Package 'GenomicDistributions'

October 14, 2021

Version 1.0.0 **Date** 2021-2-9

Title Produces Summaries and Plots of Features Distributed Across Genomes

Description If you have a set of genomic ranges, this package can help you with visualization and comparison. It produces several kinds of plots, for example: Chromosome distribution plots, which visualize how your regions are distributed over chromosomes; feature distance distribution plots, which visualizes how your regions are distributed relative to a feature of interest, like Transcription Start Sites (TSSs); genomic partition plots, which visualize how your regions overlap given genomic features such as promoters, introns, exons, or intergenic regions. It also makes it easy to compare one set of ranges to another.

Depends R (>= 4.0), IRanges, GenomicRanges

Imports data.table, ggplot2, reshape2, methods, utils, Biostrings,

Suggests AnnotationFilter, rtracklayer, testthat, knitr, BiocStyle, rmarkdown

Enhances BSgenome, extrafont, ensembldb, GenomicFeatures

LazyData true

VignetteBuilder knitr

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biocViews Software, GenomeAnnotation, GenomeAssembly,
DataRepresentation, Sequencing, Coverage, FunctionalGenomics,
Visualization

RoxygenNote 7.1.1

URL http://code.databio.org/GenomicDistributions

BugReports http://github.com/databio/GenomicDistributions

Encoding UTF-8

git_url https://git.bioconductor.org/packages/GenomicDistributions

git_branch RELEASE_3_13

git_last_commit 84c4876
git_last_commit_date 2021-05-19
Date/Publication 2021-10-14
Author Nathan C. Sheffield [aut],
 Kristyna Kupkova [aut, cre],
 Jose Verdezoto [aut],
 Tessa Danehy [ctb],
 John Lawson [ctb],
 Jose Verdezoto [ctb],
 Michal Stolarczyk [ctb],
 Jason Smith [ctb]

Maintainer Kristyna Kupkova <kristynakupkova@gmail.com>

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Description

Checks to make sure a package object is installed, and if so, returns it. If the library is not installed, it issues a warning and returns NULL.

Usage

.requireAndReturn(BSgenomeString)

Arguments

BSgenomeString A BSgenome compatible genome string.

Value

A BSgenome object if installed.

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.validateInputs

Checks class of the list of variables. To be used in functions

Description

Checks class of the list of variables. To be used in functions

Usage

```
.validateInputs(checkList)
```

Arguments

checkList

list of object to check, e.g. list(varname=c("data.frame", "numeric")). Multiuple strings in the vector are treated as OR.

Value

A warning if the wrong input class is provided.

Examples

```
x <- function(var1) {
    cl = list(var1=c("numeric","character"))
    .validateInputs(cl)
    return(var1^2)
}</pre>
```

binBSGenome

Bins a BSgenome object.

Description

Given a BSgenome object (to be loaded via loadBSgenome), and a number of bins, this will bin that genome. It is a simple wrapper of the binChroms function

Usage

```
binBSGenome(genome, binCount)
```

Arguments

genome A UCSC-style string denoting reference assembly (e.g. 'hg38')

binCount number of bins per chromosome

binChroms 5

Value

A data.table object showing the region and bin IDs of the reference genome.

Examples

```
## Not run:
binCount = 1000
refGenomeBins = binBSGenome("hg19", binCount)
## End(Not run)
```

binChroms

Naively splits a chromosome into bins

Description

Given a list of chromosomes with corresponding sizes, this script will produce (roughly) evenly-sized bins across the chromosomes. It does not account for assembly gaps or the like.

Usage

```
binChroms(binCount, chromSizes)
```

Arguments

binCount number of bins (total; *not* per chromosome)

chromSizes a named list of size (length) for each chromosome.

Value

A data.table object assigning a bin ID to each chromosome region.

```
chromSizes = c(chr1=249250621, chr2=243199373, chr3=198022430)
cBins = binChroms(1000, chromSizes)
```

6 binRegion

Ł	inRegion	Divide regions into roughly equal bins

Description

Given a start coordinate, end coordinate, and number of bins to divide, this function will split the regions into that many bins. Bins will be only approximately the same size, due to rounding. (they should not be more than 1 different).

Usage

```
binRegion(start, end, binSize = NULL, binCount = NULL, indicator = NULL)
```

Arguments

start	The starting coordinate
end	The ending coordinate
binSize	The size of bin to divide the genome into. You must supply either binSize (priority) or binCount.
binCount	The number of bins to divide. If you do not supply binSize, you must supply binCount, which will be used to calculate the binSize.
indicator	A vector with identifiers to keep with your bins, in case you are doing this on a long table with multiple segments concatenated

Details

Use case: take a set of regions, like CG islands, and bin them; now you can aggregate signal scores across the bins, giving you an aggregate signal in bins across many regions of the same type.

In theory, this just runs on 3 values, but you can run it inside a data.table j expression to divide a bunch of regions in the same way.

Value

A data.table, expanded to nrow = number of bins, with these id columns: id: region ID binID: repeating ID (this is the value to aggregate across) ubinID: unique bin IDs

```
Rbins = binRegion(1, 3000, 100, 1000)
```

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BSdtToGRanges

Converts a list of data.tables (From BSreadbeds) into GRanges.

Description

Converts a list of data.tables (From BSreadbeds) into GRanges.

Usage

```
BSdtToGRanges(dtList)
```

Arguments

dtList

A list of data.tables

Value

A GRangesList object.

calcChromBins

Calculates the distribution of a query set over the genome

Description

Returns a data.table showing counts of regions from the query that overlap with each bin. In other words, where on which chromosomes are the ranges distributed? You must provide binned regions. Only the midpoint of each query region is used to test for overlap with the bin regions.

Usage

```
calcChromBins(query, bins)
```

Arguments

query A GenomicRanges or GenomicRangesList object with query regions

bins Pre-computed bins (as a GRangesList object) to aggregate over; for example,

these could be genome bins

Value

A data table showing where on which chromosomes ranges are distributed.

```
vistaSftd = GenomicRanges::shift(vistaEnhancers, 100000)
vistaSftd2 = GenomicRanges::shift(vistaEnhancers, 200000)
calcChromBins(vistaEnhancers, GRangesList(vistaSftd, vistaSftd2))
```

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calcChromBinsRef

Returns the distribution of query over a reference assembly Given a query set of elements (a GRanges object) and a reference assembly (*e.g. 'hg38'), this will aggregate and count the distribution of the query elements across bins of the reference genome. This is a helper function to create features for common genomes. It is a wrapper of calcChromBins, which is more general.

Description

Returns the distribution of query over a reference assembly Given a query set of elements (a GRanges object) and a reference assembly (*e.g. 'hg38'), this will aggregate and count the distribution of the query elements across bins of the reference genome. This is a helper function to create features for common genomes. It is a wrapper of calcChromBins, which is more general.

Usage

```
calcChromBinsRef(query, refAssembly, binCount = 10000)
```

Arguments

query A GenomicRanges or GenomicRangesList object with query regions

refAssembly A character vector that will be used to grab chromosome sizes with getChromSizes

binCount Number of bins to divide the chromosomes into

Value

A data.table showing the distribution of regions across bins of the reference genome.

Examples

```
ChromBins = calcChromBinsRef(vistaEnhancers, "hg19")
```

calcCumulativePartitions

Calculates the cumulative distribution of overlaps between query and arbitrary genomic partitions

Description

Takes a GRanges object, then assigns each element to a partition from the provided partitionList, and then tallies the number of regions assigned to each partition. A typical example of partitions is promoter, exon, intron, etc; this function will yield the number of each for a query GRanges object There will be a priority order to these, to account for regions that may overlap multiple genomic partitions.

calcCumulativePartitionsRef

Usage

```
calcCumulativePartitions(query, partitionList, remainder = "intergenic")
```

Arguments

query GRanges or GRangesList with regions to classify.

partitionList An ORDERED and NAMED list of genomic partitions GRanges. This list must

be in priority order; the input will be assigned to the first partition it overlaps.

remainder Which partition do you want to account for 'everything else'?

Value

A data.frame assigning each element of a GRanges object to a partition from a previously provided partitionList.

Examples

calcCumulativePartitionsRef

Calculates the cumulative distribution of overlaps for a query set to a reference assembly

Description

This function is a wrapper for calcCumulativePartitions that uses built-in partitions for a given reference genome assembly.

Usage

```
calcCumulativePartitionsRef(query, refAssembly)
```

Arguments

query A GenomicRanges or GenomicRangesList object with query regions

refAssembly A character vector specifying the reference genome assembly (*e.g.* 'hg19').

This will be used to grab chromosome sizes with getTSSs.

Value

A data frame indicating the number of query region overlaps in several genomic partitions.

Examples

```
calcCumulativePartitionsRef(vistaEnhancers, "hg19")
```

```
calcExpectedPartitions
```

Calculates the distribution of overlaps between query and arbitrary genomic partitions

Description

Takes a GRanges object, then assigns each element to a partition from the provided partitionList, and then tallies the number of regions assigned to each partition. A typical example of partitions is promoter, exon, intron, etc; this function will yield the number of each for a query GRanges object There will be a priority order to these, to account for regions that may overlap multiple genomic partitions.

Usage

```
calcExpectedPartitions(
  query,
  partitionList,
  genomeSize = NULL,
  remainder = "Intergenic"
)
```

Arguments

query GRanges or GRangesList with regions to classify.

partitionList An ORDERED and NAMED list of genomic partitions GRanges. This list must

be in priority order; the input will be assigned to the first partition it overlaps.

genomeSize The number of bases in the query genome. In other words, the sum of all chro-

mosome sizes.

remainder Which partition do you want to account for 'everything else'?

Value

A data.frame assigning each element of a GRanges object to a partition from a previously provided partitionList.

calcExpectedPartitionsRef

Calculates the distribution of observed versus expected overlaps for a query set to a reference assembly

Description

This function is a wrapper for calcExpectedPartitions that uses built-in partitions for a given reference genome assembly.

Usage

```
calcExpectedPartitionsRef(query, refAssembly)
```

Arguments

query A GenomicRanges or GenomicRangesList object with query regions

refAssembly A character vector specifying the reference genome assembly (*e.g.* 'hg19').

This will be used to grab chromosome sizes with getTSSs.

Value

A data frame indicating the number of query region overlaps in several genomic partitions.

Examples

```
calcExpectedPartitionsRef(vistaEnhancers, "hg19")
```

calcFeatureDist	Find the distance to the neare.	st genomic feature

Description

For a given query set of genomic regions, and a given feature set of regions, this function will return the distance for each query region to its closest feature. It ignores strand and returns the distance as positive or negative, depending on whether the feature is upstream or downstream

Usage

```
calcFeatureDist(query, features)
```

Arguments

query A GRanges or GRangesList object with query sets features A GRanges object with features to test distance to 12 calcFeatureDistRefTSS

Details

This function is similar to the bioconductor distanceToNearest function, but returns negative values for downstream distances instead of absolute values. This allows you to assess the relative location.

Value

A vector of genomic distances for each query region relative to its closest feature.

Examples

```
vistaSftd = GenomicRanges::shift(vistaEnhancers, 100000)
calcFeatureDist(vistaEnhancers, vistaSftd)
```

calcFeatureDistRefTSS Calculates the distribution of distances from a query set to closest TSS

Description

Given a query GRanges object and an assembly string, this function will grab the TSS list for the given reference assembly and then calculate the distance from each query feature to the closest TSS. It is a wrapper of calcFeatureDist that uses built-in TSS features for a reference assembly

Usage

```
calcFeatureDistRefTSS(query, refAssembly)
```

Arguments

query A GenomicRanges or GenomicRangesList object with query regions

refAssembly A character vector specifying the reference genome assembly (*e.g.* 'hg19').

This will be used to grab chromosome sizes with getTSSs.

Value

A vector of distances for each query region relative to TSSs.

```
calcFeatureDistRefTSS(vistaEnhancers, "hg19")
```

calcGCContent 13

calcGCContent Calculate GC content over genomic ranges	calcGCContent	Calculate GC content over genomic ranges	
--	---------------	--	--

Description

Given a reference genome as a BSgenome object and some ranges on that reference, this function will return a vector of the same length as the granges object, with percent of Cs and Gs.

Usage

```
calcGCContent(query, ref)
```

Arguments

query A GenomicRanges or GenomicRangesList object with query regions.

ref Reference genome BSgenome object.

Value

A numeric vector of list of vectors with the GC percentage of the query regions.

Examples

```
## Not run:
bsg = loadBSgenome('hg19')
gcvec = calcGCContent(vistaEnhancers, bsg)
## End(Not run)
```

calcGCContentRef

Calculate GC content over genomic ranges

Description

Given a reference genome as a BSgenome object and some ranges on that reference, this function will return a vector of the same length as the granges object, with percent of Cs and Gs.

Usage

```
calcGCContentRef(query, refAssembly)
```

Arguments

query A GenomicRanges or GenomicRangesList object with query regions

refAssembly A character vector specifying the reference genome assembly (*e.g.* 'hg19').

This will be used to grab chromosome sizes with getTSSs.

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Value

A numeric vector or list of vectors with the GC percentage of the query regions.

Examples

```
## Not run:
refAssembly = 'hg19'
GCcontent = calcGCContentRef(vistaEnhancers, refAssembly)
## End(Not run)
```

calcNeighborDist

Group regions from the same chromosome together and calculate the distances between neighboring regions. Distances are then lumped into a numeric vector.

Description

Group regions from the same chromosome together and calculate the distances between neighboring regions. Distances are then lumped into a numeric vector.

Usage

```
calcNeighborDist(query)
```

Arguments

query

A GRanges or GRangesList object.

Value

A numeric vector or list with different vectors containing the distances within neighboring regions.

```
dist = calcNeighborDist(vistaEnhancers)
```

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calcOpenSignal	The function calcOpenSignal takes the input BED file(s) in form of GRanges or GRangesList object, overlaps it with all defined open chromatin regions across cell types and returns a matrix, where each row is the input genomic region (if overlap was found), each column is a cell type, and the value is a normalized ATAC-seq signal.
----------------	---

Description

The function calcOpenSignal takes the input BED file(s) in form of GRanges or GRangesList object, overlaps it with all defined open chromatin regions across cell types and returns a matrix, where each row is the input genomic region (if overlap was found), each column is a cell type, and the value is a normalized ATAC-seq signal.

Usage

```
calcOpenSignal(query, cellMatrix)
```

Arguments

allow.	Conomia ragions to be analyzed	Can be Changes or Changes List object
query	denomic regions to be analyzed.	. Can be GRanges or GRangesList object.

cellMatrix Matrix with open chromatin signal values, rows are genomic regions, columns

are cell types. First column contains information about the genomic region in following form: chr_start_end. Can be either data.frame or data.table object.

Value

A list with named components: signalMatrix - data.table with cell specific open chromatin signal values for query regions matrixStats - data.frame containing boxplot stats for individual cell type

Examples

openRegionSummary = calcOpenSignal(vistaEnhancers, exampleOpenSignalMatrix_hg19)

calcPartitions	Calculates the distribution of overlaps between query and arbitrary
	genomic partitions

Description

Takes a GRanges object, then assigns each element to a partition from the provided partitionList, and then tallies the number of regions assigned to each partition. A typical example of partitions is promoter, exon, intron, etc; this function will yield the number of each for a query GRanges object There will be a priority order to these, to account for regions that may overlap multiple genomic partitions.

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Usage

```
calcPartitions(query, partitionList, remainder = "intergenic")
```

Arguments

query GRanges or GRangesList with regions to classify

partitionList an ORDERED and NAMED list of genomic partitions GRanges. This list must

be in priority order; the input will be assigned to the first partition it overlaps

remainder A character vector to assign any query regions that do not overlap with anything

in the partitionList. Defaults to "intergenic"

Value

A data frame assigning each element of a GRanges object to a partition from a previously provided partition List.

Examples

calcPartitionsRef

Calculates the distribution of overlaps for a query set to a reference assembly

Description

This function is a wrapper for calcPartitions and calcPartitionPercents that uses built-in partitions for a given reference genome assembly.

Usage

```
calcPartitionsRef(query, refAssembly)
```

Arguments

query A GenomicRanges or GenomicRangesList object with query regions

refAssembly A character vector specifying the reference genome assembly (*e.g.* 'hg19').

This will be used to grab chromosome sizes with getTSSs.

Value

A data frame indicating the number of query region overlaps in several genomic partitions.

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Examples

```
calcPartitionsRef(vistaEnhancers, "hg19")
```

calcWidth

Calculate the widths of regions

Description

The length of a genomic region (the distance between the start and end) is called the width When given a query set of genomic regions, this function returns the width

Usage

```
calcWidth(query)
```

Arguments

query

A GRanges or GRangesList object with query sets

Value

A vector of the widths (end-start coordinates) of GRanges objects.

Examples

```
TSSdist = calcFeatureDistRefTSS(vistaEnhancers, "hg19")
plotFeatureDist(TSSdist, featureName="TSS")
```

cellTypeMetadata

Table the maps cell types to tissues and groups

Description

Table the maps cell types to tissues and groups

Usage

```
data(cellTypeMetadata)
```

Format

data.table with 3 columns (cellType, tissue and group) and 74 rows (one per cellType)

Source

self-curated dataset

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chromSizes_hg19

hg19 chromosome sizes

Description

A dataset containing chromosome sizes for Homo Sapiens hg38 genome assembly

Usage

```
data(chromSizes_hg19)
```

Format

A named vectors of lengths with one item per chromosome

Source

BSgenome. Hsapiens. UCSC. hg19 package

dtToGr

Converts a data.table (DT) object to a GenomicRanges (GR) object. Tries to be intelligent, guessing chr and start, but you have to supply end or other columns if you want them to be carried into the GR.

Description

Converts a data.table (DT) object to a GenomicRanges (GR) object. Tries to be intelligent, guessing chr and start, but you have to supply end or other columns if you want them to be carried into the GR.

Usage

```
dtToGr(
  DT,
  chr = "chr",
  start = "start",
  end = NA,
  strand = NA,
  name = NA,
  splitFactor = NA,
  metaCols = NA
```

dtToGrInternal 19

Arguments

DT A data.table representing genomic regions.

chr A string representing the chromosome column.

start A string representing the name of the start column.

end A string representing the name of the end column.

strand A string representing the name of the strand column.

name A string representing the name of the name column.

splitFactor A string representing the name of the column to use to split the data.table into

multiple data.tables.

metaCols A string representing the name of the metadata column(s) to include in the re-

turned GRanges object.

Value

A GRanges object.

Examples

dtToGrInternal

Two utility functions for converting data.tables into GRanges objects

Description

Two utility functions for converting data.tables into GRanges objects

Usage

```
dtToGrInternal(DT, chr, start, end = NA, strand = NA, name = NA, metaCols = NA)
```

Arguments

DT	A data.table representing genomic regions.
chr	A string representing the chromosome column.
start	A string representing the name of the start column.
end	A string representing the name of the end column.
strand	A string representing the name of the strand column.
name	A string representing the name of the name column.

metaCols A string representing the name of the metadata column(s) to include in the re-

turned GRanges object.

Value

A GRanges object.

exampleOpenSignalMatrix_hg19

A dataset containing a subset of open chromatin regions across all cell types defined by ENCODE for Homo Sapiens hg19

Description

Preparation steps:

- made a universe of regions by merging regions across cell types defined as opened in EN-CODE
- 2. took bigwig files from ENCODE for individual cell types, merged replicates, filtered out blacklisted sites
- 3. evaluated the signal above regions defined by previous step
- 4. performed quantile normalization
- 5. subsetted it

Usage

data(exampleOpenSignalMatrix_hg19)

Format

data.frame, rows represent whole selection of open chromatin regions across all cell types defined by ENCODE, columns are individual cell types and values are normalized open chromatin signal values.

Source

http://big.databio.org/open_chromatin_matrix/openSignalMatrix_hg19_quantileNormalized_ round4.txt.gz geneModels_hg19 21

|--|

Description

A dataset containing gene models for Homo Sapiens hg38 genome assembly.

Usage

```
data(geneModels_hg19)
```

Format

A list of two GRanges objects, with genes and exons locations

Source

EnsDb.Hsapiens.v75 package

genomePartitionList	Create a basic genome partition list of genes, exons, introns, UTRs,
	and intergenic

Description

Given GRanges for genes, and a GRanges for exons, returns a list of GRanges corresponding to various breakdown of the genome, based on the given annotations; it gives you proximal and core promoters, exons, and introns.

Usage

```
genomePartitionList(genesGR, exonsGR, threeUTRGR = NULL, fiveUTRGR = NULL)
```

Arguments

fiveUTRGR

```
genesGR a GRanges object of gene coordinates
exonsGR a GRanges object of exons coordinates
threeUTRGR a GRanges object of 3' UTRs
```

a GRanges object of 5' UTRs

Details

To be used as a partitionList for calcPartitions.

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Value

A list of GRanges objects, each corresponding to a partition of the genome. Partitions include proximal and core promoters, exons and introns.

Examples

GenomicDistributions

Produces summaries and plots of features distributed across genomes

Description

If you have a set of genomic ranges, the GenomicDistributions R package can help you with some simple visualizations. Currently, it can produce two kinds of plots: First, the chromosome distribution plot, which visualizes how your regions are distributed over chromosomes; and second, the feature distribution plot, which visualizes how your regions are distributed relative to a feature of interest, like Transcription Start Sites (TSSs).

Author(s)

Nathan C. Sheffield

References

http://github.com/databio/GenomicDistributions

getChromSizes

Returns built-in chrom sizes for a given reference assembly

Description

Returns built-in chrom sizes for a given reference assembly

Usage

```
getChromSizes(refAssembly)
```

Arguments

refAssembly

A string identifier for the reference assembly

getGeneModels 23

Value

A vector with the chromosome sizes corresponding to a specific genome assembly.

Examples

```
getChromSizes("hg19")
```

getGeneModels

Returns built-in gene models for a given reference assembly

Description

Some functions require gene models, which can obtained from any source. This function allows you to retrieve a few common built-in ones.

Usage

```
getGeneModels(refAssembly)
```

Arguments

refAssembly

A string identifier for the reference assembly

Value

A list containing the gene models corresponding to a specific reference assembly.

Examples

```
getGeneModels("hg19")
```

getReferenceData

Get reference data for a specified assembly

Description

This is a generic getter function that will return a data object requested, if it is included in the built-in data with the GenomicDistributions package or GenomicDistributionsData package (if installed). Data objects can be requested for different reference assemblies and data types (specified by a tagline, which is a unique string identifying the data type).

Usage

```
getReferenceData(refAssembly, tagline)
```

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Arguments

refAssembly Reference assembly string (e.g. 'hg38')

tagline The string that was used to identify data of a given type in the data building

step. It's used for the filename so we know what to load, and is what makes this

function generic (so it can load different data types).

Value

A requested and included package data object.

grToDt

Convert a GenomicRanges into a data.table.

Description

Convert a GenomicRanges into a data.table.

Usage

```
grToDt(GR)
```

Arguments

GR

A Granges object

Value

A data.table object.

labelCuts

Creates labels based on a discretization definition.

Description

If you are building a histogram of binned values, you want to have labels for your bins that correspond to the ranges you used to bin. This function takes the breakpoints that define your bins and produces nice-looking labels for your histogram plot.

Usage

```
labelCuts(
  breakPoints,
  round_digits = 1,
  signif_digits = 3,
  collapse = "-",
  infBins = FALSE
)
```

loadBSgenome 25

Arguments

breakPoints The exact values you want as boundaries for your bins

round_digits Number of digits to cut round labels to.
signif_digits Number of significant digits to specify.

collapse Character to separate the labels infBins use >/< as labels on the edge bins

Details

labelCuts will take a cut group, (e.g., a quantile division of some signal), and give you clean labels (similar to the cut method).

Value

A vector of histogram axis labels.

1oadBSgenome

Loads BSgenome objects from UCSC-style character vectors.

Description

This function will let you use a simple character vector (e.g. 'hg19') to load and then return BSgenome objects. This lets you avoid having to use the more complex annotation for a complete BSgenome object (e.g. BSgenome.Hsapiens.UCSC.hg38.masked)

Usage

```
loadBSgenome(genomeBuild, masked = TRUE)
```

Arguments

genomeBuild One of 'hg19', 'hg38', 'mm10', 'mm9', or 'grch38' masked Should we used the masked version? Default:TRUE

Value

A BSgenome object corresponding to the provided genome build.

```
## Not run:
bsg = loadBSgenome('hg19')
## End(Not run)
```

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loadEnsDb

Load selected EnsDb library

Description

Load selected EnsDb library

Usage

```
loadEnsDb(genomeBuild)
```

Arguments

```
genomeBuild string, genome identifier
```

Value

loaded library

Examples

```
## Not run:
loadEnsDb("hg19")
## End(Not run)
```

neighbordt

Internal helper function to calculate distance between neighboring regions.

Description

Internal helper function to calculate distance between neighboring regions.

Usage

```
neighbordt(querydt)
```

Arguments

querydt

A data table with regions grouped according to chromosome.

Value

A numeric vector with the distances in bp

nlist 27

nlist

Nathan's magical named list function. This function is a drop-in replacement for the base list() function, which automatically names your list according to the names of the variables used to construct it. It seamlessly handles lists with some names and others absent, not overwriting specified names while naming any unnamed parameters. Took me awhile to figure this out.

Description

Nathan's magical named list function. This function is a drop-in replacement for the base list() function, which automatically names your list according to the names of the variables used to construct it. It seamlessly handles lists with some names and others absent, not overwriting specified names while naming any unnamed parameters. Took me awhile to figure this out.

Usage

```
nlist(...)

Arguments
... arguments passed to list()
```

Value

A named list object.

Examples

```
x=5
y=10
nlist(x,y) # returns list(x=5, y=10)
list(x,y) # returns unnamed list(5, 10)
```

plotChromBins

Plot distribution over chromosomes

Description

Plots result from genomicDistribution calculation

Usage

```
plotChromBins(
   genomeAggregate,
   binCount = 10000,
   plotTitle = "Distribution over chromosomes"
)
```

Arguments

genomeAggregate

The output from the genomicDistribution function

binCount Number of bins (should match the call to genomicDistribution)

plotTitle Title for plot.

Value

A ggplot object showing the distribution of the query regions over bins of the reference genome.

Examples

plotCumulativePartitions

Plot the cumulative distribution of regions in features

Description

This function plots the cumulative distribution of regions across a feature set.

Usage

```
plotCumulativePartitions(assignedPartitions, feature_names = NULL)
```

Arguments

assignedPartitions

Results from calcCumulativePartitions

feature_names An optional character vector of feature names, in the same order as the Genom-

icRanges or GenomicRangesList object.

Value

A ggplot object of the cumulative distribution of regions in features.

```
p = calcCumulativePartitionsRef(vistaEnhancers, "hg19")
cumuPlot = plotCumulativePartitions(p)
```

plotExpectedPartitions 29

```
plotExpectedPartitions
```

Produces a barplot showing how query regions of interest are distributed relative to the expected distribution across a given partition list

Description

Produces a barplot showing how query regions of interest are distributed relative to the expected distribution across a given partition list

Usage

```
plotExpectedPartitions(expectedPartitions, feature_names = NULL)
```

Arguments

expectedPartitions

A data frame holding the frequency of assignment to each of the partitions, the expected number of each partition, and the log10 of the observed over expected. Produced by calcExpectedPartitions.

feature_names

Character vector with labels for the partitions (optional). By default it will use the names from the first argument.

Value

A ggplot object using a barplot to show the distribution of the query regions across a given partition list.

Examples

```
p = calcExpectedPartitionsRef(vistaEnhancers, "hg19")
expectedPlot = plotExpectedPartitions(p)
```

plotFeatureDist

Plots a histogram of distances to genomic features

Description

Given the results from featureDistribution, plots a histogram of distances surrounding the features of interest

30 plotGCContent

Usage

```
plotFeatureDist(
  dists,
  bgdists = NULL,
  featureName = "features",
  numbers = FALSE,
  nbins = 50,
  size = 1e+05,
  infBins = FALSE,
  tile = FALSE
)
```

Arguments

dists Results from featureDistribution

bgdists Background distances. If provided, will plot a background distribution of ex-

pected distances

featureName Character vector for plot labels (optional).

numbers a logical indicating whether the raw numbers should be displayed, rather than

percentages (optional).

nbins Number of bins on each side of the center point.

size Number of bases to include in plot on each side of the center point.

infBins Include catch-all bins on the sides?

tile Turn on a tile mode, which plots a tiled figure instead of a histogram.

Value

A ggplot2 plot object

Examples

```
TSSdist = calcFeatureDistRefTSS(vistaEnhancers, "hg19")
f = plotFeatureDist(TSSdist, featureName="TSS")
```

plotGCContent Plots a density distribution of GC vectors Give results from the

calcGCContent function, this will produce a density plot

Description

Plots a density distribution of GC vectors Give results from the calcGCContent function, this will produce a density plot

Usage

```
plotGCContent(gcvectors)
```

plotNeighborDist 31

Arguments

gcvectors

A numeric vector or list of numeric vectors of GC contents.

Value

A ggplot object plotting distribution of GC content in query regions.

Examples

plotNeighborDist

Plot the distances between neighboring regions. The distance in the x axis is log10 transformed for ease of comparison between different regionsets and to account for outliers.

Description

Plot the distances between neighboring regions. The distance in the x axis is log10 transformed for ease of comparison between different regionsets and to account for outliers.

Usage

```
plotNeighborDist(dcvec)
```

Arguments

dcvec

A numeric vector or list with vectors containing distances between neighbor regions. Produced by calcNeighborDist

Value

A ggplot density object showing the distribution of log10 transformed distances.

```
numVector = rnorm(400, mean=5, sd=0.1)
d = plotNeighborDist(numVector)
```

32 plotOpenSignal

plotOpenSignal	The function plotOpenSignal visualizes the signalMatrix obtained
	from calcOpenSignal.

Description

The function plotOpenSignal visualizes the signalMatrix obtained from calcOpenSignal.

Usage

Arguments

openRegionSummary

Output list from calcOpenSignal function.

plotType

Options are: jitter - jitter plot with box plot on top boxPlot - box plot without individual points and outliers barPlot (default) - bar height represents the median signal value for a given cell type.

cellGroup

- This option allows to selcet a tissue type to be plotted, if NA (default) all available tissue types are ploted, available options: "blood", "bone", "CNS", "embryonic", "eye", "foreskin", "gastrointestinal", "heart", "liver", "lymphatic", "mammaryGland", "mouth", "respiratorySystem", "skeletalMuscle", "skin", "urinarySystem", "vasculature", can be passed as a character string or vector of strings.

cellTypeMetadata

Metadata for cell type - tissue association. This option is for users, who provide their own open region signal matrix. The cellTypeMetadata matrix must contain two columns called cellType and tissueType. cellType column containes the cell type names in the provided signalMatrix column names. The tissueType columns provides an information, which tissue the cell type comes from.

colorScheme

Provide color values for each tissue Type if you want to change the default colors.

Value

A ggplot object.

plotPartitions 33

Examples

```
## Not run:
openRegionSummary = calcOpenSignal(vistaEnhancers, exampleOpenSignalMatrix_hg19)
plotSignal = plotOpenSignal(openRegionSummary)
plotSignal = plotOpenSignal(openRegionSummary, plotType = "jitter",
cellGroup = "blood")
## End(Not run)
```

plotPartitions

Produces a barplot showing how query regions of interest are distributed across a given partition list

Description

This function can be used to test a GRanges object against any arbitrary list of genome partitions. The partition list is a priority-ordered list of GRanges objects. Each region in the query will be assigned to a given partition that it overlaps with the highest priority.

Usage

```
plotPartitions(assignedPartitions, numbers = FALSE)
```

Arguments

assignedPartitions

A table holding the frequency of assignment to each of the partitions. Produced

by calcPartitions

numbers logical indicating whether raw overlaps should be plotted instead of the default

percentages

Value

A ggplot object using a barplot to show the distribution of the query regions across a given partition list.

```
p = calcPartitionsRef(vistaEnhancers, "hg19")
partPlot = plotPartitions(p)
partCounts = plotPartitions(p, numbers=TRUE)
```

34 plotQTHist

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Plot quantile-trimmed histogram

Description

Given the results from calcWidth, plots a histogram with outliers trimmed.

Usage

```
plotQTHist(
    x,
    EndBarColor = "gray57",
    MiddleBarColor = "gray27",
    quantile = NULL,
    bins = NULL,
    indep = FALSE,
    numbers = FALSE
)
```

Arguments

x Data values to plot

EndBarColor Color for the quantile bars on both ends of the graph (optional)

MiddleBarColor Color for the bars in the middle of the graph (optional)

quantile Quantile of data to be contained in each end bar (optional) Quantiles must be

under .2, optimal size is under .1

bins The number of bins for the histogram to allocate data to. (optional)

indep logical value which returns a list of plots that have had their bins calculated

independently; the normal version will plot them on the same x and y axis.

numbers a logical indicating whether the raw numbers should be displayed, rather than

percentages (optional).

Details

x-axis breaks for the frequency calculations are based on the "divisions" results from helper function calcDivisions.

Value

A ggplot2 plot object

```
plotQTHist(runif(500)*1000)
plotQTHist(list(q1=runif(500)*1000, q2=runif(500)*1000))
```

setB_100 35

setB_100

Example BED file read with rtracklayer::import

Description

Example BED file read with rtracklayer::import

Usage

```
data(setB_100)
```

Format

GenomicRanges::GRanges

splitDataTable

Efficiently split a data.table by a column in the table

Description

Efficiently split a data.table by a column in the table

Usage

```
splitDataTable(DT, split_factor)
```

Arguments

DT Data.table to split

split_factor Column to split, which can be a character vector or an integer.

Value

List of data.table objects, split by column

36 TSS_hg19

```
theme_blank_facet_label
```

Clear ggplot face label.

Description

Usually ggplot2 facets are labeled with boxes surrounding the label. This function removes the box, so it's a simple label for each facet.

Usage

```
theme_blank_facet_label()
```

Value

A ggplot theme

TSS_hg19

hg19 TSS locations

Description

A dataset containing chromosome sizes for Homo Sapiens hg38 genome assembly

Usage

```
data(TSS_hg19)
```

Format

A named vectors of lengths with one item per chromosome

Source

EnsDb.Hsapiens.v75 package

vistaEnhancers 37

 ${\tt vistaEnhancers}$

 ${\it Example BED file read with rtracklayer:: import}$

Description

Example BED file read with rtracklayer::import

Usage

data(vistaEnhancers)

Format

GenomicRanges::GRanges

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