Package ‘soGGi’

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Type Package

Title Visualise ChIP-seq, MNase-seq and motif occurrence as aggregate plots Summarised Over Grouped Genomic Intervals

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Description The soGGi package provides a toolset to create genomic interval aggregate/summary plots of signal or motif occurrence from BAM and bigWig files as well as PWM, rlelist, GRanges and GAlignments Bioconductor objects. soGGi allows for normalisation, transformation and arithmetic operation on and between summary plot objects as well as grouping and subsetting of plots by GRanges objects and user supplied metadata. Plots are created using the GGplot2 library to allow user defined manipulation of the returned plot object. Coupled together, soGGi features a broad set of methods to visualise genomics data in the context of groups of genomic intervals such as genes, superenhancers and transcription factor binding events.

biocViews Sequencing, ChIPSeq, Coverage

License GPL (>= 3)

LazyLoad yes

Depends R (>= 3.2.0), BiocGenerics, SummarizedExperiment

Imports methods, reshape2, ggplot2, S4Vectors, IRanges, GenomeInfoDb, GenomicRanges, Biostats, Rsamtools, GenomicAlignments, rtracklayer, preprocessCore, chipseq, BiocParallel

Collate 'allClasses.r' 'motifTools.R' 'peakTransforms.r' 'plots.R' 'soGGi.R'

VignetteBuilder knitr

Suggests testthat, BiocStyle, knitr

NeedsCompilation no
Join, subset and manipulate ChIPprofile objects

Description

Join, subset and manipulate ChIPprofile objects

Usage

```r
## S4 method for signature 'ChIPprofile'
c(x, ..., recursive = FALSE)
```

```r
## S4 method for signature 'ChIPprofile'
rbind(x, ..., deparse.level = 1)
```

```r
## S4 method for signature 'ChIPprofile'
cbind(x, ..., deparse.level = 1)
```

```r
## S4 method for signature 'ChIPprofile,ANY,missing'
```
x[[i, j, ...]]

## S4 method for signature 'ChIPprofile'
x$name

### Arguments

- **j**: Should be missing
- **...**: objects to be concatenated.
- **recursive**: logical. If recursive = TRUE, the function recursively descends through lists (and pairlists) combining all their elements into a vector.
- **deparse.level**: See ?base::cbind for a description of this argument.
- **x**: object from which to extract element(s) or in which to replace element(s).
- **i**: indices specifying elements to extract or replace. Indices are numeric or character vectors or empty (missing) or NULL. Numeric values are coerced to integer as by as.integer (and hence truncated towards zero). Character vectors will be matched to the names of the object (or for matrices/arrays, the dimnames): see ‘Character indices’ below for further details.
  - For [-indexing only: i, j, ... can be logical vectors, indicating elements/slices to select. Such vectors are recycled if necessary to match the corresponding extent. i, j, ... can also be negative integers, indicating elements/slices to leave out of the selection.
  - When indexing arrays by [ a single argument i can be a matrix with as many columns as there are dimensions of x; the result is then a vector with elements corresponding to the sets of indices in each row of i.
  - An index value of NULL is treated as if it were integer(0).
- **name**: A literal character string or a name (possibly backtick quoted). For extraction, this is normally (see under ‘Environments’) partially matched to the names of the object.

### Value

A ChIPprofile object

### Examples

data(chipExampleBig)
x <- c(chipExampleBig[[1]], chipExampleBig[[2]])
y <- rbind(chipExampleBig[[1]], chipExampleBig[[2]])
**Description**

This dataset contains peaks from ChIP-signal over genes

**Usage**

data(chipExampleBig)

**Details**

- ChIPprofiles

**Value**

A ChIPprofile object

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**ChIPprofile-class**

The soggi function and ChIPprofile object.

**Description**

Manual for soggi and ChIPprofile object

The soggi function is the constructor for ChIPprofile objects.

**Usage**

regionPlot(bamFile, testRanges, samplename = NULL, nOfWindows = 100, FragmentLength = 150, style = "point", distanceAround = NULL, distanceUp = NULL, distanceDown = NULL, distanceInRegionStart = NULL, distanceOutRegionStart = NULL, distanceInRegionEnd = NULL, distanceOutRegionEnd = NULL, paired = FALSE, normalize = "RPM", plotBy = "coverage", removeDup = FALSE, verbose = TRUE, format = "bam", seqlengths = NULL, forceFragment = NULL, method = "bin", genome = NULL, cutoff = 80, downSample = NULL, minFragmentLength = NULL, maxFragmentLength = NULL)
Arguments

- **bamFile**: Character vector for location of BAM file or bigWig, an rleList or PWM matrix.
- **testRanges**: GRanges object or character vector of BED file location of regions to plot.
- **samplename**: Character vector of sample name. Default is NULL.
- **nOfWindows**: Number of windows to bin regions into for coverage calculations (Default 100)
- **FragmentLength**: Integer vector Predicted or expected fragment length.
- **style**: "Point" for per base pair plot, "percentOfRegion" for normalised length and "region" for combined plot
- **distanceAround**: Distance around centre of region to be used for plotting
- **distanceUp**: Distance upstream from centre of region to be used for plotting
- **distanceDown**: Distance downstream from centre of region to be used for plotting
- **distanceInRegionStart**: Distance into region start (5’ for Watson/positive strand or not specified strand Regions, 3’ for Crick/negative strand regions) for plotting.
- **distanceOutRegionStart**: Distance out from region start (5’ for Watson/positive strand or not specified strand Regions, 3’ for Crick/negative strand regions) for plotting.
- **distanceInRegionEnd**: Distance into region end (3’ for Watson/positive strand or not specified strand Regions, 5’ for Crick/negative strand regions) for plotting.
- **distanceOutRegionEnd**: Distance out from region end (3’ for Watson/positive strand or not specified strand Regions, 5’ for Crick/negative strand regions) for plotting.
- **paired**: Is data paired end
- **normalize**: Calculate coverage as RPM. Presently only RPM available.
- **plotBy**: Score to be used for plotting. Presently only coverage.
- **removeDup**: Remove duplicates before calculating coverage.
- **verbose**: TRUE or FALSE
- **format**: Character vector of "BAM", "BigWig", "RleList" or "PWM"
- **seqLengths**: Chromosomes to be used. If missing will report all.
- **forceFragment**: Centre fragment and force consistent fragment width.
- **method**: Character vector of value "bp","bin" or "spline". The bin method divides a region of interest into equal sized bins of number specified in nOfWindows. Coverage or counts are then summarised within these windows. The spline method creates a spline with the number of spline points as specified in nOfWindows argument.
- **downSample**: Down sample BAM reads to this proportion of original.
- **genome**: BSGenome object to be used when using PWM input.
- **cutoff**: Cut-off for idnetifying motifs when using PWM input.
- **minFragmentLength**: Remove fragments smaller than this.
- **maxFragmentLength**: Remove fragments larger than this.
findconsensusRegions

Value
ChIPprofile

References
See [http://bioinformatics.csc.mrc.ac.uk](http://bioinformatics.csc.mrc.ac.uk) for more details on soGGi workflows

Examples
```r
data(chipExampleBig)
chipExampleBig
```

Description
Plot coverage of points or regions. Returns summits and summmit scores after optional fragment length prediction and read extension

Usage
```r
findconsensusRegions(testRanges, bamFiles = NULL, method = "majority", summit = "mean", resizepeak = "asw", overlap = "any", fragmentLength = NULL, NonPrimaryPeaks = list(withinsample = "drop", betweensample = "mean"))
```

Arguments
- `testRanges`: Named character vector of region locations
- `bamFiles`: Named character vector of bamFile locations
- `method`: Method to select reproducible summits to merge.
- `summit`: Only mean available
- `resizepeak`: Only asw available
- `overlap`: Type of overlap to consider for finding consensus sites
- `fragmentLength`: Predicted fragment length. Set to NULL to auto-calculate
- `NonPrimaryPeaks`: A list of parameters to deal with non primary peaks in consensus regions.
- `peakfile`: GRanges of genomic intervals to summit.
- `reads`: Character vector of bamFile location or GAlignments object
- `readlength`: Read length of alignments.
**groupByOverlaps**

**Value**

Consensus A GRanges object of consensus regions with consensus summits.
Summits A GRanges object of summits and summit scores.

---

**Description**

Create GRangeslist from all combinations of GRanges

**Usage**

`groupByOverlaps(testRanges)`

**Arguments**

- `testRanges`: A named list of GRanges or a named GRangesList

**Value**

- `groupedGRanges`: A named GRangesList object.

**Examples**

```r
data(ik_Example)
gts <- groupByOverlaps(ik_Example)
```

---

**ik_Example**

*Example Ikaros peaksets*

---

**Description**

This dataset contains peaks from Ikaros ChIP by two antibodies

**Usage**

`data(ik_Example)`

**Details**

- Ikpeaksets

**Value**

A list containing two GRanges objects
ik_Profiles

Example Ikaros signal over peaksets

Description

This dataset contains signal over peaks from Ikaros ChIP by two antibodies

Usage

data(ik_Profiles)

Details

• ik_Profiles

Value

A ChIPprofile object

normalise

Normalise ChIPprofiles

Description

Various normalisation methods for ChIPprofile objects

Usage

## S4 method for signature 'ChIPprofile'
normalise(object)

## S4 method for signature 'ChIPprofile,character,numeric'
normalise(object = "ChIPprofile",
  method = "rpm", normFactors = NULL)

Arguments

object A ChIPprofile object

method A character vector specifying normalisation method. Currently "rpm" for normalising signal for BAM to total reads, "quantile" to quantile normalise across samples, "signalInRegion" to normalise to proportion of signal within intervals, "normaliseSample" to normalise across samples and "normaliseRegions" to apply a normalisation across intervals.

normFactors A numeric vector used to scale columns or rows.
normaliseQuantiles

Value
A ChIPprofile object

Author(s)
Thomas Carroll

Examples
data(chipExampleBig)
normalise(chipExampleBig, method="quantile", normFactors=1)

normaliseQuantiles
Normalise quantile

Description
Quantile normalisation across bins/regions.

Usage
## S4 method for signature 'ChIPprofile'
normaliseQuantiles(object)

## S4 method for signature 'ChIPprofile'
normaliseQuantiles(object = "ChIPprofile")

Arguments
object A ChIPprofile object

Value
A ChIPprofile object containing normalised data

Author(s)
Thomas Carroll

Examples
data(chipExampleBig)
normaliseQuantiles(chipExampleBig)
Description
Arithmetic operations

Usage

```r
## S4 method for signature 'ChIPprofile,ChIPprofile'
Ops(e1, e2)

## S4 method for signature 'ChIPprofile,numeric'
Ops(e1, e2)

## S4 method for signature 'numeric,ChIPprofile'
Ops(e1, e2)

## S4 method for signature 'ChIPprofile'
mean(x, ...)

## S4 method for signature 'ChIPprofile'
log2(x)

## S4 method for signature 'ChIPprofile'
log(x, base = exp(1))
```

Arguments

- `e1`: ChIPprofile object
- `e2`: ChIPprofile object
- `x`: objects.
- `...`: further arguments passed to methods.
- `base`: a positive or complex number: the base with respect to which logarithms are computed. Defaults to `exp(1)`.

Value

A ChIPprofile object of result of arithmetic operation.

Examples

```r
data(chipExampleBig)
chipExampleBig[[1]] + chipExampleBig[[2]]
```
orientBy

Set strand by overlapping or nearest anchor GRanges

Description

Set strand by overlapping or nearest anchor GRanges

Usage

orientBy(testRanges, anchorRanges)

Arguments

testRanges  The GRanges object to anchor.
anchorRanges  A GRanges object by which to anchor strand orientation.

Value

newRanges A GRanges object.

Examples

data(iK_Example)
strand(iK_Example[[1]]) <- "+"
anchoredGRanges <- orientBy(iK_Example[[2]], iK_Example[[1]])

plotRegion

Plot regions

Description

A function to plot regions

Usage

## S4 method for signature 'ChIPprofile'
plotRegion(object = "ChIPprofile", gts = NULL,
  sampleData = NULL, groupData = NULL, summariseBy = NULL,
  colourBy = NULL, lineBy = NULL, groupBy = NULL,
  plotregion = "full", outliers = NULL, freeScale = FALSE)

## S4 method for signature 'ChIPprofile'
plotRegion(object = "ChIPprofile", gts = NULL,
  sampleData = NULL, groupData = NULL, summariseBy = NULL,
  colourBy = NULL, lineBy = NULL, groupBy = NULL,
  plotregion = "full", outliers = NULL, freeScale = FALSE)
Arguments

object A ChIPprofile object

gts A list of character vectors or GRangesList

plotregion region to plot. For combined plots with style "region", may be "start" or "end" to show full resolution of plot of edges.

groupData Dataframe of metadata for groups

sampleData Dataframe of metadata for sample

summariseBy Column names from GRanges element metadata. Formula or character vector of column names to use to collapse genomic ranges to summarised profiles. summariseBy can not be used in unjust with groups specified by gts argument.

colourBy Character vector or formula of either column names from colData(object) containing sample metadata or character vector "group" to colour by groups in gts

lineBy Character vector or formula of either column names from colData(object) containing sample metadata or character vector "group" to set line type by groups in gts

groupBy Character vector or formula of either column names from colData(object) containing sample metadata or character "group" to colour by groups in gts

outliers A numeric vector of length 1 containing proportion from limits to windsorise.

freeScale TRUE or FALSE to set whether ggplot 2 facets have their own scales. Useful for comparing multiple samples of differing depths without normalisation. Default is FALSE.

Value

A gg object from ggplot2

Author(s)

Thomas Carroll

Examples

data(chipExampleBig)
plotRegion(chipExampleBig[[2]])

data(pwmCov)

Description

This dataset contains an rlelist of motif coverage

Usage

data(pwmCov)
pwmToCoverage

Details

- pwmCov

Value

A rlelist of motif coverage

pwmToCoverage | PWM hits and motif scores as an RLElist

Description

Creates rlelist of pwm hits.
Motif score as an RLElist

Usage

pwmToCoverage(pwm, genome, min = "70\%", removeRand = FALSE, chrsOfInterest = NULL)

makeMotifScoreRle(pwm, regions, genome, extend, removeRand = FALSE, strandScore = "mean", atCentre = FALSE)

Arguments

- pwm: A PWM matrix object.
- genome: A BSgenome object
- min: pwm score (as percentage of maximum score) cutoff
- removeRand: Remove contigs with rand string
- chrsOfInterest: Chromosomes to use
- regions: GRanges object to include in pwm rlelist
- extend: bps to extend regions by
- strandScore: Method for averaging strand. Options are max, mean, sum, bothstrands
- atCentre: TRUE/FALSE. TRUE assigns score onto 1bp position at centre of motif. FALSE assigns every basepair the sum of scores of all overlapping motifs.

Value

A RLElist of motif density per base pair to be used as input to main soggi function.

Author(s)

Thomas Carroll
Examples

```r
data(pwmCov)
data(singleGRange)
```

---

| singleGRange | A single GRange |

Description

This dataset contains an rlelist of motif coverage

Usage

```r
data(singleGRange)
```

Details

- `singleGRange`

Value

A single GRanges used in motif coverage example/
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