Package ‘DChIPRep’

March 22, 2017

Title DChIPRep - Analysis of chromatin modification ChIP-Seq data with replication

Version 1.4.0

Description The DChIPRep package implements a methodology to assess differences between chromatin modification profiles in replicated ChIP-Seq studies as described in Chabbert et. al - http://www.dx.doi.org/10.15252/msb.20145776. A detailed description of the method is given in the software paper at https://doi.org/10.7717/peerj.1981

Depends R (>= 3.3), DESeq2

Imports methods, stats, utils, ggplot2, fdrtool, reshape2,
GenomicRanges, SummarizedExperiment, smoothenest, plyr, tidyr,
assertthat, S4Vectors, purrr, soGGi, ChIPpeakAnno

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LazyData true

Suggests mgcv, testthat, BiocStyle, knitr, rmarkdown

Collate 'AllClasses.R' 'AllGenerics.R' 'DChIPRep.R' 'dataImport.R'
'dataImportsoGGi.R' 'documentData.R' 'methods.R'
'plottingFunctions.R' 'runTesting.R'

VignetteBuilder knitr

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SystemRequirements Python 2.7, HTSeq (>= 0.6.1), numpy, argpase, sys

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R topics documented:

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Another example ChIP data set that can be used with `importDataFromMatrices` from rom Galonska et. al., 2015.

Description

The containes the H3Kme3 data in the serum and 24h_2i conditions from Galonska et. al., 2015 as well as the whole cell extract data, which is treated as input for all four samples. The data were downloaded from the SRA at the european nucleotide archive (ENA, accession PRJNA242892). The reads were aligned to the mm9 reference genome using bowtie2 (Langmead and Salzberg, 2012) with default options. Then, filtering of unmapped, low mapping quality (< 10), duplicated and multi-mapping reads was performed with Picard tools. The fragment length was inferred using cross correlation plots from SPP (Kharchenko, et. al., 2008).

Usage

data(chip_galonska)

Format

a matrix

Value

a matrix

References

DChIPRep


See Also

sample_table_galonska input_galonska TSS_galonska

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**DChIPRep**

*DChIPRep: A package for differential analysis of histone modification ChIP-Seq profiles*

**Description**

The DChIPRep package provides functions to perform a differential analysis of histone modification profiles at base-pair resolution.

**DChIPRep functions**

The DChIPRep packages provides functions for data import `importData` and performing position wise tests. After data import, a `DChIPRepResults` object on which the function `runTesting` is run to perform the tests and add the result to the object. Then, plots can be created from this object. See the vignette for additional details: `vignette("DChIPRepVignette")`

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**DChIPRepResults-class**

*DChIPRepResults object and constructor*

**Description**

The DChIPRepResults contains a DESeqDataSet as obtained after the initial import.

**Usage**

`DChIPRepResults(object)`

**Arguments**

- `object` A DESeqDataSet

**Value**

A DChIPRepResult object.

**Examples**

```r
data(testData)
dcr <- DChIPRepResults(testData)
```
DESeq2Data

Accessors for the 'DESeq2Data' slot of a DChIPRepResults object.

Description

The slot contains the DESeqDataSet as it is obtained after the initial data import. The DESeqDataSet contains the counts per position and the normalization factors as computed using the input counts.

Usage

```r
## S4 method for signature 'DChIPRepResults'
DESeq2Data(object)

## S4 replacement method for signature 'DChIPRepResults,DESeqDataSet'
DESeq2Data(object) <- value
```

Arguments

- `object`: a DChIPRepResults object
- `value`: A DESeqDataSet object

Value

the DESeq2Data object contained in the DChIPRepResults object

Examples

```r
data(testData)
dcr <- DChIPRepResults(testData)
DESeq2Data(dcr)
```

exampleChipData

An example ChIP data.

Description

An example ChIP data set that can be used with `importDataFromMatrices`. Its associated sample table can be accessed via `data(data(exampleSampleTable))`.

Usage

```r
data(exampleChipData)
```

Format

- a matrix
exampleInputData

Value

a matrix

See Also

eexampleSampleTable exampleInputData

An example input data.

Description

An example input data set that can be used with importDataFromMatrices. Its associated sample table can be accessed via data(data(exampleSampleTable)).

Usage

data(exampleInputData)

Format

a matrix

Value

a matrix

See Also

eexampleSampleTable exampleChipData

exampleSampleTable An example sample table data.frame

Description

An example sample table

Usage

data(exampleSampleTable)

Format

a data.frame

Value

a data.frame

See Also

eexampleChipData exampleInputData
FDRresults accessor and setter for the 'FDRresults' slot of a DChIPRepResults object.

Description
The slot contains the results of the FDR estimation as performed within the function runTesting. It is the complete output of the fdrtool function.

Usage

```r
## S4 method for signature 'DChIPRepResults'
FDRresults(object)

## S4 replacement method for signature 'DChIPRepResults,list'
FDRresults(object) <- value
```

Arguments

- `object`: a DChIPRepResults object
- `value`: A DESeqDataSet object

Value

A list containing the estimated false discovery rates

Examples

```r
data(testData)
dcr <- DChIPRepResults(testData)
dcr <- runTesting(dcr)
str(FDRresults(dcr))
```

getMATfromDataFrame helper function to turn a data.frame into a matrix and remove the ID column.

Description
This function takes a data.frame, with the genomic features (e.g. transcripts or genes) in the rows and the positions upstream and downstream of the TSS in the columns as well as a column ID containing a genomic feature ID and returns the data.frame with the ID column removed. The input for this function are tables obtained after running the Python import script.

Usage

```r
getMATfromDataFrame(df, ID = "name")
```
Arguments

df    the input data frame with positions in the columns and the genomic features in the rows.
ID    the name of the ID column to be removed.

Value

a matrix with the ID column removed

Examples

data(exampleSampleTable)
directory <- file.path(system.file("extdata", package="DChIPRep"))
df <- lapply(file.path(directory, exampleSampleTable$Input),
            read.delim)[[1]]
mat <- getMATfromDataFrame(df)

importData Import the data after running the Python script

Description

This function imports the data from the count table files as returned by the accompanying Python script.

Usage

importData(sampleTable, directory = "", ID = "name", ...)

Arguments

sampleTable a data.frame that has to contain the columns ChiP, Input, sampleID, upstream, downstream and condition. Each row of the table describes one experimental sample. Each row of the table describes one experimental sample. See data(exampleSampleTable) for an example table. and the vignette for further information.
directory the directory relative to which the filenames are specified given as a character.
ID character giving the name of the feature identifier column in the count tables. Defaults to "name"
... parameters passed to summarizeCountsPerPosition

Value

a DChIPRepResults object containg the imported data as a DESeqDataSet.

Examples

data(exampleSampleTable)
directory <- file.path(system.file("extdata", package="DChIPRep"))
importedData <- importData(exampleSampleTable, directory)
importDataFromMatrices

Import the data from ChiP and input matrices

**Description**

This function imports the data from two matrices that contain counts summarized per position. It computes the normalization factors from the input (one per position) and creates a DChIPRepResults object.

**Usage**

```r
importDataFromMatrices(inputData, chipData, sampleTable)
```

**Arguments**

- `inputData`: a matrix containing the counts for the input per position.
- `chipData`: a matrix containing the counts for the ChIP per position.
- `sampleTable`: a data.frame that has to contain the columns sampleID, upstream, downstream and condition. Each row of the table describes one experimental sample. See `data(exampleSampleTable)` for an example table, and the vignette for further information.

**Details**

The normalization factors are computed as $t(t(inputData) * (covC/covI))$, Where covC and covI contain the total sum of the ChIP and the input samples. Zero normalization factors can arise if the input has zero counts for certain positions. That’s why input values equal to zero are set to 1 in order to always obtain valid normalizationFactors.

**Value**

A DChIPRepResults object containing the imported data as a DESeqDataSet.

**Examples**

```r
data(exampleSampleTable)
data(exampleInputData)
data(exampleChipData)
imDataFromMatrices <- importDataFromMatrices(inputData = exampleInputData,
chipData = exampleChipData,
sampleTable = exampleSampleTable)
```
importData_soGGi

**Import the data from bam files directly**

**Description**

This function imports the data from .bam files directly. It will return a matrix with one column per .bam file and the respective counts per position in the rows. It uses the function `regionPlot` from the package soGGi.

**Usage**

```r
importData_soGGi(bam_paths, TSS, fragment_lengths, sample_ids, 
distanceUp = 1000, distanceDown = 1500, ...)
```

**Arguments**

- `bam_paths` a character vector of paths to the bam file(s) to be imported.
- `TSS` a GRanges (GenomicRanges-class) (or a class that inherits from it) object containing the TSS of interest.
- `fragment_lengths` an integer vector of fragment lengths.
- `sample_ids` a character vector of sample ids for the .bam files. This can also be a factor.
- `distanceUp` Distance upstream from centre of the TSS provided.
- `distanceDown` Distance downstream from centre of the TSS provided.
- `...` additional arguments passed to `regionPlot`.

**Details**

In the example below, we use a subsampled .bam file (0.1 % of the reads) from the Galonska et. al. WCE (whole cell extract) H3Kme3 data and associated TSS near identified peaks. For additional details on the data, see input_galonska and TSS_galonska.

**Value**

a matrix that contains the postion-wise profiles per .bam file in the columns.

**See Also**

`regionPlot` input_galonska TSS_galonska sample_table_galonska

**Examples**

```r
## Not run:
data(sample_table_galonska)
data(TSS_galonska)
bam_dir <- file.path(system.file("extdata", package="DChIPRep"))
wce_bam <- "subsampled_0001_pc_SRR2144628_WCE_bowtie2_mapped-only_XS-filt_no-dups.bam"
mat_wce <- importData_soGGi(bam_paths = file.path(bam_dir, wce_bam), 
TSS = TSS_galonska, 
fragment_lengths = sample_table_galonska$input_fragment_length[1], 
sample_ids = sample_table_galonska$input[1],
```
```r
plotProfiles

pair = FALSE,
removeDup=FALSE

head(mat_wce)
```

```r
input_galonska
Another example Input data set that can be used with importDataFromMatrices from rom Galonska et. al., 2015.
```

**Description**

The matrix contains the whole cell extract (WCE) data for H3Kme3 from the paper in each of the four columns, since this is the only input data provided for all 4 samples. For additional information see the documentation of chip_galonska.

**Usage**

```r
data(input_galonska)
```

**Format**

a matrix

**Value**

a matrix

**See Also**

chip_galonska sample_table_galonska TSS_galonska

```r
plotProfiles

Produce a TSS plot of the two conditions in the data
```

**Description**

This function plots the positionwise mean of the log2 of the normalized counts of the two conditions after runTesting has been run on a DChIPRepResults object.

**Usage**

```r
## S4 method for signature 'DChIPRepResults'
plotProfiles(object, meanFunction = function(x) {
  smhuber(x)$mu }, ...)
```
plotSignificance

Arguments

object a DChIPRepResults object after runTesting
meanFunction a function to compute the positionwise mean per group, defaults to a Huber estimator of the mean.
... additional parameters for plotting (NOT YET IMPLEMENTED)

Value

a ggplot2 object

Examples

if (requireNamespace("mgcv", quietly=TRUE)) {
  data(testData)
  dcr <- DChIPRepResults(testData)
  dcr <- runTesting(dcr)
  plotProfiles(dcr)
}

plotSignificance Produce a plot that colors the positions identified as significant

Description

This function plots the positionwise mean of the two conditions after runTesting has been run on a DChIPRepResults object. The points corresponding to significant positions are colored black in both of the conditions. The function returns the plot as a ggplot2 object that can be modified afterwards.

Usage

## S4 method for signature 'DChIPRepResults'
plotSignificance(object, 
  meanFunction = function(x) { smhuber(x)$mu }, lfdrThresh = 0.2, ...)

Arguments

object a DChIPRepResults object after runTesting
meanFunction a function to compute the positionwise mean per group, defaults to a Huber estimator of the mean.
lfdrThresh Threshold for the local FDR
... additional parameters for plotting (NOT YET IMPLEMENTED)

Value

a ggplot2 object

Examples

data(testData)
  dcr <- DChIPRepResults(testData)
  dcr <- runTesting(dcr)
  plotSignificance(dcr)
resultsDChIPRep  

**Description**

The slot contains the results of the position wise tests in a data.frame after running the function `runTesting`. It is a modified output of the `results` function of the `DESeq2` package.

**Usage**

```r
## S4 method for signature 'DChIPRepResults'
resultsDChIPRep(object)

## S4 replacement method for signature 'DChIPRepResults,list'
resultsDChIPRep(object) <- value
```

**Arguments**

- `object`: a `DChIPRepResults` object
- `value`: A `DESeqDataSet` object

**Value**

A data.frame containing the results of the position wise tests

**Examples**

```r
data(testData)
dcr <- DChIPRepResults(testData)
dcr <- runTesting(dcr)
head(resultsDChIPRep(dcr))
```

---

**runTesting**

**Run the tests on a DChIPRepResults object.**

**Description**

This function runs the testing on a `DChIPRepResults` object. It adds the FDR calculations and the result table to the `DChIPRepResults` object.

**Usage**

```r
## S4 method for signature 'DChIPRepResults'
runTesting(object, lfcThreshold = 0.05,
plotFDR = FALSE, ...)
```
Arguments

object A DChIPRepResults object.

lfcThreshold A non-negative threshold value, which determines the null hypothesis. The null hypothesis is \( H_0 : |\log_2(FC)| > \text{lfcThreshold} \)

plotFDR If set to TRUE a plot showing the estimated FDRs will be displayed

Value

A modified DChIPRepResults object containing the testing results

See Also

resultsDChIPRep

Examples

data(testData)
dcr <- DChIPRepResults(testData)
dcr <- runTesting(dcr)

Description

This table contains the sample annotation for the H3Kme3 data from Galonska et. al., 2015. For additional information see the documentation of chip_galonska.

Usage

data(sample_table_galonska)

Format

a data.frame

Value

a data.frame

See Also

chip_galonska input_galonska TSS_galonska
show

prints the DESeq2Data slot of the DChIPRepResults object

Description

prints the data

Usage

## S4 method for signature 'DChIPRepResults'
show(object)

Arguments

object A DChIPRepResults object

Value

A compact representation of the DChIPRepResults object

Examples

data(testData)
dcr <- DChIPRepResults(testData)
dcr
dcr <- runTesting(dcr)
dcr

summarizeCountsPerPosition

Helper function to summarize the counts per position

Description

This function takes a matrix of counts, with the genomic features (e.g. transcripts or genes) in the rows and the positions upstream and downstream of the TSS in the columns and returns a vector with the summarized counts per position.

Usage

summarizeCountsPerPosition(mat, ct = 0, trim = 0.15)

Arguments

mat the input matrix with positions in the columns and the genomic features in the rows.
ct the count threshold to use.
trim the trimming percentage for the trimmed mean.
Details

The summary per condition is computed as a trimmed mean per position. First, counts greater than \( ct \) are removed and then a trimmed mean with a trimming percentage of \( \text{trim} \) is computed on the log scale. This mean is then exponentiated again and multiplied by the total number of features passing the threshold \( ct \) per position. If a position contains only zero counts, its mean is returned as zero.

Value

a vector containing the summarized counts per condition

Examples

data(exampleSampleTable)
directory <- file.path(system.file("extdata", package="DChIPRep"))
df <- lapply(file.path(directory, exampleSampleTable$Input), read.delim)[[1]]
mat <- getMATfromDataFrame(df)
summaryPerPos <- summarizeCountsPerPosition(mat)
**Description**

This data contains mouse mm9 TSS close to called peak regions for H3Kme3 from Galonska et al. The original peak lists are from GEO (GSE56312) and have been merged into a common peaklist and then annotated to the closest mm9 TSS using `annotatePeakInBatch`.

**Usage**

```r
data(exampleChipData)
```

**Format**

an annoGR object from the package ChIPpeakAnno.

**Value**

an annoGR object from the package ChIPpeakAnno.

**See Also**

`chip_galonska` `input_galonska` `sample_table_galonska`
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