Package 'ssviz'

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Type Package

Title A small RNA-seq visualizer and analysis toolkit

Version 1.22.0

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Description Small RNA sequencing viewer

License GPL-2

Depends R (>=

2.15.1), methods, Rsamtools, Biostrings, reshape, ggplot2, RColorBrewer, stats

biocViews ImmunoOncology,

Sequencing, RNASeq, Visualization, MultipleComparison, Genetics

Collate AllClasses.R AllGenerics.R helper.R

VignetteBuilder knitr

Suggests knitr

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counts

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ssviz-package ssviz

Description

A package for short RNA seq visualization and quantification.

Details

| Package: | ssviz |
|----------|------------|
| Type: | Package |
| Version: | 0.99 |
| Date: | 2014-05-08 |
| License: | GPL-2 |

Author(s)

Diana H.P. Low Maintainer: Diana Low <dlow@imcb.a-star.edu.sg>

counts

counts data

Description

counts is an example total read count for bam reads

Usage

data(ssviz)

ctrlbam

Source

internal

ctrlbam

ctrlbam data

Description

ctrlbam is an example control dataset from bam file read in with readBam

Usage

data(ssviz)

Source

internal

getCountMatrix getCountMatrix

Description

returns the bam data.frame with an additional column counts. Only relevant if the fasta file used for mapping input was previously collapsed via fastx_toolkit to return a fasta read name in the format of readnumber-totalcounts

Usage

getCountMatrix(bam_file,pseudo=FALSE)

Arguments

| bam_file | An object of class DataFrame (from IRanges). Can be generated from readBam. |
|----------|---|
| pseudo | Logical. If TRUE, assume the reads in the bam file does not have a count record |
| | and sets all counts to 1. |

Value

An object of class data.frame having the values from the original bam file with an additional 'count' column.

Author(s)

Diana H.P. Low

See Also

readBam

Examples

data(ssviz)
getCountMatrix(ctrlbam)

getCountMatrix-methods

getCountMatrix

Description

returns the bam data.frame with an additional column counts. Only relevant if the fasta file used for mapping input was previously collapsed via fastx_toolkit to return a fasta read name in the format of readnumber-totalcounts

Methods

signature(object="DataFrame") Returns and object of class data.frame having the values from the original bam file with an additional 'count' column.

logicalORmissing-class

Class "logicalORmissing"

Description

Class union of logical and missing object.

Author(s)

Diana H.P. Low

Examples

showClass("logicalORmissing")

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ntfreq

Description

Calculates nucleotide frequency of reads in bam file

Usage

```
ntfreq(bam_file, ntlength, toRNA = TRUE, count_type = "total")
```

Arguments

| bam_file | An object of class data.frame or DataFrame |
|------------|---|
| ntlength | An integer specifying the length of the sequence to quantify |
| toRNA | A logical value on whether to translate the DNA sequence to RNA |
| count_type | A character string on how to count the nucleotides. Can be either "total" or "unique". If total is selected, the function will look for the countcolumn and multiply the reads by its number of occurence when calculating the frequency. |

Value

Returns a data.frame of the frequency of nucleotides (either A/C/G/T or A/C/G/U) at each position up to the specified ntlength

Author(s)

Diana H.P. Low

Examples

```
data(ssviz)
freq<-ntfreq(pctrlbam,ntlength=10)</pre>
```

ntfreq-methods *ntfreq*

Description

Calculates nucleotide frequency of reads in bam file

Methods

ntfreq(bam_file, ntlength, toRNA = TRUE, count_type = "total") Returns a data frame of nucleotide frequences along length of sequence provided.

Author(s)

Diana H.P. Low

pctrlbam

Description

pctrlbam is an example control dataset from bam file read in with readBam

Usage

data(ssviz)

Source

internal

pingpong

pingpong

Description

piRNA ping-pong analysis of complementary sequences

Usage

```
pingpong(bam_file)
```

Arguments

bam_file An object of class data.frame or DataFrame

Details

The ping-pong mechanism is a proposed method for the amplification of primary piRNAs, which leads to the production of new primary piRNAs from their precursor transcripts, which eventually amplifies the pool of both primary and secondary piRNAs. This positive feedback loop is a secondary biogenesis mechanism that requires complementary transcripts to a pre-existing pool of piRNAs.

Value

This function returns a data.frame object with frequency of overlapping complementary piRNAs.

Author(s)

Diana H.P. Low

References

Brennecke J. et al. Cell 128, 1089-1103, March 23, 2007

pingpong-methods

Examples

```
data(ssviz)
pp<-pingpong(pctrlbam)</pre>
```

pingpong-methods pingpong

Description

piRNA ping-pong analysis of complementary sequences

Methods

pingpong(bam_file) Returns a data.frame object with frequency of overlapping complementary piRNAs.

Author(s)

Diana H.P. Low

plotDistro

plotDistro

Description

Plots distribution of reads in the bam file based on length, direction (strand) or location (rname)

Usage

```
plotDistro(bamlist, type = "qwidth", samplenames = NULL, unique = FALSE, ncounts = NULL, norm = FALSE
```

Arguments

| bamlist | An object of type list, giving a list of bam files. If you only have 1 file, use list(bam_file) |
|-------------|---|
| type | An object of type character. Can be qwidth, rname or strand. In theory, any column property existing in the bam file can be used, but these 3 would be most meaningful. |
| samplenames | Labels for the plot. |
| unique | Logical value to use unique reads (TRUE) or all reads (FALSE) |
| ncounts | Number of total counts in the bam file, used if unique is set to FALSE. |
| norm | Logical value to determine if plot will be normalised. |
| yname | y axis label. |

Author(s)

Diana H.P. Low

plotFreq

Examples

```
data(ssviz)
plotDistro(list(ctrlbam))
```

plotDistro-methods plotDistro

Description

Plots distribution of reads in the bam file based on length, direction (strand) or location (rname)

Methods

```
plotDistro(bamlist, type = "qwidth", samplenames = NULL, unique = FALSE, ncounts = 1e+06, norm = FALSE,
Returns a distribution plot.
```

Author(s)

Diana H.P. Low

plotFreq

```
plotFreq
```

Description

Plots nucleotide frequency generated by ntfreq

Usage

```
plotFreq(freqvector, percentage = TRUE)
```

Arguments

| freqvector | data.frame object generated by ntfreq |
|------------|---|
| percentage | Logical value to represent y-axis as percentage or frequency. |

Author(s)

Diana H.P. Low

See Also

ntfreq

Examples

```
data(ssviz)
freq<-ntfreq(pctrlbam,ntlength=10)
plotFreq(freq)</pre>
```

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plotFreq-methods plotFreq

Description

Plots nucleotide frequency generated by ntfreq

Methods

plotFreq(freqvector, percentage = TRUE) Returns a frequency bar plot.

Author(s)

Diana H.P. Low

plotPP

plotPP

Description

Plots the ping-pong frequency of piRNA amplification

Usage

plotPP(pout, samplenames = NULL)

Arguments

| pout | An object of type data.frame generated by pingpong |
|-------------|--|
| samplenames | An object of type character for sample labels. |

Author(s)

Diana H.P. Low

References

Brennecke J. et al. Cell 128, 1089-1103, March 23, 2007

See Also

pingpong

Examples

```
data(ssviz)
pp<-pingpong(pctrlbam)
plotPP(list(pp))</pre>
```

plotPP-methods plotPP

Description

Plots the ping-pong frequency of piRNA amplification

Methods

plotPP(pout, samplenames = NULL) Returns the pingpong amplification plot.

Author(s)

Diana H.P. Low

plotRegion plotRegion

Description

Plots the read density given a chromosome region.

Usage

```
plotRegion(bamlist, region, howsmooth = 2, ncounts = NULL, samplenames = NULL)
```

Arguments

| bamlist | An object of type list, giving a list of bam files. If you only have 1 file, use list(bam_file) |
|-------------|---|
| region | An object of type character defining the region to plot. Eg. chr1:1000-2000 |
| howsmooth | Numeric value controlling smoothness of the plot. |
| ncounts | Total number of reads for plot normalization. |
| samplenames | Sample names |

Value

Returns the x and y components of the region's reads and plots the density.

Author(s)

Diana H.P. Low

Examples

```
data(ssviz)
region<-'chr1:3015526-3080526'
plotRegion(list(ctrlbam), region=region)</pre>
```

plotRegion-methods plotRegion

Description

Plots the read density given a chromosome region.

Methods

plotRegion(bamlist, region, howsmooth = 2, ncounts = NULL, samplenames = NULL) Returns
 the x and y components of the region's reads and plots the density.

Author(s)

Diana H.P. Low

ptreatbam

ptreatbam data

Description

ptreatbam is an example treatment dataset from bam file read in with readBam

Usage

data(ssviz)

Source

internal

| readBam | readBam | | |
|---------|---------|--|--|
| | | | |

Description

Reads a bam file through RSamtools, and converts it into a data frame of class DataFrame

Usage

readBam(file_name, tags = character(0))

Arguments

| file_name | Character string of bam file location |
|-----------|---|
| tags | Bam tags to import into the data frame. By default it only takes the standard |
| | values if none are given. |

Details

This function formalizes what had been described in the RSamtools documentation and makes it easier to compute the downstream functions in this package.

Value

Returns the bam file contents in a readable dataframe format.

Author(s)

Diana H.P. Low

References

RSamtools package

Examples

```
bam.files <- dir(system.file("extdata", package = "ssviz"), full = TRUE, patt = "bam$")
ctrlbam <- readBam(bam.files[1])</pre>
```

readBam-methods readBam

Description

Reads a bam file through RSamtools, and converts it into a data frame of class DataFrame

Methods

readBam(bam_file, tags = character(0)) Returns the bam file contents in a readable dataframe format.

Author(s)

Diana H.P. Low

treatbam

treatbam data

Description

treatbam is an example treatment dataset from bam file read in with readBam

Usage

data(ssviz)

Source

internal

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