The chromstaR user's guide

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1 Introduction

ChIP-seq has become the standard technique for assessing the genome-wide chromatin state of DNA. *chromstaR* provides functions for the joint analysis of multiple ChIP-seq samples. It allows peak calling for transcription factor binding and histone modifications with a narrow (e.g. H3K4me3, H3K27ac, ...) or broad (e.g. H3K36me3, H3K27me3, ...) profile. All analysis can be performed on each sample individually (=univariate), or in a joint analysis considering all samples simultaneously (=multivariate).

2 Outline of workflow

Every analysis with the chromstaR package starts from aligned reads in either BAM or BED format. In the first step, the genome is partitioned into non-overlapping, equally sized bins and the reads that fall into each bin are counted. These read counts serve as the basis for both the univariate and the multivariate peak- and broad-region calling. Univariate peak calling is done by fitting a three-state Hidden Markov Model to the binned read counts. Multivariate peak calling for S samples is done by fitting a 2^S -state Hidden Markov Model to all binned read counts.

3 Univariate analysis

3.1 Task 1: Peak calling for a narrow histone modification

Examples of histone modifications with a narrow profile are H3K4me3, H3K9ac and H3K27ac in most human tissues. For such peak-like modifications, the bin size should be set to a value between 200bp and 1000bp.

```
library(chromstaR)
```

```
## === Step 1: Binning ===
# Get an example BED file
bedfile <- system.file("extdata","euratrans","lv-H3K4me3-BN-male-bio2-tech1.bed.gz",</pre>
                        package="chromstaRData")
# Get the chromosome lengths (see ?GenomeInfoDb::fetchExtendedChromInfoFromUCSC)
# This is only necessary for BED files. BAM files are handled automatically.
data(rn4_chrominfo)
head(rn4_chrominfo)
##
    UCSC_seqlevel UCSC_seqlength NCBI_seqlevel
## 1
             chrM
                            16300
                                              MT
## 2
             chr12
                          46782294
                                              12
```

```
## 3 chr20 55268282
## 4
                    59218465
           chr19
                                       19
## 5
           chr18
                      87265094
                                       18
## 6
           chr11
                      87759784
                                       11
# We use bin size 1000bp and chromosome 12 to keep the example quick
binned.data <- binReads(bedfile, assembly=rn4_chrominfo, binsizes=1000,</pre>
                     chromosomes='chr12')
print(binned.data)
## GRanges object with 46782 ranges and 1 metadata column:
                              ranges strand | counts
##
        seqnames
##
             <Rle>
                             <IRanges> <Rle> | <integer>
                         [ 1, 1000]
       [1] chr12
##
                                       * | 0
       [2] chr12
                         [1001, 2000]
                                        * |
       [3] chr12
                         [2001, 3000]
##
                                         * |
                                                   0
       [4] chr12
[5] chr12
                                         * |
##
                         [3001, 4000]
                                                    0
##
                          [4001, 5000]
                                         * |
                                                    0
##
                                        . . . .
##
   [46778] chr12 [46777001, 46778000]
##
    [46779] chr12 [46778001, 46779000]
                                          * |
                                                    1
    [46780] chr12 [46779001, 46780000]
##
                                          * |
                                                    0
             chr12 [46780001, 46781000]
##
    [46781]
                                                     1
             chr12 [46781001, 46782000]
##
    [46782]
                                                    1
##
## seqinfo: 1 sequence from an unspecified genome
```

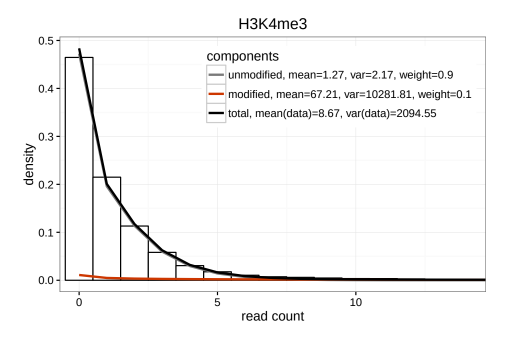
```
## === Step 2: Peak calling ===
# We restrict the peak calling to 60 seconds to keep this example quick.
model <- callPeaksUnivariate(binned.data, max.time=60, verbosity=0)

## Replaced read counts > 500 by 500 in 111 bins. Set option 'read.cutoff=FALSE' to disable
this filtering. This filtering was done to increase the speed of the HMM.

## Calculating states from posteriors ...
## 0.13s

## Making segmentation ...
## 0.14s

## === Step 3: Checking the fit ===
# For a narrow modification, the fit should look something like this,
# with the 'modified'-component near the bottom of the figure
plot(model) + ggtitle('H3K4me3')
```



```
## === Step 4: Export to genome browser ===
# We can export peak calls and binned read counts with
exportUnivariates(list(model), filename=tempfile(), what=c('peaks','counts'))
```

3.2 Task 2: Peak calling for a broad histone modification

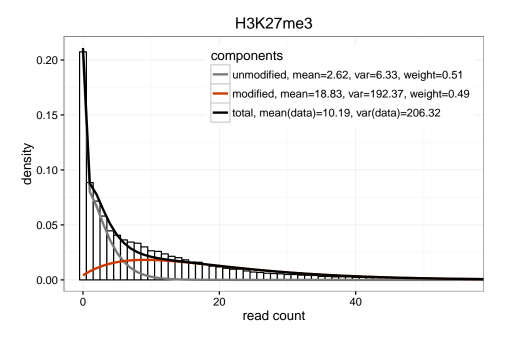
library(chromstaR)

Examples of histone modifications with a broad profile are H3K9me3, H3K27me3, H3K36me3, H4K20me1 in most human tissues. These modifications usually cover broad domains of the genome, and the enrichment is best captured with a bin size between 500bp and 2000bp.

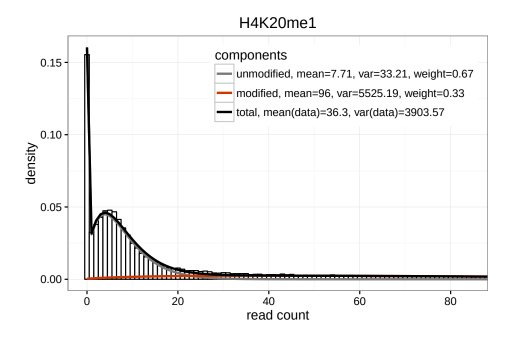
```
## === Step 2: Peak calling ===
# We restrict the peak calling to 60 seconds to keep this example quick.
model <- callPeaksUnivariate(binned.data, max.time=60, verbosity=0)

## Calculating states from posteriors ...
## 0.17s
## Making segmentation ...
## 0.13s</pre>
```

```
## === Step 3: Checking the fit ===
# For a broad modification, the fit should look something like this,
# with a 'modified'-component that fits the thick tail of the distribution.
plot(model) + ggtitle('H3K27me3')
```



```
## === Step 4: Export to genome browser ===
# We can export peak calls and binned read counts with
exportUnivariates(list(model), filename=tempfile(), what=c('peaks','counts'))
```



3.3 Task 3: Peak calling for ATAC-seq, DNase-seq, FAIRE-seq, ...

Peak calling for ATAC-seq and DNase-seq is similar to the peak calling of a narrow histone modification (see section 3.1). FAIRE-seq experiments seem to exhibit a broad profile with our model, so the procedure is similar to the domain calling of a broad histone modification (see section 3.2).

4 Multivariate analysis

4.1 Task 1: Integrating multiple replicates

chromstaR can be used to call peaks with multiple replicates, without the need of prior merging. The underlying statistical model integrates information from all replicates to identify common peaks. It is, however, important to note that replicates with poor quality can affect the joint peak calling negatively. It is therefore recommended to first check the replicate quality and discard poor-quality replicates. The necessary steps for peak calling for an example ChIP-seq experiment with 4 replicates are detailed below.

library(chromstaR)

```
## === Step 1: Binning ===
# Let's get some example data with 3 replicates in spontaneous hypertensive rat (SHR)
file.path <- system.file("extdata","euratrans", package='chromstaRData')</pre>
bedfiles.good <- list.files(file.path, pattern="H3K27me3.*SHR", full.names=TRUE)[1:3]
# We fake a replicate with poor quality by taking a different mark entirely
bedfiles.poor <- list.files(file.path, pattern="H4K2Ome1.*SHR", full.names=TRUE)[1]
bedfiles <- c(bedfiles.good, bedfiles.poor)</pre>
# Obtain chromosome lengths. This is only necessary for BED files. BAM files are
# handled automatically.
data(rn4_chrominfo)
head(rn4_chrominfo)
## UCSC_seqlevel UCSC_seqlength NCBI_seqlevel
## 1
                           16300
             chrM
## 2
                        46782294
                                             12
             chr12
                                            20
## 3
             chr20
                        55268282
                                            19
## 4
                        59218465
            chr19
## 5
           chr18
                        87265094
## 6
                                             11
           chr11
                        87759784
# Define experiment structure
exp <- data.frame(file=bedfiles, mark='H3K27me3', condition='SHR', replicate=1:4,</pre>
                  pairedEndReads=FALSE)
# We use bin size 1000bp and chromosome 12 to keep the example quick
binned.data <- list()</pre>
for (bedfile in bedfiles) {
  binned.data[[basename(bedfile)]] <- binReads(bedfile, binsize=1000,</pre>
                                         assembly=rn4_chrominfo, chromosomes='chr12',
                                         experiment.table=exp)
## === Step 2: Univariate peak calling ===
# The univariate fit is obtained for each replicate
models <- list()</pre>
for (i1 in 1:length(binned.data)) {
  models[[i1]] <- callPeaksUnivariate(binned.data[[i1]], max.time=60)</pre>
## === Step 3: Check replicate correlation ===
\hbox{\it \# We run a multivariate peak calling on all 4 replicates}
# A warning is issued because replicate 4 is very different from the others
multi.model <- callPeaksReplicates(models, max.time=60, eps=1)</pre>
## HMM: number of states = 16
## HMM: number of bins = 46782
## HMM: maximum number of iterations = none
## HMM: maximum running time = 60 sec
## HMM: epsilon = 1
## HMM: number of experiments = 4
                         log(P)
## Iteration
                                               dlog(P)
                                                          Time in sec
##
           0
                             -inf
## HMM: Precomputing densities ...
```

```
log(P)
                                             dlog(P)
## Iteration
                                                        Time in sec
##
          0
                            -inf
                                                                 0
##
           1
                 -548036.299631
                                                 inf
                                                                 0
##
           2
                  -543428.085670
                                         4608.213961
                                                                 1
##
           3
                 -543327.002716
                                         101.082954
                                                                 1
##
           4
                 -543306.772386
                                          20.230330
                                                                 1
##
           5
                  -543300.996034
                                          5.776353
                                           2.040704
                  -543298.955330
##
           6
                                                                 1
##
           7
                  -543298.122400
                                           0.832930
## HMM: Convergence reached!
## HMM: Recoding posteriors ...
## Warning in callPeaksReplicates(models, max.time = 60, eps = 1): Your replicates cluster
in 2 groups. Consider redoing your analysis with only the group with the highest average
coverage:
## H3K27me3-SHR-rep4
## Replicates from groups with lower coverage are:
## H3K27me3-SHR-rep1
## H3K27me3-SHR-rep2
## H3K27me3-SHR-rep3
# Checking the correlation confirms that Rep4 is very different from the others
print(multi.model$replicateInfo$correlation)
                    H3K27me3-SHR-rep1 H3K27me3-SHR-rep2 H3K27me3-SHR-rep3
## H3K27me3-SHR-rep1
                        1.0000000 0.9997432 0.9993580
## H3K27me3-SHR-rep2
                           0.9997432
                                             1.0000000
                                                               0.9994435
## H3K27me3-SHR-rep3
                           0.9993580
                                             0.9994435
                                                               1.0000000
                          -0.3685248
## H3K27me3-SHR-rep4
                                            -0.3683620
                                                              -0.3683201
                 H3K27me3-SHR-rep4
## H3K27me3-SHR-rep1
                          -0.3685248
## H3K27me3-SHR-rep2
                           -0.3683620
## H3K27me3-SHR-rep3
                           -0.3683201
## H3K27me3-SHR-rep4
                           1.0000000
## === Step 4: Peak calling with replicates ===
# We redo the previous step without the "bad" replicate
# Also, we force all replicates to agree in their peak calls
multi.model <- callPeaksReplicates(models[1:3], force.equal=TRUE, max.time=60)</pre>
## === Step 5: Export to genome browser ===
# Finally, we can export the results as BED file
exportMultivariate(multi.model, filename=tempfile(), what=c('peaks','counts'))
```

4.2 Task 2: Detecting differentially modified regions

chromstaR is extremely powerful in detecting differentially modified regions in two or more samples. The following example illustrates this on ChIP-seq data for H4K20me1 in brown norway (BN) and spontaneous hypertensive

rat (SHR) in left-ventricle (lv) heart tissue. The mode of analysis is called *condition*, because all conditions are analyzed simultaneously.

```
library(chromstaR)
#=== Step 1: Preparation ===
## Prepare the file paths. Exchange this with your input and output directories.
inputfolder <- system.file("extdata","euratrans", package="chromstaRData")</pre>
outputfolder <- file.path(tempdir(), 'H4K20me1-example')</pre>
## Define experiment structure
data(experiment_table_H4K20me1)
print(experiment_table_H4K20me1)
##
                                                  mark condition replicate
                                         file
## 19 lv-H4K20me1-BN-male-bio1-tech1.bed.gz H4K20me1 BN 1
## 20 lv-H4K20me1-BN-male-bio2-tech1.bed.gz H4K20me1
                                                              BN
                                                                          2
## 22 lv-H4K20me1-SHR-male-bio1-tech1.bed.gz H4K20me1 SHR ## 22 lv-H4K20me1-SHR-male-bio2-tech1.bed.gz H4K20me1 SHR ## 23 lv-H4K20me1-SHR-male-bio2-tech1.bed.gz H4K20me1 SHR
                                                                          1
## pairedEndReads
         FALSE
## 19
## 20
               FALSE
           FALSE
## 22
## 23
             FALSE
## Define assembly
# This is only necessary if you have BED files, BAM files are handled automatically.
# For common assemblies you can also specify them as 'hq19' for example.
data(rn4_chrominfo)
head(rn4_chrominfo)
## UCSC_seqlevel UCSC_seqlength NCBI_seqlevel
             chrM
## 1
                           16300 MT
## 2
                        46782294
                                             12
             chr12
## 3
           chr20
                        55268282
                                             20
## 4 chr19 59218465
## 5 chr18 87265094
## 6 chr11 87759784
                                              19
                                              18
#=== Step 2: Run Chromstar ===
## Run ChromstaR
Chromstar(inputfolder, experiment.table=experiment_table_H4K20me1,
          outputfolder=outputfolder, numCPU=2, binsize=1000, assembly=rn4_chrominfo,
          prefit.on.chr='chr12', mode='condition')
## Results are stored in 'outputfolder' and can be loaded for further processing
list.files(outputfolder)
## [1] "binned"
                               "browserfiles" "chrominfo.tsv"
                             "experiment_table.tsv" "multivariate"
## [4] "chromstaR.config"
## [7] "plots"
                             "univariate"
model <- get(load(file.path(outputfolder,'multivariate',</pre>
                            'multivariate_mode-condition_mark-H4K20me1.RData')))
```

```
## === Step 3: Construct differential and common states ===
diff.states <- stateBrewer(experiment_table_H4K20me1, mode='condition',</pre>
                         differential.states=TRUE)
print(diff.states)
## combination state
## 1 [SHR] 3
## 2
          [BN]
common.states <- stateBrewer(experiment_table_H4K20me1, mode='condition',</pre>
                            common.states=TRUE)
print(common.states)
## combination state
## 1 [] 0
## 2
       [BN+SHR]
                   15
## === Step 5: Export to genome browser ===
# Export only differential states
exportMultivariate(model, filename=tempfile(),
                  what=c('peaks','counts','combinations'),
```

4.3 Task 3: Finding combinatorial chromatin states

include.states=diff.states)

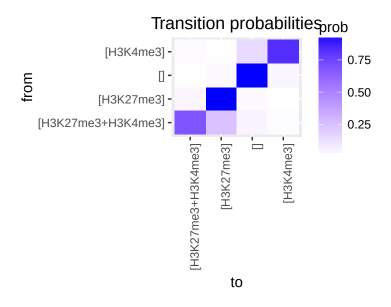
Most experimental studies that probe several histone modifications are interested in combinatorial chromatin states. An example of a simple combinatorial state would be [H3K4me3+H3K27me3], which is also frequently called "bivalent promoter", due to the simultaneous occurrence of the promoter marking H3K4me3 and the repressive H3K27me3. Finding combinatorial states with *chromstaR* is equivalent to a multivariate peak calling. The following code chunks demonstrate how to find bivalent promoters and do some simple analysis. The mode of analysis is called *mark*, because all marks are analyzed simultaneously.

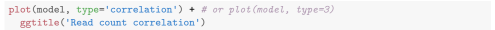
```
library(chromstaR)

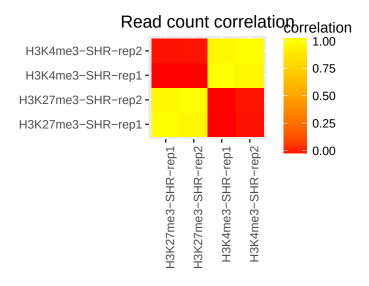
#=== Step 1: Preparation ===
## Prepare the file paths. Exchange this with your input and output directories.
inputfolder <- system.file("extdata","euratrans", package="chromstaRData")
outputfolder <- file.path(tempdir(), 'SHR-example')

## Define experiment structure (SHR = spontaneous hypertensive rat)
data(experiment_table_SHR)
print(experiment_table_SHR)</pre>
```

```
##
                                      file mark condition replicate
## 4 lv-H3K27me3-SHR-male-bio2-tech1.bed.gz H3K27me3 SHR
                                                                   1
## 5 lv-H3K27me3-SHR-male-bio2-tech2.bed.gz H3K27me3
                                                          SHR
                                                                     2
## 16 lv-H3K4me3-SHR-male-bio2-tech1.bed.gz H3K4me3
                                                          SHR
                                                                     1
## 17 lv-H3K4me3-SHR-male-bio3-tech1.bed.gz H3K4me3
                                                        SHR
                                                                     2
## pairedEndReads
## A
              FALSE
## 5
              FALSE
## 16
              FALSE
## 17
              FALSE
## Define assembly
# This is only necessary if you have BED files, BAM files are handled automatically.
# For common assemblies you can also specify them as 'hg19' for example.
data(rn4_chrominfo)
head(rn4_chrominfo)
## UCSC_seqlevel UCSC_seqlength NCBI_seqlevel
## 1
                         16300
            chrM
## 2
                        46782294
            chr12
                       55268282
## 3
            chr20
                                           20
## 4
            chr19
                       59218465
                                           19
## 5
            chr18
                        87265094
                                           18
                      87759784
## 6
            chr11
#=== Step 2: Run Chromstar ===
## Run ChromstaR
Chromstar(inputfolder, experiment.table=experiment_table_SHR,
         outputfolder=outputfolder, numCPU=2, binsize=1000, assembly=rn4_chrominfo,
         prefit.on.chr='chr12', mode='mark')
## Results are stored in 'outputfolder' and can be loaded for further processing
list.files(outputfolder)
## [1] "binned"
                             "browserfiles"
                                                   "chrominfo.tsv"
## [4] "chromstaR.config"
                             "experiment_table.tsv" "multivariate"
## [7] "plots"
                             "univariate"
model <- get(load(file.path(outputfolder,'multivariate',</pre>
                           'multivariate_mode-mark_condition-SHR.RData')))
# Obtain genomic frequencies for combinatorial states
genomicFrequencies(model)
##
                                               [H3K27me3]
##
                  []
                             [H3K4me3]
##
          0.42995169
                             0.09356163
                                               0.41840879
## [H3K27me3+H3K4me3]
          0.05807789
# Plot transition probabilities and read count correlation
plot(model, type='transitionMatrix') + # or plot(model, type=1)
ggtitle('Transition probabilities')
```



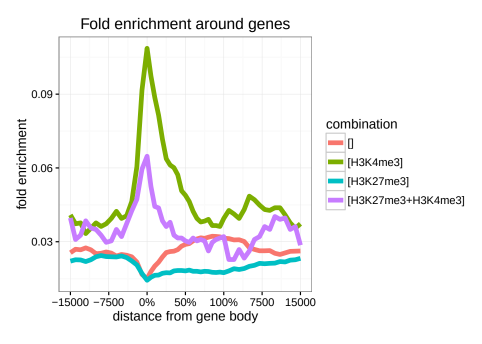




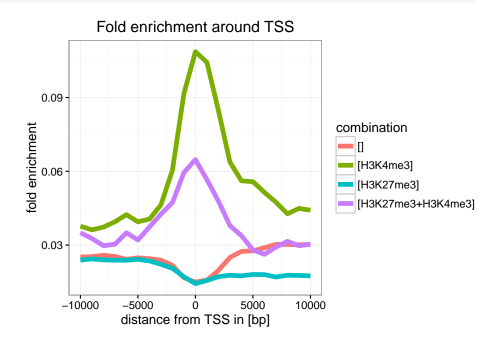
=== Step 3: Enrichment analysis ===
Annotations can easily be obtained with the biomaRt package. Of course, you can
also load them from file if you already have annotation files available.
library(biomaRt)
ensembl <- useMart('ENSEMBL_MART_ENSEMBL', host='may2012.archive.ensembl.org',</pre>

```
dataset='rnorvegicus_gene_ensembl')
genes <- getBM(attributes=c('ensembl_gene_id', 'chromosome_name', 'start_position',</pre>
                            'end_position', 'strand', 'external_gene_id',
                             'gene_biotype'),
               mart=ensembl)
# Transform to GRanges for easier handling
genes <- GRanges(seqnames=paste0('chr',genes$chromosome_name),</pre>
                 ranges=IRanges(start=genes$start, end=genes$end),
                 strand=genes$strand,
                 name=genes$external_gene_id, biotype=genes$gene_biotype)
print(genes)
## GRanges object with 29516 ranges and 2 metadata columns:
##
             seqnames
                                      ranges strand |
                <Rle>
##
                                   <IRanges> <Rle> | <character>
##
         [1]
                chr13
                          [1120899, 1121213]
                                               - |
- |
                                                        L0C682397
##
         [2]
               chr13
                          [1192186, 2293551]
                                                        L0C304725
##
         [3]
               chr13
                          [3174383, 3175216]
                                                  + |
##
         [4]
              chr13
                          [4377731, 4379174]
                                                  - | D3ZPH4_RAT
##
         [5]
               chr13
                          [4866302, 4866586]
                                                  - | F1LZC7_RAT
##
                 . . .
                                                . . . .
                chr6 [134310258, 134310338]
     [29512]
                                                         SNORD113
##
                                                 + |
                chr9 [ 6920889, 6921049]
                                                  - |
##
     [29513]
                                                          U1
             chr11 [ 40073746, 40073816]
chr2 [233090372, 233090478]
##
     [29514]
                                                  - | SNORD19B
                                                  - |
##
     [29515]
                                                         U6
                chr6 [ 92917449, 92917541]
##
     [29516]
##
                  biotype
##
                <character>
##
        [1] protein_coding
##
         [2] protein_coding
##
         [3]
                pseudogene
##
         [4] protein_coding
##
         [5] protein_coding
##
##
     [29512]
                     snoRNA
##
     [29513]
                     snRNA
##
     [29514]
                     snoRNA
##
     [29515]
                     snRNA
                      miRNA
##
     [29516]
##
     seqinfo: 22 sequences from an unspecified genome; no seqlengths
##
```

```
# We expect promoter [H3K4me3] and bivalent-promoter signatures [H3K4me3+H3K27me3]
# to be enriched at transcription start sites.
plotEnrichment(hmm = model, annotation = genes, bp.around.annotation = 15000) +
    ggtitle('Fold enrichment around genes') +
    xlab('distance from gene body')
```

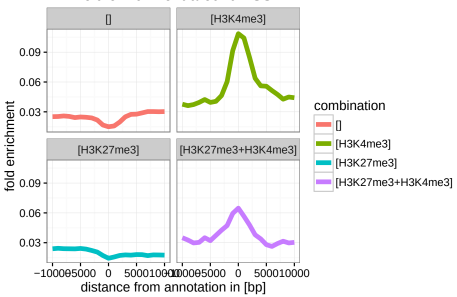


Plot enrichment only at TSS. We make use of the fact that TSS is the start of a gene.
plotEnrichment(model, genes, region = 'start') +
 ggtitle('Fold enrichment around TSS') +
 xlab('distance from TSS in [bp]')



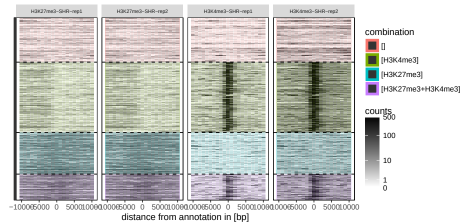
```
# Note: If you want to facet the plot because you have many combinatorial states you
# can do that with
plotEnrichment(model, genes, region = 'start') +
   facet_wrap(~ combination) + ggtitle('Fold enrichment around TSS')
```

Fold enrichment around TSS

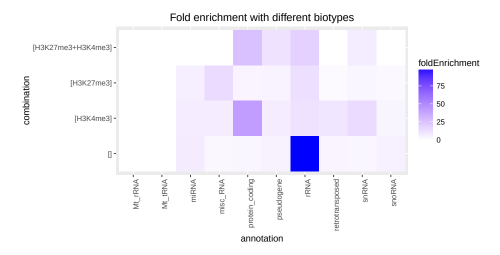


```
# Another form of visualization that shows every TSS in a heatmap
tss <- resize(genes, width = 3, fix = 'start')
plotHeatmap(model, tss) +
  theme(strip.text.x = element_text(size=6)) +
  ggtitle('Read count around TSS')</pre>
```

Read count around TSS



```
# Fold enrichment with different biotypes, showing that protein coding genes are
# enriched with (bivalent) promoter combinations [H3K4me3] and [H3K4me3+H3K27me3],
# while rRNA is enriched with the empty [] combination.
biotypes <- split(tss, tss$biotype)
plotMultipleEnrichment(model, annotations=biotypes) + coord_flip() +
    ggtitle('Fold enrichment with different biotypes')</pre>
```



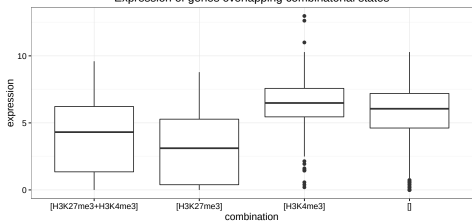
```
# === Step 4: Expression analysis ===
# Suppose you have expression data as well for your experiment. The following code
# shows you how to get the expression values for each combinatorial state.
data(expression_lv)
head(expression_lv)
##
        ensembl_gene_id expression_BN expression_SHR
## 1 ENSRNOGO0000000001
                                  8.8
                                                 7.4
## 2 ENSRNOG00000000007
                                 20.0
                                                 13.0
## 3 ENSRNOG00000000008
                                  1.8
                                                 3.4
## 4 ENSRNOGOOOOOOOO10
                                  6.2
                                                506.8
## 5 ENSRNOG0000000012
                                 48.0
                                                36.4
## 6 ENSRNOGOOOOOOO014
                                 18.2
                                                 15.2
# We need to get coordinates for each of the genes
library(biomaRt)
ensembl <- useMart('ENSEMBL_MART_ENSEMBL', host='may2012.archive.ensembl.org',</pre>
                   dataset='rnorvegicus_gene_ensembl')
genes <- getBM(attributes=c('ensembl_gene_id', 'chromosome_name', 'start_position',</pre>
                             'end_position', 'strand', 'external_gene_id',
                             'gene_biotype'),
               mart=ensembl)
expr <- merge(genes, expression_lv, by='ensembl_gene_id')</pre>
# Transform to GRanges
expression.SHR <- GRanges(seqnames=paste0('chr',expr$chromosome_name),
                          ranges=IRanges(start=expr$start, end=expr$end),
                          strand=expr$strand, name=expr$external_gene_id,
                          biotype=expr$gene_biotype,
```

```
expression=expr$expression_SHR)

# We apply an asinh transformation to reduce the effect of outliers
expression.SHR$expression <- asinh(expression.SHR$expression)

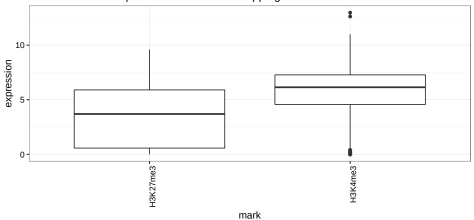
## Plot
plotExpression(model, expression.SHR) +
    theme(axis.text.x=element_text(angle=0, hjust=0.5)) +
    ggtitle('Expression of genes overlapping combinatorial states')
```

Expression of genes overlapping combinatorial states



```
plotExpression(model, expression.SHR, return.marks=TRUE) +
    ggtitle('Expression of marks overlapping combinatorial states')
```

Expression of marks overlapping combinatorial states



4.4 Task 4: Finding differences between combinatorial chromatin states

Consider bivalent promoters defined by [H3K4me3+H3K27me3] at two different developmental stages, or in two different strains or tissues. This is an example where one is interested in *differences* between *combinatorial states*. The following example demonstrates how such an analysis can be done with *chromstaR*. We use a data set from the Euratrans project (downsampled to chr12) to find differences in bivalent promoters between brown norway (BN) and spontaneous hypertensive rat (SHR) in left-ventricle (lv) heart tissue.

Chromstar can be run in three different modes:

- full: Recommended mode if your (number of marks) * (number of conditions) is less or equal to 8. With 8 ChIP-seq experiments there are already 2⁸ = 256 combinatorial states which is the maximum that most computers can handle computationally for a human-sized genome at bin size 1000bp.
- condition: Choose this mode if you are interested in highly significant differences between conditions. The computational limit for the number of conditions is ~ 8 for a human-sized genome. Combinatorial states are not as accurate as in mode mark or full.
- **DEFAULT** *mark*: This mode will yield good combinatorial chromatin state calls for any number of marks and conditions. However, differences between conditions have more false positives than in mode *condition* or *full*.

```
library(chromstaR)
```

```
#=== Step 1: Preparation ===
## Prepare the file paths. Exchange this with your input and output directories.
inputfolder <- system.file("extdata","euratrans", package="chromstaRData")</pre>
outputfolder <- file.path(tempdir(), 'SHR-BN-example')</pre>
## Define experiment structure
data(experiment_table)
print(experiment_table)
                                        file
                                                 mark condition replicate
## 1 lv-H3K27me3-BN-male-bio2-tech1.bed.gz H3K27me3 BN
                                                                       1
## 2 lv-H3K27me3-BN-male-bio2-tech2.bed.gz H3K27me3
                                                             BN
## 4 lv-H3K27me3-SHR-male-bio2-tech1.bed.gz H3K27me3
                                                            SHR
                                                                        1
## 5 lv-H3K27me3-SHR-male-bio2-tech2.bed.gz H3K27me3
                                                            SHR
```

```
## 13 lv-H3K4me3-BN-female-bio1-tech1.bed.gz H3K4me3
                                                       BN
## 14 lv-H3K4me3-BN-male-bio2-tech1.bed.gz H3K4me3
                                                         BN
                                                                    2
## 16 lv-H3K4me3-SHR-male-bio2-tech1.bed.gz H3K4me3
                                                         SHR
                                                                    1
## 17 lv-H3K4me3-SHR-male-bio3-tech1.bed.gz H3K4me3
                                                         SHR
                                                                    2
## pairedEndReads
## 1
            FALSE
             FALSE
## 2
## 4
             FALSE
             FALSE
## 5
## 13
            FALSE
## 14
            FALSE
            FALSE
## 16
## 17
             FALSE
## Define assembly
# This is only necessary if you have BED files, BAM files are handled automatically.
\mbox{\# For common assemblies you can also specify them as 'hg19' for example.}
data(rn4_chrominfo)
head(rn4_chrominfo)
## UCSC_seqlevel UCSC_seqlength NCBI_seqlevel
## 1 chrM 16300 MT
## 2
            chr12
                       46782294
## 3
                      55268282
                                         20
           chr20
## 4
           chr19
                      59218465
## 5
                      87265094
           chr18
                                          18
          chr11
                     87759784
#=== Step 2: Run Chromstar ===
## Run ChromstaR
Chromstar(inputfolder, experiment.table=experiment_table,
         outputfolder=outputfolder, numCPU=2, binsize=1000, assembly=rn4_chrominfo,
         prefit.on.chr='chr12', mode='mark')
## Results are stored in 'outputfolder' and can be loaded for further processing
list.files(outputfolder)
## [1] "binned"
                             "browserfiles"
                                                    "chrominfo.tsv"
                             "experiment_table.tsv" "multivariate"
## [4] "chromstaR.config"
## [7] "multivariate-combined" "plots"
                                                    "univariate"
model <- get(load(file.path(outputfolder,'multivariate-combined',</pre>
                         'combined_mode-mark.RData')))
#=== Step 3: Analysis and export ===
## Obtain all genomic regions where the two tissues have different states
segments <- model$segments</pre>
diff.segments <- segments[segments$combination.SHR != segments$combination.BN]
# Let's be strict with the differences and get only clear ones
diff.segments <- diff.segments[diff.segments$differential.score > 0.999]
exportGRanges(diff.segments, trackname='differential_chromatin_states',
```

```
filename=tempfile(), scorecol='differential.score')
## Obtain all genomic regions where we find a bivalent promoter in BN but not in SHR
bivalent.BN <- segments[segments$BN == '[H3K27me3+H3K4me3]' &
                        segments$SHR != '[H3K27me3+H3K4me3]']
## Obtain all genomic regions where BN and SHR have promoters
promoter.BN <- segments[segments$transition.group == 'constant [H3K4me3]']</pre>
## Get transition frequencies
print(model$frequencies)
##
                                         SHR
                                                frequency cumulative.frequency
## 1
                       [] 4.107135e-01
                                                                     0.4107135
                                   [H3K4me3] 3.956436e-01
## 2
               [H3K4me3]
                                                                     0.8063571
## 3
              [H3K27me3]
                                          [] 5.752212e-02
                                                                     0.8638793
      [H3K27me3+H3K4me3]
                                   [H3K4me3] 3.591125e-02
## 4
                                                                     0.8997905
## 5
              [H3K27me3]
                                          [] 3.428669e-02
                                                                     0.9340772
## 6
               [H3K4me3]
                                   [H3K4me3] 2.037108e-02
                                                                     0.9544483
## 7
                                         [] 1.718610e-02
                      []
                                                                     0.9716344
## 8
      [H3K27me3+H3K4me3]
                                   [H3K4me3] 1.697234e-02
                                                                     0.9886067
## 9
              [H3K27me3]
                                          [] 2.907101e-03
                                                                     0.9915138
## 10 [H3K27me3+H3K4me3] [H3K27me3+H3K4me3] 2.351332e-03
                                                                      0.9938652
## 11
              [H3K27me3]
                                          [] 2.287204e-03
                                                                     0.9961524
## 12 [H3K27me3+H3K4me3]
                                   [H3K4me3] 1.496302e-03
                                                                     0.9976487
## 13
              [H3K27me3]
                                  [H3K27me3] 1.496302e-03
                                                                     0.9991450
## 14
              [H3K27me3]
                                  [H3K27me3] 5.557693e-04
                                                                      0.9997007
## 15 [H3K27me3+H3K4me3]
                                   [H3K4me3] 2.565089e-04
                                                                      0.9999572
## 16 [H3K27me3+H3K4me3] [H3K27me3+H3K4me3] 4.275149e-05
                                                                      1.0000000
##
                                   group
## 1
                        zero transition
## 2
                    constant [H3K27me3]
## 3
                     constant [H3K4me3]
## 4
            constant [H3K27me3+H3K4me3]
## 5
              stage-specific [H3K4me3]
## 6
              stage-specific [H3K27me3]
## 7
              stage-specific [H3K27me3]
## 8
                                   other
## 9
      stage-specific [H3K27me3+H3K4me3]
## 10
                                   other
## 11
                                   other
## 12
                                   other
## 13
               stage-specific [H3K4me3]
## 14 stage-specific [H3K27me3+H3K4me3]
## 15
## 16
                                   other
```

5 FAQ

5.1 The peak calls are too lenient. Can I adjust the strictness of the peak calling?

The strictness of the peak calling can be controlled with a posterior cutoff. The Hidden Markov Model gives posterior probabilities for each peak, and based on these probabilites the model decides if a peak is present or not by picking the state with the highest probability. This way of peak calling leads to very lenient peak calls, and for some applications it may be desirable to obtain only very clear peaks. This can be achieved by setting a posterior cutoff. To follow the below example, please first run step 1 and 2 from section 4.4. !!Note that in general more peaks are obtained with a stricter cutoff, as this will lead to the split-up of previously broader peaks into several smaller, more well defined peaks.

5.2 The combinatorial differences that chromstaR gives me are not convincing. Is there a way to restrict the results to a more convincing set?

You were interested in combinatorial state differences as in section 4.4 and checked the results in a genome browser. You found that some differences are convincing by eye and some are not. There are several possibilities to explore:

- 1. You can play with the "differential score" (see section 4.4, step 3) and export only differences with a score close to 1.
- 2. Run Chromstar with mode='mark' or mode='condition' and see if the results improve.
- 3. Check for bad replicates that are very different from the rest and exclude them prior to the analysis.

6 Session Info

```
sessionInfo()
## R Under development (unstable) (2016-02-17 r70182)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 14.04.4 LTS
## locale:
## [1] LC_CTYPE=en_US.UTF-8
                                LC_NUMERIC=C
## [3] LC_TIME=nl_NL.UTF-8
                                LC_COLLATE=C
## [5] LC_MONETARY=nl_NL.UTF-8 LC_MESSAGES=en_US.UTF-8
## [7] LC_PAPER=n1_NL.UTF-8
                                 LC_NAME=C
## [9] LC_ADDRESS=C
                                 LC_TELEPHONE=C
## [11] LC_MEASUREMENT=n1_NL.UTF-8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] stats4 parallel stats
                                 graphics grDevices utils
                                                                datasets
## [8] methods base
##
## other attached packages:
## [1] biomaRt_2.27.2 chromstaR_0.99.1
                                             chromstaRData_0.99.0
## [4] ggplot2_2.1.0
## [7] IRanges_2.6.0
                          GenomicRanges_1.24.0 GenomeInfoDb_1.8.0
                         S4Vectors_0.10.0 BiocGenerics_0.18.0
## loaded via a namespace (and not attached):
## [1] Rcpp_0.12.4.5
                          compiler_3.3.0
##
    [3] formatR_1.3
                                 plyr_1.8.3
                                XVector_0.12.0
## [5] highr_0.5.1
## [7] bitops_1.0-6
                                iterators_1.0.8
## [9] tools_3.3.0
                                zlibbioc_1.18.0
## [11] digest_0.6.9
                                RSQLite_1.0.0
## [13] evaluate_0.8.3
                                gtable_0.2.0
                                DBI_0.3.1
## [15] foreach_1.4.3
## [17] stringr_1.0.0
                                knitr_1.12.3
                               grid_3.3.0
## [19] Biostrings_2.40.0
## [21] Biobase_2.32.0
                                 AnnotationDbi_1.33.7
                                BiocParallel_1.6.0
## [23] XML_3.98-1.4
## [25] reshape2_1.4.1
                                magrittr_1.5
                            Rsamtools_1.24.0
GenomicAlignments_1.8.0
## [27] scales_0.4.0
## [29] codetools_0.2-14
## [31] SummarizedExperiment_1.2.0 colorspace_1.2-6
## [33] labeling_0.3
                                stringi_1.0-1
## [35] RCurl 1.95-4.8
                                 doParallel_1.0.10
## [37] munsell_0.4.3
warnings()
## NULL
```