

Package ‘genomation’

November 19, 2025

Type Package

Title Summary, annotation and visualization of genomic data

Version 1.43.0

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Description A package for summary and annotation of genomic intervals. Users can visualize and quantify genomic intervals over pre-defined functional regions, such as promoters, exons, introns, etc. The genomic intervals represent regions with a defined chromosome position, which may be associated with a score, such as aligned reads from HT-seq experiments, TF binding sites, methylation scores, etc. The package can use any tabular genomic feature data as long as it has minimal information on the locations of genomic intervals. In addition, It can use BAM or BigWig files as input.

License Artistic-2.0

LazyLoad yes

VignetteBuilder knitr

biocViews Annotation, Sequencing, Visualization, CpGIsland

Encoding latin1

URL <http://bioinformatics.mdc-berlin.de/genomation/>

BugReports <https://github.com/BIMSBbioinfo/genomation/issues>

Depends R (>= 3.0.0), grid

Imports Biostrings (>= 2.47.6), BSgenome (>= 1.47.3), data.table, Seqinfo, GenomicRanges (>= 1.31.8), GenomicAlignments (>= 1.15.6), S4Vectors (>= 0.17.25), ggplot2, gridBase, impute, IRanges (>= 2.13.12), matrixStats, methods, parallel, plotrix, plyr, readr, reshape2, Rsamtools (>= 1.31.2), seqPattern, rtracklayer (>= 1.39.7), Rcpp (>= 0.12.14)

Suggests BiocGenerics, genomationData, knitr, RColorBrewer, rmarkdown, RUnit

Collate 'combineScoreMatrixList.R' 'documentData.R'
 'genomation-classes.R' 'getRandomEnrichment.R'
 'findFeatureComb.R' 'patternMatrix.R' 'plotMatrix.R'
 'randomizeFeature.R' 'readAnnotate.R' 'readData.R'
 'scoreMatrix.R' 'scoreMatrixBin.R' 'scoreMatrixList.R' 'Ops.R'
 'test_genomation_package.R' 'deprecated_defunct.R'
 'enrichmentMatrix.R' 'RcppExports.R'

LinkingTo Rcpp

RoxygenNote 6.0.1.9000

git_url <https://git.bioconductor.org/packages/genomation>

git_branch devel

git_last_commit c618839

git_last_commit_date 2025-10-29

Repository Bioconductor 3.23

Date/Publication 2025-11-18

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annotateWithFeature	<i>Function to annotate given GRanges object with a given genomic feature</i>
---------------------	---

Description

Function to annotate given GRanges object with a given genomic feature

Usage

```
annotateWithFeature(target, feature, strand = FALSE, extend = 0,
  feature.name = NULL, intersect.chr = FALSE)

## S4 method for signature 'GRanges,GRanges'
annotateWithFeature(target, feature,
  strand = FALSE, extend = 0, feature.name = NULL,
  intersect.chr = FALSE)
```

Arguments

target	a GRanges object storing chromosome locations to be annotated
feature	a GRanges object storing chromosome locations of a feature (can be CpG islands, ChIP-seq peaks, etc)
strand	If set to TRUE, annotation features and target features will be overlapped based on strand (def:FAULT)
extend	specifying a positive value will extend the feature on both sides as much as extend

feature.name	name of the annotation feature. For example: H3K4me1,CpGisland etc. by default the name is taken from the given variable
intersect.chr	boolean, whether to select only chromosomes that are common to feature and target. FALSE by default

Value

returns an AnnotationByFeature object

Examples

```
data(cpgi)
data(promoters)
annot = annotateWithFeature(cpgi, promoters)
```

annotateWithFeatureFlank

Function to annotate a given GRanges object with promoter,exon,intron & intergenic values

Description

Function to annotate a given GRanges object with promoter,exon,intron & intergenic values

Usage

```
annotateWithFeatureFlank(target, feature, flank, feature.name = NULL,
  flank.name = "flank", strand = FALSE, intersect.chr = FALSE)

## S4 method for signature 'GRanges,GRanges,GRanges'
annotateWithFeatureFlank(target, feature,
  flank, feature.name = NULL, flank.name = "flank", strand = FALSE,
  intersect.chr = FALSE)
```

Arguments

target	a granges object storing chromosome locations to be annotated
feature	a granges object storing chromosome locations of a feature (can be CpG islands, ChIP-seq peaks, etc)
flank	a granges object storing chromosome locations of the flanks of the feature
feature.name	string for the name of the feature
flank.name	string for the name of the flanks
strand	If set to TRUE, annotation features and target features will be overlapped based on strand (def:FAULT)
intersect.chr	boolean, whether to select only chromosomes that are common to feature and target. FALSE by default

Value

returns an AnnotationByFeature object

Examples

```
data(cpgi)
data(cage)
cpgi.flanks = getFlanks(cpgi)
flank.annot = annotateWithFeatureFlank(cage, cpgi, cpgi.flanks)
```

annotateWithFeatures *Annotate given ranges with genomic features*

Description

The function annotates a target GRangesList or GRanges object as overlapping or not with the elements of a named GRangesList. This is useful to annotate your regions of interest with genomic features with arbitrary categories such as repeat classes or families, or output from genome segmentation algorithms such as chromHMM.

Usage

```
annotateWithFeatures(target, features, strand.aware = FALSE,
  intersect.chr = FALSE)

## S4 method for signature 'GRanges,GRangesList'
annotateWithFeatures(target, features,
  strand.aware = FALSE, intersect.chr = FALSE)

## S4 method for signature 'GRangesList,GRangesList'
annotateWithFeatures(target, features,
  strand.aware = FALSE, intersect.chr = FALSE)
```

Arguments

target	GRanges or GRangesList object storing chromosome locations to be annotated (e.g. chipseq peaks)
features	a named GRangesList object containing GRanges objects different set of features. The function calculates percent overlaps with and without precedence at the same time. The order of objects in GRangesList defines their precedence. If a range in target overlaps with a more precedent range in an element of features, the other overlaps from other less precedent elements will be discarded. This is useful for getting piecharts where percentages should add up to 100.
strand.aware	if set to TRUE, annotation features and target features will be overlapped based on strand (def:FALSE)

`intersect.chr` logical value, whether to select only chromosomes that are common to feature and target. FALSE by default

Value

returns an `AnnotationByFeature` object or if target is a `GRangesList`, a list of `AnnotationByFeature` objects.

See Also

see [getMembers](#), [heatTargetAnnotation](#), [plotTargetAnnotation](#)

Examples

```
library(GenomicRanges)
data(cage)
data(cpgi)
cage$tpm=NULL
gl = GRangesList(cage=cage, cpgi=cpgi)

bed.file = system.file("extdata/chr21.refseq.hg19.bed", package = "genomation")
gene.parts = readTranscriptFeatures(bed.file)
annot = annotateWithFeatures(gl, gene.parts, intersect.chr=TRUE)
```

`annotateWithGeneParts` *Annotate given object with promoter, exon, intron and intergenic regions*

Description

The function annotates `GRangesList` or `GRanges` object as overlapping with promoter,exon,intron or intergenic regions.

Usage

```
annotateWithGeneParts(target, feature, strand = FALSE,
  intersect.chr = FALSE)

## S4 method for signature 'GRanges,GRangesList'
annotateWithGeneParts(target, feature,
  strand = FALSE, intersect.chr = FALSE)

## S4 method for signature 'GRangesList,GRangesList'
annotateWithGeneParts(target, feature,
  strand = FALSE, intersect.chr = FALSE)
```

Arguments

target	GRanges or GRangesList object storing chromosome locations to be annotated (e.g. chipseq peaks)
feature	GRangesList object containing GRanges object for promoter, exons, introns and transcription start sites, or simply output of readTranscriptFeatures function
strand	If set to TRUE, annotation features and target features will be overlapped based on strand (def:FALSE)
intersect.chr	boolean, whether to select only chromosomes that are common to feature and target. FALSE by default

Value

[AnnotationByGeneParts](#) object or a list of [AnnotationByGeneParts](#) objects if target is a [GRangesList](#) object.

Examples

```
data(cage)
bed.file = system.file("extdata/chr21.refseq.hg19.bed", package = "genomation")
gene.parts = readTranscriptFeatures(bed.file)
cage.annot = annotateWithGeneParts(cage, gene.parts, intersect.chr=TRUE)
cage.annot
```

annotatGrWithGeneParts

annotatGrWithGeneParts function

Description

annotatGrWithGeneParts function

Usage

```
annotatGrWithGeneParts(gr, prom, exon, intron, strand = FALSE)
```

Arguments

gr	target object
prom	promoters
exon	exons
intron	introns
strand	logical

AnnotationByFeature-class

An S4 class that information on overlap of target features with annotation features

Description

This object is desgined to hold statistics and information about genomic feature overlaps

Slots

members a matrix showing overlap of target features with annotation genomic features

annotation a named vector of percentages

precedence a named vector of percentages

num.annotation vector

num.precedence vector

no.of.OlapFeat vector

perc.of.OlapFeat vector

AnnotationByGeneParts-class

An S4 class that information on overlap of target features with annotation features

Description

This object is desgined to hold statistics and information about genomic feature overlaps

Slots

members a matrix showing overlap of target features with annotation genomic features

annotation a named vector of percentages

precedence a named vector of percentages

num.annotation vector

num.precedence vector

no.of.OlapFeat vector

perc.of.OlapFeat vector

dist.to.TSS a data frame showing distances to TSS and gene/TSS names and strand

bed12ToExons	<i>bed12ToExons function</i>
--------------	------------------------------

Description

extracts exons from a bed12 file and puts them into GRanges object

Usage

```
bed12ToExons(ref)
```

Arguments

ref	data.frame object
-----	-------------------

bed12ToIntrons	<i>bed12ToIntrons function</i>
----------------	--------------------------------

Description

extracts introns from a bed12 file and puts them into GRanges object

Usage

```
bed12ToIntrons(ref)
```

Arguments

ref	data.frame object
-----	-------------------

binMatrix	<i>Bins the columns of a matrix using a user provided function</i>
-----------	--

Description

Bins the columns of a matrix using a user provided function

Usage

```
binMatrix(x, bin.num = NULL, fun = "mean")

## S4 method for signature 'ScoreMatrix'
binMatrix(x, bin.num = NULL, fun = "mean")

## S4 method for signature 'ScoreMatrixList'
binMatrix(x, bin.num = NULL, fun = "mean")
```

Arguments

x	ScoreMatrix or a ScoreMatrixList object
bin.num	integer number of bins in the final matrix
fun	character vector or an anonymous function that will be used for binning

Value

ScoreMatrix or ScoreMatrixList object

Examples

```
# binning the columns in a ScoreMatrix object
library(GenomicRanges)
target = GRanges(rep(c(1,2),each=7), IRanges(rep(c(1,1,2,3,7,8,9), times=2), width=5),
weight = rep(c(1,2),each=7),
strand=c('-', '-', '-', '-', '+', '-', '+', '-', '-', '-', '-', '-', '-', '+'))
windows = GRanges(rep(c(1,2),each=2), IRanges(rep(c(1,2), times=2), width=5),
strand=c('-', '+', '-', '+'))
sm = ScoreMatrix(target, windows)
bin = binMatrix(sm, bin.num=2)
```

binMax	<i>Function that computes a maximum value for each bin</i>
--------	--

Description

Function that computes a maximum value for each bin

Usage

```
binMax(x, n)
```

Arguments

x	NumericVector
n	intiger - number of bins

binMean	<i>Function that computes a mean value for each bin</i>
---------	---

Description

Function that computes a mean value for each bin

Usage

```
binMean(x, n)
```

Arguments

x	NumericVector
n	intiger - number of bins

binMedian	<i>Function that computes a median value for each bin</i>
-----------	---

Description

Function that computes a median value for each bin

Usage

```
binMedian(x, n)
```

Arguments

x	NumericVector
n	intiger - number of bins

binMin	<i>Function that computes a minimum value for each bin</i>
--------	--

Description

Function that computes a minimum value for each bin

Usage

```
binMin(x, n)
```

Arguments

x	NumericVector
n	intiger - number of bins

binner	<i>binner function</i>
--------	------------------------

Description

given a vector and length smooths the vector to a given size

Usage

```
binner(start, end, nbins)
```

Arguments

start	start position
end	end position
nbins	number of bins

binSum	<i>Function that computes a sum of values in a bin</i>
--------	--

Description

Function that computes a sum of values in a bin

Usage

```
binSum(x, n)
```

Arguments

x	NumericVector - vector of values of a bin
n	integer - number of bins

`c.ScoreMatrix``c.ScoreMatrix`

Description

Combine a scoreMatrix into a scoreMatrixList object - when a ScoreMatrix is a first argument

Usage

```
## S3 method for class 'ScoreMatrix'  
c(..., recursive = FALSE, use.names = TRUE)
```

Arguments

<code>...</code>	contains scoreMatrix and scoreMatrixList objects
<code>recursive</code>	logical
<code>use.names</code>	logical

Value

returns a scoreMatrixList object

`c.ScoreMatrixList``c.ScoreMatrixList`

Description

Combine a scoreMatrix into a scoreMatrixList object - when a ScoreMatrixList is a first argument

Usage

```
## S3 method for class 'ScoreMatrixList'  
c(..., recursive = FALSE, use.names = TRUE)
```

Arguments

<code>...</code>	contains scoreMatrix and scoreMatrixList objects
<code>recursive</code>	logical
<code>use.names</code>	logical

Value

returns a scoreMatrixList object

cage	<i>Example CAGE data set.</i>
------	-------------------------------

Description

Location and tag per million values for CAGE TSS clusters on chr21 and chr22 of human genome (hg19 assembly). The clusters are downloaded from ENCODE project downloads for NHEK cells.

Format

[GRanges](#) object

calculateOverlapSignificance	<i>function that calculates the significance of overlaps of two sets of features using randomization</i>
------------------------------	--

Description

This function calculates the significance of overlaps of two sets of features using randomization. #' It returns a distribution of overlaps of a target set with a given randomized feature set. The randomization can be constrained by supplied arguments. The function is still in Beta mode - the regions can overlap excluded regions, and the randomized regions are not disjoint. Please take care that the excluded and included regions are not too strict when compared to the total width of the ranges.

Usage

```
calculateOverlapSignificance(target, feature, chrom.sizes = NULL,
  stranded = TRUE, keep.strand.prop = TRUE, keep.chrom = TRUE,
  exclude = NULL, include = NULL, seed = NULL, nrand = 1)
```

```
## S4 method for signature 'GRanges,GRanges'
calculateOverlapSignificance(target, feature,
  chrom.sizes = NULL, stranded = TRUE, keep.strand.prop = TRUE,
  keep.chrom = TRUE, exclude = NULL, include = NULL, seed = NULL,
  nrand = 1)
```

Arguments

target	a GRanges object for which the overlap needs to be calculates
feature	a GRanges object to be randomized
chrom.sizes	sizes of chromosomes as a named vector (names are chromosomes names and elements of the vectors are lengths). , if not given sizes in GRanges object will be used if no sizes there the end of each chr will be the end last feature on each chr

stranded	if FALSE, all of the returned features will be strandless (will have "*" in the strand slot)
keep.strand.prop	If TRUE strands will have the same proportion as the features
keep.chrom	If TRUE, number of features and randomized features for a chromosome will match. Currently setting this to FALSE is not supported.
exclude	A GRanges object where no randomized feature should overlap, can be gaps or unmappable regions in the genome as an example.
include	A GRanges object which defines the boundaries of randomized features
seed	random number generator seed
nrand	number of randomizations (default:1)

Value

returns a GRanges object which is randomized version of the feature

checkBedValidity	<i>checkBedValidity function</i>
------------------	----------------------------------

Description

checks the validity of the bed data.frame if it is a legitimate bed columns

Usage

```
checkBedValidity(bed.df, type = "none")
```

Arguments

bed.df	data.frame object
type	type

checkClass	<i>checkClass function</i>
------------	----------------------------

Description

check whether the x object corresponds to the given class

Usage

```
checkClass(x, class.name, var.name = deparse(substitute(x)))
```


Arguments

x	object
class.name	class name
var.name	uses x object

compressedAndUrl2temp	<i>compressedAndUrl2temp function</i>
-----------------------	---------------------------------------

Description

compressedAndUrl2temp function

Usage

compressedAndUrl2temp(filename)

Arguments

filename	file name
----------	-----------

constrainRanges	<i>constrainRanges function</i>
-----------------	---------------------------------

Description

removes ranges that fell of the rle object does not check for the correspondence of the chromosome names - always check before using this function

Usage

constrainRanges(target, windows)

Arguments

target	target file
windows	windows

convertBed2Exons	<i>convert a data frame read-in from a bed file to a GRanges object for exons</i>
------------------	---

Description

convert a data frame read-in from a bed file to a GRanges object for exons

Usage

```
convertBed2Exons(bed.df)

## S4 method for signature 'data.frame'
convertBed2Exons(bed.df)
```

Arguments

bed.df	a data.frame where column order and content resembles a bed file with 12 columns
--------	--

Value

GRanges object

Note

one bed track per file is only accepted, the bed files with multiple tracks will cause an error

Examples

```
file = system.file('extdata/chr21.refseq.hg19.bed', package='genomation')
bed12 = read.table(file)
exons = convertBed2Exons(bed12)
head(exons)
```

convertBed2Introns	<i>convert a data frame read-in from a bed file to a GRanges object for introns</i>
--------------------	---

Description

convert a data frame read-in from a bed file to a GRanges object for introns

Usage

```
convertBed2Introns.bed.df)

## S4 method for signature 'data.frame'
convertBed2Introns.bed.df)
```

Arguments

bed.df a data.frame where column order and content resembles a bed file with 12 columns

Value

[GRanges](#) object

Note

one bed track per file is only accepted, the bed files with multiple tracks will cause an error

Examples

```
file = system.file('extdata/chr21.refseq.hg19.bed', package='genomation')
bed12 = read.table(file)
introns = convertBed2Introns.bed12)
head(introns)
```

convertBedDf	<i>convert a data frame read-in from a bed file to a GRanges object</i>
--------------	---

Description

convert a data frame read-in from a bed file to a GRanges object

Usage

```
convertBedDf.bed)

## S4 method for signature 'data.frame'
convertBedDf.bed)
```

Arguments

bed a data.frame where column order and content resembles a bed file with 12 columns

Value

[GRanges](#) object

Note

one bed track per file is only accepted, the bed files with multiple tracks will cause an error
bed files are expected not to have header lines

cpgi	<i>Example CpG island data set.</i>
------	-------------------------------------

Description

CpG islands of hg19 assembly of human genome on chr21 and chr22. Downloaded from UCSC genome browser.

Format

GRanges object

detectUCSCheader	<i>detectUCSCheader function</i>
------------------	----------------------------------

Description

detects UCSC header (and first track)

Usage

detectUCSCheader(filename)

Arguments

filename	file name
----------	-----------

distance2NearestFeature	<i>distance2NearestFeature function</i>
-------------------------	---

Description

distance2NearestFeature function

Usage

distance2NearestFeature(g.idh, tss)

Arguments

g.idh	target object
tss	TSSes

enrichmentMatrix	<i>Compute an enrichment of IP over control both stored in ScoreMatrix objects</i>
------------------	--

Description

This is an enrichmentMatrix function for ScoreMatrix objects, that enables to normalize ChIP-seq signals with respect to IgG or input DNA control.

Usage

```
\S4method{enrichmentMatrix}{ScoreMatrix,ScoreMatrix}(IP, control)
```

Arguments

IP	ScoreMatrix object storing an IP sample
control	ScoreMatrix object storing a control sample

Value

ScoreMatrix object

Note

The function computes an enrichment of IP over control as follow: Suppose both IP and control are ScoreMatrix objects that have same dimensions. Then, the enrichment is calculated usign a formula: $\log_2((IP + 1) / (control + 1))$.

See Also

[ScoreMatrix](#)

Examples

```
#load IP and control BAM files and create ScoreMatrix objects
library('genomationData')
bam.file_IP <- system.file("extdata",
  "wgEncodeBroadHistoneH1hesoSuz12051317AlnRep1.chr21.bam", package = "genomationData")
bam.file_c <- system.file("extdata",
  "wgEncodeBroadHistoneH1hesoSctcfStdAlnRep1.chr21.bam", package = "genomationData")
data(promoters)
IP <- ScoreMatrix(target = bam.file_IP, windows = promoters, type = 'bam')
control <- ScoreMatrix(target = bam.file_c, windows = promoters, type = 'bam')

# compute an enrichment of IP over control
enrichmentMatrix(IP, control)
```

enrichmentMatrix, ScoreMatrixList, ScoreMatrix-method

Compute an enrichment of IP (stored in ScoreMatrixList object) over control (stored in ScoreMatrix object)

Description

This is an enrichmentMatrix function for IP ScoreMatrixList object and control ScoreMatrix object, that enables to normalize ChIP-seq signals with respect to IgG or input DNA control.

Usage

```
\S4method{enrichmentMatrix}{ScoreMatrixList, ScoreMatrix}(IP, control)
```

Arguments

IP [ScoreMatrixList](#) object storing IP samples
control [ScoreMatrix](#) storing control sample

Value

ScoreMatrixList object

Note

The function computes an enrichment of IP over control as follow: Suppose both IP and control are ScoreMatrix objects that have same dimensions. Then, the enrichment is calculated using a formula: $\log_2((IP + 1) / (control + 1))$.

See Also

[ScoreMatrixList](#), [ScoreMatrix](#)

Examples

```
#load IP and control BAM files and create ScoreMatrix objects
library('genomationData')
data(promoters)
bam.file_IP_1 <- system.file("extdata",
  "wgEncodeSydhTfbsH1hesCZnf143IggrabAlnRep1.chr21.bam", package = "genomationData")
IP_1 <- ScoreMatrix(target = bam.file_IP_1, windows = promoters, type = 'bam')

bam.file_IP_2 <- system.file("extdata",
  "wgEncodeBroadHistoneH1hesCZnf143IggrabAlnRep1.chr21.bam", package = "genomationData")
IP_2 <- ScoreMatrix(target=bam.file_IP_2, windows = promoters, type = 'bam')

bam.file_c <- system.file("extdata",
  "wgEncodeBroadHistoneH1hesCZnf143IggrabAlnRep1.chr21.bam", package = "genomationData")
control <- ScoreMatrix(target = bam.file_c, windows = promoters, type = 'bam')
```

```
# create a ScoreMatrixList object storing IP ScoreMatrix objects
sml_IP <- ScoreMatrixList(list(IP1 = IP_1, IP2 = IP_2))

# compute an enrichment of IP over control
enrichmentMatrix(sml_IP, control)
```

enrichmentMatrix, ScoreMatrixList, ScoreMatrixList-method

Compute an enrichment of IP over control both stored in ScoreMatrixList objects

Description

This is an enrichmentMatrix function for ScoreMatrixList objects, that enables to normalize ChIP-seq signals with respect to IgG or input DNA control.

Usage

```
\S4method{enrichmentMatrix}{ScoreMatrixList, ScoreMatrixList}(IP, control)
```

Arguments

IP	ScoreMatrixList object storing IP samples
control	ScoreMatrixList storing control samples

Value

ScoreMatrixList object

Note

The function computes an enrichment of IP over control as follow: Suppose both IP and control are ScoreMatrix objects that have same dimensions. Then, the enrichment is calculated using a formula: $\log_2((IP + 1) / (control + 1))$.

See Also

[ScoreMatrixList](#)

Examples

```
#load IP and control BAM files and create ScoreMatrix objects
library('genomationData')
data(promoters)
bam.file_IP_1 <- system.file("extdata",
  "wgEncodeSydhTfbsH1hescZnf143IggrabAlnRep1.chr21.bam", package = "genomationData")
IP_1 <- ScoreMatrix(target = bam.file_IP_1, windows = promoters, type = 'bam')
```

```
bam.file_IP_2 <- system.file("extdata",
"wgEncodeBroadHistoneH1hesoSuz12051317AlnRep1.chr21.bam", package = "genomationData")
IP_2 <- ScoreMatrix(target=bam.file_IP_2, windows = promoters, type = 'bam')

bam.file_c <- system.file("extdata",
"wgEncodeBroadHistoneH1hesoSctcfStdAlnRep1.chr21.bam", package = "genomationData")
control <- ScoreMatrix(target = bam.file_c, windows = promoters, type = 'bam')

# create a ScoreMatrixList object storing IP ScoreMatrix objects
sml_IP <- ScoreMatrixList(list(IP1 = IP_1, IP2 = IP_2))

# create a ScoreMatrixList object storing control ScoreMatrix objects
sml_control <- ScoreMatrixList(list(c1 = control, c2 = control))

# compute an enrichment of IP over control
enrichmentMatrix(sml_IP, sml_control)
```

file.ext	<i>file.ext function</i>
----------	--------------------------

Description

Extract file extension from file path

Usage

```
file.ext(x)
```

Arguments

x file

findFeatureComb	<i>Find combinations of genomic features</i>
-----------------	--

Description

Provided a GRangesList, finds the combinations of sets of ranges. It is mostly used to look at the combinatorics of transcription factor binding. The function works by, firstly, constructing a union of all ranges in the list, which are then designated by the combinatorics of overlap with the original sets. A caveat of this approach is that the number of possible combinations increases exponentially, so we would advise you to use it with up to 6 data sets. If you wish to take a look at a greater number of factors, methods like self organizing maps or ChromHMM might be more appropriate.

Usage

```
findFeatureComb(gl, width=0, use.names=FALSE, collapse.char=':')

## S4 method for signature 'GRangesList'
findFeatureComb(gl, width = 0, use.names = FALSE,
  collapse.char = ":")
```

Arguments

<code>gl</code>	a <code>GRangesList</code> object, containing ranges for which represent regions enriched for transcription factor binding
<code>width</code>	integer is the requested width of each enriched region. If 0 the ranges are not resized, if a positive integer, the width of all ranges is set to that number. Ranges are resized relative to the center of original ranges.
<code>use.names</code>	a boolean which tells the function whether to return the resulting ranges with a numeric vector which designates each class (the default), or to construct the names of each class using the names from the <code>GRangesList</code>
<code>collapse.char</code>	a character which will be used to separate the class names if <code>use.names=TRUE</code> . The default is ':'

Value

a `GRanges` object

Examples

```
library(GenomicRanges)
g = GRanges(paste('chr', rep(1:2, each=3), sep=''), IRanges(rep(c(1,5,9), times=2), width=3))
gl = GRangesList(g1=g, g2=g[2:5], g3=g[3:4])
findFeatureComb(gl)
findFeatureComb(gl, use.names=TRUE)
```

galpTo2Ranges

galpTo2Ranges function

Description

galpTo2Ranges function

Usage

```
galpTo2Ranges(x)
```

Arguments

<code>x</code>	object
----------------	--------

genes

Example RefSeq genes data set.

Description

RefSeq genes of hg19 assembly of human genome on chr21 and chr22. Downloaded from UCSC genome browser.

Format

[GRanges](#) object

getAssociationWithTSS *Get distance to nearest TSS and gene id from AnnotationByGeneParts*

Description

This accessor function gets the nearest TSS, its distance to target feature, strand and name of TSS/gene from AnnotationByGeneParts object

Usage

```
getAssociationWithTSS(x)
```

```
## S4 method for signature 'AnnotationByGeneParts'
getAssociationWithTSS(x)
```

Arguments

x a AnnotationByGeneParts object

Value

RETURNS a data.frame containing row number of the target features, distance of target to nearest TSS, TSS/Gene name, TSS strand

Examples

```
data(cage)
bed.file = system.file("extdata/chr21.refseq.hg19.bed", package = "genomation")
gene.parts = readTranscriptFeatures(bed.file)
cage.annot = annotateWithGeneParts(cage, gene.parts, intersect.chr=TRUE)
head(getAssociationWithTSS(cage.annot))
```

getColors

*getColors function***Description**

gets colors for a factor variable

Usage

```
getColors(n)
```

Arguments

n n

getFeatsWithTargetsStats

Get the percentage/count of annotation features overlapping with target features from AnnotationByFeature

Description

This function retrieves percentage/number of annotation features overlapping with targets. For example, if AnnotationByFeature object is containing statistics of differentially methylated regions overlapping with gene annotation. This function will return number/percentage of introns,exons and promoters overlapping with differentially methylated regions.

Usage

```
getFeatsWithTargetsStats(x,percentage=TRUE)
```

```
## S4 method for signature 'AnnotationByFeature'
getFeatsWithTargetsStats(x, percentage = TRUE)
```

Arguments

x a AnnotationByFeature object

percentage TRUE|FALSE. If TRUE percentage of annotation features will be returned. If FALSE, number of annotation features will be returned

Value

RETURNS a vector of percentages or counts showing quantity of annotation features overlapping with target features

Examples

```
data(cage)
bed.file=system.file("extdata/chr21.refseq.hg19.bed", package = "genomation")
gene.parts = readTranscriptFeatures(bed.file)
cage.annot = annotateWithGeneParts(cage, gene.parts, intersect.chr=TRUE)
getFeatsWithTargetsStats(cage.annot)
```

getFlanks	<i>Function to get upstream and downstream adjacent regions to a genomic feature such as CpG islands</i>
-----------	--

Description

Function to get upstream and downstream adjacent regions to a genomic feature such as CpG islands

Usage

```
getFlanks(grange, flank=2000, clean=TRUE)

## S4 method for signature 'GRanges'
getFlanks(grange, flank = 2000, clean = TRUE)
```

Arguments

grange	GRanges object for the feature
flank	number of basepairs for the flanking regions
clean	If set to TRUE, flanks overlapping with other main features will be trimmed, and overlapping flanks will be removed. This will remove multiple counts when other features overlap with flanks

Value

GRanges object for flanking regions

Examples

```
data(cpgi)
cpgi.flanks = getFlanks(cpgi)
head(cpgi.flanks)
```

getMembers	<i>Get the membership slot of AnnotationByFeature</i>
------------	---

Description

Membership slot defines the overlap of target features with annotation features. For example, if a target feature overlaps with an exon

Usage

```
getMembers(x)

## S4 method for signature 'AnnotationByFeature'
getMembers(x)
```

Arguments

x a AnnotationByFeature object

Value

matrix showing overlap of target features with annotation features. 1 for overlap, 0 for non-overlap

getRandomEnrichment	<i>get enrichment based on randomized feature overlap</i>
---------------------	---

Description

This function measures the association between two genomic features by randomizing one feature and counting the overlaps in randomized sets. That is to say, query feature will be randomly distributed over the genome (constrained by provided options), and the overlap of target with these randomized features will be measured.

Usage

```
getRandomEnrichment(target, query, randomizations = 1000, rand.set = NULL,
  ...)

## S4 method for signature 'GRanges,GRanges'
getRandomEnrichment(target, query,
  randomizations = 1000, rand.set = NULL, ...)
```

Arguments

target	a GRanges object to be overlapped with query
query	a GRanges object that will be randomly placed across the genome and overlap of these random regions with target will be the background distribution of association between target and query.
randomizations	number of times the features to be shuffled
rand.set	instead of randomly placing features in query one can supply an already shuffled set of query genomic features.
...	other parameters to be passed to randomizeFeature function. These parameters ccontrol how randomization is done.

Value

returns a RandomEnrichment object

See Also

[randomizeFeature](#)

Examples

```
data(cage)
data(cpgi)

enr = getRandomEnrichment(cage, cpgi, randomizations=50)
```

getTargetAnnotationStats

Get the percentage of target features overlapping with annotation from AnnotationByFeature

Description

This function retrieves percentage/number of target features overlapping with annotation

Usage

```
getTargetAnnotationStats(x,percentage=TRUE,precedence=TRUE)

## S4 method for signature 'AnnotationByFeature'
getTargetAnnotationStats(x, percentage = TRUE,
  precedence = TRUE)
```

Arguments

x	a AnnotationByFeature object
percentage	TRUE FALSE. If TRUE percentage of target features will be returned. If FALSE, number of target features will be returned
precedence	TRUE FALSE. If TRUE there will be a hierarchy of annotation features when calculating numbers (with promoter>exon>intron precedence) That means if a feature overlaps with a promoter it will be counted as promoter overlapping only, or if it is overlapping with a an exon but not a promoter, #' it will be counted as exon overlapping only whether or not it overlaps with an intron.

Value

a vector of percentages or counts showing quantity of target features overlapping with annotation

Examples

```
data(cage)
bed.file=system.file("extdata/chr21.refseq.hg19.bed", package = "genomation")
gene.parts = readTranscriptFeatures(bed.file)
cage.annot=annotateWithGeneParts(cage, gene.parts, intersect.chr=TRUE)
getTargetAnnotationStats(cage.annot)
```

gffToGRanges	<i>Converts a gff formatted data.frame into a GenomicRanges object. The GenomicRanges object needs to be properly formatted for the function to work.</i>
--------------	---

Description

Converts a gff formatted data.frame into a GenomicRanges object. The GenomicRanges object needs to be properly formatted for the function to work.

Usage

```
gffToGRanges(gff.file, filter = NULL, zero.based = FALSE, ensembl = FALSE)
```

Arguments

gff.file	path to a gff formatted file. The file can end in .gz, .bz2, .xz, or .zip and/or start with http:// or ftp://. If the file is not compressed it can also start with https:// or ftps://.
filter	a character designating which elements to retain from the gff file (e.g. exon, CDS, ...)
zero.based	boolean whether the coordinates are 0 or 1 based. 0 is the default
ensembl	boolean if TRUE, add the chr prefix to seqlevels. FALSE by default

Value

returns a GenomicRanges object

Examples

```
gff.file = system.file('extdata/chr21.refseq.hg19.gtf', package='genomation')
gff = gffToGRanges(gff.file)
```

heatMatrix

Draw a heatmap of a given ScoreMatrix object

Description

The function makes a heatmap out of given ScoreMatrix object. If desired it can use clustering using given clustering function (e.g. k-means) and plot cluster color codes as a sidebar. In addition, user can define groups of rows using 'group' argument.

Usage

```
heatMatrix(mat, grid = FALSE, col = NULL, xcoords = NULL, group = NULL,
  group.col = NULL, order = FALSE, user.order = FALSE, winsorize = c(0,
  100), clustfun = NULL, main = "", legend.name = NULL, cex.legend = 1,
  xlab = NULL, cex.main = 1, cex.lab = 1, cex.axis = 1,
  newpage = TRUE)
```

Arguments

mat	a ScoreMatrix object
grid	if TRUE, grid graphics will be used. if FALSE, base graphics will be used on the top level, so users can use par(mfrow) or par(mfcol) prior to calling the function. Default:FALSE
col	a vector of colors, such as the ones created by heat.colors(10). If NULL (which is default), jet color scheme (common in matlab plots) will be used.
xcoords	a vector of numbers showing relative positions of the bases or windows. It must match the number of columns in the ScoreMatrix. Alternatively, it could be a numeric vector of two elements. Such as c(0,100) showing the relative start and end coordinates of the first and last column of the ScoreMatrix object.
group	a list of vectors of row numbers or a factor. This grouping is used for row side colors of the heatmap. If it is a list, each element of the list must be a vector of row numbers. Names of the elements of the list will be used as names of groups. If group is a factor, it's length must match the number of rows of the matrix, and factor levels will be used as the names of the groups in the plot.
group.col	a vector of color names to be used at the row side colors if group argument is given or clustfun function is given.

order	Logical indicating if the rows should be ordered or not (Default:FALSE). If order=TRUE the matrix will be ordered with rowSums(mat) values in descending order. If group argument is provided, first the groups will be ordered in descending order of sums of rows then, everything within the clusters will be ordered by sums of rows. If clustfun is given then rows within clusters will be order in descending order of sums of rows.
user.order	a numerical vector indicating the order of groups/clusters (it works only when group or clustfun argument is given).
winsorize	Numeric vector of two, defaults to c(0,100). This vector determines the upper and lower percentile values to limit the extreme values. For example, c(0,99) will limit the values to only 99th percentile, everything above the 99 percentile will be equalized to the value of 99th percentile. This is useful for visualization of matrices that have outliers.
clustfun	a function for clustering rows of mat that returns a vector of integers indicating the cluster to which each point is allocated (a vector of cluster membership), e.g. k-means algorithm with 3 centers: function(x) kmeans(x, centers=3)\$cluster. By default FALSE.
main	a character string for the plot title
legend.name	a character label plotted next to the legend
cex.legend	A numerical value giving the amount by which legend axis marks should be magnified relative to the default
xlab	label a character string for x-axis of the heatmap
cex.main	A numerical value giving the amount by which plot title should be magnified
cex.lab	A numerical value giving the amount by which axis labels (including 'legend.name') should be magnified relative to the default.
cex.axis	A numerical value giving the amount by which axis marks should be magnified relative to the default
newpage	logical indicating if grid.newpage() function should be invoked if grid=TRUE.

Value

returns clustering result invisibly, if clustfun is defined

Examples

```
data(cage)
data(promoters)
scores1=ScoreMatrix(target=cage, windows=promoters, strand.aware=TRUE,
                     weight.col="tpm")

set.seed(1000)

heatMatrix(mat=scores1, legend.name="tpm", winsorize=c(0,99), xlab="region around TSS",
           xcoords=-1000:1000,
           cex.legend=0.8, main="CAGE clusters on promoters", cex.lab=1,
           cex.axis=0.9, grid=FALSE)
```

```
## examples using clustering functions
## k-means
cl1 <- function(x) kmeans(x, centers=3)$cluster
set.seed(1000)
heatMatrix(mat=scores1, legend.name="tpm", winsorize=c(0,99), xlab="region around TSS",
           xcoords=-1000:1000, clustfun=cl1,
           cex.legend=0.8, main="CAGE clusters on promoters", cex.lab=1,
           cex.axis=0.9, grid=FALSE,
           user.order=c(1,3,2))

## hierarchical clustering
cl2 <- function(x) cutree(hclust(dist(x), method="complete"), k=3)
set.seed(1000)
heatMatrix(mat=scores1, legend.name="tpm", winsorize=c(0,99), xlab="region around TSS",
           xcoords=-1000:1000, clustfun=cl2,
           cex.legend=0.8, main="CAGE clusters on promoters", cex.lab=1,
           cex.axis=0.9, grid=FALSE)
```

heatMeta

Heatmap for meta-region profiles

Description

Function calculates meta-profile(s) from a `ScoreMatrix` or a `ScoreMatrixList`, then produces a heatmap or a set of stacked heatmaps for meta-region profiles

Usage

```
heatMeta(mat, centralTend = "mean", profile.names = NULL, xcoords = NULL,
         col = NULL, meta.rescale = FALSE, winsorize = c(0, 100),
         legend.name = NULL, cex.legend = 1, xlab = NULL, main = "",
         cex.lab = 1, cex.axis = 1)
```

Arguments

<code>mat</code>	<code>ScoreMatrix</code> or <code>ScoreMatrixList</code> to be plotted
<code>centralTend</code>	a character that determines central tendency of meta-profile(s). It takes "mean" (default) or "median".
<code>profile.names</code>	a character vector for names of profiles. If <code>NULL</code> , the names will be taken from <code>names(mat)</code> if <code>mat</code> is a <code>ScoreMatrixList</code> object.
<code>xcoords</code>	a vector of numbers showing relative positions of the bases or windows. It must match the number of columns in the <code>ScoreMatrix</code> . For example: if there are 2001 elements in the matrices which are base-pair resolution and they are centered around an anchor point like TSS, the <code>xcoords</code> argument should be -1000:1000. This argument is used to plot accurate x-axis labels for the plots. If <code>NULL</code> it will be equal to <code>1:ncol(mat)</code> .

col	a vector of color palette. color scheme to be used. If NULL, a version of jet colors will be used.
meta.rescale	if TRUE meta-region profiles are scaled to 0 to 1 range by subtracting the min from profiles and dividing them by max-min.
winsorize	Numeric vector of two, defaults to c(0,100). This vector determines the upper and lower percentile values to limit the extreme values. For example, c(0,99) will limit the values to only 99th percentile, everything above the 99 percentile will be equalized to the value of 99th percentile. This is useful for visualization of matrices that have outliers.
legend.name	a character label plotted next to the legend
cex.legend	A numerical value giving the amount by which legend axis marks should be magnified relative to the default
xlab	label a character string for x-axis
main	a character string for the plot title
cex.lab	A numerical value giving the amount by which axis labels (including 'legend.name') should be magnified relative to the default.
cex.axis	A numerical value giving the amount by which axis marks should be magnified relative to the default

Value

returns meta-profile matrix invisibly.

Examples

```
data(cage)
data(promoters)
scores1=ScoreMatrix(target=cage, windows=promoters, strand.aware=TRUE)
data(cpgi)
scores2=ScoreMatrix(target=cpgi, windows=promoters, strand.aware=TRUE)

x=new("ScoreMatrixList", list(scores1, scores2))

heatMeta(mat=x, legend.name="fg", cex.legend=0.8, main="fdf", cex.lab=6,
         cex.axis=0.9)
```

heatTargetAnnotation	<i>Plots the percentage of overlapping intervals with genomic features in a heatmap</i>
----------------------	---

Description

This function plots a heatmap for percentage of overlapping ranges with provided genomic features. The input object is a list of AnnotationByFeature objects, which contains necessary information about overlap statistics to make the plot.

Usage

```
heatTargetAnnotation(l, cluster = FALSE, col = c("white", "blue"),
  precedence = FALSE, plot = TRUE)
```

Arguments

<code>l</code>	a list of <code>AnnotationByFeature</code> objects. This could be returned by annotateWithFeatures function.
<code>cluster</code>	TRUE/FALSE. If TRUE the heatmap is going to be clustered using a default hierarchical clustering scheme.
<code>col</code>	a vector of two colors that will be used for interpolation. The first color is the lowest one, the second is the highest one
<code>precedence</code>	TRUE/FALSE. If TRUE the precedence of annotation features will be used when plotting. The precedence will be taken from the <code>GRangesList</code> used as annotation. The first <code>GRanges</code> will be treated as most important, and the second as the second most important and so on. Such that, if an interval overlaps with features on that is part of the first <code>GRanges</code> object in the annotation <code>GRangesList</code> , the rest of its overlaps with other elements in the annotation <code>GRangesList</code> will be ignored. This feature is important to have if the user desires that percentage of overlaps adds up to 100. This is only possible when the annotation features are non-overlapping with each other or there is a hierarchy/precedence among them such as (with promoter>exon>intron precedence). In this case, anything that overlaps with a promoter annotation will only be counted as promoter even if it overlaps with exons or introns.
<code>plot</code>	If FALSE, does not plot the heatmap just returns the matrix used to make the heatmap

Value

returns the matrix used to make the heatmap when `plot` FALSE, otherwise returns `ggplot2` object which can be modified further.

See Also

see [getMembers](#), [annotateWithFeatures](#)

Examples

```
library(GenomicRanges)
data(cage)
data(cpgi)
cage$tpm=NULL
gl = GRangesList(cage=cage, cpgi=cpgi)

bed.file = system.file("extdata/chr21.refseq.hg19.bed", package = "genomation")
gene.parts = readTranscriptFeatures(bed.file)
annot = annotateWithFeatures(gl, gene.parts, intersect.chr=TRUE)
```

```
heatTargetAnnotation(annot)
```

```
intersectScoreMatrixList
```

Get common rows from all matrices in a ScoreMatrixList object

Description

Returns a intersection of rows for each matrix in a ScoreMatrixList object. This is done using the rownames of each element in the list.

Usage

```
intersectScoreMatrixList(sml, reorder = FALSE)
```

```
## S4 method for signature 'ScoreMatrixList'  
intersectScoreMatrixList(sml, reorder = FALSE)
```

Arguments

sml	a ScoreMatrixList object
reorder	if TRUE ScoreMatrix objects in the list are sorted based on their common row ids.

Value

ScoreMatrixList object

Examples

```
library(GenomicRanges)  
target = GRanges(rep(c(1,2),each=7),  
                  IRanges(rep(c(1,1,2,3,7,8,9), times=2), width=5),  
                  weight = rep(c(1,2),each=7))  
  
windows1 = GRanges(rep(c(1,2),each=2),  
                   IRanges(rep(c(1,2), times=2), width=5),  
                   strand=c('-', '+', '-', '+'))  
windows2 = windows1[c(1,3)]  
sml = as(list(ScoreMatrix(target, windows1),  
             ScoreMatrix(target, windows2)), 'ScoreMatrixList')  
sml  
intersectScoreMatrixList(sml)
```

listSliceMax	<i>Function creates a matrix storing data with desirable number of bins for each window</i>
--------------	---

Description

listSliceMax() function calls the binMax() function

Usage

```
listSliceMax(xlist, n)
```

Arguments

xlist	List of vectors storing values of a bin
n	intiger - number of bins

listSliceMean	<i>Function creates a matrix storing data with desirable number of bins for each window</i>
---------------	---

Description

listSliceMean() function calls the binMean() function

Usage

```
listSliceMean(xlist, n)
```

Arguments

xlist	List of vectors storing values of a bin
n	intiger - number of bins

listSliceMedian	<i>Function creates a matrix storing data with desirable number of bins for each window</i>
-----------------	---

Description

listSliceMean() function calls the binMedian() function

Usage

```
listSliceMedian(xlist, n)
```

Arguments

xlist	List of vectors storing values of a bin
n	intiger - number of bins

listSliceMin	<i>Function creates a matrix storing data with desirable number of bins for each window</i>
--------------	---

Description

listSliceMin() function calls the binMin() function

Usage

```
listSliceMin(xlist, n)
```

Arguments

xlist	List of vectors storing values of a bin
n	intiger - number of bins

listSliceSum	<i>Function creates a matrix storing data with desirable number of bins for each window</i>
--------------	---

Description

listSliceSum() function calls the binSum() function

Usage

```
listSliceSum(xlist, n)
```

Arguments

xlist	List of vectors storing values of a bin
n	integer - number of bins

Max_c	<i>Function that computes a max value</i>
-------	---

Description

Function that computes a max value

Usage

```
Max_c(x)
```

Arguments

x	NumericVector
---	---------------

Mean_c	<i>Function that computes a mean value</i>
--------	--

Description

Function that computes a mean value

Usage

```
Mean_c(x)
```

Arguments

x	NumericVector
---	---------------

Median_c	<i>Function that computes a median value</i>
----------	--

Description

Function that computes a median value

Usage

Median_c(x)

Arguments

x	NumericVector
---	---------------

Min_c	<i>Function that computes a min value</i>
-------	---

Description

Function that computes a min value

Usage

Min_c(x)

Arguments

x	NumericVector
---	---------------

multiHeatMatrix	<i>Draw multiple heatmaps from a ScoreMatrixList object</i>
-----------------	---

Description

The function plots multiple heatmaps for a ScoreMatrixList object side by side. Each matrix can have different color schemes but it is essential that each matrix is obtained from same regions or neighbouring regions.

Usage

```
multiHeatMatrix(sml, grid = TRUE, col = NULL, xcoords = NULL,
  group = NULL, group.col = NULL, order = FALSE, user.order = FALSE,
  winsorize = c(0, 100), clustfun = FALSE, clust.matrix = NULL,
  column.scale = TRUE, matrix.main = NULL, common.scale = FALSE,
  legend = TRUE, legend.name = NULL, cex.legend = 0.8, xlab = NULL,
  cex.lab = 1, cex.main = 1, cex.axis = 0.8, newpage = TRUE)
```

Arguments

sml	a ScoreMatrixList object
grid	if TRUE, grid graphics will be used. if FALSE, base graphics will be used on the top level, so users can use par(mfrow) or par(mfcol) prior to calling the function. Default:FALSE
col	a color palette or list of color palettes, such as list(heat.colors(10),topo.colors(10)). If it is a list, its length must match the number of matrices to be plotted. If it is a single palette every heatmap will have the same colors.
xcoords	a vector of numbers showing relative positions of the bases or windows or a list of vectors. The elements of the list must match the number of columns in the corresponding ScoreMatrix. Alternatively, the elements could be a numeric vector of two elements. Such as c(0,100) showing the relative start and end coordinates of the first and last column of the ScoreMatrix object. The remaining coordinates will be automatically matched in this case. If the argument is not a list but a single vector, then all heatmaps will have the same coordinate on their x-axis.
group	a list of vectors of row numbers or a factor. The rows will be reordered to match their grouping. The grouping is used for row side colors of the heatmap. If it is a list, each element of the list must be a vector of row numbers. Names of the elements of the list will be used as names of groups. If group is a factor, its length must match the number of rows of the matrix, and factor levels will be used as the names of the groups in the plot.
group.col	a vector of color names to be used at the row side colors if group and clustfun arguments are given
order	Logical indicating if the rows should be ordered or not (Default:FALSE). If order=TRUE the matrix will be ordered with rowSums(mat) values in descending order. If group argument is provided, first the groups will be ordered in descending order of sums of rows then, everything within the clusters will be ordered by sums of rows. If clustfun is given then rows within clusters will be ordered in descending order by sums of rows.
user.order	a numerical vector indicating the order of groups/clusters (it works only when group or clustfun argument is given).
winsorize	Numeric vector of two, defaults to c(0,100). This vector determines the upper and lower percentile values to limit the extreme values. For example, c(0,99) will limit the values to only 99th percentile for a matrix, everything above the 99th percentile will be equalized to the value of 99th percentile. This is useful for visualization of matrices that have outliers.

<code>clustfun</code>	a function for clustering rows of <code>mat</code> that returns a vector of integers indicating the cluster to which each point is allocated (a vector of cluster membership), e.g. k-means algorithm with 3 centers: <code>function(x) kmeans(x, centers=3)\$cluster</code> . By default <code>FALSE</code> .
<code>clust.matrix</code>	a numerical vector of indexes or a character vector of names of the <code>ScoreMatrix</code> objects in <code>'sml'</code> to be used in clustering (if <code>clustfun</code> argument is provided). By default all matrices are clustered. Matrices that are not indicated in <code>clust.matrix</code> are ordered according to result of clustering algorithm.
<code>column.scale</code>	Logical indicating if matrices should be scaled or not, prior to clustering or ordering. Setting this to <code>TRUE</code> scales the columns of the matrices using, <code>scale()</code> function. scaled columns are only used for clustering or ordering. Original scores are displayed for heatmaps.
<code>matrix.main</code>	a vector of strings for the titles of the heatmaps. If <code>NULL</code> titles will be obtained from names of the <code>ScoreMatrix</code> objects in the <code>ScoreMatrixList</code> objects.
<code>common.scale</code>	if <code>TRUE</code> (Default: <code>FALSE</code>) all the heatmap colors will be coming from the same score scale, although each heatmap color scale can be different. The color intensities will be coming from the same scale. The scale will be determined by minimum of all matrices and maximum of all matrices. This is useful when all matrices are on the same score scale. If <code>FALSE</code> , the color scale will be determined by minimum and maximum of each matrix individually.
<code>legend</code>	if <code>TRUE</code> and color legend for the heatmap is drawn.
<code>legend.name</code>	a vector of legend labels to be plotted with legends of each heatmap. If it is a length 1 vector, all heatmaps will have the same legend label.
<code>cex.legend</code>	A numerical value giving the amount by which legend axis marks should be magnified relative to the default
<code>xlab</code>	a vector of character strings for x-axis labels of the heatmaps. if it is length 1, all heatmaps will have the same label.
<code>cex.lab</code>	A numerical value giving the amount by which axis labels (including <code>'legend.name'</code>) should be magnified relative to the default.
<code>cex.main</code>	A numerical value giving the amount by which plot title should be magnified
<code>cex.axis</code>	A numerical value giving the amount by which axis marks should be magnified relative to the default
<code>newpage</code>	logical indicating if <code>grid.newpage()</code> function should be invoked if <code>grid=TRUE</code> .

Value

invisibly returns the order of rows, if `clustfun` is provided and/or `order=TRUE`

Examples

```
data(cage)
data(promoters)
scores1=ScoreMatrix(target=cage, windows=promoters, strand.aware=TRUE)

data(cpgi)
scores2=ScoreMatrix(target=cpgi, windows=promoters, strand.aware=TRUE)
```

```

sml=new("ScoreMatrixList",list(a=scores1,b=scores2))

# use with k-means
multiHeatMatrix(sml,
                 clustfun=function(x) kmeans(x, centers=2)$cluster,
                 cex.axis=0.8,xcoords=c(-1000,1000),
                 winsorize=c(0,99),
                 legend.name=c("tpm","coverage"),xlab="region around TSS")

# use with hierarchical clustering
cl2 <- function(x) cutree(hclust(dist(x), method="complete"), k=2)
multiHeatMatrix(sml,legend.name="tpm",winsorize=c(0,99),xlab="region around TSS",
                 xcoords=-1000:1000,clustfun=cl2,
                 cex.legend=0.8,cex.lab=1,
                 cex.axis=0.9,grid=FALSE)

# use different colors
require(RColorBrewer)
col.cage= brewer.pal(9,"Blues")
col.cpgi= brewer.pal(9,"YlGn")
multiHeatMatrix(sml,
                 clustfun=function(x) kmeans(x, centers=2)$cluster,
                 cex.axis=0.8,xcoords=c(-1000,1000),
                 winsorize=c(0,99),col=list(col.cage,col.cpgi),
                 legend.name=c("tpm","coverage"),xlab="region around TSS")

```

Ops,numeric,ScoreMatrixList-method

Ops method for a ScoreMatrixList object. It enables to use arithmetic, indicator and logic operations on ScoreMatrixList objects.

Description

Arithmetic method for ScoreMatrixList

Usage

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

Arguments

e1	the numeric value
e2	the ScoreMatrixList object

Value

ScoreMatrixList

Ops,ScoreMatrix,ScoreMatrix-method*Ops method for a ScoreMatrix object. It enables to use arithmetic, indicator and logic operations on ScoreMatrix objects.*

Description

Arithmetic method for ScoreMatrix and ScoreMatrixList classes

Usage

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

Arguments

e1 the [ScoreMatrix](#) object or numeric value
e2 the [ScoreMatrix](#) object or numeric value

Value

ScoreMatrix

Ops,ScoreMatrixList,numeric-method*Ops method for a ScoreMatrixList object. It enables to use arithmetic, indicator and logic operations on ScoreMatrixList objects.*

Description

Arithmetic method for ScoreMatrixList

Usage

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

Arguments

e1 the [ScoreMatrixList](#) object
e2 the numeric value

Value

ScoreMatrixList

Ops, ScoreMatrixList, ScoreMatrixList-method

Ops method for a ScoreMatrixList object. It enables to use arithmetic, indicator and logic operations on ScoreMatrixList objects.

Description

Arithmetic methods for ScoreMatrixList

Usage

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

Arguments

e1	the ScoreMatrixList object
e2	the ScoreMatrixList object

Value

ScoreMatrixList

orderBy

Reorder all elements of a ScoreMatrixList to a given ordering vector

Description

Reorder all elements of a ScoreMatrixList to a given ordering vector

Usage

```
orderBy(sml, ord.vec)

## S4 method for signature 'ScoreMatrixList'
orderBy(sml, ord.vec)
```

Arguments

sml	ScoreMatrixList object
ord.vec	an integer vector

Value

ScoreMatrixList object

Examples

```

library(GenomicRanges)
data(cage)
data(cpgi)
data(promoters)

cage$tpm = NULL
targets = GRangesList(cage=cage, cpgi=cpgi)
sml = ScoreMatrixList(targets, promoters, bin.num=10)
kmeans.clust = kmeans(sml$cage,3)

sml.ordered = orderBy(sml, kmeans.clust$cluster)

multiHeatMatrix(sml.ordered)

```

patternMatrix	<i>Get scores that correspond to k-mer or PWM matrix occurrence for bases in each window</i>
---------------	--

Description

The function produces a base-pair resolution matrix or matrices of scores that correspond to k-mer or PWM matrix occurrence over predefined windows that have equal width. It finds either positions of pattern hits above a specified threshold and creates score matrix filled with 1 (presence of pattern) and 0 (its absence) or matrix with scores themselves. If pattern is a character of length 1 or PWM matrix then the function returns a ScoreMatrix object, if character of length more than 1 or list of PWMs then ScoreMatrixList.

Usage

```

patternMatrix(pattern, windows, genome = NULL, min.score = 0.8,
  asPercentage = FALSE, cores = 1)

\S4method{patternMatrix}{character,DNAStringSet}(pattern, windows,
  asPercentage, cores)

\S4method{patternMatrix}{character,GRanges,BSgenome}(pattern, windows, genome,
  cores)

\S4method{patternMatrix}{matrix,DNAStringSet}(pattern, windows,
  min.score, asPercentage,
  cores)

\S4method{patternMatrix}{matrix,GRanges,BSgenome}(pattern, windows, genome,
  min.score, asPercentage,
  cores)

```

```
\S4method{patternMatrix}{list,DNAStringSet}(pattern, windows,
                                              min.score, asPercentage,
                                              cores)

\S4method{patternMatrix}{list,GRanges,BSgenome}(pattern, windows, genome,
                                              min.score, asPercentage,
                                              cores)
```

Arguments

pattern	matrix (a PWM matrix), list of matrices or a character vector of length 1 or more. A matrix is a PWM matrix that needs to have one row for each nucleotide ("A","C","G" and "T" respectively). IUPAC ambiguity codes can be used and it will match any letter in the subject that is associated with the code.
windows	GRanges object or DNAStringSet object that have equal width of ranges or sequences.
genome	BSgenome object
min.score	numeric or character indicating minimum score to count a match. It can be given as a character string containing a percentage of the highest possible score or a single number (by default "80%" or 0.8). If min.score is set to NULL then patternMatrix returns scores themselves (default).
asPercentage	boolean telling whether scores represent percentage of the maximal motif PWM score (default: TRUE) or raw scores (FALSE).
cores	the number of cores to use (default: 1). It is supported only on Unix-like platforms.

Details

patternMatrix is based on functions from the seqPattern package: getPatternOccurrenceList function to find position of pattern that is a character vector in a list of sequences (a DNAStringSet object) and adapted function motifScanHits to find pattern that is a PWM matrix in sequences (a DNAStringSet object).

If cores > 1 is provided then for every window occurrence of pattern is counted in parallel.

Value

returns a scoreMatrix object or a scoreMatrixList object

See Also

[ScoreMatrix](#), [ScoreMatrixList](#)

Examples

```
library(Biostrings)

# consensus sequence of the ctcf motif
motif = "CCGCGNGGNGGCAG"
```



```
# Creates 10 random DNA sequences
seqs = sapply(1:10,
  function(x) paste(sample(c("A","T","G","C"), 180, replace=TRUE), collapse=""))
windows = DNASTringSet(seqs)
p = patternMatrix(pattern=motif, windows=windows, min.score=0.8)
p
```

plotMeta

*Line plot(s) for meta-region profiles***Description**

Function calculates meta-profile(s) from a ScoreMatrix or a ScoreMatrixList, then produces a line plot or a set of line plots for meta-region profiles

Usage

```
plotMeta(mat, centralTend = "mean", overlay = TRUE, winsorize = c(0, 100),
  profile.names = NULL, xcoords = NULL, meta.rescale = FALSE,
  smoothfun = NULL, line.col = NULL, dispersion = NULL,
  dispersion.col = NULL, ylim = NULL, ylab = "average score",
  xlab = "bases", ...)
```

Arguments

mat	ScoreMatrix or ScoreMatrixList object. If it is a ScoreMatrixList object, all matrices in the ScoreMatrixList should have the same number of columns.
centralTend	a character that determines central tendency of meta-profile(s). It takes "mean" (default) or "median".
overlay	If TRUE multiple profiles will be overlayed in the same plot (Default:TRUE). If FALSE, and mat is a ScoreMatrixList, consider using par(mfrow=c(1,length(mat))) to see the plots from all matrices at once.
winsorize	Numeric vector of two, defaults to c(0,100). This vector determines the upper and lower percentile values to limit the extreme values. For example, c(0,99) will limit the values to only 99th percentile, everything above the 99 percentile will be equalized to the value of 99th percentile. This is useful for visualization of matrices that have outliers.
profile.names	a character vector for names of the profiles. The order should be same as the order of ScoreMatrixList.
xcoords	a numeric vector which designates relative base positions of the meta-region profiles. For example, for a 2001 column ScoreMatrix, xcoord=-1000:1000 indicates relative positions of each column in the score matrix. If NULL (Default), xcoords equals to 1:ncol(mat)
meta.rescale	if TRUE meta-region profiles are scaled to 0 to 1 range by subtracting the min from profiles and dividing them by max-min. If dispersion is not NULL, then dispersion will be scaled as well.

smoothfun	a function to smooth central tendency and dispersion bands (Default: NULL), e.g. stats::lowess.
line.col	color of lines for centralTend of meta-region profiles. Defaults to colors from rainbow() function.
dispersion	shows dispersion interval bands around centralTend (default:NULL). It takes one of the character: • "se" shows standard error of the mean and 95 percent confidence interval for the mean • "sd" shows standard deviation and 2*(standard deviation) • "IQR" shows 1st and 3rd quartile and confidence interval around the median based on the median $\pm 1.57 * \text{IQR}/\sqrt{n}$ (notches)
dispersion.col	color of bands of dispersion. Defaults to colors from rainbow() and transparency is set to 0.5 (rainbow(length(mat), alpha = 0.5)).
ylim	same as ylim at plot function. if NULL ylim is estimated from all meta-region profiles.
ylab	same as ylab at plot function. Default: "average score"
xlab	same as xlab at plot function. Default: "bases"
...	other options to plot

Value

returns the meta-region profiles invisibly as a matrix.

Note

Score matrices are plotted according to ScoreMatrixList order. If ScoreMatrixList contains more than one matrix then they will overlap each other on a plot, i.e. the first one is plotted first and every next one overlays previous one(s) and the last one is the topmost.

Missing values in data slow down plotting dispersion around central tendency. The reason is that dispersion is plotted only for non-missing values, for each segment that contains numerical values graphics::polygon function is used to plot dispersion bands. There might be a situation, when in a column of ScoreMatrix is only one numeric number and the rest are NAs, then at corresponding position only central tendency will be plotted.

Notches show the 95 percent confidence interval for the median according to an approximation based on the normal distribution. They are used to compare groups - if notches corresponding to adjacent base pairs on the plot do not overlap, this is strong evidence that medians differ. Small sample sizes (5-10) can cause notches to extend beyond the interquartile range (IQR) (Martin Krzywinski *et al. Nature Methods* 11, 119-120 (2014))

Examples

```
data(cage)
data(promoters)
scores1=ScoreMatrix(target=cage, windows=promoters, strand.aware=TRUE)

data(cpgi)
```

```

scores2=ScoreMatrix(target=cpgi, windows=promoters, strand.aware=TRUE)

# create a new ScoreMatrixList
x=new("ScoreMatrixList", list(scores1, scores2))

plotMeta(mat=x, overlay=TRUE, main="my plotowski")

# plot dispersion and smooth central tendency and variation interval bands
plotMeta(mat=x, centralTend="mean", dispersion="se", winsorize=c(0,99),
         main="Dispersion as interquartile band", lwd=4,
         smoothfun=function(x) stats::lowess(x, f = 1/5))

```

plotTargetAnnotation *Plot annotation categories from AnnotationByGeneParts or AnnotationByFeature*

Description

This function plots a pie or bar chart for showing percentages of targets annotated by genic parts or other query features

Usage

```

plotTargetAnnotation(x, precedence = TRUE,
  col = getColors(length(x@annotation)), cex.legend = 1, ...)

## S4 method for signature 'AnnotationByFeature'
plotTargetAnnotation(x, precedence = TRUE,
  col = getColors(length(x@annotation)), cex.legend = 1, ...)

```

Arguments

x	a AnnotationByFeature or AnnotationByGeneParts object
precedence	TRUE FALSE. If TRUE there will be a hierarchy of annotation features when calculating numbers (with promoter>exon>intron precedence). This option is only valid when x is a AnnotationByGeneParts object
col	a vector of colors for piechart or the bar plot
cex.legend	a numeric value of length 1 to specify the size of the legend. By default 1.
...	graphical parameters to be passed to pie or barplot functions

Value

plots a piechart or a barplot for percentage of the target features overlapping with annotation

Examples

```
data(cage)

bed.file = system.file("extdata/chr21.refseq.hg19.bed", package = "genomation")
gene.parts = readTranscriptFeatures(bed.file)
annot = annotateWithGeneParts(cage, gene.parts, intersect.chr=TRUE)

plotTargetAnnotation(annot)
```

promoters

Example promoter data set.

Description

promoters of hg19 assembly of human genome on chr21 and chr22. Promoter set is derived from refseq TSS.

Format

[GRanges](#) object

RandomEnrichment-class

An S4 class for storing getRandomEnrichment function results

Description

The resulting object stores the results of getRandomEnrichment function

Slots

orig.cnt: number of features overlapping with query at getRandomEnrichment
rand.olap.dist: set of number of features overlapping with randomized queries at getRandomEnrichment
log2fc: log2 fold change calculated by dividing orig.cnt by mean(rand.olap.dist) and taking log2 of that result
p.value: P-value assuming rand.olap.dist has a normal distribution and comparing orig.cnt with that distribution
rand.p.value: p-value from randomization by calculation the proportion of how many times a random number of overlap exceeds the original number of overlap

See Also

[getRandomEnrichment](#)

randomizeFeature	<i>function that randomizes the genomic coordinates</i>
------------------	---

Description

This function randomly distributes the coordinates of genomic features which is stored in a GRanges object. The randomization can be constrained by supplied arguments. The function is still in Beta mode - the regions can overlap excluded regions, and the randomized regions are not disjoint. Please take care that the excluded and included regions are not too strict when compared to the total width of the ranges.

Usage

```
randomizeFeature(feature, chrom.sizes = NULL, stranded = TRUE,
  keep.strand.prop = TRUE, keep.chrom = TRUE, exclude = NULL,
  include = NULL, seed = NULL, nrand = 1)
```

```
## S4 method for signature 'GRanges'
randomizeFeature(feature, chrom.sizes = NULL,
  stranded = TRUE, keep.strand.prop = TRUE, keep.chrom = TRUE,
  exclude = NULL, include = NULL, seed = NULL, nrand = 1)
```

Arguments

feature	a GRanges object to be randomized
chrom.sizes	sizes of chromosomes as a named vector (names are chromosomes names and elements of the vectors are lengths). , if not given sizes in GRanges object will be used if no sizes there the end of each chr will be the end last feature on each chr
stranded	if FALSE, all of the returned features will be strandless (will have "*" in the strand slot)
keep.strand.prop	If TRUE strands will have the same proportion as the features
keep.chrom	If TRUE, number of features and randomized features for a chromosome will match. Currently setting this to FALSE is not supported.
exclude	A GRanges object where no randomized feature should overlap, can be gaps or unmappable regions in the genome as an example.
include	A GRanges object which defines the boundaries of randomized features. If not provided the whole genome is used, as defined using the chrom.sizes parameter.
seed	random number generator seed
nrand	number of randomizations (default:1)

Value

returns a GRanges object which is randomized version of the feature, along with a "set" column in the metadata which designates to which iteration of the randomization the range belong.

read.zip	<i>read.zip function</i>
----------	--------------------------

Description

read.zip function

Usage

read.zip(file, ...)

Arguments

file	zip file
...	additional objects and parameters

readBam	<i>readBam function</i>
---------	-------------------------

Description

given a big bam path reads the big wig file into a RleList to be used by ScoreMatrix:char,GRanges

Usage

readBam(target, windows, rpm = FALSE, unique = FALSE, extend = 0,
param = NULL, paired.end = FALSE, library.size = NULL, ...)

Arguments

target	target object
windows	windows
rpm	logical
unique	logical
extend	numeric
param	ScanBamParam object
paired.end	llogical
library.size	numeric
...	additional parameters

readBed	<i>Read a BED file and convert it to GRanges.</i>
---------	---

Description

The function reads a BED file that contains location and other information on genomic features and returns a [GRanges](#) object. The minimal information that the BED file has to have is chromosome, start and end columns. it can handle all BED formats up to 12 columns.

Usage

```
readBed(file, track.line = FALSE, remove.unusual = FALSE,  
        zero.based = TRUE)
```

Arguments

file	location of the file, a character string such as: "/home/user/my.bed" or the input itself as a string (containing at least one \n). The file can end in .gz, .bz2, .xz, or .zip and/or start with http:// or ftp://. If the file is not compressed it can also start with https:// or ftps://.
track.line	the number of track lines to skip, "auto" to detect them automatically or FALSE(default) if the bed file doesn't have track lines
remove.unusual	if TRUE remove the chromosomes with unusual names, such as chrX_random (Default:FALSE)
zero.based	a boolean which tells whether the ranges in the bed file are 0 or 1 base encoded. (Default: TRUE)

Value

[GRanges](#) object

Examples

```
my.file=system.file("extdata", "chr21.refseq.hg19.bed", package="genomation")  
refseq = readBed(my.file, track.line=FALSE, remove.unusual=FALSE)  
head(refseq)
```

readBigWig	<i>readBigWig function</i>
------------	----------------------------

Description

given a big wig path reads the big wig file into a RleList to be used by ScoreMatrix:char,GRanges

Usage

```
readBigWig(target, windows = NULL, ...)
```

Arguments

target	target object
windows	windows
...	additional parameters

readBroadPeak	<i>A function to read the Encode formatted broad peak file into a GRanges object</i>
---------------	--

Description

A function to read the Encode formatted broad peak file into a GRanges object

Usage

```
readBroadPeak(file, track.line=FALSE, zero.based=TRUE)
```

Arguments

file	an absolute or relative path to a bed file formatted by the Encode broadPeak standard. The file can end in .gz, .bz2, .xz, or .zip and/or start with http:// or ftp://. If the file is not compressed it can also start with https:// or ftps://.
track.line	the number of track lines to skip, "auto" to detect them automatically or FALSE(default) if the bed file doesn't have track lines
zero.based	a boolean which tells whether the ranges in the bed file are 0 or 1 base encoded. (Default: TRUE)

Value

a GRanges object

Examples

```

broad.peak.file = system.file('extdata',"ex.broadPeak", package='genomation')

broad.peak = readBroadPeak(broad.peak.file)
head(broad.peak)

```

readFeatureFlank	<i>A function to read-in genomic features and their upstream and downstream adjacent regions such as CpG islands and their shores</i>
------------------	---

Description

A function to read-in genomic features and their upstream and downstream adjacent regions such as CpG islands and their shores

Usage

```

readFeatureFlank(location,remove.unusual=TRUE,flank=2000,
                  clean=TRUE,feature.flank.name=NULL)

## S4 method for signature 'character'
readFeatureFlank(location, remove.unusual = TRUE,
                  flank = 2000, clean = TRUE, feature.flank.name = NULL)

```

Arguments

location	for the bed file of the feature.
remove.unusual	remove chromosomes with unusual names random, Un and anything with "_" character
flank	number of basepairs for the flanking regions
clean	If set to TRUE, flanks overlapping with other main features will be trimmed
feature.flank.name	the names for feature and flank ranges, it should be a character vector of length 2. example: c("CpGi","shores")

Value

a GRangesList object containing one GRanges object for flanks and one for GRanges object for the main feature. NOTE: This can not return a CompressedGRangesList at the moment because flanking regions do not have to have the same column name as the feature. CompressedGRangesList elements should resemble each other in the column content. We can not satisfy that criteria for the flanks

Examples

```
cgi.path = system.file('extdata/chr21.CpG.hg19.bed', package='genomation')
cgi.shores = readFeatureFlank(cgi.path)
cgi.shores
```

readGeneric	<i>Read a tabular file and convert it to GRanges.</i>
-------------	---

Description

The function reads a tabular text file that contains location and other information on genomic features and returns a [GRanges](#) object. The minimal information that the file has to have is chromosome, start and end columns. Strand information is not compulsory.

Usage

```
readGeneric(file, chr = 1, start = 2, end = 3, strand = NULL,
  meta.cols = NULL, keep.all.metadata = FALSE, zero.based = FALSE,
  remove.unusual = FALSE, header = FALSE, skip = 0, sep = "\t")
```

Arguments

file	location of the file, a character string such as: "/home/user/my.bed" or the input itself as a string (containing at least one \n).
chr	number of the column that has chromosomes information in the table (Def:1)
start	number of the column that has start coordinates in the table (Def:2)
end	number of the column that has end coordinates in the table (Def:3)
strand	number of the column that has strand information, only -/+ is accepted (Default:NULL)
meta.cols	named list that maps column numbers to meta data columns. e.g. list(name=5, score=10), which means 5th column will be named "name", and 10th column will be named "score" and their contents will be a part of the returned GRanges object. If header = TRUE, meta.cols parameter will over-write the column names given by the header line of the data frame.
keep.all.metadata	logical determining if the extra columns (the ones that are not designated by chr,start,end,strand and meta.cols arguments) should be kept or not. (Default:FALSE)
zero.based	a boolean which tells whether the ranges in the bed file are 0 or 1 base encoded. (Default: FALSE)
remove.unusual	if TRUE(default) remove the chromosomes with unusual names, such as chrX_random (Default:FALSE)
header	whether the original file contains a header line which designates the column names. If TRUE header will be used to construct column names. These names can be over written by meta.cols argument.

skip	number of lines to skip. If there is a header line(s) you do not wish to include you can use skip argument to skip that line.
sep	a single character which designates the separator in the file. The default value is tab.

Value

GRanges object

Examples

```
my.file=system.file("extdata", "chr21.refseq.hg19.bed", package="genomation")
refseq = readGeneric(my.file, chr=1, start=2, end=3, strand=NULL,
                    meta.cols=list(score=5, name=4),
                    keep.all.metadata=FALSE, zero.based=TRUE)
head(refseq)
```

readNarrowPeak	<i>A function to read the Encode formatted narrowPeak file into a GRanges object</i>
----------------	--

Description

A function to read the Encode formatted narrowPeak file into a GRanges object

Usage

```
readNarrowPeak(file, track.line=FALSE, zero.based=TRUE)
```

Arguments

file	an absolute or relative path to a bed file formatted by the Encode narrowPeak standard. The file can end in .gz, .bz2, .xz, or .zip and/or start with http:// or ftp://. If the file is not compressed it can also start with https:// or ftps://.
track.line	the number of track lines to skip, "auto" to detect them automatically or FALSE(default) if the bed file doesn't have track lines
zero.based	a boolean which tells whether the ranges in the bed file are 0 or 1 base encoded. (Default: TRUE)

Value

a GRanges object

Examples

```
narrow.peak.file = system.file('extdata',"ex.narrowPeak", package='genomation')

narrow.peak = readBroadPeak(narrow.peak.file)
head(narrow.peak)
```

readTableFast	<i>readTableFast function</i>
---------------	-------------------------------

Description

fast reading of big tables chr indicates index of column of chromosomes

Usage

```
readTableFast(filename, header = TRUE, skip = 0, sep = "\t", chr = 1)
```

Arguments

filename	file name
header	logical
skip	numeric
sep	character
chr	numeric

readTranscriptFeatures	<i>Function for reading exon intron and promoter structure from a given bed file</i>
------------------------	--

Description

Function for reading exon intron and promoter structure from a given bed file

Usage

```
readTranscriptFeatures(location,remove.unusual=TRUE,
                        up.flank=1000,down.flank=1000,unique.prom=TRUE)

## S4 method for signature 'character'
readTranscriptFeatures(location, remove.unusual = TRUE,
  up.flank = 1000, down.flank = 1000, unique.prom = TRUE)
```

Arguments

location	location of the bed file with 12 or more columns. The file can end in .gz, .bz2, .xz, or .zip and/or start with http:// or ftp://. If the file is not compressed it can also start with https:// or ftps://.
remove.unusual	remove the chromosomes with unusual names, mainly random chromosomes etc
up.flank	up-stream from TSS to detect promoter boundaries
down.flank	down-stream from TSS to detect promoter boundaries
unique.prom	get only the unique promoters, promoter boundaries will not have a gene name if you set this option to be TRUE

Value

a [GRangesList](#) containing locations of exon/intron/promoter/TSS

Note

one bed track per file is only accepted, the bed files with multiple tracks will cause an error

Examples

```
my.bed12.file = system.file("extdata/chr21.refseq.hg19.bed", package = "genomation")
my.bed12.file
feats = readTranscriptFeatures(my.bed12.file)
names(feats)
sapply(feats, head)
```

scaleScoreMatrix	<i>Scales the values in the matrix by rows and/or columns</i>
------------------	---

Description

Scales the values in the matrix by rows and/or columns

Usage

```
scaleScoreMatrix(mat, columns = FALSE, rows = TRUE, scalefun = NULL)

## S4 method for signature 'ScoreMatrix'
scaleScoreMatrix(mat, columns = FALSE, rows = TRUE,
  scalefun = NULL)
```

Arguments

mat	ScoreMatrix object
columns	columns whether to scale the matrix by columns. Set by default to FALSE.
rows	rows Whether to scale the matrix by rows. Set by default to TRUE
scalefun	function object that takes as input a matrix and returns a matrix. By default the argument is set to $(x - \text{mean}(x))/(\text{max}(x)-\text{min}(x)+1)$

Value

ScoreMatrix object

Examples

```
# scale the rows of a scoreMatrix object
library(GenomicRanges)
target = GRanges(rep(c(1,2),each=7), IRanges(rep(c(1,1,2,3,7,8,9), times=2), width=5),
              weight = rep(c(1,2),each=7),
              strand=c('-', '-', '-', '-', '+', '-', '+', '-', '-', '-', '-', '-', '-', '+'))
windows = GRanges(rep(c(1,2),each=2), IRanges(rep(c(1,2), times=2), width=5),
              strand=c('-', '+', '-', '+'))
sm = ScoreMatrix(target, windows)
ssm = scaleScoreMatrix(sm, rows=TRUE)
```

scaleScoreMatrixList *Scale the ScoreMatrixList*

Description

Scales each ScoreMatrix in the ScoreMatrixList object, by rows and/or columns

Usage

```
scaleScoreMatrixList(sml, columns, rows, scalefun)
```

```
## S4 method for signature 'ScoreMatrixList'
scaleScoreMatrixList(sml, columns = FALSE,
  rows = TRUE, scalefun = NULL)
```

Arguments

sml	a ScoreMatrixList object
columns	a columns whether to scale the matrix by columns. Set by default to FALSE
rows	a rows Whether to scale the matrix by rows. Set by default to TRUE
scalefun	a function object that takes as input a matrix and returns a matrix. By default the argument is set to the R scale function with center=TRUE and scale=TRUE

Value

ScoreMatrixList object

Examples

```
library(GenomicRanges)
data(cage)
data(cpgi)
data(promoters)

cage$tpm = NULL
targets = GRangesList(cage=cage, cpgi=cpgi)
sml = ScoreMatrixList(targets, promoters, bin.num=10, strand.aware=TRUE)
sml.scaled = scaleScoreMatrixList(sml, rows=TRUE)
sml.scaled

multiHeatMatrix(sml)
```

ScoreMatrix

Get base-pair score for bases in each window

Description

The function produces a base-pair resolution matrix of scores for given equal width windows of interest. The returned matrix can be used to draw meta profiles or heatmap of read coverage or wig track-like data. The windows argument can be a predefined region around transcription start sites or other regions of interest that have equal lengths. The function removes all window that fall off the Rle object - have the start coordinate < 1 or end coordinate > length(Rle). The function takes the intersection of names in the Rle and GRanges objects. On Windows OS the function will give an error if the target is a file in .bigWig format.

Usage

```
ScoreMatrix(target, windows, strand.aware = FALSE, weight.col = NULL,
  is.noCovNA = FALSE, type = "auto", rpm = FALSE, unique = FALSE,
  extend = 0, param = NULL, bam.paired.end = FALSE, library.size = NULL)

\S4method{ScoreMatrix}{RleList,GRanges}(target, windows, strand.aware)

\S4method{ScoreMatrix}{GRanges,GRanges}(target, windows, strand.aware,
  weight.col, is.noCovNA)

\S4method{ScoreMatrix}{character,GRanges}(target, windows, strand.aware,
  weight.col=NULL, is.noCovNA=FALSE,
  type='auto', rpm=FALSE,
  unique=FALSE, extend=0, param=NULL,
```

```
bam.paired.end=FALSE,
library.size=NULL)
```

Arguments

target	RleList , GRanges, a BAM file or a BigWig to be overlapped with ranges in windows
windows	GRanges object that contains the windows of interest. It could be promoters, CpG islands, exons, introns. However the sizes of windows have to be equal.
strand.aware	If TRUE (default: FALSE), the strands of the windows will be taken into account in the resulting ScoreMatrix. If the strand of a window is -, the values of the bins for that window will be reversed
weight.col	if the object is GRanges object a numeric column in meta data part can be used as weights. This is particularly useful when genomic regions have scores other than their coverage values, such as percent methylation, conservation scores, GC content, etc.
is.noCovNA	(Default:FALSE) if TRUE, and if 'target' is a GRanges object with 'weight.col' provided, the bases that are uncovered will be preserved as NA in the returned object. This useful for situations where you can not have coverage all over the genome, such as CpG methylation values.
type	(Default:"auto") if target is a character vector of file paths, then type designates the type of the corresponding files (bam or bigWig).
rpm	boolean telling whether to normalize the coverage to per million reads. FALSE by default. See library.size.
unique	boolean which tells the function to remove duplicated reads based on chr, start, end and strand
extend	numeric which tells the function to extend the reads to width=extend
param	ScanBamParam object
bam.paired.end	boolean indicating whether given BAM file contains paired-end reads (default:FALSE). Paired-reads will be treated as fragments.
library.size	numeric indicating total number of mapped reads in a BAM file (rpm has to be set to TRUE). If is not given (default: NULL) then library size is calculated using the Rsamtools idxstatsBam function: sum(idxstatsBam(target)\$mapped).

Value

returns a ScoreMatrix object

Note

We assume that a paired-end BAM file contains reads with unique ids and we remove both mates of reads if they are repeated. Due to the fact that ScoreMatrix uses the GenomicAlignments::readGAlignmentPairs function to read paired-end BAM files a duplication of reads occurs when mates of one pair map into two different windows.

Strands of reads in a paired-end BAM are inferred depending on strand of first alignment from the pair. This is a default setting in the GenomicAlignments::readGAlignmentPairs function (see a

strandMode argument). This mode should be used when the paired-end data was generated using one of the following stranded protocols: Directional Illumina (Ligation), Standard SOLiD.

See Also

[ScoreMatrixBin](#)

Examples

```
# When target is GRanges
data(cage)
data(promoters)
scores1=ScoreMatrix(target=cage, windows=promoters, strand.aware=TRUE,
                    weight.col="tpm")

# When target is RleList
library(GenomicRanges)
covs = coverage(cage)
scores2 = ScoreMatrix(target=covs, windows=promoters, strand.aware=TRUE)
scores2

# When target is a bam file
bam.file = system.file('unitTests/test.bam', package='genomation')
windows = GRanges(rep(c(1,2), each=2), IRanges(rep(c(1,2), times=2), width=5))
scores3 = ScoreMatrix(target=bam.file, windows=windows, type='bam')
scores3
```

ScoreMatrix-class	<i>An S4 class for storing ScoreMatrix function results</i>
-------------------	---

Description

The resulting object is an extension of a matrix object, and stores values (typically genome-wide scores) for a predefined set of regions. Each row on the ScoreMatrix is a predefined region (Ex: CpG islands, promoters) and columns are values across those regions.

Constructors

see [ScoreMatrix](#)

Coercion

as(from, "matrix"): Creates a matrix from ScoreMatrix object. You can also use S3Part() function to extract the matrix from ScoreMatrix object.

Subsetting

In the code snippets below, `x` is a `ScoreMatrix` object. `'x[i,j]'`: Get or set elements from row `i` and column `j` and return a subset `ScoreMatrix` object.

See Also

[ScoreMatrix](#)

ScoreMatrixBin	<i>Get bin score for bins on each window</i>
----------------	--

Description

The function first bins each window to equal number of bins, and calculates the a summary matrix for scores of each bin (currently, mean, max and min supported) A `scoreMatrix` object can be used to draw average profiles or heatmap of read coverage or wig track-like data. windows can be a predefined region such as CpG islands, gene bodies, transcripts or CDS (coding sequences) that are not necessarily equi-width. Each window will be chopped to equal number of bins based on `bin.num` option.

Usage

```
ScoreMatrixBin(target, windows, bin.num = 10, bin.op = "mean",
  strand.aware = FALSE, weight.col = NULL, is.noCovNA = FALSE,
  type = "auto", rpm = FALSE, unique = FALSE, extend = 0,
  param = NULL, bam.paired.end = FALSE, library.size = NULL)

\S4method{ScoreMatrixBin}{RleList,GRanges}(target, windows, bin.num, bin.op,
  strand.aware)

\S4method{ScoreMatrixBin}{GRanges,GRanges}(target, windows,
  bin.num, bin.op,
  strand.aware, weight.col,
  is.noCovNA)

\S4method{ScoreMatrixBin}{character,GRanges}(target, windows, bin.num=10,
  bin.op='mean', strand.aware,
  is.noCovNA=FALSE, type='auto',
  rpm, unique, extend, param,
  bam.paired.end=FALSE,
  library.size=NULL)

\S4method{ScoreMatrixBin}{RleList,GRangesList}(target, windows,
  bin.num, bin.op,
  strand.aware)

\S4method{ScoreMatrixBin}{GRanges,GRangesList}(target, windows,
```

```

        bin.num,bin.op,
        strand.aware,weight.col,
        is.noCovNA)

\S4method{ScoreMatrixBin}{character,GRangesList}(target, windows, bin.num=10,
        bin.op='mean',strand.aware,
        weight.col=NULL,
        is.noCovNA=FALSE, type='auto',
        rpm, unique, extend, param,
        bam.paired.end=FALSE,
        library.size=NULL)

```

Arguments

target	RleList, GRanges, a BAM file or a bigWig file object to be overlapped with ranges in windows
windows	GRanges or GRangesList object that contains the windows of interest. It could be promoters, CpG islands, exons, introns as GRanges object or GrangesList object representing exons of each transcript. Exons must be ordered by ascending rank by their position in transcript. The sizes of windows does NOT have to be equal.
bin.num	single integer value denoting how many bins there should be for each window
bin.op	bin operation that is either one of the following strings: "max","min","mean","median","sum". The operation is applied on the values in the bin. Defaults to "mean"
strand.aware	If TRUE (default: FALSE), the strands of the windows will be taken into account in the resulting scoreMatrix. If the strand of a window is -, the values of the bins for that window will be reversed
weight.col	if the object is GRanges object a numeric column in meta data part can be used as weights. This is particularly useful when genomic regions have scores other than their coverage values, such as percent methylation, conservation scores, GC content, etc.
is.noCovNA	(Default:FALSE) if TRUE,and if 'target' is a GRanges object with 'weight.col' provided, the bases that are uncovered will be preserved as NA in the returned object. This useful for situations where you can not have coverage all over the genome, such as CpG methylation values.
type	(Default:"auto") if target is a character vector of file paths, then type designates the type of the corresponding files (bam or bigWig)
rpm	boolean telling whether to normalize the coverage to per milion reads. FALSE by default. See library.size.
unique	boolean which tells the function to remove duplicated reads based on chr, start, end and strand
extend	numeric which tells the function to extend the reads to width=extend
param	ScanBamParam object
bam.paired.end	boolean indicating whether given BAM file contains paired-end reads (default:FALSE). Paired-reads will be treated as fragments.

`library.size` numeric indicating total number of mapped reads in a BAM file (rpm has to be set to TRUE). If is not given (default: NULL) then library size is calculated using the Rsamtools `idxstatsBam` function: `sum(idxstatsBam(target)$mapped)`.

Value

returns a `scoreMatrix` object

See Also

[ScoreMatrix](#)

Examples

```
data(cage)
data(cpgi)
data(promoters)
myMat=ScoreMatrixBin(target=cage,
                     windows=cpgi,bin.num=10,bin.op="mean",weight.col="tpm")

plot(colMeans(myMat,na.rm=TRUE),type="l")

myMat2=ScoreMatrixBin(target=cage,
                     windows=promoters,bin.num=10,bin.op="mean",
                     weight.col="tpm",strand.aware=TRUE)

plot(colMeans(myMat2,na.rm=TRUE),type="l")

# Compute transcript coverage of a set of exons.
library(GenomicRanges)
bed.file = system.file("extdata/chr21.refseq.hg19.bed",
                       package = "genomation")
gene.parts = readTranscriptFeatures(bed.file)
transcripts = split(gene.parts$exons, gene.parts$exons$name)
transcripts = transcripts[]
myMat3 = ScoreMatrixBin(target=cage, windows=transcripts[1:250],
                       bin.num=10)
myMat3
```

ScoreMatrixList

Make ScoreMatrixList from multiple targets

Description

The function constructs a list of `ScoreMatrix` objects in the form of `ScoreMatrixList` object. This object can be visualized using `multiHeatMatrix`, `heatMeta` or `plotMeta`

Usage

```
ScoreMatrixList(targets, windows = NULL, bin.num = NULL, bin.op = "mean",
  strand.aware = FALSE, weight.col = NULL, is.noCovNA = FALSE,
  type = "auto", rpm = FALSE, unique = FALSE, extend = 0,
  param = NULL, library.size = NULL, cores = 1)
```

Arguments

targets	can be a list of scoreMatrix objects, that are coerced to the ScoreMatrixList, a list of RleList objects, or a character vector specifying the locations of multiple bam files or bigWig files that are used to construct the scoreMatrixList. If it is either a RleList object or a character vector of files, it is obligatory to give a windows argument.
windows	GenomicRanges containing viewpoints for the scoreMatrix or ScoreMatrixList functions
bin.num	an integer telling the number of bins to bin the score matrix
bin.op	an name of the function that will be used for smoothing windows of ranges
strand.aware	a boolean telling the function whether to reverse the coverage of ranges that come from - strand (e.g. when plotting enrichment around transcription start sites)
weight.col	if the object is GRanges object a numeric column in meta data part can be used as weights. This is particularly useful when genomic regions have scores other than their coverage values, such as percent methylation, conservation scores, GC content, etc.
is.noCovNA	(Default:FALSE) if TRUE, and if 'targets' is a GRanges object with 'weight.col' provided, the bases that are uncovered will be preserved as NA in the returned object. This useful for situations where you can not have coverage all over the genome, such as CpG methylation values.
type	(Default:"auto") if targets is a character vector of file paths, then type designates the type of the corresponding files (bam or bigWig)
rpm	boolean telling whether to normalize the coverage to per million reads. FALSE by default. See library.size.
unique	boolean which tells the function to remove duplicated reads based on chr, start, end and strand
extend	numeric which tells the function to extend the features (i.e aligned reads) to total length of width+extend
param	ScanBamParam object
library.size	a numeric vector of the same length as targets indicating total number of mapped reads in BAM files (targets). If is not given (default: NULL) then library sizes for every target is calculated using the Rsamtools idxstatsBam function: sum(idxstatsBam(target)\$mapped). rpm argument has to be set to TRUE.
cores	the number of cores to use (default: 1)

Value

returns a ScoreMatrixList object

Examples

```
# visualize the distribution of cage clusters and cpg islands around promoters
library(GenomicRanges)
data(cage)
data(cpgi)
data(promoters)

cage$tpm = NULL
targets = GRangesList(cage=cage, cpgi=cpgi)
sml = ScoreMatrixList(targets, promoters, bin.num=10, strand.aware=TRUE)
sml

multiHeatMatrix(sml)
```

ScoreMatrixList-class *An S4 class for storing a set of ScoreMatrixList*

Description

The resulting object is an extension of a list object, where each element corresponds to a score matrix object

Constructors

see [ScoreMatrixList](#)

Coercion

`as(from, "ScoreMatrixList")`: Creates a `ScoreMatrixList` object from a list containing [ScoreMatrix](#) or [ScoreMatrixBin](#) objects.

Subsetting

In the code snippets below, `x` is a `ScoreMatrixList` object.

`x[[i]],x[[i]]`: Get or set elements `i`, where `i` is a numeric or character vector of length 1.

`x$name, x$name: value`: Get or set element name, where `name` is a name or character vector of length 1.

See Also

[ScoreMatrixList](#)

show,RandomEnrichment-method	<i>show method for some of the genomation classes</i>
------------------------------	---

Description

show method for some of the genomation classes

Usage

```
## S4 method for signature 'RandomEnrichment'
show(object)

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

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

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

## S4 method for signature 'ScoreMatrixList'
show(object)
```

Arguments

object object of class RandomEnrichment

Value

Shows the dimension of the ScoreMatrix
Shows the number of matrices and their sizes

Sum_c	<i>Function that computes a sum value</i>
-------	---

Description

Function that computes a sum value

Usage

```
Sum_c(x)
```

Arguments

x	NumericVector
target.type	<i>target.type function</i>

Description

Check if a target file is in bam and bigWig formats by looking at the file extension

Usage

```
target.type(target, type = "")
```

Arguments

target	target file
type	type name
[,ScoreMatrix,ANY,ANY,ANY-method	
<i>Extract method for a ScoreMatrix object.</i>	

Description

Extract method for a ScoreMatrix object.

Usage

```
## S4 method for signature 'ScoreMatrix,ANY,ANY,ANY'
x[i, j]
```

Arguments

x	the ScoreMatrix object
i	numeric value
j	numeric value

[,ScoreMatrixList,ANY,ANY,ANY-method

Extract method for a ScoreMatrixList object.

Description

Extract method for a ScoreMatrixList object.

Usage

```
## S4 method for signature 'ScoreMatrixList,ANY,ANY,ANY'  
x[i]
```

Arguments

x	the ScoreMatrixList object
i	numeric value

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