Package 'cliProfiler'

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Type Package

Title A package for the CLIP data visualization

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Description An easy and fast way to visualize and profile the high-throughput IP data. This package generates the meta gene profile and other profiles. These profiles could provide valuable information for understanding the IP experiment results.

License Artistic-2.0

VignetteBuilder knitr

Encoding UTF-8

LazyData TRUE

URL https://github.com/Codezy99/cliProfiler

BugReports https://github.com/Codezy99/cliProfiler/issues

biocViews Sequencing, ChIPSeq, Visualization, Epigenetics, Genetics

RoxygenNote 7.1.1

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exonProfile exonProfile for the GRanges objects

Description

An function to check the position of peaks in the exonic region.

Usage

```
exonProfile(
    object,
    annotation,
    title = "Exon Profile",
    group = NA,
    exlevel = NA,
    extranscript_support_level = NA,
    maxLength = NA,
    minLength = NA,
    nomap = FALSE
)
```

Arguments

| object | A GRanges object which should contains all the peaks that you want to check |
|------------|--|
| annotation | A path way to the annotation file. The format of the annotation file should be gff3 and downloaded from https://www.gencodegenes.org/ |
| title | The main title for the output meta gene profile plot. |
| group | The column name which contains the information of grouping for making the comparison plot. NA means all the peaks belongs to the same catagory. |
| exlevel | A parameter for the annotation filtering. exlevel represents the level that you would like to exclude. NA means no level filtering for the annotation file. The level from the annotations refers to how reliable this annotation is. For more information about level please check https://www.gencodegenes.org/pages/data_format.html. |

exonProfile

extranscript_support_level

| | A parameter for the annotation filtering. extranscript_support_level represents the transcript_support_level that you would like to exclude (e.g. 4 and 5). NA means no transcript_support_level filtering for the annotation file. Transcripts are scored according to how well mRNA and EST alignments match over its full length. Here the number 6 means the transcript_support_level NA. For more in- formation about level please check https://www.gencodegenes.org/pages/data_format.html. |
|-----------|---|
| maxLength | A numeric value which indicate the maximum value of exon length for the anno- tation filtering. Or a NA which will turn off the max length annotation filtering. |
| minLength | A numeric value which indicate the minimum value of exon length for the anno- tation filtering. Or a NA which will turn off the min length annotation filtering. |
| nomap | A logical vector (TRUE or FALSE). It indicates whether you would like to ex- clude peaks that cannot assign to annotations in the plot. |

Details

Here is an explanation of output meta data in the list 1:

center: The center position of each peaks. This center position is used for calculating the position of peaks within the genomic regions.

- exon_S and exon_E: The location of 5' and 3' splice sites (SS) of the exon.
- exon_length: The length of the exon that peak assigned.
- exon_transcript_id: The transcript ID for the exon
- exon_map: The relative position of each peak. This value close to 0 means this peak located close to the 3' SS. The position value close to one means the peak close to the 5' SS. Value 3 means this peaks can not map to any annotation.

Value

A list object, the list 1 contains the information of the assignment of the peaks and their position value within the exon. The value close to 1 means the peak close to the 5' splice site. The list 2 includes the plot of exonProfile.

Author(s)

You Zhou, Kathi Zarnack

```
## Load the test data and get the path to the test gff3 file
testpath <- system.file("extdata", package = "cliProfiler")
test <- readRDS(file.path(testpath, "test.rds"))
test_gff3 <- file.path(testpath, "annotation_test.gff3")</pre>
```

```
output <- exonProfile(test, test_gff3)</pre>
```

geneTypeProfile

Description

An function to check the gene type belonging for the peaks.

Usage

```
geneTypeProfile(
    object,
    annotation,
    title = "Gene Type Profile",
    exlevel = NA,
    extranscript_support_level = NA
)
```

Arguments

| object | A GRanges object which should contains all the peaks that you want to check |
|-----------------|--|
| annotation | A path way to the annotation file. The format of the annotation file should be gff3 and downloaded from https://www.gencodegenes.org/ |
| title | The main title for the output meta gene profile plot. |
| exlevel | A parameter for the annotation filtering. exlevel represents the level that you would like to exclude. NA means no level filtering for the annotation file. The level from the annotations refers to how reliable this annotation is. For more information about level please check https://www.gencodegenes.org/pages/data_format.html. |
| extranscript_su | pport_level |
| | A parameter for the annotation filtering. extranscript_support_level represents the transcript_support_level that you would like to exclude (e.g. 4 and 5). NA means no transcript_support_level filtering for the annotation file. Transcripts are scored according to how well mRNA and EST alignments match over its full length. Here the number 6 means the transcript_support_level NA. For more information about level please check https://www.gencodegenes.org/pages/data_format.html. |

Details

Here is an explanation of output meta data in the list 1:

center: The center position of each peaks. This center position is used for calculating the position of peaks within the genomic regions.

- geneType: The gene type of the gene that input peak belongs to.
- Gene_ID: The gene ID of the gene that input peak belongs to.

intronProfile

Value

A list object, the list 1 contains the information of the assignment of the peaks and the gene type of their located genes. The list 2 includes the plot of geneTypeProfile

Author(s)

You Zhou, Kathi Zarnack

Examples

```
## Load the test data and get the path to the test gff3 file
testpath <- system.file("extdata", package = "cliProfiler")
test <- readRDS(file.path(testpath, "test.rds"))
test_gff3 <- file.path(testpath, "annotation_test.gff3")</pre>
```

```
output <- geneTypeProfile(test, test_gff3)</pre>
```

intronProfile *intronProfile for the GRanges objects*

Description

An function to check the position of peaks in the intronic region.

Usage

```
intronProfile(
    object,
    annotation,
    title = "Intron Profile",
    group = NA,
    exlevel = NA,
    extranscript_support_level = NA,
    maxLength = NA,
    minLength = NA,
    nomap = FALSE
```

```
)
```

Arguments

| object | A GRanges object which should contains all the peaks that you want to check |
|------------|---|
| annotation | A path way to the annotation file. The format of the annotation file should be gff3 and downloaded from https://www.gencodegenes.org/ |
| title | The main title for the output meta gene profile plot. |
| group | The column name which contains the information of grouping for making the comparison plot. NA means all the peaks belongs to the same catagory. |

| exlevel | A parameter for the annotation filtering. exlevel represents the level that you would like to exclude. NA means no level filtering for the annotation file. The level from the annotations refers to how reliable this annotation is. For more information about level please check https://www.gencodegenes.org/pages/data_format.html. |
|-----------------|--|
| extranscript_su | |
| | A parameter for the annotation filtering. extranscript_support_level represents the transcript_support_level that you would like to exclude (e.g. 4 and 5). NA means no transcript_support_level filtering for the annotation file. Transcripts are scored according to how well mRNA and EST alignments match over its full length. Here the number 6 means the transcript_support_level NA. For more information about level please check https://www.gencodegenes.org/pages/data_format.html. |
| maxLength | A numeric value which indicate the maximum value of exon length for the anno- tation filtering. Or a NA which will turn off the max length annotation filtering. |
| minLength | A numeric value which indicate the minimum value of exon length for the anno- tation filtering. Or a NA which will turn off the min length annotation filtering. |
| nomap | A logical vector (TRUE or FALSE). It indicates whether you would like to exclude peaks that cannot assign to annotations in the plot. |

Details

Here is an explanation of output meta data in the list 1:

center: The center position of each peaks. This center position is used for calculating the position of peaks within the genomic regions.

- Intron_S and Intron_E: The location of 5' and 3' splice sites (SS) of the intron.
- Intron_length: The length of the intron that peak assigned.
- Intron_transcript_id: The transcript ID for the intron.
- Intron_map: The relative position of each peak. This value close to 0 means this peak located close to the 5' SS. The position value close to one means the peak close to the 3' SS. Value 3 means this peaks can not map to any annotation.

Value

A list object, the list 1 contains the information of the assignment of the peaks and their position value within the intron. The value close to 1 means the peak close to the 3' splice site. The list 2 includes the plot of intronProfile.

Author(s)

You Zhou, Kathi Zarnack

Examples

```
## Load the test data and get the path to the test gff3 file
testpath <- system.file("extdata", package = "cliProfiler")
test <- readRDS(file.path(testpath, "test.rds"))
test_gff3 <- file.path(testpath, "annotation_test.gff3")</pre>
```

output <- intronProfile(test, test_gff3)</pre>

metaGeneProfile *metaGeneProfile for the GRanges objects*

Description

An function for calculating the genomic position and generate the meta gene profile plot of the input peaks.

Usage

```
metaGeneProfile(
    object,
    annotation,
    include_intron = FALSE,
    title = "Meta Gene Profile",
    group = NA,
    split = FALSE,
    exlevel = NA,
    extranscript_support_level = NA,
    adjust = 1,
    nomap = FALSE
)
```

Arguments

| object | A GRanges object which should contains all the peaks that you want to check |
|-----------------|---|
| annotation | A path way to the annotation file. The format of the annotation file should be gff3 and downloaded from https://www.gencodegenes.org/ |
| include_intron | A logical vector TRUE or FALSE that define whether the intronic region should be included in the position calculation or not. |
| title | The main title for the output meta gene profile plot. |
| group | The column name which contains the information of grouping for making the comparison plot. NA means all the peaks belongs to the same catagory. |
| split | A logical vector which indicates whether the plot should show the density curve for 3'UTR, CDS 5'UTR, respectively. |
| exlevel | A parameter for the annotation filtering. exlevel represents the level that you would like to exclude. NA means no level filtering for the annotation file. The level from the annotations refers to how reliable this annotation is. For more information about level please check https://www.gencodegenes.org/pages/data_format.html. |
| extranscript_su | upport_level |
| | A parameter for the annotation filtering. extranscript_support_level represents the transcript_support_level that you would like to exclude (e.g. 4 and 5). NA means no transcript_support_level filtering for the annotation file. Transcripts are scored according to how well mRNA and EST alignments match over its full length. Here the number 6 means the transcript_support_level NA. For more in- formation about level please check https://www.gencodegenes.org/pages/data_format.html. |

| adjust | A parameter inherit from ggplot2. A multiplicate bandwidth adjustment. This |
|--------|---|
| | makes it possible to adjust the bandwidth while still using the a bandwidth esti- |
| | mator. For example, $adjust = 1/2$ means use half of the default bandwidth. |
| nomap | A logical vector. It indicates whether you would like to exclude peaks that |
| | cannot assign to annotations in the plot. |

Details

Here is an explanation of output meta data in the list 1:

center: The center position of each peaks. This center position is used for calculating the position of peaks within the genomic regions.

- location: Which genomic region this peak belongs to.
- Gene ID: Which gene this peak belongs to.
- Position: The relative position of each peak. This value close to 0 means this peak located close to the 5' end of the genomic feature. The position value close to one means the peak close to the 3' end of the genomic feature. Value 5 means this peaks can not map to any annotation.

Value

A list object, the list 1 contains the information of the assignment of the peaks and their position value. The position value between 0 to 1 means it located at the 5' UTR, the value close to the 1 means the position of this peak close to the 3' end of the 5' UTR. Peaks located at CDS would have a number between 1 and 2. Postion value between 2 to 3 means this peak assigned to the 3' UTR. For the peaks which can not be assignment to any annotations, they have the value 5. The list 2 includes the plot of meta gene profile.

Author(s)

You Zhou, Kathi Zarnack

```
## Load the test data and get the path to the test gff3 file
testpath <- system.file("extdata", package = "cliProfiler")
test <- readRDS(file.path(testpath, "test.rds"))
test_gff3 <- file.path(testpath, "annotation_test.gff3")
output <- metaGeneProfile(
    object = test, annotation = test_gff3,
    include_intron = FALSE
)
```

motifProfile

Description

An function to plot the frequency or fraction of the interested motif around the center of input peaks.

Usage

```
motifProfile(
    object,
    motif = NA,
    genome = NA,
    fraction = TRUE,
    title = "Motif Profile",
    flanking = 10
)
```

Arguments

| object | A GRanges object which should contains all the peaks that you want to check |
|----------|---|
| motif | A character string which use the IUPAC nucleotide code, e.g. DRACH, TTAGGG. |
| genome | The name of the full genome sequences package in the Bioconductor, e.g. "BSgenome.Mmusculus.UCSC You should install the package before running this function. |
| fraction | A logical vector (TRUE or FALSE) that the result should be presented in fraction or number. |
| title | The main title for the output meta gene profile plot. |
| flanking | The size of the flanking windows that you would like to check. Flanking=5 will give you the result of the 10+1nt windows around the center of peaks. |

Value

A list object, the list 1 contains the information of the frequency of specified motif around the center of peaks. The list 2 includes the plot of motifProfile.

Author(s)

You Zhou, Kathi Zarnack

```
## Load the test data and get the path to the test gff3 file
testpath <- system.file("extdata", package = "cliProfiler")
test <- readRDS(file.path(testpath, "test.rds"))
test_gff3 <- file.path(testpath, "annotation_test.gff3")</pre>
```

```
## Please make sure that the correct BSgenome package have installed before
## running motifProfile. For example,library("BSgenome.Mmusculus.UCSC.mm10")
## would be required for the mouse data.
output <- motifProfile(test,
    motif = "DRACH",
    genome = "BSgenome.Mmusculus.UCSC.mm10",
    flanking = 20
)</pre>
```

spliceSiteProfile spliceSiteProfile for the GRanges objects

Description

An function to check the enrichment of peaks around the splice sites in a absolute distance.

Usage

```
spliceSiteProfile(
    object,
    annotation,
    title = "Splice Site Profile",
    exlevel = NA,
    extranscript_support_level = NA,
    exon_length_filtering = TRUE,
    intron_length_filtering = TRUE,
    flanking = 150,
    bin = 30
)
```

Arguments

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windowProfile

| exon_length_filtering | |
|-----------------------|--|
| | The exon_length_filtering should be a logical value which indicated whether user would like to exclude the exons that have a length less than flanking value. |
| | Set this parameter to TRUE to turn on this filtering step. |
| intron_length_ | filtering |
| | The intron_length_filtering should be a logical value which indicated whether user would like to exclude the introns that have a length less than flanking value. Set this parameter to TRUE to turn on this filtering step. |
| flanking | The size of the flanking windows that you would like to check. Flanking=5 will give you the result of the 10+1nt windows around the center of peaks. |
| bin | A number that indicates how many bins would you like to use in the histogram. |

Value

A list object, the list 1 contains the information of the position of peaks around 5' or 3' splice sites. The list 2 includes the plot of spliceSiteProfile

Author(s)

You Zhou, Kathi Zarnack

Examples

```
## Load the test data and get the path to the test gff3 file
testpath <- system.file("extdata", package = "cliProfiler")
test <- readRDS(file.path(testpath, "test.rds"))
test_gff3 <- file.path(testpath, "annotation_test.gff3")
output <- spliceSiteProfile(test, test_gff3,
      flanking = 200, bin = 40
)
```

windowProfile windowProfile for the GRanges objects

Description

An function to check the position of peaks within the given GRanges windows.

Usage

```
windowProfile(
    object,
    annotation,
    title = "Window Profile",
    group = NA,
    nomap = FALSE
)
```

Arguments

| object | A GRanges object which contains all the peaks that you want to check |
|------------|---|
| annotation | A GRanges object that includes the customised genomic region. |
| title | The main title for the output meta gene profile plot. |
| group | The column name which contains the information of grouping for making the comparison plot. NA means all the peaks belongs to the same catagory. |
| nomap | A logical vector (TRUE or FALSE). It indicates whether you would like to exclude peaks that cannot assign to annotations in the plot. |

Details

Here is an explanation of output meta data in the list 1:

center: The center position of each peaks. This center position is used for calculating the position of peaks within the genomic regions.

- window_S and window_E: The boundary of the annotation that peaks are assigned.
- window_length: The length of the annotation feature that peak assigned.
- window_map: The relative position of each peak. This value close to 0 means this peak located close to the 5' end of the annotation. The position value close to one means the peak close to the 3' end. Value 3 means this peaks can not map to any annotation.

Value

A list object, the list 1 contains the information of the assignment of the peaks and their position value within the given region. The value close to 1 means the peak close to the end of region in 3' end direction. The list 2 includes the ggplot of windowProfile.

Author(s)

You Zhou, Kathi Zarnack

```
## Load the test data and get the path to the test gff3 file
testpath <- system.file("extdata", package = "cliProfiler")
test <- readRDS(file.path(testpath, "test.rds"))
test_gff3 <- file.path(testpath, "annotation_test.gff3")
test_gff3 <- rtracklayer::import.gff3(test_gff3)</pre>
```

```
output <- windowProfile(test, test_gff3)</pre>
```

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