPeakCall_Porse.txt
```

## Cebp ChIPseq in from Benner

```
Script/jahmmPeakCalling.sh -c 2 -g mm10 -r 300 -i Input/PeakCalls/Benner/\
-o Output/PeakCalls/Benner/ 2> Log/jahmmPeakCall_Benner.txt
```

## 2.4 Component Calls

## Cebp from Samuel

```
Script/mmComponentCalls.R --model='GMM' --ncomp=3 --oneComp=TRUE\
--input_dir='/home/ricky/Rlim/ChromatinConformation/Input/ComponentCalls/Macs/Samuel/'\
--output_dir='/home/ricky/Rlim/ChromatinConformation/Output/ComponentCalls/Macs/Samuel/'\
2> Log/mmComponentCall_Samuel.txt\
```

## Cebp from Porse

```
Rscript Script/mmComponentCalls.R --model='GMM' --ncomp=5 --oneComp=TRUE\
--input_dir='Input/ComponentCalls/Jahmm/Porse/'\
--output_dir='Output/ComponentCalls/Jahmm/Porse/' 2> Log/mmComponentCall_Porse.txt
```

## Cebp from Benner

```
Script/mmComponentCalls.R --model='GMM' --ncomp=3 --oneComp=TRUE\
--input_dir='Input/ComponentCalls/Jahmm/Benner/'\
--output_dir='Output/ComponentCalls/Jahmm/Benner/' 2> Log/mmComponentCall_Benner.txt
```

## 2.5 BiClustering: Direct and Indirect

## Cebp from Samuel

```
Rscript Script/assignLocalClustering.R --distance=3000 --filter=TRUE\
--input_dir='/home/ricky/Rlim/ChromatinConformation/Input/LocalClusters/Macs/Samuel/'\
--output_dir='/home/ricky/Rlim/ChromatinConformation/Output/LocalClusters/Macs/Samuel/'\
2> Log/assignLocalCluster_Samuel.txt
```

## Cebp from Porse

```
Script/assignLocalClustering.R --distance=3000 --filter=TRUE\
--input_dir='/home/ricky/Rlim/ChromatinConformation/Input/LocalClusters/Jahmm/Porse/'\
--output_dir='/home/ricky/Rlim/ChromatinConformation/Output/LocalClusters/Jahmm/Porse/'
```

## Cebp from Benner

```
Script/assignLocalClustering.R --distance=3000 --filter=TRUE\
--input_dir='/home/ricky/Rlim/ChromatinConformation/Input/LocalClusters/Jahmm/Benner/'\
--output_dir='/home/ricky/Rlim/ChromatinConformation/Output/LocalClusters/Jahmm/Benner/'\
2> Log/assignLocalCluster_Benner.txt
```

## 2.6 Motif Calls from Component Calls

#### Cebp from Samuel

```
# extend bed peak summit from 1 bp resolution to 300 bp bin (150 bp in both strand directions)
ls Input/MotifCalls/Macs/Samuel/ComponentCalls/*.bed |\
parallel -j 2 ``bedtools slop -i {}\
-g /home/ricky/Rlim/Biotools/Genomes/mm10/refGenome/mm10.genome\ -b 150 >{.}_150bp_extended.bed''

# bed to fasta
ls Input/MotifCalls/Macs/Samuel/ComponentCalls/*extended.bed |\
parallel -j 2 ``bedtools getfasta\
-fi /home/ricky/Rlim/Biotools/Genomes/mm10/refGenome/mm10.fa -bed {} -fo {.}.fa''
```

```
ls Input/MotifCalls/Macs/Samuel/ComponentCalls/*.fa |\
parallel -j 2 ``meme-chip\
-db /home/ricky/Rlim/Biotools/motif_databases/JASPAR_CORE_2014_vertebrates.meme
-oc \ /home/ricky/Rlim/ChromatinConformation/Output/MotifCalls/Macs/Samuel/ComponentCalls/\{/.\} \\
-index-name {/.} -meme-mod zoops -meme-minw 4 -meme-maxw 10 -meme-nmotifs 10 {}''
Cebp from Porse
# convert the bed to fasta
ls Input/MotifCalls/Jahmm/Porse/ComponentCalls/*.bed |\
parallel -j 2 "bedtools getfasta\
-fi /home/ricky/Rlim/Biotools/Genomes/mm10/refGenome/mm10.fa -bed {} -fo {.}.fa"
# meme-chip
ls Input/MotifCalls/Jahmm/Porse/ComponentCalls/*.fa |\
parallel -j 2 "meme-chip -db ~/Rlim/Biotools/motif_databases/JASPAR_CORE_2014_vertebrates.meme
-oc Output/MotifCalls/Jahmm/Porse/ComponentCalls{/.}\
-index-name {/.} -meme-mod zoops -meme-minw 4 -meme-maxw 10 -meme-nmotifs 10 {}"
Cebp from Benner
# convert the bed to fasta
ls Input/MotifCalls/Jahmm/Benner/ComponentCalls/*.bed |\
parallel -j 2 "bedtools getfasta\
-fi /home/ricky/Rlim/Biotools/Genomes/mm10/refGenome/mm10.fa -bed {} -fo {.}.fa"
# meme-chip
ls Input/MotifCalls/Jahmm/Benner/ComponentCalls/*.fa |\
parallel -j 2 "meme-chip -db ~/Rlim/Biotools/motif_databases/JASPAR_CORE_2014_vertebrates.meme
-oc Output/MotifCalls/Jahmm/Benner/ComponentCalls/{/.}
-index-name {/.} -meme-mod zoops -meme-minw 4 -meme-maxw 10 -meme-nmotifs 10 {}"
     Motif Calls from Local Clusters
2.7
Cebp from Samuel
# extend bed peak summit from 1 bp resolution to 300 bp bin (150 bp in both strand directions)
ls Input/MotifCalls/Macs/Samuel/LocalClusters/*.bed |\
parallel -j 2 ``bedtools slop -i \{\}\
-g /home/ricky/Rlim/Biotools/Genomes/mm10/refGenome/mm10.genome\ -b 150 >{.}_150bp_extended.bed''
# bed to fasta
ls Input/MotifCalls/Macs/Samuel/LocalClusters/*extended.bed |\
parallel -j 2 ``bedtools getfasta\
-fi /home/ricky/Rlim/Biotools/Genomes/mm10/refGenome/mm10.fa -bed {} -fo {.}.fa''
ls Input/MotifCalls/Macs/Samuel/LocalClusters/*.fa |\
parallel -j 2 ``meme-chip\
-db /home/ricky/Rlim/Biotools/motif_databases/JASPAR_CORE_2014_vertebrates.meme
-oc\ /home/ricky/Rlim/ChromatinConformation/Output/MotifCalls/Macs/Samuel/LocalClusters/\{/.\} \\
-index-name {/.} -meme-mod zoops -meme-minw 4 -meme-maxw 10 -meme-nmotifs 10 {}''
Cebp from Porse
# convert the bed to fasta
ls Input/MotifCalls/Jahmm/Porse/LocalClusters/*.bed |\
```

parallel -j 2 "bedtools getfasta\

```
-fi /home/ricky/Rlim/Biotools/Genomes/mm10/refGenome/mm10.fa -bed {} -fo {.}.fa"
# meme-chip
ls Input/MotifCalls/Jahmm/Porse/LocalClusters/*.fa |\
parallel -j 2 "meme-chip -db ~/Rlim/Biotools/motif_databases/JASPAR_CORE_2014_vertebrates.meme
-oc Output/MotifCalls/Jahmm/Porse/LocalClusters{/.}
-index-name {/.} -meme-mod zoops -meme-minw 4 -meme-maxw 10 -meme-nmotifs 10 {}"
Cebp from Benner
# convert the bed to fasta
ls Input/MotifCalls/Jahmm/Benner/LocalClusters/*.bed |\
parallel -j 2 "bedtools getfasta\
-fi /home/ricky/Rlim/Biotools/Genomes/mm10/refGenome/mm10.fa -bed {} -fo {.}.fa"
# meme-chip
ls Input/MotifCalls/Jahmm/Benner/LocalClusters/*.fa |\
parallel -j 2 "meme-chip -db ~/Rlim/Biotools/motif_databases/JASPAR_CORE_2014_vertebrates.meme
-oc Output/MotifCalls/Jahmm/Benner/LocalClusters/{/.}
-index-name \{/.\} -meme-mod zoops -meme-minw 4 -meme-maxw 10 -meme-nmotifs 10 \{\}"
  Filename: chromatinConformation.Rnw
  Working directory: /home/ricky/Rlim/ChromatinConformation
```

## 3 Metainfo

```
sessionInfo()
## R version 3.2.0 (2015-04-16)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 14.04.1 LTS
## locale:
## [1] LC_CTYPE=en_SG.UTF-8
                                LC_NUMERIC=C
## [3] LC_TIME=en_SG.UTF-8
                                 LC_COLLATE=en_SG.UTF-8
## [5] LC_MONETARY=en_SG.UTF-8
                               LC_MESSAGES=en_SG.UTF-8
## [7] LC_PAPER=en_SG.UTF-8
                                 LC_NAME=C
## [9] LC_ADDRESS=C
                                 LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_SG.UTF-8 LC_IDENTIFICATION=C
## attached base packages:
## [1] stats graphics grDevices utils
                                             datasets methods base
##
## other attached packages:
## [1] knitr_1.10.5
## loaded via a namespace (and not attached):
## [1] magrittr_1.5 formatR_1.2 tools_3.2.0
                                              stringi_0.5-2 highr_0.5
## [6] digest_0.6.8 stringr_1.0.0 evaluate_0.7
```