Package ‘GenomicPlot’

March 29, 2024

Type Package

Title Plot profiles of next generation sequencing data in genomic features

Version 1.0.6

Description Visualization of next generation sequencing (NGS) data is essential for interpreting high-throughput genomics experiment results. 'GenomicPlot' facilitates plotting of NGS data in various formats (bam, bed, wig and bigwig); both coverage and enrichment over input can be computed and displayed with respect to genomic features (such as UTR, CDS, enhancer), and user defined genomic loci or regions. Statistical tests on signal intensity within user defined regions of interest can be performed and represented as boxplots or bar graphs. Parallel processing is used to speed up computation on multicore platforms. In addition to genomic plots which is suitable for displaying of coverage of genomic DNA (such as ChIPseq data), metagenomic (without introns) plots can also be made for RNAseq or CLIPseq data as well.

License GPL-2

Encoding UTF-8

LazyData FALSE

Collate `` DrawingFunctions.R" " GenomicPlot.R" " HandleDataMatrix.R"
`` HandleFeatures.R" " Parallel.R" " ReadData.R" " Setup.R"
`` Plot_5parts_metagene.R" " Plot_start_end.R"
`` Plot_start_end_with_random.R" " Plot_region.R" " Plot_locus.R"
`` data.R" " Plot_locus_with_random.R" " Plot_peak_annotation.R"
`` Plot_bam_correlation.R"

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Perform one-way ANOVA and post hoc TukeyHSD tests

Description

This is a helper function for performing one-way ANOVA analysis and post hoc Tukey’s Honest Significant Differences tests

Usage

aov_TukeyHSD(df, xc = "Group", yc = "Intensity", op = NULL, verbose = FALSE)

Arguments

df a dataframe
xc a string denoting column name for grouping
yc a string denoting column name for numeric data to be plotted
op output prefix for statistical analysis results
verbose logical, to indicate whether a file should be produced to save the test results

Value

a list of two elements, the first is the p-value of ANOVA test and the second is a matrix of the output of TukeyHSD tests

Note

used in plot_locus

Author(s)

Shuye Pu

Examples

stat_df <- data.frame(
  Feature = rep(c("A", "B"), c(20, 30)),
  Intensity = c(rnorm(20, mean = 2, sd = 1), rnorm(30, mean = 3, sd = 1))
)

out <- aov_TukeyHSD(stat_df, xc = "Feature")
out
check_constraints

Check constraints of genomic ranges

Description

Make sure the coordinates of GRanges are within the boundaries of chromosomes, and trim anything that goes beyond. Also, remove entries whose seqname is not in the seqname of a query GRanges.

Usage

check_constraints(gr, genome, queryRle = NULL)

Arguments

- gr: a GenomicRanges object
- genome: genomic version name such as "hg19"
- queryRle: a RleList object used as a query against gr

Value

a GRanges object

Author(s)

Shuye Pu

Examples

```r
subject <- GRanges("chr19",
    IRanges(rep(c(10, 15), 2), width = c(1, 20, 400, 2e+8)),
    strand = c("+", "+", "-", "-")
)

g <- check_constraints(gr = subject, genome = "hg19")
identical(g, subject)

subject1 <- GRanges("chr19",
    IRanges(rep(c(10, 15), 2), width = c(1, 20, 400, 28)),
    strand = c("+", "+", "-", "-")
)

g1 <- check_constraints(gr = subject1, genome = "hg19")
identical(g1, subject1)
```
custom_TxDB_from_GTF  Make custom TxDB object from a GTF/GFF file

Description
This is a helper function for creating custom TxDB object from a GTF/GFF file. Mitochondrial chromosome is excluded.

Usage

custom_TxDB_from_GTF(gtfFile, genome = "hg19")

Arguments

*gtfFile*  path to a gene annotation gtf file

*genome*  a string denoting the genome name and version

Value

a TxDB object defined in the GenomicFeatures package.

Author(s)

Shuye Pu

Examples

```r
gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", package = "GenomicPlot")
txdb <- custom_TxDB_from_GTF(gtfFile, genome = "hg19")
```

draw_boxplot_by_factor

*Plot boxplot with two factors*

Description

Plot violin plot with boxplot components for data with one or two factors, p-value significance levels are displayed. "****" = 0.001, "***" = 0.01, "**" = 0.05.
Usage

draw_boxplot_by_factor(
  stat_df,
  xc = "Feature",
  yc = "Intensity",
  fc = xc,
  comp = list(c(1, 2)),
  stats = "wilcox.test",
  Xlab = xc,
  Ylab = yc,
  nf = 1
)

Arguments

  stat_df a dataframe with column names c(xc, yc)
  xc a string denoting column name for grouping
  yc a string denoting column name for numeric data to be plotted
  fc a string denoting column name for sub-grouping based on an additional factor
  comp a list of vectors denoting pair-wise comparisons to be performed between groups
  stats the name of pair-wise statistical tests, like t.test or wilcox.test
  Xlab a string for x-axis label
  Ylab a string for y-axis label
  nf a integer normalizing factor for correct count of observations when the data table
       has two factors, such as those produced by 'pivot_longer', equals to the number
       of factors

Value

  a ggplot object

Note

  used by plot_locus, plot_locus_with_random, plot_region

Author(s)

  Shuye Pu

Examples

  stat_df <- data.frame(
    Feature = rep(c("A", "B"), c(20, 30)),
    Intensity = c(rnorm(20, 2, 0.5), rnorm(30, 3, 0.6))
  )
  p <- draw_boxplot_by_factor(stat_df,
    xc = "Feature", yc = "Intensity",
    }
draw_boxplot_wo_outlier

Plot boxplot without outliers

Description

Plot boxplot without outliers, useful when outliers have a wide range and the median is squeezed at the bottom of the plot. The p-value significance level is the same as those in draw_boxplot_by_factor, but not displayed.

Usage

draw_boxplot_wo_outlier(
    stat_df,  
    xc = "Feature",  
    yc = "Intensity",  
    fc = xc,  
    comp = list(c(1, 2)),  
    stats = "wilcox.test",  
    Xlab = xc,  
    Ylab = yc,  
    nf = 1
)

Arguments

stat_df a dataframe with column names c(xc, yc)
xc a string denoting column name for grouping
yc a string denoting column name for numeric data to be plotted
fc a string denoting column name for sub-grouping
comp a list of vectors denoting pair-wise comparisons to be performed between groups
stats the name of pair-wise statistical tests, like t.test or wilcox.test
Xlab a string for x-axis label
Ylab a string for y-axis label
nf a integer normalizing factor for correct count of observations when the data table has two factors, such as those produced by ‘pivot_longer’, equals to the number of factors

Value

a ggplot object
Examples

```r
stat_df <- data.frame(
  Feature = rep(c("A", "B"), c(20, 30)),
  Intensity = c(rnorm(20, 2), rnorm(30, 3))
)

p <- draw_boxplot_wo_outlier(stat_df,
  xc = "Feature", yc = "Intensity",
  Ylab = "Signal Intensity"
)
p
```

---

**draw_combo_plot**

Make combo plot for statistics plots

**Description**

Place violin plot, boxplot without outliers, mean+se barplot and quantile plot on the same page

**Usage**

```r
draw_combo_plot(
  stat_df,
  xc = "Feature",
  yc = "Intensity",
  comp = list(c(1, 2)),
  Xlab = xc,
  Ylab = yc,
  stats = "wilcox.test",
  fc = xc,
  Ylim = NULL,
  title = "",
  nf = 1
)
```

**Arguments**

- `stat_df`: a dataframe with column names `c(xc, yc)`
- `xc`: a string denoting column name for grouping
- `yc`: a string denoting column name for numeric data to be plotted
- `comp`: a list of vectors denoting pair-wise comparisons to be performed between groups
- `Xlab`: a string for x-axis label
- `Ylab`: a string for y-axis label
- `stats`: the name of pair-wise statistical tests, like `t.test` or `wilcox.test`
draw_locus_profile

Plot signal profile around genomic loci

Description

Plot lines with standard error as the error band

Usage

draw_locus_profile(
plot_df,
xc = "Position",
yc = "Intensity",
cn = "Query",
fn = NULL,
fc = NULL,
Ylim = NULL,
title = NULL,
nf = NULL
)
Arguments

- **plot_df**: a dataframe with column names c(xc, yc, cn, "lower", "upper")
- **xc**: a string denoting column name for values on x-axis
- **yc**: a string denoting column name for numeric data to be plotted
- **cn**: a string denoting column name for sample grouping, like 'Query' or 'Reference'
- **sn**: a string denoting column name for the subject of sample grouping, if 'cn' is 'Query', then 'sn' will be 'Reference'
- **Xlab**: a string for x-axis label
- **Ylab**: a string for y-axis label
- **shade**: logical indicating whether to place a shaded rectangle around the loci bounded by hl
- **hl**: a vector of two integers defining upstream and downstream boundaries of the rectangle

Value

a ggplot object

Note

used by `plot_locus, plot_locus_with_random`

Author(s)

Shuye Pu

Examples

```r
library(dplyr)
Reference <- rep(rep(c("Ref1", "Ref2"), each = 100), 2)
Query <- rep(c("Query1", "Query2"), each = 200)
Position <- rep(seq(-50, 49), 4)
Intensity <- rlnorm(400)
se <- runif(400)
df <- data.frame(Intensity, se, Position, Query, Reference) %>%
  mutate(lower = Intensity - se, upper = Intensity + se) %>%
  mutate(Group = paste(Query, Reference, sep = ":"))

p <- draw_locus_profile(df, cn = "Group", shade = TRUE, hl = c(-10, 20))
p
```
draw_matrix_heatmap  
Display matrix as a heatmap

Description

Make a complex heatmap with column annotations

Usage

draw_matrix_heatmap(
  fullMatrix,
  dataName = "geneData",
  labels_col = NULL,
  levels_col = NULL,
  ranking = "Sum",
  ranges = NULL,
  verbose = FALSE
)

Arguments

fullMatrix  a numeric matrix
dataName  the nature of the numeric data
labels_col  a named vector for column annotation
levels_col  factor levels for names of labels_col, specifying the order of labels_col
ranking  method for ranking the rows of the input matrix, options are c("Sum", "Max",
  "Hierarchical", "None")
ranges  a numeric vector with three elements, defining custom range for color ramp, default=NULL, i.e. the range is defined automatically based on the c(minimum, median, maximum) of fullMatrix
verbose  logical, whether to output the input matrix for inspection

Value

a grob object

Author(s)

Shuye Pu
draw_mean_se_barplot

Examples

```r
fullMatrix <- matrix(rnorm(10000), ncol = 100)
for (i in seq_len(80)) {
  fullMatrix[i, 16:75] <- runif(60) + i
}
labels_col <- as.character(seq_len(100))
levels_col <- c("start", "center", "end")
names(labels_col) <- rep(levels_col, c(15, 60, 25))
draw_matrix_heatmap(fullMatrix, dataName = "test", labels_col, levels_col,
  ranges = c(-2, 0, 20))
```

draw_mean_se_barplot  Plot barplot for mean with standard error bars

Description

Plot barplot for mean with standard error bars, no p-value significance levels are displayed, but ANOVA p-value is provided as tag and TukeyHSD test are displayed as caption.

Usage

```r
draw_mean_se_barplot(
  stat_df,
  xc = "Feature",
  yc = "Intensity",
  fc = xc,
  comp = list(c(1, 2)),
  Xlab = xc,
  Ylab = yc,
  Ylim = NULL,
  nf = 1
)
```

Arguments

- **stat_df**: a dataframe with column names c(xc, yc)
- **xc**: a string denoting column name for grouping
- **yc**: a string denoting column name for numeric data to be plotted
- **fc**: a string denoting column name for sub-grouping based on an additional factor
- **comp**: a list of vectors denoting pair-wise comparisons to be performed between groups
- **Xlab**: a string for x-axis label
- **Ylab**: a string for y-axis label
- **Ylim**: a numeric vector of two elements, defining custom limits of y-axis
draw_quantile_plot

nf

a integer normalizing factor for correct count of observations when the data table has two factors, such as those produced by pivot_longer, equals to the number of factors

Value

a ggplot object

Note

used by plot_locus, plot_locus_with_random

Author(s)

Shuye Pu

Examples

```r
stat_df <- data.frame(
  Feature = rep(c("A", "B"), c(20, 30)),
  Intensity = c(rnorm(20, 2), rnorm(30, 3))
)
p <- draw_mean_se_barplot(stat_df,
  xc = "Feature",
  yc = "Intensity",
  Ylab = "Intensity"
)
p
```

---

draw_quantile_plot | Plot quantile over value

Description

Plot quantiles as y-axis, and values as x-axis. Same as 'geom_ecdf', but allows sub-grouping by a second factor.

Usage

draw_quantile_plot(
  stat_df, 
  xc = "Feature", 
  yc = "Intensity", 
  Ylab = yc, 
  fc = xc
)
Arguments

- `stat_df`: a dataframe with column names (xc, yc)
- `xc`: a string denoting column name for grouping
- `yc`: a string denoting column name for numeric data to be plotted
- `Ylab`: a string for y-axis label
- `fc`: a string denoting column name for sub-grouping based on an additional factor

Value

- a ggplot object

Note

used by `plot_locus, plot_locus_with_random`

Author(s)

Shuye Pu

Examples

```r
stat_df <- data.frame(
  Feature = rep(c("A", "B"), c(20, 30)),
  Intensity = c(rnorm(20, 2, 5), rnorm(30, 3, 5)),
  Height = c(rnorm(20, 5, 5), rnorm(30, 1, 5))
)
stat_df_long <- tidyr::pivot_longer(stat_df, 
  cols = c(Intensity, Height), names_to = "type", 
  values_to = "value"
)
print(draw_quantile_plot(stat_df, xc = "Feature", yc = "Intensity"))
print(draw_quantile_plot(stat_df, xc = "Feature", yc = "Height"))
print(draw_quantile_plot(stat_df_long, 
  xc = "Feature", yc = "value", 
  fc = "type", Ylab = "value"
))
```

---

**draw_rank_plot**  
Plot fraction of cumulative sum over rank

Description

Plot cumulative sum over rank as line plot, both cumulative sum and rank are scaled between 0 and 1. This is the same as the fingerprint plot of the deepTools.
Usage

draw_rank_plot(stat_df, xc = "Feature", yc = "Intensity", Ylab = yc)

Arguments

stat_df a dataframe with column names c(xc, yc)
xc a string denoting column name for grouping
yc a string denoting column name for numeric data to be plotted
Ylab a string for y-axis label

Value

a ggplot object

Author(s)

Shuye Pu

Examples

stat_df <- data.frame(
  Feature = rep(c("A", "B"), c(20, 30)),
  Intensity = c(rlnorm(20, 5, 5), rlnorm(30, 1, 5))
)
stat_df1 <- data.frame(
  Feature = rep(c("A", "B"), c(20, 30)),
  Height = c(rnorm(20, 5, 5), rnorm(30, 1, 5))
)

print(draw_rank_plot(stat_df,
  xc = "Feature", yc = "Intensity",
  Ylab = "Intensity"
))
print(draw_rank_plot(stat_df1,
  xc = "Feature", yc = "Height",
  Ylab = "Height"
))

---

draw_region_landmark  Plot genomic region landmark indicator

Description

Plot a gene centered polygon for demarcating gene and its upstream and downstream regions

Usage

draw_region_landmark(featureNames, vx, xmax)
**draw_region_name**

Plot genomic region names

---

**Arguments**
- **featureNames**: a string vector giving names of sub-regions
- **vx**: a vector on integers denoting the x coordinates of start of each sub-region
- **xmax**: an integer denoting the left most boundary

**Value**
a ggplot object

**Note**
used by `plot_5parts_metagene, plot_region`

**Author(s)**
Shuye Pu

**Examples**
```r
fn <- c("5'UTR", "CDS", "3'UTR")
mark <- c(1, 5, 20)
xmax <- 25

p <- draw_region_landmark(featureNames = fn, vx = mark, xmax = xmax)
```

---

**draw_region_name**  
Plot genomic region names

**Description**
Plot sub-region labels under the landmark

**Usage**

```r
draw_region_name(featureNames, scaled_bins, xmax)
```

**Arguments**
- **featureNames**: a string vector giving names of sub-regions
- **scaled_bins**: a vector of integers denoting the lengths of each sub-region
- **xmax**: an integer denoting the right most boundary

**Value**
a ggplot object
draw_region_profile

Note
used by plot_5parts_metagene, plot_region

Author(s)
Shuye Pu

Examples

```rn <- c("5'UTR", "CDS", "3'UTR")
bins <- c(5, 15, 5)
xmax <- 25

p <- draw_region_name(featureNames = fn, scaled_bins = bins, xmax = xmax)
```

---

draw_region_profile  Plot signal profile in genomic regions

Description

Plot lines with standard error as the error band

Usage

```r
draw_region_profile(
  plot_df,
  xc = "Position",
  yc = "Intensity",
  cn = "Query",
  sn = NULL,
  Ylab = "Signal Intensity",
  vx
)
```

Arguments

plot_df a dataframe with column names c(xc, yc, cn, "lower", "upper")
xc a string denoting column name for values on x-axis
yc a string denoting column name for numeric data to be plotted
cn column name in plot_df for query samples grouping
sn column name in plot_df for subject name to be shown in the plot title
Ylab a string for Y-axis label
vx a vector on integers denoting the x coordinates of start of each sub-region
**Value**

a ggplot object

**Note**

used by `plot_5parts_metagene, plot_region`

**Author(s)**

Shuye Pu

**Examples**

```r
library(dplyr)
Reference <- rep(rep(c("Ref1", "Ref2"), each = 100), 2)
Query <- rep(c("Query1", "Query2"), each = 200)
Position <- rep(seq_len(100), 4)
Intensity <- rlnorm(400)
se <- runif(400)
df <- data.frame(Intensity, se, Position, Query, Reference) %>%
  mutate(lower = Intensity - se, upper = Intensity + se) %>%
  mutate(Group = paste(Query, Reference, sep = ":"))
vx <- c(1, 23, 70)

p <- draw_region_profile(df, cn = "Group", vx = vx)
p
```

**Description**

Plot profile on top of heatmap, and align feature labels.

**Usage**

```r
draw_stacked_plot(plot_list, heatmap_list)
```

**Arguments**

- `plot_list` a list of profile plots
- `heatmap_list` a list of heatmaps

**Value**

a null value
draw_stacked_profile

**Note**

used by `plot_locus, plot_5parts_metagene, plot_region`

**Author(s)**

Shuye Pu

---

**draw_stacked_profile**  *Plot signal profile around start, center, and end of genomic regions*

**Description**

Plot lines with standard error as the error band, also plots number of regions having non-zero signals

**Usage**

```r
draw_stacked_profile(
  plot_df,  
  xc = "Position",  
  yc = "Intensity",  
  cn = "Query",  
  ext = c(0, 0, 0, 0),  
  hl = c(0, 0, 0, 0),  
  atitle = "title",  
  insert = 0,  
  Ylab = "Signal Intensity",  
  shade = FALSE,  
  stack = TRUE
)
```

**Arguments**

- `plot_df`: a dataframe with column names `(xc, yc, cn, "Interval", "lower", "upper")`
- `xc`: a string denoting column name for values on x-axis
- `yc`: a string denoting column name for numeric data to be plotted
- `cn`: a string denoting column name for grouping
- `ext`: a vector of 4 integers denoting upstream and downstream extension around start and end, the range of extensions must be within the range of `xc` of the `plot_df`
- `hl`: a vector of 4 integers defining upstream and downstream boundaries of the rectangle for start and end
- `atitle`: a string for the title of the plot
- `insert`: a integer denoting the width of the center region
- `Ylab`: a string for y-axis label
- `shade`: logical, indicating whether to place a shaded rectangle around the point of interest
- `stack`: logical, indicating whether to plot the number of valid (non-zero) data points in each bin
**draw_stacked_profile**

**Value**

a ggplot object

**Note**

used by `plot_start_end, plot_start_end_with_random`

**Author(s)**

Shuye Pu

**Examples**

```r
library(dplyr)
Reference <- rep(rep(c("Ref1", "Ref2"), each = 100), 2)
Query <- rep(c("Query1", "Query2"), each = 200)
Position <- rep(seq(-50, 49), 4)
Intensity <- rlnorm(400)
se <- runif(400)
start_df <- data.frame(Intensity, se, Position, Query, Reference) %>%
  mutate(lower = Intensity - se, upper = Intensity + se) %>%
  mutate(Group = paste(Query, Reference, sep = ":")) %>%
  mutate(Location = rep("Start", 400)) %>%
  mutate(Interval = sample.int(1000, 400))
Intensity <- rlnorm(400, meanlog = 1.5)
se <- runif(400)
center_df <- data.frame(Intensity, se, Position, Query, Reference) %>%
  mutate(lower = Intensity - se, upper = Intensity + se) %>%
  mutate(Group = paste(Query, Reference, sep = ":")) %>%
  mutate(Location = rep("Center", 400)) %>%
  mutate(Interval = sample.int(600, 400))
Intensity <- rlnorm(400, meanlog = 2)
se <- runif(400)
end_df <- data.frame(Intensity, se, Position, Query, Reference) %>%
  mutate(lower = Intensity - se, upper = Intensity + se) %>%
  mutate(Group = paste(Query, Reference, sep = ":")) %>%
  mutate(Location = rep("End", 400)) %>%
  mutate(Interval = sample.int(2000, 400))

df <- rbind(start_df, center_df, end_df)
p <- draw_stacked_profile(df, cn = "Group", shade = TRUE,
  ext = c(-50, 50, -50, 50),
  hl = c(-20, 20, -25, 25), insert = 100)
p
```
effective_size  Normalize sample library size to effective size

Description

This is a helper function for handle_input. edgeR::calcNormFactors function is used to estimate normalizing factors, which is used to multiply library sizes.

Usage

effective_size(outlist, outRle, genome = "hg19", nc = 2, verbose = FALSE)

Arguments

outlist  a list of list objects with four elements, 'query' is a GRanges object, 'size' is the library size, 'type' is the input file type, 'weight' is the name of the metadata column
outRle   logical, indicating whether the 'query' element of the output should be an RleList object or a GRanges object
genome   a string denoting the genome name and version
nc       integer, number of cores for parallel processing
verbose  logical, whether to output additional information

Value

a list of list objects with four elements ('query', 'size', 'type', 'weight'), with the 'size' element modified.

Author(s)

Shuye Pu

Examples

queryFiles <- system.file("extdata", "chip_treat_chr19.bam", package = "GenomicPlot"
)names(queryFiles) <- "query"

inputFiles <- system.file("extdata", "chip_input_chr19.bam", package = "GenomicPlot"
)names(inputFiles) <- "input"

chipImportParams <- setImportParams(
  offset = 0, fix_width = 150, fix_point = "start", norm = TRUE, useScore = FALSE, outRle = FALSE, useSizeFactor = FALSE, genome = "hg19"
out_list <- handle_input(
  inputFiles = c(queryFiles, inputFiles),
  importParams = chipImportParams, verbose = TRUE, nc = 2
)

out <- effective_size(out_list, outRle = TRUE)

---

**Description**

The data files in the extdata directory contain data for next generation sequencing read alignments, MACS2 peaks and gene annotation, which are used to test the package and generate plots in the package vignettes. To meet the package file size limit, all data are restricted to chr19:58000-507000 of the human genome version hg19. Details for each file are as follows.

**Details**

- "gencode.v19.annotation_chr19.gtf" is an excerpt of a gene annotation file by limiting to chr19:58000-507000 of the human genome.
- "gencode.v19.annotation_chr19.gtf.granges.rds" is a GRanges object produced by importing the above gtf file using RCAS::importGtf.
- "chip_treat_chr19.bam(.bai)" and "chip_input_chr19.bam(.bai)" are paired-end read alignment data from ChIPseq experiments.
- "treat_chr19.bam(.bai)" and "input_chr19.bam(.bai)" are single-end read alignment data from iCLIP experiments.
- "test_wig_chr19_+(-).wig", "test_wig_chr19_+(-).bw" are iCLIP alignment data in WIG and BIGWIG format, respectively; '+' and '-' represent forward and reverse strand, respectively.
- "test_clip_peak_chr19.bed" contains strand-specific iCLIP peak in BED format.
- "test_chip_peak_chr19.bed" and "test_chip_peak_chr19.narrowPeak" contain ChIPseq peaks generated with MACS2, in summit peak and narrow peak format, respectively. "test_chr19.bedGraph" contains the same data in bedGraph format.
- "test_file1.txt", "test_file2.txt", "test_file3.txt" and "test_file4.txt" are tab-delimited text files, each contains various human gene names in different columns.

**Value**

Various files used as inputs to run examples and tests

**Author(s)**

Shuye Pu
Source
The original gene annotation (gtf) file is downloaded from https://www.gencodegenes.org/human/. Except for the gtf file, all other files are derived from experimental data produced in-house at the Greenblatt Lab, University of Toronto, Canada.

extract_longest_tx Extract the longest transcript for each protein-coding genes

Description
Gene level computations require selecting one transcript per gene to avoid bias by genes with multiple isoforms. In ideal case, the most abundant transcript (principal or canonical isoform) should be chosen. However, the most abundant isoform may vary depending on tissue type or physiological condition, the longest transcript is usually the principal isoform, and alternatively spliced isoforms are not. This method get the longest transcript for each gene. The longest transcript is defined as the isoform that has the longest transcript length. In case of tie, the one with longer CDS is selected. If the lengths of CDS tie again, the transcript with smaller id is selected arbitrarily.

Usage
extract_longest_tx(txdb)

Arguments
taxdb a TxDb object defined in the GenomicFeatures package

Value
a dataframe of transcript information with the following columns: "tx_id tx_name gene_id nexon tx_len cds_len utr5_len utr3_len"

Author(s)
Shuye Pu

Examples

```
rown = system.file("extdata", "gencode.v19.annotation_chr19.gtf", package = "GenomicPlot")

taxdb <- custom_TxDb_from_GTF(rown, genome = "hg19")
longestTx <- extract_longest_tx(txdb)
```
filter_by_nonoverlaps_stranded

Filter GRanges by nonoverlaps in a stranded way

Description
This function reports all query GRanges that do not overlaps GRanges in subject. Strand information is used to define overlap.

Usage

filter_by_nonoverlaps_stranded(
  query,
  subject,
  maxgap = -1L,
  minoverlap = 0L,
  ignore.order = TRUE
)

Arguments

query a GRanges object
subject a GRanges object
maxgap an integer denoting the distance that define overlap
minoverlap The minimum amount of overlap between intervals as a single integer greater than 0. If you modify this argument, maxgap must be held fixed.
ignore.order logical, indicating whether the order of query and subject can be switched, default = TRUE. This parameter is used to avoid the situation that the size of overlaps is bigger than the size of subject, which will produce an error when plotting Venn diagrams.

Value

a GRanges object

Author(s)

Shuye Pu

Examples

query <- GRanges("chr19",
  IRanges(rep(c(10, 15), 2), width = c(1, 20, 40, 50)),
  strand = c("+", "+", "-", "-")
)

subject <- GRanges("chr19",

filter_by_overlaps_nonstranded

Filter GRanges by overlaps in a nonstranded way

Description

This function reports all query GRanges that have overlaps in subject GRanges. Strand information is not required.

Usage

filter_by_overlaps_nonstranded(
  query,
  subject,
  maxgap = -1L,
  minoverlap = 0L,
  ignore.order = TRUE
)

Arguments

query a GRanges object
subject a GRanges object
maxgap an integer denoting the distance that define overlap
minoverlap The minimum amount of overlap between intervals as a single integer greater than 0. If you modify this argument, maxgap must be held fixed.
ignore.order logical, indicating whether the order of query and subject can be switched, default = TRUE. This parameter is used to avoid the situation that the size of overlaps is bigger than the size of subject, which will produce an error when plotting Venn diagrams.

Value

a GRanges object

Author(s)

Shuye Pu
Examples

query <- GRanges("chr19",
  IRanges(rep(c(10, 15), 2), width = c(1, 20, 40, 50)),
  strand = c("+", "+-", "-", "+-")
)

subject <- GRanges("chr19",
  IRanges(rep(c(13, 150), 2), width = c(10, 14, 20, 28)),
  strand = c("+", "-", "-", "+")
)

res <- filter_by_overlaps_nonstranded(query, subject, ignore.order = TRUE)
res

filter_by_overlaps_stranded

Filter GRanges by overlaps in a stranded way

Description
This function reports all query GRanges that have overlaps in subject GRanges. Strand information
is used to define overlap.

Usage

filter_by_overlaps_stranded(
  query,
  subject,
  maxgap = -1L,
  minoverlap = 0L,
  ignore.order = TRUE
)

Arguments

query a GRanges object
subject a GRanges object
maxgap an integer denoting the distance that define overlap
minoverlap The minimum amount of overlap between intervals as a single integer greater
  than 0. If you modify this argument, maxgap must be held fixed.
ignore.order logical, indicating whether the order of query and subject can be switched, de-
  fault = TRUE. Overlaps in query and subject often have different sizes. This
  parameter will make the function use whichever is smaller to avoid errors when
  making Venn diagrams.
find_mate

Value
a GRanges object

Author(s)
Shuye Pu

Examples

query <- GRanges("chr19",
    IRanges(rep(c(10, 15), 2), width = c(1, 20, 40, 50)),
    strand = c("+", "+", "-", "-")
)

subject <- GRanges("chr19",
    IRanges(rep(c(13, 150), 2), width = c(10, 14, 20, 28)),
    strand = c("+", "+", "+", "+")
)

res <- filter_by_overlaps_stranded(query, subject)
res
resf <- filter_by_overlaps_stranded(query, subject, ignore.order = FALSE)
resf

find_mate  Find wig/bw file for the negative strand

Description
Find the file name of the negative strand, if a .wig/bw file for positive strand if provided, by looking
for file names with one character difference. If no negative strand file is found, assume the input
.wig/bw file is non-stranded

Usage
find_mate(inputFile, verbose = FALSE)

Arguments
inputFile path to a .wig/bw file, presumably for positive strand
verbose logical, whether to output additional information

Value
path to the negative .wig/bw file or NULL
gene2tx

Author(s)
Shuye Pu

Examples

queryFile <- system.file("extdata", "test_wig_chr19_+.wig",  
   package = "GenomicPlot"  
)  
names(queryFile) <- "test_wig"  

out <- GenomicPlot:::find_mate(inputFile = queryFile, verbose = TRUE)

gene2tx <- function(gtfFile, geneList, geneCol = 1) {
  # Translate gene names to transcript ids using a GTF file for a subset of genes
  # Given a list of gene names in a file or in a character vector, turn them into a vector of transcript ids.
  # Usage
  # gene2tx(gtfFile, geneList, geneCol = 1)
  # Arguments
  # gtfFile: path to a GTF file
  # geneList: path to a tab-delimited text file with one gene name on each line, or a character vector of gene names (eg. RPRD1B)
  # geneCol: the position of the column that containing gene names in the case that geneList is a file
  # Value
  # a vector of transcript ids (eg. ENST00000577222.1)
  # Author(s)
  # Shuye Pu
  # Examples
  # gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf",  
  #    package = "GenomicPlot"  
  #)
  # genes <- c("RPRD1A", "RPAP2", "RPRD1B", "RPRD2", "ZNF281", "YTHDF2")
  # tx <- gene2tx(gtfFile = gtfFile, geneList = genes)
Description

An R package for efficient and flexible visualization of genome-wide NGS coverage profiles

Details

The goal of ‘GenomicPlot‘ is to provide an efficient visualization tool for next generation sequencing (NGS) data with rich functionality and flexibility. ‘GenomicPlot‘ enables plotting of NGS data in various formats (bam, bed, wig and bigwig); both coverage and enrichment over input can be computed and displayed with respect to genomic features (such as UTR, CDS, enhancer), and user defined genomic loci or regions. Statistical tests on signal intensity within user defined regions of interest can be performed and presented as box plots or pie charts. Parallel processing is enabled to speed up computation on multi-core platforms. Main functions are as follows:

- **plot_5parts_metagene** generates genomic (with introns) or metagenomic (without introns) plots around gene body and its upstream and downstream regions, the gene body can be further segmented into 5’UTR, CDS and 3’UTR.
- **plot_start_end** plots genomic profiles around the start and end of genomic features (like exons or introns), or user defined genomic regions. A center region with user defined width can be plotted simultaneously.
- **plot_locus** plots distance between sample peaks and genomic features, or distance from one set of peaks to another set of peaks.
- **plot_region** plots signal profiles within and around genomic features, or user defined genomic regions.
- **plot_peak_annotation** plots peak annotation statistics (distribution in different type of genes, and in different parts of genes).
- **plot_overlap_bed** plots peak overlaps as Venn diagrams.
- Random features can be generated and plotted to serve as contrast to real features in plot_locus_with_random and plot_start_end_with_random.
- All profile line plots have error bands.
- Statistical analysis results on user defined regions of interest are plotted along with the profile plots in plot_region, plot_locus and plot_locus_with_random.

Author(s)

Shuye Pu

PACKAGE
get_genomic_feature_coordinates

Extract genomic features from TxDb object

Description

Extract genomic coordinates and make bed or bed 12 files from a TxDb object for a variety of annotated genomic features. The output of this function is a list. The first element of the list is a GRanges object that provide the start and end information of the feature. The second element is a GRangesList providing information for sub-components. The third element is the name of a bed file.

Usage

get_genomic_feature_coordinates(
  txdb,  
  featureName,  
  featureSource = NULL,  
  export = FALSE,  
  longest = FALSE,  
  protein_coding = FALSE
)

Arguments

txdb a TxDb object defined in the GenomicFeatures package

featureName one of the genomic feature in c("utr3", "utr5", "cds", "intron", "exon", "transcript", "gene")

featureSource the name of the gtf/gff3 file or the online database from which txdb is derived, used as name of output file

export logical, indicating if the bed file should be produced

longest logical, indicating whether the output should be limited to the longest transcript of each gene

protein_coding logical, indicating whether to limit to protein_coding genes

Details

For "utr3", "utr5", "cds" and "transcript", the GRanges object denotes the start and end of the feature in one transcript, and the range is named by the transcript id and may span introns; the GrangesList object is a list of exons comprising each feature and indexed on transcript id. The bed file is in bed12 format. For "exon" and "intron", the GRanges object denotes unnamed ranges of individual exon and intron, and the GrangesList object is a list of exons or introns belonging to one transcript and indexed on transcript id. The bed file is in bed6 format. For "gene", both GRanges object and GRangesList object have the same ranges and names. The bed file is in bed6 format.
get_targeted_genes

Value

a list of three objects, the first is a GRanges object, the second is a GRangesList object, the last is
the output file name if export is TRUE.

Author(s)

Shuye Pu

Examples

gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf",
    package = "GenomicPlot"
)

txdb <- custom_TxDb_from_GTF(gtfFile, genome = "hg19")

output <- get_genomic_feature_coordinates(txdb,
    featureName = "cds", featureSource = "gencode",
    export = FALSE, longest = TRUE, protein_coding = TRUE
)

get_targeted_genes

Get the number of peaks overlapping each feature of all protein-coding
genes

Description

Annotate each peak with genomic features based on overlap, and produce summary statistics for
distribution of peaks in features of protein-coding genes. If a peak overlap multiple features, a
feature is assigned to the peak in the following order of precedence: "5'UTR", "3'UTR", "CDS",
"Intron", "Promoter", "TTS".

Usage

get_targeted_genes(peak, features, stranded = TRUE)

Arguments

peak a GRanges object defining query ranges
features a GRangesList object representing genomic features
stranded logical, indicating whether the overlap should be strand-specific

Value

a list object
get_txdb_features

Note

used in plot_peak_annotation

Author(s)

Shuye Pu

Examples

gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", package = "GenomicPlot")

_txdb <- custom_TxDb_from_GTF(gtfFile, genome = "hg19")
f <- get_txdb_features(_txdb, dsTSS = 100, fiveP = 0, threeP = 1000)

p <- RCAS::importBed(system.file("extdata", "test_chip_peak_chr19.bed", package = "GenomicPlot")
ann <- get_targeted_genes(peak = p, features = f, stranded = FALSE)

get_txdb_features

Get genomic coordinates of features of protein-coding genes

Description

Get genomic coordinates of promoter, 5'UTR, CDS, 3'UTR, TTS and intron for the longest transcript of protein-coding genes. The range of promoter is defined by fiveP and dsTSS upstream and downstream TSS, respectively, the TTS ranges from the 3' end of the gene to threeP downstream, or the start of a downstream gene, whichever is closer.

Usage

get_txdb_features(txdb, fiveP = -1000, dsTSS = 300, threeP = 1000, nc = 2)

Arguments

txdb a TxDb object defined in the GenomicFeatures package
fiveP extension upstream of the 5' boundary of genes
dsTSS range of promoter extending downstream of TSS
threeP extension downstream of the 3' boundary of genes
nc number of cores for parallel processing

Value

a GRangesList object
Author(s)
Shuye Pu

Examples
```r
gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", 
    package = "GenomicPlot"
)

txdb <- custom_TxDb_from_GTF(gtfFile, genome = "hg19")

f <- get_txdb_features(txdb, dsTSS = 100, fiveP = -100, threeP = 100)
```

Description
Genomic coordinates of 72 transcripts in hg19 for genomic features promoter, 5'UTR, CDS, 3'UTR, TTS, as well as user inputs for processing these features. See `prepare_5parts_genomic_features` for details.

Value
A named list with the following elements:

- **windowRs** a list of 5 GrangesList objects for the 5 genomic features
- **nbins** a positive integer
- **scaled_bins** a vector of 5 integers
- **fiveP** a negative integer
- **threeP** a positive integer
- **meta** logical
- **longest** logical

Author(s)
Shuye Pu

Source
The data is produced by running the following code:
```r
txdb <- AnnotationDbi::loadDb(system.file("extdata", "txdb.sql", package = "GenomicPlot"))
gf5_genomic <- GenomicPlot::prepare_5parts_genomic_features(txdb, meta = FALSE, nbins = 100, 
    fiveP = -2000, threeP = 1000, longest = TRUE)
```
Toy data for examples and testing of the ‘GenomicPlot’ package

Description

Metagenomic coordinates of 72 transcripts in hg19 for genomic features promoter, 5′ UTR, CDS, 3′ UTR, TTS, as well as user inputs for processing these features. See `prepare_5parts_genomic_features` for details.

Value

A named list with the following elements:

- `windowRs` a list of 5 `GrangesList` objects for the 5 genomic features
- `nbins` a positive integer
- `scaled_bins` a vector of 5 integers
- `fiveP` a negative integer
- `threeP` a positive integer
- `meta` logical
- `longest` logical

Author(s)

Shuye Pu

Source

The data is produced by running the following code:

```r
txdb <- AnnotationDbi::loadDb(system.file("extdata", "txdb.sql", package = "GenomicPlot"))
gf5_meta <- GenomicPlot::prepare_5parts_genomic_features(txdb, meta = TRUE, nbins = 100, fiveP = -2000, threeP = 1000, longest = TRUE)
```

Convert GRanges to dataframe

Description

Convert a GRanges object with meta data columns to a dataframe, with the first 6 columns corresponding those of BED6 format, and the meta data as additional columns

Usage

```r
gr2df(gr)
```
handle_bam

Arguments

gr       a GRanges object

Value

a dataframe

Author(s)

Shuye Pu

Examples

gr2 <- GenomicRanges::GRanges(c("chr1", "chr1"),
    IRanges::IRanges(c(7, 13), width = 3),
    strand = c("+", "-"))
GenomicRanges::mcols(gr2) <- data.frame(
    score = c(0.3, 0.9),
    cat = c(TRUE, FALSE))
df2 <- gr2df(gr2)

Description

This is a function for read NGS reads data in bam format, store the input data in a list of GRanges objects or RleList objects. For paired-end reads, only take the second read in a pair, assuming which is the sense read for strand-specific RNAseq.

Usage

handle_bam(inputFile, importParams = NULL, verbose = FALSE)

Arguments

inputFile       a string denoting path to the input file
importParams    a list of parameters, refer to handle_input for details
verbose         logical, whether to output additional information

Details

The reads are filtered using mapq score >= 10 by default, only mapped reads are counted towards library size.
**Value**

A list object with four elements, 'query' is a list GRanges objects or RleList objects, 'size' is the library size, 'type' is the input file type, 'weight' is the name of the metadata column to be used as weight for coverage calculation.

**Author(s)**

Shuye Pu

**Examples**

```r
queryFiles <- system.file("extdata", "treat_chr19.bam", package = "GenomicPlot"
)  
names(queryFiles) <- "query"

bamimportParams <- setImportParams(
  offset = -1, fix_width = 0, fix_point = "start", norm = TRUE,
  useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)  
out <- handle_bam(
  inputFile = queryFiles, importParams = bamimportParams, verbose = TRUE
)
```

**Description**

This is a function for read peaks data in bed format, store the input data in a list of GRanges objects or RleList objects.

**Usage**

`handle_bed(inputFile, importParams = NULL, verbose = FALSE)`

**Arguments**

- `inputFile`: a string denoting path to the input file
- `importParams`: a list of parameters, refer to `handle_input` for details
- `verbose`: logical, whether to output additional information

**Value**

A list object with four elements, 'query' is a list GRanges objects or RleList objects, 'size' is the library size, 'type' is the input file type, 'weight' is the name of the metadata column to be used as weight for coverage calculation.
handle_bedGraph

Handle files in bedGraph format

Description

This is a function for read peaks data in bedGraph format, store the input data in a list of GRanges objects or RleList objects.

Usage

handle_bedGraph(inputFile, importParams = NULL, verbose = FALSE)

Arguments

inputFile a string denoting path to the input file
importParams a list of parameters, refer to handle_input for details
verbose logical, whether to output additional information

Value

a list object with four elements, 'query' is a list GRanges objects or RleList objects, 'size' is the library size, 'type' is the input file type, 'weight' is the name of the metadata column to be used as weight for coverage calculation

Author(s)

Shuye Pu
Examples

queryFiles <- system.file("extdata", "test_chr19.bedGraph", 
    package = "GenomicPlot"
)
names(queryFiles) <- "chipPeak"

importParams <- setImportParams(
    offset = 0, fix_width = 0, fix_point = "start", norm = FALSE,
    useScore = TRUE, outRle = FALSE, useSizeFactor = FALSE, genome = "hg19",
    val = 4, skip = 1
)

out <- handle_bedGraph(queryFiles, importParams, verbose = TRUE)
out$query

handle_bw

Handle files in bw|bigwig|bigWig|BigWig|BW|BIGWIG format

Description

This is a function for read NGS coverage data in bigwig format, store the input data in a list of 
GRanges objects or RleList objects. The input bw file can be stranded or non-stranded. Library size 
is calculate as the sum of all coverage.

Usage

handle_bw(inputFile, importParams, verbose = FALSE)

Arguments

inputFile a string denoting path to the input file
importParams a list of parameters, refer to handle_input for details
verbose logical, whether to output additional information

Details

For stranded files, forward and reverse strands are stored in separate files, with '+' or 'p' in the forward strand file name and '-' or 'm' in the reverse strand file name.

Value

a list object with four elements, 'query' is a list GRanges objects or RleList objects, 'size' is the estimated library size, 'type' is the input file type, weight' is the name of the metadata column to be used as weight for coverage calculation

Author(s)

Shuye Pu
Examples

```r
queryFiles <- system.file("extdata", "test_wig_chr19_.bw", package = "GenomicPlot"
)
names(queryFiles) <- "test_bw"

wigimportParams <- setImportParams(
    offset = 0, fix_width = 0, fix_point = "start", norm = FALSE,
    useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

out <- handle_bw(queryFiles, wigimportParams, verbose = TRUE)
```

Description

This is a wrapper function for read NGS data in different file formats, store the input data in a list of GRanges objects or RleList objects. File names end in bed|bam|bw|bigwig|bigWig|BigWig|BW|BIGWIG are recognized, and a named list of files with mixed formats are allowed.

Usage

```r
handle_input(inputFiles, importParams = NULL, verbose = FALSE, nc = 2)
```

Arguments

- **inputFiles**: a vector of strings denoting file names
- **importParams**: a list with the 9 elements: list(offset, fix_width, fix_point, useScore, outRle, norm, genome, useSizeFactor). Details are described in the documentation of `setImportParams` function
- **verbose**: logical, whether to output additional information
- **nc**: integer, number of cores for parallel processing

Details

When 'useScore' is TRUE, the score column of the bed file will be used in the metadata column 'score' of the GRanges object, or the 'Values' field of the RleList object. Otherwise the value 1 will be used instead. When the intended use of the input bed is a reference feature, both 'useScore' and 'outRle' should be set to FALSE.

Value

A list object with four elements, 'query' is a list GRanges objects or RleList objects, 'size' is the library size, 'type' is the input file type, 'weight' is the name of the metadata column to be used as weight for coverage calculation.
Examples

```r
queryFiles1 <- system.file("extdata", "treat_chr19.bam",
    package = "GenomicPlot"
)
names(queryFiles1) <- "query"

inputFiles1 <- system.file("extdata", "input_chr19.bam",
    package = "GenomicPlot"
)
names(inputFiles1) <- "input"

bamimportParams <- setImportParams(
    offset = -1, fix_width = 0, fix_point = "start", norm = TRUE,
    useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

out_list <- handle_input(
    inputFiles = c(queryFiles1, inputFiles1),
    importParams = bamimportParams, verbose = TRUE, nc = 2
)

queryFiles2 <- system.file("extdata", "test_wig_chr19_+.wig",
    package = "GenomicPlot"
)
names(queryFiles2) <- "test_wig"

wigimportParams <- setImportParams(
    offset = 0, fix_width = 0, fix_point = "start", norm = FALSE,
    useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

out <- handle_input(queryFiles2, wigimportParams, verbose = TRUE)

queryFiles3 <- system.file("extdata", "test_wig_chr19_+.bw",
    package = "GenomicPlot"
)
names(queryFiles3) <- "test_bw"

out <- handle_input(c(queryFiles1, queryFiles2, queryFiles3),
    wigimportParams,
    verbose = TRUE
)
```

---

**handle_wig**

Handle files in wig format
Description

This is a function for read NGS coverage data in wig format, store the input data in a list of GRanges objects or RleList objects. The input wig file can be stranded or non-stranded. Library size is calculate as the sum of all coverage.

Usage

handle_wig(inputFile, importParams, verbose = FALSE)

Arguments

inputFile a string denoting path to the input file
importParams a list of parameters, refer to handle_input for details
verbose logical, whether to output additional information

Details

For stranded files, forward and reverse strands are stored in separate files, with '+' or 'p' in the forward strand file name and '-' or 'm' in the reverse strand file name.

Value

a list object with four elements, 'query' is a list GRanges objects or RleList objects, 'size' is the library size, 'type' is the input file type, 'weight' is the name of the metadata column to be used as weight for coverage calculation

Author(s)

Shuye Pu

Examples

queryFiles <- system.file("extdata", "test_wig_chr19_.wig",
                          package = "GenomicPlot"
) names(queryFiles) <- "test_wig"

wigimportParams <- setImportParams(
  offset = 0, fix_width = 0, fix_point = "start", norm = FALSE,
  useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

out <- handle_wig(queryFiles, wigimportParams, verbose = TRUE)
**Description**

Replace 0 and missing values in a sparse non-negative matrix with half of minimum of non-zero values, to avoid use of arbitrary pseudo numbers, and to allow computing ratios and log transformation of matrices. When a matrix is sparse (assuming it has many all-zero rows and few all-zero columns), the half of minimum of non-zero values is a number that is small enough so that is will not distort the data too much (comparing to a pseudo count = 1), but large enough to avoid huge ratios when used as a denominator.

**Usage**

```r
impute_hm(fullmatrix, verbose = FALSE)
```

**Arguments**

- `fullmatrix`: a numeric matrix
- `verbose`: logical, whether to output additional information

**Value**

a numeric matrix

**Author(s)**

Shuye Pu

**Examples**

```r
fullMatrix <- matrix(rlnorm(100), ncol = 10)
for (i in 5:6) {
  fullMatrix[i - 1, 4:7] <- 0
}
imp <- GenomicPlot:::impute_hm(fullMatrix, verbose = TRUE)
```
inspect_matrix

Inspect a numeric matrix

Description
Check the matrix for NA, NaN, INF, -INF and 0 values

Usage
inspect_matrix(fullmatrix, verbose = FALSE)

Arguments
fullmatrix a numeric matrix
verbose logical, indicating whether to print out the stats in the console

Value
a numerical matrix summarizing the unusual values

Author(s)
Shuye Pu

Examples
fullMatrix <- matrix(rnorm(100), ncol = 10)
for (i in 5:6) {
  fullMatrix[i, 4:7] <- NaN
  fullMatrix[i + 1, 4:7] <- NA
  fullMatrix[i + 2, 4:7] <- -Inf
  fullMatrix[i - 1, 4:7] <- 0
  fullMatrix[i - 2, 1:3] <- Inf
}
GenomicPlot:::inspect_matrix(fullMatrix, verbose = TRUE)
**Description**

Make a partial TxDb object given a GTF file and a list of gene names in a file or in a character vector.

**Usage**

```r
make_subTxDb_from_GTF(gtfFile, genome = "hg19", geneList, geneCol = 1)
```

**Arguments**

- `gtfFile`: path to a GTF file
- `genome`: version of genome, like "hg19"
- `geneList`: path to a tab-delimited text file with one gene name on each line, or a character vector of gene names
- `geneCol`: the position of the column that containing gene names in the case that `geneList` is a file

**Value**

a TxDb object

**Author(s)**

Shuye Pu

**Examples**

```r
gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", package = "GenomicPlot")
genesis <- c("RPRD1A", "RPAP2", "RPRD1B", "RPRD2", "ZNF281", "YTHDF2")
txdb <- make_subTxDb_from_GTF(gtfFile = gtfFile, geneList = genesis)
```
Description

This is a helper function for Venn diagram plot. A Venn diagram is plotted as output. For GRanges, as A overlap B may not be the same as B overlap A, the order of GRanges in a list matters, certain order may produce an error.

Usage

```r
overlap_pair(apair, overlap_fun, title = NULL)
```

Arguments

- **apair**: a list of two vectors
- **overlap_fun**: the name of the function that defines overlap, depending on the type of object in the vectors. For GRanges, use `filter_by_overlaps_stranded` or `filter_by_nonoverlaps_stranded`, for gene names, use `intersect`.
- **title**: main title of the figure

Value

a `VennDiagram` object

Author(s)
Shuye Pu

Examples

```r
test_list <- list(A = c(1, 2, 3, 4, 5), B = c(4, 5, 7))
overlap_pair(test_list, intersect, title = "test")

## GRanges overlap
query <- GRanges("chr19",
  IRanges(rep(c(10, 15), 2), width = c(1, 20, 40, 50)),
  strand = c("+", "+", "+", "-"))

subject <- GRanges("chr19",
  IRanges(rep(c(13, 150), 2), width = c(10, 14, 20, 28)),
  strand = c("+", "+", "+", "+"))

overlap_pair(
  list(query = query, subject = subject),
  filter_by_overlaps_stranded
```
Description
This is a helper function for Venn diagram plot. A Venn diagram is plotted as output. For GRanges, as A overlap B may not be the same as B overlap A, the order of GRanges in a list matters, certain order may produce an error.

Usage
overlap_quad(aquad, overlap_fun, title = NULL)

Arguments
- aquad: a list of four vectors
- overlap_fun: the name of the function that defines overlap, depending on the type of object in the vectors. For GRanges, use `filter_by_overlaps_stranded` or `filter_by_nonoverlaps_stranded`, for gene names, use `intersect`.
- title: main title of the figure

Value
a VennDiagram object

Author(s)
Shuye Pu

Examples
```
test_list <- list(A = c(1, 2, 3, 4, 5), B = c(4, 5, 7), C = c(1, 3), D = 6)
overlap_quad(test_list, intersect)

## GRanges overlap
query1 <- GRanges("chr19",
    IRanges(rep(c(10, 15), 2), width = c(1, 20, 40, 50)),
    strand = c("+", "+", "+", "+")
)

query2 <- GRanges("chr19",
    IRanges(rep(c(1, 15), 2), width = c(1, 20, 40, 50)),
    strand = c("+", "+", "+", "+")
)
```
subject1 <- GRanges("chr19",
    IRanges(rep(c(13, 150), 2), width = c(10, 14, 20, 28)),
    strand = c("+", ",", ",", "+")
)

subject2 <- GRanges("chr19",
    IRanges(rep(c(13, 50), 2), width = c(10, 14, 20, 21)),
    strand = c("+", ",", ",", "+")
)

overlap_quad(list(
    subject1 = subject1, subject2 = subject2, query1 = query1,
    query2 = query2
), filter_by_overlaps_stranded)

---

**overlap_triple**  
*Plot three-sets Venn diagram*

**Description**

This is a helper function for Venn diagram plot. A Venn diagram is plotted as output. For GRanges, as A overlap B may not be the same as B overlap A, the order of GRanges in a list matters, certain order may produce an error.

**Usage**

```r
overlap_triple(atriple, overlap_fun, title = NULL)
```

**Arguments**

- **atriple**: a list of three vectors
- **overlap_fun**: the name of the function that defines overlap, depending on the type of object in the vectors. For GRanges, use `filter_by_overlaps_stranded` or `filter_by_nonoverlaps_stranded`.
- **title**: main title of the figure

**Value**

- A VennDiagram object

**Author(s)**

- Shuye Pu
Examples

test_list <- list(A = c(1, 2, 3, 4, 5), B = c(4, 5, 7), C = c(1, 3))
overlap_triple(test_list, intersect, title = "test")

## GRanges overlap
query <- GRanges("chr19",
  IRanges(rep(c(10, 15), 2), width = c(1, 20, 40, 50)),
  strand = c("+", "+", "-", "-"))

subject1 <- GRanges("chr19",
  IRanges(rep(c(13, 150), 2), width = c(10, 14, 20, 28)),
  strand = c("+", "-", "-", "+"))

subject2 <- GRanges("chr19",
  IRanges(rep(c(13, 50), 2), width = c(10, 14, 20, 21)),
  strand = c("+", "-", "+", "-"))

overlap_triple(
  list(subject1 = subject1, subject2 = subject2, query = query),
  filter_by_overlaps_stranded
)

\[\text{parallel\_countOverlaps}\]

---

**Parallel execution of countOverlaps**

---

**Description**

Function for parallel computation of countOverlaps function in the GenomicRanges package

**Usage**

```r
parallel_countOverlaps(grange_list, tileBins, nc = 2, switch = FALSE)
```

**Arguments**

- `grange_list` a list of GRanges objects.
- `tileBins` a GRanges object of tiled genome
- `nc` integer, number of cores for parallel processing
- `switch` logical, switch the order of query and feature

**Value**

a list of numeric vectors
Author(s)
Shuye Pu

Examples

```r
bedQueryFiles <- c(
  system.file("extdata", "test_chip_peak_chr19.narrowPeak", package = "GenomicPlot"),
  system.file("extdata", "test_chip_peak_chr19.bed", package = "GenomicPlot"),
  system.file("extdata", "test_clip_peak_chr19.bed", package = "GenomicPlot")
)
names(bedQueryFiles) <- c("NarrowPeak", "SummitPeak", "iCLIPPeak")

bedimportParams <- setImportParams(
  offset = 0, fix_width = 100, fix_point = "center", norm = FALSE,
  useScore = FALSE, outRle = FALSE, useSizeFactor = FALSE, genome = "hg19"
)

out_list <- handle_input(
  inputFiles = bedQueryFiles,
  importParams = bedimportParams, verbose = TRUE, nc = 2
)

chromInfo <- circlize::read.chromInfo(species = "hg19")$df
seqi <- Seqinfo(seqnames = chromInfo$chr, seqlengths = chromInfo$end, 
isCircular = rep(FALSE, nrow(chromInfo)), genome = "hg19")
grange_list <- lapply(out_list, function(x) x$query)
tilewidth <- 100000
tileBins <- tileGenome(seqi, 
tilewidth = tilewidth, 
cut.last.tile.in.chrom = TRUE)

score_list1 <- parallel_countOverlaps(grange_list, tileBins, nc = 2)
dplyr::glimpse(score_list1)
```

**parallel_scoreMatrixBin**

*Parallel execution of scoreMatrixBin on a huge target windows object split into chunks*
Description

Function for parallel computation of scoreMatrixBin. The ‘windows’ parameter of the scoreMatrixBin method is split into nc chunks, and scoreMatrixBin is called on each chunk simultaneously to speed up the computation.

Usage

```
parallel_scoreMatrixBin(
  queryRegions,
  windowRs,
  bin_num,
  bin_op,
  weight_col,
  stranded,
  nc = 2
)
```

Arguments

- **queryRegions**: a RleList object or Granges object providing input for the 'target' parameter of the scoreMatrixBin method.
- **windowRs**: a single GRangesList object.
- **bin_num**: number of bins the windows should be divided into.
- **bin_op**: operation on the signals in a bin, a string in c("mean", "max", "min", "median", "sum") is accepted.
- **weight_col**: if the queryRegions is a GRanges object, a numeric column in meta data part can be used as weights.
- **stranded**: logical, indicating if the strand of the windows should be considered to determine upstream and downstream.
- **nc**: an integer denoting the number of cores requested, 2 is the default number that is allowed by CRAN but 5 gives best trade-off between speed and space.

Value

a numeric matrix

Author(s)

Shuye Pu

Examples

```
queryFiles <- system.file("extdata", "chip_treat_chr19.bam",
                         package = "GenomicPlot")

names(queryFiles) <- "query"

chipimportParams <- setImportParams(
```
plot_5parts_metagene

offset = 0, fix_width = 150, fix_point = "start", norm = TRUE,
useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

queryRegion <- handle_input(queryFiles, chipimportParams,
verbose = TRUE
)[1]$query

windowFiles <- system.file("extdata", "test_chip_peak_chr19.narrowPeak",
package = "GenomicPlot"
)
names(windowFiles) <- "narrowPeak"

importParams <- setImportParams(
    offset = 0, fix_width = 0, fix_point = "start", norm = FALSE,
    useScore = FALSE, outRle = FALSE, useSizeFactor = FALSE, genome = "hg19"
)

windowRegion <- handle_bed(windowFiles, importParams, verbose = TRUE)$query

out <- parallel_scoreMatrixBin(
    queryRegions = queryRegion,
    windowRs = windowRegion,
    bin_num = 50,
    bin_op = "mean",
    weight_col = "score",
    stranded = TRUE,
    nc = 2
)

#

plot_5parts_metagene  Plot promoter, 5' UTR, CDS, 3' UTR and TTS

Description
Plot reads or peak Coverage/base/gene of samples given in the query files around genes. The up-
stream and downstream windows flanking genes can be given separately, metagene plots are gener-
ated with 5'UTR, CDS and 3'UTR segments. The length of each segments are prorated according
to the median length of each segments. If Input files are provided, ratio over Input is computed and
displayed as well.

Usage
plot_5parts_metagene(
    queryFiles,
gFeatures_list,
    inputFiles = NULL,
    importParams = NULL,
plot_5parts_metagene

verbose = FALSE,
transform = NA,
smooth = FALSE,
scale = FALSE,
stranded = TRUE,
outPrefix = NULL,
heatmap = FALSE,
heatRange = NULL,
rmOutlier = 0,
Ylab = "Coverage/base/gene",
hw = c(10, 10),
nc = 2
)

Arguments

queryFiles a vector of sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed

gFeatures_list a list of genomic features as output of the function prepare_5parts_genomic_features

inputFiles a vector of input sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed

importParams a list of parameters for handle_input

verbose logical, indicating whether to output additional information (data used for plotting or statistical test results)

transform logical, whether to log2 transform the matrix

smooth logical, indicating whether the line should smoothed with a spline smoothing algorithm

scale logical, indicating whether the score matrix should be scaled to the range 0:1, so that samples with different baseline can be compared

stranded logical, indicating whether the strand of the feature should be considered

outPrefix a string specifying output file prefix for plots (outPrefix.pdf)

heatmap logical, indicating whether a heatmap of the score matrix should be generated

heatRange a numeric vector with three elements, defining custom range for color ramp, default=NULL, i.e. the range is defined automatically based on the c(minimum, median, maximum) of a data matrix

rmOutlier a numeric value serving as a multiplier of the MAD in Hampel filter for outliers identification, 0 indicating not removing outliers. For Gaussian distribution, use 3, adjust based on data distribution.

Ylab a string for y-axis label

hw a vector of two elements specifying the height and width of the output figures

nc integer, number of cores for parallel processing

Value

a dataframe containing the data used for plotting
Author(s)
Shuye Pu

Examples

```r
data(gf5_meta)
queryfiles <- system.file("extdata", "treat_chr19.bam", package = "GenomicPlot")
names(queryfiles) <- "clip_bam"
inputfiles <- system.file("extdata", "input_chr19.bam", package = "GenomicPlot")
names(inputfiles) <- "clip_input"

bamimportParams <- setImportParams(
  offset = -1, fix_width = 0, fix_point = "start", norm = TRUE,
  useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

plot_5parts_metagene(
  queryFiles = queryfiles,
  gFeatures_list = list("metagene" = gf5_meta),
  inputFiles = inputfiles,
  scale = FALSE,
  verbose = FALSE,
  transform = NA,
  smooth = TRUE,
  stranded = TRUE,
  outPrefix = NULL,
  importParams = bamimportParams,
  heatmap = TRUE,
  rmOutlier = 0,
  nc = 2
)
```

Description

Plot correlation in reads coverage distributions along the genome for bam files. Generates a fingerprint plot, a heatmap of correlation coefficients with hierarchical clustering, a pairwise correlation plot and a PCA plot.

Usage

```r
plot_bam_correlation(
  bamFiles,
  binSize = 1e+06,
)```
plot_bam_correlation

outPrefix = NULL,
importParams = NULL,
grouping = NULL,
verbose = FALSE,
hw = c(8, 8),
nc = 2
)

Arguments

bamFiles       a named vector of strings denoting file names
binSize        an integer denoting the tile width for tiling the genome, default 1000000
outPrefix      a string denoting output file name in pdf format
importParams   a list of parameters for handle_input
grouping       a named vector for bamFiles group assignment
verbose        logical, indicating whether to output additional information
hw             a vector of two elements specifying the height and width of the output figures
nc             integer, number of cores for parallel processing

Value

a dataframe of read counts per bin per sample

Examples

bamQueryFiles <- c(
  system.file("extdata", "chip_input_chr19.bam", package = "GenomicPlot"),
  system.file("extdata", "chip_treat_chr19.bam", package = "GenomicPlot")
)
grouping <- c(1, 2)
names(bamQueryFiles) <- names(grouping) <- c("chip_input", "chip_treat")

bamImportParams <- setImportParams(
  offset = 0, fix_width = 150, fix_point = "start", norm = FALSE,
  useScore = FALSE, outRle = FALSE, useSizeFactor = FALSE, genome = "hg19"
)

plot_bam_correlation(
  bamFiles = bamQueryFiles, binSize = 100000, outPrefix = NULL,
  importParams = bamImportParams, nc = 2, verbose = FALSE
)


**plot_locus**  
*Plot signal around custom genomic loci*

**Description**
Plot reads or peak Coverage/base/gene of samples given in the query files around reference locus (start, end or center of a genomic region) defined in the centerFiles. The upstream and downstream windows flanking loci can be given separately, a smaller window can be defined to allow statistical comparisons between samples for the same reference, or between references for a given sample. If Input files are provided, ratio over Input is computed and displayed as well.

**Usage**

```r
plot_locus(
  queryFiles,
  centerFiles,
  txdb = NULL,
  ext = c(-100, 100),
  hl = c(0, 0),
  shade = TRUE,
  smooth = FALSE,
  importParams = NULL,
  verbose = FALSE,
  binSize = 10,
  refPoint = "center",
  Xlab = "Center",
  Ylab = "Coverage/base/gene",
  inputFiles = NULL,
  stranded = TRUE,
  heatmap = TRUE,
  scale = FALSE,
  outPrefix = NULL,
  rmOutlier = 0,
  transform = NA,
  statsMethod = "wilcox.test",
  heatRange = NULL,
  hw = c(8, 8),
  nc = 2
)
```

**Arguments**

- `queryFiles` a vector of sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed
- `centerFiles` a named vector of reference file names or genomic features in c("utr3", "utr5", "cds", "intron", "exon", "transcript", "gene"). The file should be in .bed format only
txdb  a TxDb object defined in the GenomicFeatures package. Default NULL, needed only when genomic features are used as centerFiles.

ext  a vector of two integers defining upstream and downstream boundaries of the plot window, flanking the reference locus

hl  a vector of two integers defining upstream and downstream boundaries of the highlight window, flanking the reference locus

shade  logical indicating whether to place a shaded rectangle around the point of interest

smooth  logical, indicating whether the line should smoothed with a spline smoothing algorithm

importParams  a list of parameters for handle_input

verbose  logical, indicating whether to output additional information (data used for plotting or statistical test results)

binSize  an integer defines bin size for intensity calculation

refPoint  a string in c("start", "center", "end")

Xlab  a string denotes the label on x-axis

Ylab  a string for y-axis label

inputFiles  a vector of input sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed

stranded  logical, indicating whether the strand of the feature should be considered

heatmap  logical, indicating whether a heatmap of the score matrix should be generated

scale  logical, indicating whether the score matrix should be scaled to the range 0:1, so that samples with different baseline can be compared

outPrefix  a string specifying output file prefix for plots (outPrefix.pdf)

rmOutlier  a numeric value serving as a multiplier of the MAD in Hampel filter for outliers identification, 0 indicating not removing outliers. For Gaussian distribution, use 3, adjust based on data distribution.

transform  a string in c("log", "log2", "log10"), default = NA indicating no transformation of data matrix

statsMethod  a string in c("wilcox.test", "t.test"), for pair-wise group comparisons

heatRange  a numeric vector with three elements, defining custom range for color ramp, default=NULL, i.e. the range is defined automatically based on the c(minimum, median, maximum) of a data matrix

hw  a vector of two elements specifying the height and width of the output figures

nc  integer, number of cores for parallel processing

Value

a list of two dataframes containing the data used for plotting and for statistical testing

Author(s)

Shuye Pu
Examples

centerfiles <- c(
  system.file("extdata", "test_clip_peak_chr19.bed", package = "GenomicPlot"),
  system.file("extdata", "test_chip_peak_chr19.bed", package = "GenomicPlot"))

names(centerfiles) <- c("iCLIPPeak", "SummitPeak")
queryfiles <- c(
  system.file("extdata", "chip_treat_chr19.bam", package = "GenomicPlot"))

names(queryfiles) <- c("chip_bam")
inputfiles <- c(
  system.file("extdata", "chip_input_chr19.bam", package = "GenomicPlot"))
names(inputfiles) <- c("chip_input")

chipimportParams <- setImportParams(
  offset = 0, fix_width = 150, fix_point = "start", norm = TRUE,
  useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

plot_locus(
  queryFiles = queryfiles,
  centerFiles = centerfiles,
  ext = c(-500, 500),
  hl = c(-100, 100),
  shade = TRUE,
  smooth = TRUE,
  importParams = chipimportParams,
  binSize = 10,
  refPoint = "center",
  Xlab = "Center",
  inputFiles = inputfiles,
  stranded = TRUE,
  scale = FALSE,
  outPrefix = NULL,
  verbose = FALSE,
  transform = NA,
  rmOutlier = 0,
  Ylab = "Coverage/base/peak",
  statsMethod = "wilcox.test",
  heatmap = TRUE,
  nc = 2
)
Description

Plot reads or peak Coverage/base/gene of samples given in the query files around reference locus defined in the centerFiles. The upstream and downstream windows flanking loci can be given separately, a smaller window can be defined to allow statistical comparisons between reference and random loci. The loci are further divided into sub-groups that are overlapping with c("5'UTR", "CDS", "3'UTR"), "unrestricted" means all loci regardless of overlapping.

Usage

plot_locus_with_random(
  queryFiles,  
centerFiles,  
txdb,  
ext = c(-200, 200),  
hl = c(-100, 100),  
shade = FALSE,  
importParams = NULL,  
verbose = FALSE,  
smooth = FALSE,  
transform = NA,  
binSize = 10,  
refPoint = "center",  
Xlab = "Center",  
Ylab = "Coverage/base/gene",  
inputFiles = NULL,  
stranded = TRUE,  
scale = FALSE,  
outPrefix = NULL,  
rmOutlier = 0,  
n_random = 1,  
hw = c(8, 8),  
detailed = FALSE,  
statsMethod = "wilcox.test",  
nc = 2  
)

Arguments

queryFiles a vector of sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed

centerFiles a vector of reference file names. The file should be .bed format only

txdb a TxDb object defined in the ‘GenomicFeatures’ package

ext a vector of two integers defining upstream and downstream boundaries of the plot window, flanking the reference locus

hl a vector of two integers defining upstream and downstream boundaries of the highlight window, flanking the reference locus
shade logical indicating whether to place a shaded rectangle around the point of interest
importParams a list of parameters for handle_input
verbose logical, indicating whether to output additional information (data used for plotting or statistical test results)
smooth logical, indicating whether the line should smoothed with a spline smoothing algorithm
transform a string in c("log", "log2", "log10"), default = NA indicating no transformation of data matrix
binSize an integer defines bin size for intensity calculation
refPoint a string in c("start", "center", "end")
Xlab a string denotes the label on x-axis
Ylab a string for y-axis label
inputFiles a vector of input sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed
stranded logical, indicating whether the strand of the feature should be considered
scale logical, indicating whether the score matrix should be scaled to the range 0:1, so that samples with different baseline can be compared
outPrefix a string specifying output file prefix for plots (outPrefix.pdf)
rmOutlier a numeric value serving as a multiplier of the MAD in Hampel filter for outliers identification, 0 indicating not removing outliers. For Gaussian distribution, use 3, adjust based on data distribution
n_random an integer denotes the number of randomization should be performed
hw a vector of two elements specifying the height and width of the output figures
detailed logical, indicating whether to plot each parts of gene.
statsMethod a string in c("wilcox.test", "t.test"), for pair-wise groups comparisons
nc integer, number of cores for parallel processing

Value

a dataframe containing the data used for plotting

Author(s)

Shuye Pu

Examples

gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", package = "GenomicPlot"
)

fdb <- custom_TxDb_from_GTF(gtfFile, genome = "hg19")
bedQueryFiles <- c(
plot_named_list

```
system.file("extdata", "test_chip_peak_chr19.narrowPeak",
    package = "GenomicPlot"),
system.file("extdata", "test_chip_peak_chr19.bed", package = "GenomicPlot"),
system.file("extdata", "test_clip_peak_chr19.bed", package = "GenomicPlot")
)
names(bedQueryFiles) <- c("NarrowPeak", "SummitPeak", "iCLIPPeak")

bamQueryFiles <- system.file("extdata", "treat_chr19.bam",
    package = "GenomicPlot")
names(bamQueryFiles) <- "clip_bam"
bamInputFiles <- system.file("extdata", "input_chr19.bam",
    package = "GenomicPlot")
names(bamInputFiles) <- "clip_input"

bamImportParams <- setImportParams(
    offset = -1, fix_width = 0, fix_point = "start", norm = TRUE,
    useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)
plot_locus_with_random(
    queryFiles = bamQueryFiles,
    centerFiles = bedQueryFiles[3],
    txdb = txdb,
    ext = c(-200, 200),
    hl = c(-50, 50),
    shade = TRUE,
    importParams = bamImportParams,
    verbose = FALSE,
    smooth = TRUE,
    transform = NA,
    binSize = 10,
    refPoint = "center",
    Xlab = "Center",
    Ylab = "Coverage/base/peak",
    inputFiles = bamInputFiles,
    stranded = TRUE,
    scale = FALSE,
    outPrefix = NULL,
    rmOutlier = 0,
    n_random = 1,
    hw = c(8, 8),
    detailed = FALSE,
    statsMethod = "wilcox.test",
    nc = 2)
```

---

plot_named_list  plot a named list as a figure
Description

This is a helper function for displaying function arguments for a plotting function. If the runtime value of the argument is a small object, its values is displayed, otherwise, only the name of the value of the argument is displayed.

Usage

plot_named_list(params)

Arguments

params  
a list produced by as.list(environment()), with names being the arguments and values being the runtime values when the function is called.

Value

a ggplot object

Author(s)

Shuye Pu

Examples

data(gf5_genomic)

gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf",  
  package = "GenomicPlot"
)

taxdb <- custom_TxDb_from_GTF(gtfFile, genome = "hg19")

queryfiles <- system.file("extdata", "treat_chr19.bam",  
  package = "GenomicPlot"
)

names(queryfiles) <- "query"

inputfiles <- system.file("extdata", "input_chr19.bam",  
  package = "GenomicPlot"
)

names(inputfiles) <- "input"

bamimportParams <- setImportParams(  
  offset = -1, fix_width = 0, fix_point = "start", norm = TRUE,  
  useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

alist <- list(  
  "txdb" = txdb, "treat" = queryfiles, "control" = inputfiles,  
  "feature" = gf5_genomic, "param" = bamimportParams
)
**plot_overlap_bed**

Plot Venn diagrams depicting overlap of genomic regions

Description

This function takes a list of up to 4 bed file names, and produce a Venn diagram

Usage

```r
plot_overlap_bed(
  bedList,
  outPrefix = NULL,
  importParams = NULL,
  pairOnly = TRUE,
  stranded = TRUE,
  hw = c(8, 8),
  verbose = FALSE
)
```

Arguments

- `bedList`: a named list of bed files, with list length = 2, 3 or 4
- `outPrefix`: a string for plot file name
- `importParams`: a list of parameters for handle_input
- `pairOnly`: logical, indicating whether only pair-wise overlap is desirable
- `stranded`: logical, indicating whether the feature is stranded. For nonstranded feature, only "*" is accepted as strand
- `hw`: a vector of two elements specifying the height and width of the output figures
- `verbose`: logical, indicating whether to output additional information

Value

a ggplot object

Author(s)

Shuye Pu
Examples

```r
queryFiles <- c(
  system.file("extdata", "test_chip_peak_chr19.narrowPeak", package = "GenomicPlot" ),
  system.file("extdata", "test_chip_peak_chr19.bed", package = "GenomicPlot" ),
  system.file("extdata", "test_clip_peak_chr19.bed", package = "GenomicPlot" ),
)
names(queryFiles) <- c("narrowPeak", "summitPeak", "clipPeak")

bedimportParams <- setImportParams(
  offset = 0, fix_width = 100, fix_point = "center", norm = FALSE,
  useScore = FALSE, outRle = FALSE, useSizeFactor = FALSE, genome = "hg19"
)

plot_overlap_bed(
  bedList = queryFiles, importParams = bedimportParams, pairOnly = FALSE,
  stranded = FALSE, outPrefix = NULL
)
```

Description

This function takes a list of (at most 4) tab-delimited file names, and produce a Venn diagram.

Usage

```r
plot_overlap_genes(
  fileList, columnList, pairOnly = TRUE,
  hw = c(8, 8), outPrefix = NULL
)
```

Arguments

- `fileList`: a named list of tab-delimited files
- `columnList`: a vector of integers denoting the columns that have gene names in the list of files
- `pairOnly`: logical, indicating whether only pair-wise overlap is desirable
- `hw`: a vector of two elements specifying the height and width of the output figures
- `outPrefix`: a string for plot file name
plot_peak_annotation

Value
a list of vectors of gene names

Author(s)
Shuye Pu

Examples

testfile1 <- system.file("extdata", "test_file1.txt", package = "GenomicPlot")
testfile2 <- system.file("extdata", "test_file2.txt", package = "GenomicPlot")
testfile3 <- system.file("extdata", "test_file3.txt", package = "GenomicPlot")
testfile4 <- system.file("extdata", "test_file4.txt", package = "GenomicPlot")
testfiles <- c(testfile1, testfile2, testfile3, testfile4)
names(testfiles) <- c("test1", "test2", "test3", "test4")

plot_overlap_genes(testfiles, c(3, 2, 1, 1), pairOnly = FALSE)

plot_peak_annotation Annotate peaks with genomic features and genes

Description
Produce a table of transcripts targeted by peaks, and generate plots for target gene types, and peak distribution in genomic features

Usage

plot_peak_annotation(
  peakFile,
  gtfFile,
  importParams = NULL,
  fiveP = -1000,
  dsTSS = 300,
  threeP = 1000,
  simple = FALSE,
  outPrefix = NULL,
  verbose = FALSE,
  hw = c(8, 8),
  nc = 2
)
Arguments

peakFile a string denoting the peak file name, only .bed format is allowed

gtfFile path to a gene annotation gtf file with gene_biotype field

importParams a list of parameters for handle_input

fiveP extension out of the 5' boundary of genes for defining promoter: fiveP TSS + dsTSS

dsTSS extension downstream of TSS for defining promoter: fiveP TSS + dsTSS

threeP extension out of the 3' boundary of genes for defining termination region: -0 TTS + threeP

simple logical, indicating whether 5’UTR, CDS and 3’UTR are annotated in the gtfFile

outPrefix a string denoting output file name in pdf format

verbose logical, to indicate whether to write the annotation results to a file

hw a vector of two elements specifying the height and width of the output figures

nc number of cores for parallel processing

Value

a list of three dataframes, 'annotation' is the annotation of peaks into gene types, 'stat' is the summary stats for pie chart, 'simplified' is the summary stats excluding intron

Author(s)

Shuye Pu

Examples

gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", package = "GenomicPlot")

centerFile <- system.file("extdata", "test_chip_peak_chr19.bed", package = "GenomicPlot")
names(centerFile) <- c("summitPeak")

bedimportParams <- setImportParams(
  offset = 0, fix_width = 100, fix_point = "center", norm = FALSE,
  useScore = FALSE, outRle = FALSE, useSizeFactor = FALSE, genome = "hg19"
)

plot_peak_annotation(
  peakFile = centerFile, gtfFile = gtfFile, importParams = bedimportParams,
  fiveP = -2000, dsTSS = 200, threeP = 2000, simple = FALSE)
plot_region

Plot signal inside as well as around custom genomic regions

Description
Plot reads or peak Coverage/base/gene of samples given in the query files inside regions defined in the centerFiles. The upstream and downstream flanking windows can be given separately. If Input files are provided, ratio over Input is computed and displayed as well.

Usage
plot_region(
  queryFiles,
  centerFiles,
  txdb = NULL,
  regionName = "region",
  inputFiles = NULL,
  nbins = 100,
  importParams = NULL,
  verbose = FALSE,
  scale = FALSE,
  heatmap = FALSE,
  fiveP = -1000,
  threeP = 1000,
  smooth = FALSE,
  stranded = TRUE,
  transform = NA,
  outPrefix = NULL,
  rmOutlier = 0,
  heatRange = NULL,
  Ylab = "Coverage/base/gene",
  statsMethod = "wilcox.test",
  hw = c(8, 8),
  nc = 2
)

Arguments
queryFiles a named vector of sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed
centerFiles a named vector of reference file names or genomic features in c("utr3", "utr5", "cds", "intron", "exon", "transcript", "gene"). The file should be in .bed format only
txdb a TxDb object defined in the GenomicFeatures package. Default NULL, needed only when genomic features are used as centerFiles.
regionName a string specifying the name of the center region in the plots
inputFiles a named vector of input sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed

nbins an integer defines the total number of bins

importParams a list of parameters for handle_input

verbose logical, indicating whether to output additional information (data used for plotting or statistical test results)

scale logical, indicating whether the score matrix should be scaled to the range 0:1, so that samples with different baseline can be compared

heatmap logical, indicating whether a heatmap of the score matrix should be generated

fiveP an integer, indicating extension out or inside of the 5' boundary of gene by negative or positive number

threeP an integer, indicating extension out or inside of the 5' boundary of gene by positive or negative number

smooth logical, indicating whether the line should smoothed with a spline smoothing algorithm

stranded logical, indicating whether the strand of the feature should be considered

transform a string in c("log", "log2", "log10"), default = NA indicating no transformation of data matrix

outPrefix a string specifying output file prefix for plots (outPrefix.pdf)

rmOutlier a numeric value serving as a multiplier of the MAD in Hampel filter for outliers identification, 0 indicating not removing outliers. For Gaussian distribution, use 3, adjust based on data distribution

heatRange a numeric vector with three elements, defining custom range for color ramp, default=NULL, i.e. the range is defined automatically based on the c(minimun, median, maximum) of a data matrix

Ylab a string for y-axis label

statsMethod a string in c("wilcox.test", "t.test"), for pair-wise group comparisons

hw a vector of two elements specifying the height and width of the output figures

nc integer, number of cores for parallel processing

Value a dataframe containing the data used for plotting

Author(s) Shuye Pu

Examples
centerfiles <- system.file("extdata", "test_chip_peak_chr19.narrowPeak", package = "GenomicPlot")
names(centerfiles) <- c("NarrowPeak")
queryfiles <- c(
system.file("extdata", "chip_treat_chr19.bam", package = "GenomicPlot")
names(queryfiles) <- c("chip_bam")
inputfiles <- c(
  system.file("extdata", "chip_input_chr19.bam", package = "GenomicPlot"))
names(inputfiles) <- c("chip_input")

chipimportParams <- setImportParams(
  offset = 0, fix_width = 150, fix_point = "start", norm = TRUE,
  useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19")

plot_region(
  queryFiles = queryfiles,
  centerFiles = centerfiles,
  inputFiles = inputfiles,
  nbins = 100,
  heatmap = TRUE,
  scale = FALSE,
  regionName = "narrowPeak",
  importParams = chipimportParams,
  verbose = FALSE,
  fiveP = -500,
  threeP = 500,
  smooth = TRUE,
  transform = NA,
  stranded = TRUE,
  outPrefix = NULL,
  Ylab = "Coverage/base/peak",
  rmOutlier = 0,
  nc = 2
)

plot_start_end

---

**plot_start_end**

Plot signals around the start and the end of genomic features

**Description**

Plot reads or peak Coverage/base/gene of samples given in the query files around start and end of custom features. The upstream and downstream windows can be given separately, within the window, a smaller window can be defined to highlight region of interest. A line plot will be displayed for both start and end of feature. If Input files are provided, ratio over Input is computed and displayed as well.

**Usage**

```r
plot_start_end(
  queryFiles,
  inputFiles = NULL,
  centerFiles,
```
txdb = NULL,
importParams = NULL,
binSize = 10,
insert = 0,
verbose = FALSE,
ext = c(-500, 100, -100, 500),
hl = c(-50, 50, -50, 50),
stranded = TRUE,
scale = FALSE,
smooth = FALSE,
rmOutlier = 0,
outPrefix = NULL,
transform = NA,
shade = TRUE,
Ylab = "Coverage/base/gene",
hw = c(8, 8),
nc = 2
)

Arguments

queryFiles  a vector of sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed
inputFiles  a vector of input sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed
centerFiles  bed files that define the custom features, or features in c("utr3", "utr5", "cds", "intron", "exon", "transcript", "gene"), multiple features are allowed.
txdb  a TxDb object defined in the GenomicFeatures package. Default NULL, needed only when genomic features are used in the place of centerFiles.
importParams  a list of parameters for handle_input
binSize  an integer defines bin size for intensity calculation
insert  an integer specifies the length of the center regions to be included, in addition to the start and end of the feature
verbose  logical, whether to output additional information (including data used for plotting or statistical test results)
ext  a vector of four integers defining upstream and downstream boundaries of the plot window, flanking the start and end of features
hl  a vector of four integers defining upstream and downstream boundaries of the highlight window, flanking the start and end of features
stranded  logical, indicating whether the strand of the feature should be considered
scale  logical, indicating whether the score matrix should be scaled to the range 0:1, so that samples with different baseline can be compared
smooth  logical, indicating whether the line should smoothed with a spline smoothing algorithm
rmOutlier

A numeric value serving as a multiplier of the MAD in Hampel filter for outliers identification, 0 indicating not removing outliers. For Gaussian distribution, use 3, adjust based on data distribution.

outPrefix

A string specifying output file prefix for plots (outPrefix.pdf).

transform

A string in c("log", "log2", "log10"), default = NA, indicating no transformation of data matrix.

shade

Logical indicating whether to place a shaded rectangle around the point of interest.

Ylab

A string for y-axis label.

hw

A vector of two elements specifying the height and width of the output figures.

nc

An integer, number of cores for parallel processing.

Value

A list of two objects, the first is a GRanges object, the second is a GRangesList object.

Author(s)

Shuye Pu

Examples

```r
gtffile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", 
package = "GenomicPlot")

txdb <- custom_TxDB_from_GTF(gtffile, genome = "hg19")
bamQueryFiles <- system.file("extdata", "treat_chr19.bam", 
package = "GenomicPlot")
names(bamQueryFiles) <- "clip_bam"
bamInputFiles <- system.file("extdata", "input_chr19.bam", 
package = "GenomicPlot")
names(bamInputFiles) <- "clip_input"

bamimportParams <- setImportParams(
  offset = -1, fix_width = 0, fix_point = "start", norm = TRUE,
  useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

plot_start_end(
  queryFiles = bamQueryFiles, 
  inputFiles = bamInputFiles, 
  txdb = txdb, 
  centerFiles = "intron", 
  binSize = 10, 
  importParams = bamimportParams, 
  ext = c(-500, 200, -200, 500), 
  hl = c(-100, 100, -100, 100), 
  insert = 100, 
  stranded = TRUE,
```

```r
```
plot_start_end_with_random

Plot signals around the start and the end of genomic features and random regions

Description

Plot reads or peak Coverage/base/gene of samples given in the query files around start, end and center of genomic features or custom feature given in a .bed file. The upstream and downstream windows can be given separately. If Input files are provided, ratio over Input is computed and displayed as well. A random feature can be generated to serve as a background for contrasting.

Usage

plot_start_end_with_random(
    queryFiles,
    inputFiles = NULL,
    txdb = NULL,
    centerFile,
    importParams = NULL,
    binSize = 10,
    insert = 0,
    verbose = FALSE,
    ext = c(-500, 200, -200, 500),
    hl = c(-50, 50, -50, 50),
    randomize = FALSE,
    stranded = TRUE,
    scale = FALSE,
    smooth = FALSE,
    rmOutlier = 0,
    outPrefix = NULL,
    transform = NA,
    shade = TRUE,
    nc = 2,
    hw = c(8, 8),
    Ylab = "Coverage/base/gene"
)
Arguments

queryFiles  a vector of sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed
inputFiles  a vector of input sample file names. The file should be in .bam, .bed, .wig or .bw format, mixture of formats is allowed
txdb  a TxDb object defined in the GenomicFeatures package. Default NULL, needed only when genomic features are used in the place of centerFile.
centerFile  a bed file that defines the custom feature, or a feature in c("utr3", "utr5", "cds", "intron", "exon", "transcript", "gene"), multiple features are not allowed.
importParams  a list of parameters for handle_input
binSize  an integer defines bin size for intensity calculation
insert  an integer specifies the length of the center regions to be included, in addition to the start and end of the feature
verbose  logical, whether to output additional information (data used for plotting or statistical test results)
ext  a vector of four integers defining upstream and downstream boundaries of the plot window, flanking the start and end of features
hl  a vector of four integers defining upstream and downstream boundaries of the highlight window, flanking the start and end of features
randomize  logical, indicating if randomized feature should generated and used as a contrast to the real feature. The ransomized feature is generated by shifting the given feature with a random offset within the range of ext[1] and ext[4]
stranded  logical, indicating whether the strand of the feature should be considered
scale  logical, indicating whether the score matrix should be scaled to the range 0:1, so that samples with different baseline can be compared
smooth  logical, indicating whether the line should smoothed with a spline smoothing algorithm
rmOutlier  a numeric value serving as a multiplier of the MAD in Hampel filter for outliers identification, 0 indicating not removing outliers. For Gaussian distribution, use 3, adjust based on data distribution
outPrefix  a string specifying output file prefix for plots (outPrefix.pdf)
transform  a string in c("log", "log2", "log10"), default = NA indicating no transformation of data matrix
shade  logical indicating whether to place a shaded rectangle around the point of interest
nc  integer, number of cores for parallel processing
hw  a vector of two elements specifying the height and width of the output figures
Ylab  a string for y-axis label

Value

a list of two objects, the first is a GRanges object, the second is a GRangesList object
prepare_3parts_genomic_features

Demarcate genes into promoter, gene body and TTS features

Author(s)
Shuye Pu

Examples

gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf",   
   package = "GenomicPlot"
)

txdb <- custom_TxDB_from_GTF(gtfFile, genome = "hg19")

bamQueryFiles <- system.file("extdata", "treat_chr19.bam",   
   package = "GenomicPlot")

names(bamQueryFiles) <- "clip_bam"

bamInputFiles <- system.file("extdata", "input_chr19.bam",   
   package = "GenomicPlot")

names(bamInputFiles) <- "clip_input"

bamImportParams <- setImportParams(   
   offset = -1, fix_width = 0, fix_point = "start", norm = TRUE,   
   useScore = FALSE, outRle = TRUE, useSizeFactor = FALSE, genome = "hg19"
)

plot_start_end_with_random(   
   queryFiles = bamQueryFiles,   
   inputFiles = bamInputFiles,   
   txdb = txdb,   
   centerFile = "intron",   
   binSize = 10,   
   importParams = bamImportParams,   
   ext = c(-100, 100, -100, 100),   
   hl = c(-20, 20, -20, 20),   
   insert = 100,   
   stranded = TRUE,   
   scale = FALSE,   
   smooth = TRUE,   
   verbose = TRUE,   
   transform = "log2",   
   outPrefix = NULL,   
   randomize = TRUE,   
   nc = 2
)
**prepare_3parts_genomic_features**

**Description**

This is a helper function for 'plot_3parts_metagene', used to speed up plotting of multiple data sets with the same configuration. Use featureName='transcript' and meta=FALSE and longest=TRUE for genes.

**Usage**

```r
prepare_3parts_genomic_features(
  txdb,
  featureName = "transcript",
  meta = TRUE,
  nbins = 100,
  fiveP = -1000,
  threeP = 1000,
  longest = TRUE,
  protein_coding = TRUE,
  verbose = FALSE
)
```

**Arguments**

- `txdb` a TxDb object defined in the GenomicFeatures package
- `featureName` one of the gene feature in c("utr3", "utr5", "cds", "transcript")
- `meta` logical, indicating whether a metagene (intron excluded) or genomic (intron included) plot should be produced
- `nbins` an integer defines the total number of bins
- `fiveP` extension out of the 5' boundary of gene
- `threeP` extension out of the 3' boundary of gene
- `longest` logical, indicating whether the output should be limited to the longest transcript of each gene
- `protein_coding` logical, indicating whether to limit to protein_coding genes
- `verbose` logical, whether to output additional information

**Value**

a named list with the elements c("windowRs", "nbins", "scaled_bins", "fiveP", "threeP", "meta", "longest")

**Author(s)**

Shuye Pu
prepare_5parts_genomic_features

Demarcate genes into promoter, 5'UTR, CDS, 3'UTR and TTS features

Description

This is a helper function for 'plot_5parts_metagene', used to speed up plotting of multiple data sets with the same configuration. Only protein-coding genes are considered.

Usage

prepare_5parts_genomic_features(
    txdb,                  
    meta = TRUE,           
    nbins = 100,           
    fiveP = -1000,         
    threeP = 1000,         
    longest = TRUE,        
    verbose = FALSE,       
    subsetTx = NULL       
)

Arguments

taxdb a TxDb object defined in the GenomicFeatures package
meta logical, indicating whether a metagene (intron excluded) or gene (intron included) plot should be produced
nbins an integer defines the total number of bins
fiveP extension out of the 5' boundary of gene
threeP extension out of the 3' boundary of gene
longest logical, indicating whether the output should be limited to the longest transcript of each gene
verbose logical, whether to output additional information
subsetTx a vector of transcript names (eg. ENST00000587541.1) for subsetting the genome

Examples

gtffile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", 
    package = "GenomicPlot"
)

txdb <- custom_TxDb_from_GTF(gtffile, genome = "hg19")

gf <- prepare_3parts_genomic_features(txdb, 
    meta = FALSE, nbins = 100, fiveP = -1000, threeP = 1000, 
    longest = FALSE
)
Value

A named list with the elements c("windowRs", "nbins", "scaled_bins", "fiveP", "threeP", "meta", "longest")

Author(s)

Shuye Pu

Examples

gtfFile <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", package = "GenomicPlot")

)

txdb <- custom_TxDb_from_GTF(gtfFile, genome = "hg19")

gf <- prepare_5parts_genomic_features(txdb,
  meta = TRUE, nbins = 100, fiveP = -0, threeP = 0,
  longest = TRUE
)

Description

This is a helper function for manipulate the score matrix produced by ScoreMatrix or ScoreMatrin-Bin functions defined in the 'genomation' package. To facilitate downstream analysis, imputation of missing values is performed implicitly when log transformation is required, otherwise missing values are replaced with 0.

Usage

process_scoreMatrix(
  fullmatrix,
  scale = FALSE,
  rmOutlier = 0,
  transform = NA,
  verbose = FALSE
)

Arguments

fullmatrix a numeric matrix, with bins in columns and genomic windows in rows
scale logical, indicating whether the score matrix should be scaled to the range 0:1, so that samples with different baseline can be compared
process_scoreMatrix

rmOutlier a numeric value to multiple the 'mad' when detecting outliers, can be adjusted based on data. Default 0, indicating not to remove outliers.

transform a string in c("log", "log2", "log10"), default = NA indicating no transformation of data matrix

verbose logical, indicating whether to output additional information (data used for plotting or statistical test results)

Details

If inputFiles for the plotting function is null, all operations (scale, rmOutlier and transform) can be applied to the score matrix, in the order of rmOutlier -> transform -> scale. When inputFiles are provided, only rmOutlier can be applied to the score matrix, as transform and scale will affect ratio calculation, especially when log2 transformation of the ratio is intended. However, all these operations can be applied to the resulting ratio matrix. In order to avoid introducing distortion into the processed data, use caution when applying these operations.

Value

a numeric matrix with the same dimension as the fullmatrix

Author(s)

Shuye Pu

Examples

fullMatrix <- matrix(rlnorm(100), ncol = 10)
for (i in 5:6) {
    fullMatrix[i, 4:7] <- NaN
    fullMatrix[i + 1, 4:7] <- NA
    fullMatrix[i + 2, 4:7] <- Inf
    fullMatrix[i - 1, 4:7] <- 0
    fullMatrix[i - 2, 1:3] <- Inf
}
fullMatrix[9, 4:7] <- runif(4) + 90

wo <- process_scoreMatrix(fullMatrix, rmOutlier = 3, verbose = TRUE)
tf <- process_scoreMatrix(fullMatrix,
    rmOutlier = 0, transform = "log2", verbose = TRUE
)
scaled <- process_scoreMatrix(fullMatrix, scale = TRUE, verbose = TRUE)
rank_rows

**Rank rows of a matrix based on user input**

**Description**

The rows of a input numeric matrix is ordered based row sum, row maximum, or hierarchical clustering of the rows with euclidean distance and centroid linkage. This a helper function for drawing matrix heatmaps.

**Usage**

```r
description <- rank_rows(fullmatrix, ranking = "Hierarchical")
```

**Arguments**

- `fullmatrix`: a numeric matrix
- `ranking`: a string in c("Sum", "Max", "Hierarchical", "None")

**Value**

a numeric matrix

**Author(s)**

Shuye Pu

**Examples**

```r
fullMatrix <- matrix(rnorm(100), ncol = 10)
for (i in 5:8) {
  fullMatrix[i, 4:7] <- runif(4) + i
}
apply(fullMatrix, 1, sum)
ranked <- rank_rows(fullMatrix, ranking = "Sum")
apply(ranked, 1, sum)
```

ratio_over_input

**compute ratio over input**

**Description**

compute enrichment of IP samples over Input samples

**Usage**

```r
description <- ratio_over_input(IP, Input, verbose = FALSE)
```
rm_outlier

Arguments

- **IP**: a numerical matrix
- **Input**: another numerical matrix with same dimensions as the IP matrix
- **verbose**: logical, whether to output additional information

Value

a numerical matrix with same dimensions as the IP matrix

Author(s)

Shuye Pu

Examples

```r
IP <- matrix(rlnorm(100), ncol = 10)
Input <- matrix(runif(100), ncol = 10)

ratio <- GenomicPlot:::ratio_over_input(IP, Input, verbose = TRUE)
```

Description

This is a helper function for dealing with excessively high values using Hampel filter. If outliers are detected, replace the outliers with the up bound = median(rowmax) + multiplier*mad(rowmax). This function is experimental. For data with normal distribution, the multiplier is usually set at 3. As the read counts data distribution is highly skewed, it is difficult to define a boundary for outliers, try the multiplier values between 10 to 1000.

Usage

```r
rm_outlier(fullmatrix, verbose = FALSE, multiplier = 1000)
```

Arguments

- **fullmatrix**: a numeric matrix, with bins in columns and genomic windows in rows
- **verbose**: logical, whether to output the outlier information to the console
- **multiplier**: a numeric value to multiple the `mad`, default 1000, maybe adjusted based on data

Value

a numeric matrix
setImportParams

Author(s)
Shuye Pu

Examples
fullmatrix <- matrix(rnorm(100), ncol = 10)
maxm <- max(fullmatrix)
fullmatrix[3, 9] <- maxm + 1000
fullmatrix[8, 1] <- maxm + 500
rm_outlier(fullmatrix, verbose = TRUE, multiplier = 100)
r

Description
This function save as a template for setting up import parameters for reading NGS data, it provides
default values for each parameter.

Usage
setImportParams(
  offset = 0,
  fix_width = 0,
  fix_point = "start",
  norm = FALSE,
  useScore = FALSE,
  outRle = TRUE,
  useSizeFactor = FALSE,
  saveRds = FALSE,
  genome = "hg19",
  val = 4,
  skip = 0
)

Arguments

offset an integer, -1 indicating the bam reads should be shrunk to the -1 position at the 5'end of the reads, which corresponds to the cross link site in iCLIP.

fix_width an integer, for bam file, defines how long the reads should be extended from the start position, ignored when offset is not 0; for bed files, defines the width of each interval centering on the 'fix_point'.

fix_point a string in c("start", "end", "center") denoting the anchor point for extension, ignored when offset is not 0.
norm logical, indicating whether the output RleList should be normalized to RPM using library sizes.
useScore logical, indicating whether the 'score' column of the bed file should be used in calculation of coverage.
outRle logical, indicating whether the output should be RleList objects or GRanges objects.
useSizeFactor logical, indicating whether the library size should be adjusted with a size factor, using the 'calcNormFactors' function in the edgeR package, only applicable to ChIPseq data.
saveRds logical, indicating whether the results of handle_input should be saved for fast reloading
genome a string denoting the genome name and version.
val integer, indicating the column that will be used as score/value. default 4 for bedGraph.
skip integer, indicating how many rows will be skipped before reading in data, default 0.

Value

a list of nine elements

Author(s)

Shuye Pu

Examples

importParams1 <- setImportParams()
importParams2 <- setImportParams(offset = -1, saveRds = TRUE)

=set_seqinfo

Set standard chromosome size of model organisms

Description

This is a helper function for making Seqinfo objects, which is a components of GRanges and TxDb objects. It also serves to unify seqlevels between GRanges and TxDb objects. Mitochondrial chromosome is not included.

Usage

set_seqinfo(genome = "hg19")

Arguments

genome a string denoting the genome name and version
Value

A Seqinfo object defined in the GenomeInfoDb package.

Author(s)

Shuye Pu

Examples

```r
out <- set_seqinfo(genome = "hg19")
```

Description

Creating a virtual cluster for parallel processing

Usage

```r
start_parallel(nc = 2, verbose = FALSE)
```

Arguments

- `nc`: a positive integer greater than 1, denoting number of cores requested
- `verbose`: logical, whether to output additional information

Value

An object of class c("SOCKcluster", "cluster"), depending on platform

Author(s)

Shuye Pu

Examples

```r
cl <- start_parallel(2L)
stop_parallel(cl)
```
### stop_parallel

**Stop parallel processing**

**Description**

Stopping a virtual cluster after parallel processing is finished

**Usage**

```r
stop_parallel(cl)
```

**Arguments**

- `cl` a cluster or SOCKcluster object depending on platform

**Value**

0 if the cluster is stopped successfully, 1 otherwise.

**Author(s)**

Shuye Pu

**Examples**

```r
c1 <- start_parallel(2L)
stop_parallel(c1)
```

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### txdb.sql

**Toy data for examples and testing of the 'GenomicPlot' package**

**Description**

A tiny TxDb object holding genomic feature coordinates of 72 transcripts in hg19.

**Value**

A SQLite database

**Author(s)**

Shuye Pu
Source

The data is produced by running the following code:

gtf <- system.file("extdata", "gencode.v19.annotation_chr19.gtf", package = "GenomicPlot")
txdb <- custom_TxDB_from_GTF(gtf, genome = "hg19")
AnnotationDbi::saveDb(txdb, "/inst/extdata/txdb.sql")
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