Package 'riboWaltz'

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Type Package

Title Optimization of ribosome P-site positioning in ribosome profiling data

Version 1.2.0

Description riboWaltz is an R package designed for the analysis of ribosome profiling (RiboSeq) data aimed at the identification of the P-site offset. The P-site offset (PO) is specified by the localization of the P-site of ribosomes within the fragments of the RNA (reads) resulting from RiboSeq assays. It is defined as the distance of the P-site from the two ends of the reads. Determining the PO is a crucial step for a variety of RiboSeq-based analyses such as verify the so-called 3-nt periodicity of ribosomes along the coding sequence, derive translation initiation and elongation rates and reveal new translational events in unannotated open reading frames and ncRNAs. riboWaltz performs accurate computation of the PO for all the lengths of reads from single or multiple samples, taking advantage from an original two-step algorithm. Moreover, riboWaltz provides the user a variety of graphical representations, laying the groundwork for further positional analyses and new biological discoveries.

```
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LazyData TRUE
Depends R (>= 3.3.0)
Imports Biostrings (>= 2.46.0),
      data.table (>= 1.10.4.3),
      GenomicAlignments (>= 1.14.1),
      GenomicFeatures (\geq 1.24.5),
      GenomicRanges (>= 1.24.3),
      ggplot2 (>= 2.2.1),
      ggrepel (>= 0.6.5),
      IRanges (>= 2.12.0)
biocViews
RoxygenNote 7.1.1
Encoding UTF-8
Suggests knitr,
     rmarkdown
```

VignetteBuilder knitr

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bamtobed

From BAM files to BED files.

Description

This function reads one or multiple BAM files converting them into BED files that contain, for each read: i) the name of the corresponding reference sequence (i.e. of the transcript on which it aligns); ii) its leftmost and rightmost position with respect to the 1st nucleotide of the reference sequence; iii) its length (intended as the width of the reference sequence region covered by the RNA fragment. For further information about this choice please refer to section Details of function bamtolist); iv) the strand on which it aligns. Please note: this function relies on the bamtobed utility of the BEDTools suite and can be only run on UNIX, LINUX and Apple OS X operating systems. Moreover, to generate R data structures containing reads information, the function bedtolist must be run on the resulting BED files. For these reasons the authors suggest the use of bamtolist.

Usage

bamtobed(bamfolder, bedfolder = NULL)

Arguments

bamfolder Character string specifying the path to the folder storing BAM files. Please note:

the function looks for BAM files recursively starting from the specified folder.

bedfolder Character string specifying the path to the directory where BED files should be

stored. If the specified folder doesn't exist, it is automatically created. If NULL (the default), BED files are stored in a new subfolder of the working directory,

called bed.

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Examples

```
## path_bam <- "path/to/BAM/files"
## path_bed <- "path/to/output/directory"
## bamtobed(bamfolder = path_bam, bedfolder = path_bed)</pre>
```

bamtolist

From BAM files to lists of data tables or GRangesList objects.

Description

This function reads one or multiple BAM files converting them into data tables or GRanges objects, arranged in a list or a GRangesList, respectively. In both cases the list elements contain, for each read: i) the name of the corresponding reference sequence (i.e. of the transcript on which it aligns); ii) its leftmost and rightmost position with respect to the 1st nucleotide of the reference sequence; iii) its length (intended as the width of the reference sequence region covered by the RNA fragment, see parameter indel_threshold and Details); iv) the leftmost and rightmost position of the annotated CDS of the reference sequence (if any) with respect to its 1st nucleotide. Please note: start and stop codon positions for transcripts without annotated CDS are set to 0.

Usage

```
bamtolist(
  bamfolder,
  annotation,
  transcript_align = TRUE,
  name_samples = NULL,
  indel_threshold = 5,
  refseq_sep = NULL,
  output_class = "datatable"
)
```

Arguments

 ${\tt bamfolder}$

Character string specifying the path to the folder storing BAM files.

annotation

Data table as generated by create_annotation. Please make sure the name of reference transcripts in the annotation data table match those in the BAM files (see refseq_sep).

transcript_align

Logical value whether BAM files in bamfolder come from a transcriptome alignment (intended as an alignment against reference transcript sequences, see Details). If TRUE (the default), reads mapping on the negative strand should not be present and, if any, they are automatically removed.

name_samples

Named character string vector specifying the desired name for the elements of the output list of data tables. A character string for each BAM file in bamfolder is required. Plase be careful to name each element of the vector after the correct corresponding BAM file in bamfolder, leaving their path and extension out. No specific order is required. Default is NULL i.e. list elements are named after the name of the BAM files, leaving their path and extension out. See the example for additional details.

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indel_threshold

Positive integer value specifying the maximum number of indels (insertions + deletions) allowed for each read. All reads associated to more indels than specified are discorded (see Potails). Default is 5

ified are discarded (see Details). Default is 5.

refseq_sep Character specifying the separator between reference sequences' name and ad-

ditional information to discard, stored in the same field (see Details). All characters before the first occurrence of the specified separator are kept. Default is

NULL i.e. no string splitting is performed.

output_class Either "datatable" or "granges". It specifies the format of the output i.e. a list of

data tables or a GRangesList object. Default is "datatable".

Details

riboWaltz only works for read alignments based on transcript coordinates. This choice is due to the main purpose of RiboSeq assays to study translational events through the isolation and sequencing of ribosome protected fragments. Most reads from RiboSeq are supposed to map on mRNAs and not on introns and intergenic regions. Nevertheless, BAM based on transcript coordinates can be generated in two ways: i) aligning directly against transcript sequences; ii) aligning against standard chromosome sequences, requiring the outputs to be translated in transcript coordinates. The first option can be easily handled by many aligners (e.g. Bowtie), given a reference FASTA file where each sequence represents a transcript, from the beginning of the 5' UTR to the end of the 3' UTR. The second procedure is based on reference FASTA files where each sequence represents a chromosome, usually coupled with comprehensive gene annotation files (GTF or GFF). The STAR aligner, with its option –quantMode TranscriptomeSAM (see Chapter 6 of its manual), is an example of tool providing such a feature.

indel_threshold is aimed at keeping only reads showing small differences between the width of the RNA fragment and the length of the reference sequence region covered after their alignment (for additional details please read here about *qwidth* and *width*, defined in the GAlignments-class of the GenomicAlignments package). Strong differences between these values are usually due to many indels which in turn are caused by loose thresholds used in the alignment process. High numbers of consecutive deletions (nucleotides present in the reference sequence but not in the read) may be an indication of reads mapping on exon-exon junctions. Despite this eventuality is not in accordance with alignments based on transcript coordinates, stretches of deletions can be included by errors or inaccurate transcript annotation and should be discarded. In fact, **riboWaltz** deliberately identifies the length of the reference sequences covered by the reads with the reads length, referring to it as *length* in all data structures reporting reads or P-site offsets information (i.e. those generated by bamtolist itself, bamtobed, bedtolist, psite and psite_info). This value is indeed strongly connected to the identification of the P-site offsets since it determines the position of the 5' and 3' read extremities (columns *end5* and *end3* in many data structures generated by the package), used as starting points by **riboWaltz**'s core algorithm.

refseq_sep is intended to lighten the identifiers of the reference sequences included in the output list of data table or to modify them to match those in the annotation table. Many details about the reference sequence such as their version (usually dot-separated), their length, name variants, associated gene/transcript/protein names (usually pipe-separated) might indeed be stored in the FASTA file used for the alignment and automatically transferred in the BAM.

Value

A list of data tables or a GRangesList object.

```
## ## Let's suppose there are two BAM files ("Samp1.bam" and "Samp2.bam") in
```

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```
## ## "path/to/BAM/files". We want to acquire them and assign to the
## ## corresponding data tables the names "Control" and "Treated",
## ## respectively.
## ## We first define the "name_samples" character string vector as follow:
## name_of_bams <- c("Control", "Treated")
## names(name_of_bams) <- c("Samp1", "Samp2")
##
## Then, we can acquire the two files:
## path_bam <- "path/to/BAM/files"
## reads_list <- bamtolist(bamfolder = path_bam, name_samples = name_of_bams,
## annotation = annotation_dt)
##
## read_list will be a list of two data tables, named "Control" (with
## ## mapping reads from "Samp1.bam") and "Treated" (with mapping reads
## ## from"Samp2.bam")</pre>
```

bedtolist

From BED files to lists of data tables or GRangesList objects.

Description

This function reads one or multiple BED files, as generated by bamtobed, converting them into data tables or GRanges objects, arranged in a list or a GRangesList, respectively. In both cases two columns are attached to the original data containing, for each read, the leftmost and rightmost position of the annotated CDS of the reference sequence (if any) with respect to its 1st nucleotide. Please note: start and stop codon positions for transcripts without annotated CDS are set to 0.

Usage

```
bedtolist(
  bedfolder,
  annotation,
  transcript_align = TRUE,
  name_samples = NULL,
  refseq_sep = FALSE,
  output_class = "datatable"
)
```

Arguments

bedfolder

Character string specifying the path to the folder storing BED files as generated by bamtobed.

annotation

Data table as generated by create_annotation. Please make sure the name of reference transcripts in the annotation data table match those in the BED files (see also refseq_sep).

transcript_align

Logical value whether BED files in bedfolder come from a transcriptome alignment (intended as an alignment against reference transcript sequences, see Details). If TRUE (the default), reads mapping on the negative strand should not be present and, if any, they are automatically removed.

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name_samples

Named character string vector specifying the desired name for the output list elements. A character string for each BED file in bedfolder is required. Plase be careful to name each element of the vector after the correct corresponding BED file in bedfolder, leaving their path and extension out. No specific order is required. Default is NULL i.e. list elements are named after the name of the BED files, leaving their path and extension out.

refseq_sep

Character specifying the separator between reference sequences' name and additional information to discard, stored in the same field (see Details). All characters before the first occurrence of the specified separator are kept. Default is NULL i.e. no string splitting is performed.

Either "datatable" or "granges". It specifies the format of the output i.e. a list of data tables or a GRangesList object. Default is "datatable".

Details

riboWaltz only works for read alignments based on transcript coordinates. This choice is due to the main purpose of RiboSeq assays to study translational events through the isolation and sequencing of ribosome protected fragments. Most reads from RiboSeq are supposed to map on mRNAs and not on introns and intergenic regions. Nevertheless, BAM based on transcript coordinates can be generated in two ways: i) aligning directly against transcript sequences; ii) aligning against standard chromosome sequences, requiring the outputs to be translated in transcript coordinates. The first option can be easily handled by many aligners (e.g. Bowtie), given a reference FASTA file where each sequence represents a transcript, from the beginning of the 5' UTR to the end of the 3' UTR. The second procedure is based on reference FASTA files where each sequence represents a chromosome, usually coupled with comprehensive gene annotation files (GTF or GFF). The STAR aligner, with its option —quantMode TranscriptomeSAM (see Chapter 6 of its manual), is an example of tool providing such a feature.

refseq_sep is intended to lighten the identifiers of the reference sequences included in the final data table or to modify them to match those in the annotation table. Many details about the reference sequence such as their version (usually dot-separated), their length, name variants, associated gene/transcript/protein names (usually pipe-separated) might indeed be stored in the FASTA file used for the alignment and automatically transferred in the BAM.

Value

A list of data tables or a GRangesList object.

Examples

```
## path_bed <- "path/to/BED/files"
## bedtolist(bedfolder = path_bed, annotation = mm81cdna)</pre>
```

cds_coverage

Number of in-frame P-sites per coding sequence or transcript.

Description

This function generates a data table that for each transcript contains at least i) its name; ii) its length or the length of its annotated coding sequence (if any); iii) the number of P-sites falling in its annotated coding sequence or in the whole transcript for all samples. When the coverage is computed for coding sequences and not whole transcripts, a chosen number of nucleotides at the

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beginning and/or at the end of the CDSs can be excluded for restricting the analysis to a portion of the original coding sequence. In this case the output data table includes an additional column reporting the length of the selected region. Moreover, either all P-sites or only in-frame P-sites can be considered. Finally, transcripts without annotated CDS are automatically discarded.

Usage

```
cds_coverage(
  data,
  annotation,
  start_nts = 0,
  stop_nts = 0,
  in_frame = TRUE,
  whole_transcript = FALSE
)
```

Arguments

data Either list of data tables or GRangesList object from psite_info.

annotation Data table as generated by create_annotation.

start_nts Positive integer specifying the number of nucleotides at the beginning of the

coding sequences to be excluded from the analisys. Default is 0. This parameter

is considered only if whole_transcript is FALSE.

stop_nts Positive integer specifying the number of nucleotides at the end of the coding

sequences to be excluded from the analisys. Default is 0. This parameter is

considered only if whole_transcript is FALSE.

in_frame Logical value whether to consider only in-frame P-sites when computing the

coverage. Default is TRUE. This parameter is considered only if whole_transcript

is FALSE.

whole_transcript

Logical value whether to compute the coverage based on P-sites falling in any region of the transcript, i.e. CDS and UTRs. Default is FALSE.

Value

A data table.

```
## data(reads_list)
## data(mm81cdna)
##
## ## compute and add p-site datails
## psite_offset <- psite(reads_list, flanking = 6, extremity = "auto")
## reads_psite_list <- psite_info(reads_list, psite_offset)
##
## Compute the number of in-frame P-sites per whole coding sequences.
## psite_cds <- cds_coverage(reads_psite_list, mm81cdna)
##
## Compute the number of in-frame P-sites per coding sequences exluding
## ## Compute the number of in-frame P-sites per coding sequences exluding
## ## the first 15 nucleotides and the last 10 nucleotides.
## psite_cds <- cds_coverage(reads_psite_list, mm81cdna,
## start_nts = 15, stop_nts = 10)</pre>
```

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```
##
## ## Compute the number of P-sites per transcripts.
## psite_cds <- cds_coverage(reads_psite_list, mm81cdna,
## whole_transcript = TRUE)</pre>
```

codon_coverage

Number of reads per codon.

Description

This function computes transcript-specific codon coverages, defined as the number of either read footprints or P-sites mapping on each triplet of coding sequences and UTRs (see *Details*). The resulting data table contains, for each triplet: i) the name of the corresponding reference sequence (i.e. of the transcript to which it belongs); ii) its leftmost and rightmost position with respect to the 1st nucleotide of the reference sequence; iii) its position with respect to the 1st and the last codon of the annotated CDS of the reference sequence; iv) the region of the transcript (5' UTR, CDS, 3' UTR) it is in; v) the number of read footprints or P-sites falling in that region for all samples.

Usage

```
codon_coverage(
  data,
  annotation,
  sample = NULL,
  psite = FALSE,
  min_overlap = 1,
  output_class = "datatable"
)
```

Arguments

data	Either list of data tables or GRangesList object from psite_info. Data tables and GRanges objects generated by bamtolist and bedtolist can be used if psite is FALSE (the default).
annotation	Data table as generated by create_annotation.
sample	Character string vector specifying the name of the sample(s) of interest. Default is NULL i.e. all samples in data are processed.
psite	Logical value whether to return the number of P-sites per codon. Default is TRUE. If FALSE, the number of read footprints per codon is returned instead.
min_overlap	Positive integer specifying the minimum number of overlapping positions (in nucleotides) between reads and codons to be considered overlapping. If psite is TRUE this parameter must be 1 (the default).
output_class	Either "datatable" or "granges". It specifies the format of the output i.e. a list of data tables or a GRangesList object. Default is "datatable".

Details

The sequence of every transcript is divided in triplets starting from the annotated translation initiation site (if any) and proceeding towards the UTRs extremities, possibly discarding the exceeding 1 or 2 nucleotides at the extremities of the transcript. Please note: transcripts not associated to any annotated 5' UTR, CDS and 3'UTR and transcripts whose coding sequence length is not divisible by 3 are automatically discarded.

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Value

A data table or GRanges object.

Examples

```
## data(reads_list)
## data(mm81cdna)
##
## ## compute and add p-site datails
## psite_offset <- psite(reads_list, flanking = 6, extremity = "auto")
## reads_psite_list <- psite_info(reads_list, psite_offset)
##
## Compute the codon coverage based on the number of ribosome footprint per
## codon, setting the minimum overlap between reads and triplets to 3 nts:
## coverage_dt <- codon_coverage(reads_psite_list, mm81cdna, min_overlap = 3)
##
## Compute the coverage based on the number of P-sites per codon:
## coverage_dt <- codon_coverage(reads_psite_list, mm81cdna, psite = TRUE)</pre>
```

codon_usage_psite

Empirical codon usage indexes.

Description

This function computes empirical codon usage indexes based on either ribosome P-sites, A-site or E-site frequency associated to in-frame P-sites along coding sequences. Given one sample, it computes 64 codon usage indexes (one for each triplet, optionally normalized for their frequency in CDSs) and generates a bar plot of the resulting values. If two samples are specified, the function also compares the two sets of codon usage indexes returning a scatter plot. The same output is generated specifying one sample and providing 64 triplet-specific values. Scatter plots report the result of linear regressions between the two sets of values and the corresponding Pearson correlation coefficient.

Usage

```
codon_usage_psite(
  data,
  annotation,
  sample,
  site = "psite",
  fastapath = NULL,
  fasta_genome = TRUE,
 refseq_sep = NULL,
 bsgenome = NULL,
 gtfpath = NULL,
  txdb = NULL,
 dataSource = NA,
  organism = NA,
  transcripts = NULL,
  frequency_normalization = TRUE,
  codon_values = NULL,
  label_scatter = FALSE,
```

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```
label_number = 64,
label_aminoacid = FALSE
)
```

Arguments

data

Either list of data tables or GRangesList object from $psite_info$. Each data table may or may not include one or more columns among p_site_codon , a_site_codon and e_site_codon reporting the three nucleotides covered by the P-site, A-site and E-site, respectively. These columns can be previously generated by the $psite_info$ function. Otherwise, the column of interest can be specified by $psite_info$ function and it is automatically generated starting from a FASTA file or a BS genome data package.

annotation

Data table as generated by create_annotation.

sample

Either character string or character string vector specifying one or two sample names, respectively. If one sample name is specified, a bar plot displaying the 64 codon usage indexes is generated. If two sample names are specified, the function also compares the two sets of codon usage indexes returning a scatter plot. In the latter case it also performes a linear regression and computes the Pearson correlation coefficient.

site

Either "psite, "asite", "esite". It specifies if the empirical codon usage indexes should be based on ribosome P-sites ("psite"), A-sites ("asite") or E-sites ("esite"). Default is "psite".

fastapath

Character string specifying the FASTA file used in the alignment step, including its path, name and extension. This file can contain reference nucleotide sequences either of a genome assembly or of all the transcripts (see Details and fasta_genome). Please make sure the sequences derive from the same release of the annotation file used in the create_annotation function. Note: either fastapath or bsgenome is required to compute the frequency in sequences of each codon, used as normalization factors, even if data includes one or more columns among p_site_codon , a_site_codon and e_site_codon . Default is NULL.

fasta_genome

Logical value whether the FASTA file specified by fastapath contains nucleotide sequences of a genome assembly. If TRUE (the default), an annotation object is required (see gtfpath and txdb). FALSE implies nucleotide sequences of transcripts are provided instead.

refseq_sep

Character specifying the separator between reference sequences' name and additional information to discard, stored in the headers of the FASTA file specified by fastapath (if any). It might be required for matching the reference sequences' identifiers reported in the input list of data tables. All characters before the first occurrence of the specified separator are kept. Default is NULL i.e. no string splitting is performed.

bsgenome

Character string specifying the BSgenome data package with the genome sequences to be loaded. If not already present in the system, it is automatically installed through the biocLite.R script (check the list of available BSgenome data packages by running the available.genomes function of the BSgenome package). This parameter must be coupled with an annotation object (see gtfpath and txdb). Please make sure the sequences included in the specified BSgenome data pakage are in agreement with the sequences used in the alignment step. Note: either fastapath or bsgenome is required to compute the frequency in sequences of each codon, used as normalization factors, even if data includes

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one or more columns among p_site_codon , a_site_codon and e_site_codon . Default is NULL.

gtfpath

Character string specifying the location of a GTF file, including its path, name and extension. Please make sure the GTF file and the sequences specified by fastapath or bsgenome derive from the same release. Note that either gtfpath or txdb is required if and only if nucleotide sequences of a genome assembly are provided (see fastapath or bsgenome). Default is NULL.

txdb

Character string specifying the TxDb annotation package to be loaded. If not already present in the system, it is automatically installed through the biocLite.R script (check here the list of available TxDb annotation packages). Please make sure the TxDb annotation package and the sequences specified by fastapath or bsgenome derive from the same release. Note that either gtfpath or txdb is required if and only if nucleotide sequences of a genome assembly are provided (see fastapath or bsgenome). Default is NULL.

dataSource

Optional character string describing the origin of the GTF data file. This parameter is considered only if gtfpath is specified. For more information about this parameter please refer to the description of *dataSource* of the makeTxDbFromGFF function included in the GenomicFeatures package.

organism

Optional character string reporting the genus and species of the organism of the GTF data file. This parameter is considered only if gtfpath is specified. For more information about this parameter please refer to the description of *organism* of the makeTxDbFromGFF function included in the GenomicFeatures package.

transcripts

Character string vector listing the name of transcripts to be included in the analysis. Default is NULL i.e. all transcripts are used. Please note: transcripts without annotated CDS and transcripts whose coding sequence length is not divisible by 3 are automatically discarded.

frequency_normalization

Logical value whether to normalize the 64 codon usage indexes for the corresponding codon frequencies in coding sequences. Default is TRUE.

codon_values

Data table containing 64 triplet-specific values. If specified, the provided values are compared with the empirical codon usage indexes computed for the sample of interest. The data table must contain the DNA or RNA nucleotide sequence of the 64 codons and corresponding values arranged in two columns named *codon* and *value*, respectively. Default is NULL.

label_scatter

Logical value whether to label the dots in the scatter plot. Each dot is labeled using either the nucleotide sequence of the codon or the corresponding amino acid symbol (see label_aminoacid). This parameter is considered only if two sample names are specified in sample or codon_values is provided. Default is FALSE.

label_number

Integer value in [1,64] specifying how many dots in the scatter plot should be labeled. Dots farthest from the confident interval of the regression line are automatically identified and labeled. Default is 64 i.e. all dots are labeled. This parameter is considered only if label_scatter is TRUE.

label_aminoacid

Logical value whether to use amino acid symbols to label the dots of the scatter. Default is FALSE i.e. codon nucleotide sequences are used instead. This parameter is considered only if label_scatter is TRUE.

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Details

riboWaltz only works for read alignments based on transcript coordinates. This choice is due to the main purpose of RiboSeq assays to study translational events through the isolation and sequencing of ribosome protected fragments. Most reads from RiboSeq are supposed to map on mRNAs and not on introns and intergenic regions. Nevertheless, BAM based on transcript coordinates can be generated in two ways: i) aligning directly against transcript sequences; ii) aligning against standard chromosome sequences, requiring the outputs to be translated in transcript coordinates. The first option can be easily handled by many aligners (e.g. Bowtie), given a reference FASTA file where each sequence represents a transcript, from the beginning of the 5' UTR to the end of the 3' UTR. The second procedure is based on reference FASTA files where each sequence represents a chromosome, usually coupled with comprehensive gene annotation files (GTF or GFF). The STAR aligner, with its option –quantMode TranscriptomeSAM (see Chapter 6 of its manual), is an example of tool providing such a feature.

Value

A list containing ggplot2 objects and a data table with the associated data ("dt"). If only one sample name is specified in sample, one ggplot2 object is returned (a bar plot named "plot"). If two sample names are specified in sample, three ggplot2 objects are returned (two bar plots named "plot_NameSample1" and "plot_NameSample2" and a scatter plot named "plot_comparison"). If codon_values is specified, two ggplot2 objects are returned (one bar plots named "plot" and a scatter plot named "plot_comparison"). Please note: before plotting, the 64 values are scaled to make them ranging between 0 and 1. If frequency_normalization is TRUE, the data table contains raw, normalized and scaled codon usage indexes, only raw and scaled data otherwise.

Examples

```
## ## compute and add the p-site datails
## psite_offset <- psite(reads_list, flanking = 6, extremity = "auto")</pre>
## reads_psite_list <- psite_info(reads_list, psite_offset)</pre>
##
## ## codon usage from transcriptome alignment
## path_fasta <- "path/to/transcriptome/FASTA/file"</pre>
## codon_usage_psite(data = reads_psite_list, annotation = mm81cdna,
##
                      sample = "Samp1",
##
                      fastapath = path_fasta, fasta_genome = FALSE)
## ## codon usage from genome alignment
## path_fasta <- "path/to/genome/FASTA/file"</pre>
## codon_usage_psite(data = reads_psite_list, annotation = mm81cdna,
                      sample = "Samp1",
##
##
                      fastapath = path_fasta, fasta_genome = TRUE)
```

create_annotation

Annotation data table.

Description

This function generates transcript basic annotation data tables starting from GTF files or TxDb objects. Annotation data tables include a column named transcript reporting the name of the reference transcripts and four columns named l_tr , l_utr5 , l_cds and l_utr3 reporting the length of the transcripts and of their annotated 5' UTRs, CDSs and 3' UTRs, respectively. Please note: if a transcript region is not annotated its length is set to 0.

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Usage

```
create_annotation(gtfpath = NULL, txdb = NULL, dataSource = NA, organism = NA)
```

Arguments

gtfpath A character string specifying the path to a GTF file, including its name and

extension. Please make sure the GTF file derives from the same release of the sequences used in the alignment step. Note that either gtfpath or txdb must be

specified. Default is NULL.

txdb Character string specifying the TxDb annotation package to be loaded. If not al-

ready present in the system, it is automatically installed through the biocLite.R script (check here the list of available TxDb annotation packages). Please make sure the TxDb annotation package derives from the same release of the sequences used in the alignment step. Note that either gtfpath or txdb must

be specified. Default is NULL.

dataSource Optional character string describing the origin of the GTF data file. This param-

eter is considered only if gtfpath is specified. For more information about this parameter please refer to the description of dataSource of the makeTxDbFromGFF

function included in the GenomicFeatures package.

organism Optional character string reporting the genus and species of the organism of

the GTF data file. This parameter is considered only if gtfpath is specified. For more information about this parameter please refer to the description of organism of the makeTxDbFromGFF function included in the GenomicFeatures

package.

Value

A data table.

Examples

```
## gtf_file <- "path/to/GTF/file.GTF"
## create_annotation(gtfpath = gtf_file, dataSource = "gencode6", organism = "Mus musculus")</pre>
```

duplicates_filter

Duplicates filtering.

Description

This function provides multiple options for remove duplicated reads: when two or more reads are marked as duplicates, all of them are discarded but one.

Usage

```
duplicates_filter(
  data,
  sample = NULL,
  extremity = "both",
  keep = "shortest",
  output_class = "datatable",
```

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```
txt = FALSE,
  txt_file = NULL
)
```

Arguments

data Either list of data tables or GRangesList object from bamtolist, bedtolist or

length_filter.

sample Character string or character string vector specifying the name of the sample(s)

to process. Default is NULL i.e. all samples are processed.

extremity Either "both", "5end", "3end". It specifies the criterion to define which reads

should be considered duplicates. Reads are marked as duplicates if they map on the same transcript and share: both the 5' estremity and the 3' extremity ("both"), only the 5' extremity ("5end"), only the same 3' extremity ("3end"). For "5end" and "3end", reads of different lengths can be marked as duplicates.

See keep to choose which one should be kept.

keep Either "shortest" or "longest". It specifies wheter to keep the shortest or the

longest read when duplicates display different lengths. This parameter is considered only if extremity is set to "5end" or "3end". Default is "shortest".

output_class Either "datatable" or "granges". It specifies the format of the output i.e. a list of

data tables or a GRangesList object. Default is "datatable".

txt Logical value whether to write in a txt file statistics on the filtering step. Similar

information are displayed by default in the console. Default is FALSE.

txt_file Character string specifying the path, name and extension (e.g. "PATH/NAME.extension")

of the plain text file where statistics on the filtering step should be written. If the specified folder doesn't exist, it is automatically created. If NULL (the default), the information are written in "duplicates_filtering.txt", saved in the working

directory. This parameter is considered only if txt is TRUE.

Value

A list of data tables or a GRangesList object.

```
#generate an \emph{ad hoc} dataset:
library(data.table)
dt <- data.table(transcript = rep("ENSMUST0000000001.4", 6),</pre>
                  end5 = c(92, 92, 92, 94, 94, 95),
                  end3 = c(119, 119, 122, 122, 123, 123)
                  )[, length := end3 - end5 + 1
                   ][, cds_start := 14
                    ][, cds_stop := 1206]
example_reads_list <- list()</pre>
example_reads_list[["Samp_example"]] <- dt</pre>
## Reads are duplicates if they share both the 5' estremity and the
## 3' extremity:
filtered_list <- duplicates_filter(example_reads_list,</pre>
                                    extremity = "both")
## Reads are duplicates if they only share the 5' estremity. Among duplicated
## reads we keep the shortes one:
```

frame_psite 15

frame_psite

Percentage of P-sites per reading frame.

Description

This function computes the percentage of P-sites falling in the three possible translation reading frames and generates a bar plot of the resulting values. It only handles annotated 5' UTRs, coding sequences and 3' UTRs, separately.

Usage

```
frame_psite(
  data,
  sample = NULL,
  transcripts = NULL,
  region = "all",
  length_range = "all",
  plot_title = NULL
)
```

Arguments

data	Either list of data tables or GRangesList object from psite_info.
sample	Character string vector specifying the name of the sample(s) of interest. Default is NULL i.e. all samples in data are processed.
transcripts	Character string vector listing the name of transcripts to be included in the analysis. Default is NULL i.e. all transcripts are used.
region	Character string specifying the region(s) of the transcripts to be analysed. It can be either "5utr", "cds", "3utr" for 5' UTRs, CDSs and 3' UTRs, respectively. Default is "all" i.e. all regions are considered. According to this parameter the bar plots are differently arranged to optimise the organization and the visualization of the data.
length_range	Integer or an integer vector specyfying the read length(s) to be included in the analysis. Default is "all" i.e. all read lengths are used.
plot_title	Character string specifying the title of the plot. If "auto", the title of the plot reports the region specified by region (if any) and the considered read length(s). Default is NULL i.e. no title is plotted.

Value

A list containing a ggplot2 object ("plot") and the data table with the associated data ("dt").

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Examples

```
## data(reads_list)
##
## ## compute and add p-site datails
## psite_offset <- psite(reads_list, flanking = 6, extremity = "auto")
## reads_psite_list <- psite_info(reads_list, psite_offset)
##
## Generate the bar plot for all read lengths:
## frame_whole <- frame_psite(reads_psite_list, sample = "Samp1")
##
## ## Generate the bar plot restricting the analysis to coding sequences and
## ## reads of 28 nucleotides:
## frame_sub <- frame_psite(reads_psite_list, sample = "Samp1",
##
## region = "cds", length_range = 28)</pre>
```

frame_psite_length

Percentage of P-sites per reading frame stratified by read length.

Description

Similar to frame_psite, but the results are stratified by read lengths and plotted as heatmaps.

Usage

```
frame_psite_length(
  data,
  sample = NULL,
  transcripts = NULL,
  region = "all",
  cl = 95,
  length_range = "all",
  plot_title = NULL,
  colour = "#061b63"
)
```

Arguments

data	Either list of data	tables or GRange	sList object from	psite info.

sample Character string vector specifying the name of the sample(s) of interest. Default

is NULL i.e. all samples in data are processed.

transcripts Character string vector listing the name of transcripts to be included in the anal-

ysis. Default is NULL i.e. all transcripts are used.

region Character string specifying the region(s) of the transcripts to be analysed. It can

be either "5utr", "cds", "3utr" for 5' UTRs, CDSs and 3' UTRs, respectively. Default is "all" i.e. all regions are considered. According to this parameter the heatmaps are differently arranged to optimise the organization and the visual-

ization of the data.

cl Integer value in [1,100] specifying a confidence level for restricting the plot to

a sub-range of read lengths. The new range is associated to the most abundant populations of reads accounting for the cl of the sample. Default is 95. This

parameter has no effect if length_range is specified.

length_filter 17

length_range	Integer or an integer vector specyfying the read length(s) to be included in the analysis. Default is "all" i.e. all read lengths are used. If specified, this parameter prevails over c1.
plot_title	Character string specifying the title of the plot. If "auto", the title of the plot reports the region specified by region (if any) and the considered read length(s). Default is NULL i.e. no title is displayed.
colour	Character string specifying the colour of the plot. The colour scheme is as follow: tiles corresponding to the lowest signal are always white, tiles corresponding to the highest signal are of the specified colour and the progression between these two colours follows a linear gradient. Default is dark blue.

Value

A list containing a ggplot2 object ("plot") and the data table with the associated data ("dt").

Examples

```
## data(reads_list)
##
## ## compute and add p-site datails
## psite_offset <- psite(reads_list, flanking = 6, extremity = "auto")
## reads_psite_list <- psite_info(reads_list, psite_offset)
##
## Generate the heatmap for all read lengths:
## frame_len_whole <- frame_psite_length(reads_psite_list, sample = "Samp1")
##
## ## Generate the heatmap restricting the analysis to coding sequences and a
## ## sub-range of read lengths:
## frame_len_sub <- frame_psite_length(reads_psite_list, sample = "Samp1",
## region = "cds", cl = 90)</pre>
```

length_filter

Read length filtering.

Description

This function provides multiple options for filtering the reads according to their length. Read lengths to keep are either specified by the user or automatichally selected on the basis of the trinucleotide periodicity of reads mapping on the CDS.

Usage

```
length_filter(
  data,
  sample = NULL,
  length_filter_mode = "periodicity",
  periodicity_threshold = 50,
  length_range = NULL,
  output_class = "datatable",
  txt = FALSE,
  txt_file = NULL
)
```

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Arguments

data Either list of data tables or GRangesList object from bamtolist, bedtolist or

duplicates_filter.

sample Character string or character string vector specifying the name of the sample(s)

to process. Default is NULL i.e. all samples are processed.

length_filter_mode

Either "periodicity" or "custom". It specifies how read length selection should be performed. "periodicity": only read lengths satisfying a periodicity threshold (see periodicity_threshold) are kept. It ensures the removal of all reads with low or no periodicity; "custom": only read lengths specified by the user are kept (see length_range). Default is "periodicity".

periodicity_threshold

Integer in [10, 100]. Only read lengths satisfying this threshold (i.e. a higher percentage of read extremities falls in one of the three reading frames along the CDS) are kept. This parameter is considered only if length_filter_mode is

set to "periodicity". Default is 50.

length_range Integer or integer vector specifying one read length or a range of read lengths to

keep, respectively. This parameter is considered only if length_filter_mode

is set to "custom".

output_class Either "datatable" or "granges". It specifies the format of the output i.e. a list of

data tables or a GRangesList object. Default is "datatable".

txt Logical value whether to write in a txt file statistics on the filtering step. Similar

information are displayed by default in the console. Default is FALSE.

txt_file Character string specifying the path, name and extension (e.g. "PATH/NAME.extension")

of the plain text file where statistics on the filtering step should be written. If the specified folder doesn't exist, it is automatically created. If NULL (the default), the information are written in "length_filtering.txt", saved in the working

directory. This parameter is considered only if txt is TRUE.

Value

A list of data tables or a GRangesList object.

metaheatmap_psite 19

metaheatmap_psite

Ribosome occupancy metaheatmaps at single-nucleotide resolution.

Description

This function generates two heatmap-like metaprofiles (metaheatmaps) displaying the abundance of P-sites around the start and the stop codon of annotated CDSs. It works similarly to metaprofile_psite but the intensity of signal is represented by a continuous color scale rather than by the height of a line chart. This graphical output is a good option to visualize several profiles at once and compare results obtained with different read lengths or in multiple conditions.

Usage

```
metaheatmap_psite(
  data,
  annotation,
  sample,
  scale_factors = NULL,
  length_range = "all",
  transcripts = NULL,
  utr51 = 25,
  cds1 = 50,
  utr31 = 25,
  log_colour = F,
  colour = "black",
  plot_title = NULL
)
```

Arguments

data Either list of data tables or GRangesList object from psite_info.

annotation Data table as generated by create_annotation.

sample List of either character strings specifying the name of the sample(s) of inter-

est or character string vectors specifying the name of their replicates. In the latter case the final metaheatmap for each element of the list is generated by merging the corresponding replicates exploiting the scale factors specified by scale_factors. The row(s) of the final plot are labelled according to the name

of the elements of the list.

scale_factors Named numeric vector specifying the scale factors for generating metaprofiles

from multiple replicates (see sample). Scale factors can be defined for a subset of list elements of sample i.e. for all replicates of selected samples. If so, the remaining scale factors are set automatically to 1. Please be careful to name each element of the vector after the correct corresponding string in sample. No specific order is required. Default is NULL i.e. all scale factors are automatically

set to 1.

analysis. Default is "all" i.e. all read lengths are used.

transcripts Character string vector listing the name of transcripts to be included in the anal-

ysis. Default is NULL i.e. all transcripts are used. Please note: transcripts

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	with either 5' UTR, coding sequence or 3' UTR shorter than utr51, 2*cds1 and utr31, respectively, are automatically discarded.
utr5l	Positive integer specifying the length (in nucleotides) of the 5' UTR region flanking the start codon to be considered in the analysis. Default is 25.
cdsl	Positive integer specifying the length (in nucleotides) of the CDS regions flanking both the start and stop codon to be considered in the analysis. Default is 50.
utr3l	Positive integer specifying the length (in nucleotides) of the 3' UTR region flanking the stop codon to be considered in the analysis. Default is 25.
log_colour	Logical value whether to use a logarithmic colour scale (strongly suggested in case of large signal variations). Default is FALSE.
colour	Character string specifying the colour of the plot. The colour scheme is as follow: tiles corresponding to the lowest signal are always white, tiles corresponding to the highest signal are of the specified colour and the progression between these two colours follows either linear or logarithmic gradients (see log_colour). Default is "black".
plot_title	Character string specifying the title of the plot. If "auto", the title of the plot reports the number of transcripts and the read length(s) employed for generating the metaprofiles. Default is NULL i.e. no title is displayed.

Details

The intensity of signal in the metaprofiles corresponds, for each nucleotide, to the sum of the number of P-sites (defined by their leftmost position) mapping on that position for all transcripts in one or multiple replicates.

Value

A list containing a ggplot2 object ("plot") and the data table with the associated data ("dt").

```
## data(reads_list)
## data(mm81cdna)
##
## ## compute and add p-site datails
## psite_offset <- psite(reads_list, flanking = 6, extremity = "auto")</pre>
## reads_psite_list <- psite_info(reads_list, psite_offset)</pre>
## ## Generate metaheatmaps employing all read lengths:
## metaheat_whole <- metaheatmap_psite(reads_psite_list, mm81cdna,</pre>
##
                                         sample = list("Whole"=c("Samp1")))
##
\#\# \#\# Generate metaprofiles employing reads of 27, 28 and 29 nucleotides and
## ## a subset of transcripts (in this example only transcripts with at least
## ## one P-site mapping on the translation initiation site are kept):
## sample_name <- "Samp1"</pre>
## sub_reads_psite_list <- reads_psite_list[[sample_name]][psite_from_start == 0]</pre>
## transcript_names <- as.character(sub_reads_psite_list$transcript)</pre>
## metaheat_sub <- metaheatmap_psite(reads_psite_list, mm81cdna,</pre>
                                       sample = list("sub"=sample_name),
## length_range = 27:29, transcripts = transcript_names, plot_title = "auto")
```

metaprofile_psite 21

metaprofile_psite

Ribosome occupancy metaprofiles at single-nucleotide resolution.

Description

This function generates metaprofiles displaying the abundance of P-sites around the start and the stop codon of annotated CDSs. For one sample the intensity of the signal in the metaprofiles corresponds, for each nucleotide, to the sum of the number of P-sites (defined by their leftmost position) mapping on that position for all transcripts. Multiple samples can be handled in several ways.

Usage

```
metaprofile_psite(
  data,
  annotation,
  sample.
  multisamples = "separated",
  plot_style = "split",
  scale_factors = NULL,
  length_range = "all",
  transcripts = NULL,
  frequency = FALSE,
  utr51 = 25,
  cdsl = 50,
  utr31 = 25,
  colour = NULL,
  plot_title = NULL
)
```

Arguments

data Either list of data tables or GRangesList object from psite_info.

annotation Data table as generated by create_annotation.

sample

Either character string or character string vector specifying the name of the sample(s) of interest. A named list of one or more character strings and/or character string vectors can be provided. In this case i) each list element should include the name of the replicate(s) related to the sample of interest and ii) the name assigned to the elements of the list are displayed in the plot. Multiple replicates specified in character string vectors are handled according to multisample.

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multisamples

Either "separated", "average" or "sum". It specifies how to handle multiple samples and replicates. If "saparated", one metaprofile for each sample included in sample is returned as an independent ggplot object. If sample is a list, it is unlisted, coerced to character string and handled accordingly. If "average" or "sum" i) one metaprofiles is returned if sample is a character string vector or ii) one metaprofiles is built for each element of sample when it is a list. If "average", the metaprofiles display for each nucleotide the mean signal and the corresponding standard error computed among the replicates. If "sum", the metaprofiles display for each nucleotide the sum of the signal of the replicates. In both cases a single ggplot object is returned, where multiple metaprofiles are organized and displayed according to plot_style. Default is "separated".

plot_style

Either "split", "ovelaid" or "mirrored". It specifies how to organize and display multiple metaprofiles. If "split", the metaprofiles are placed one below the other, in independent boxes. If "overlaid", all metaprofiles are superimposed. If "mirrored" sample must be a list of exactly two elements and the two metaprofiles are mirrored along the x axis. Default is "split".

scale_factors

Named numeric vector the same length as sample specifying the scale factors for generating metaprofiles from multiple replicates (see sample). Please be careful to name each element of the vector after the correct corresponding string in sample. No specific order is required. When frequency is TRUE, this parameter is not considered. Default is NULL i.e. all scale factors are automatically set to 1.

length_range

Integer or integer vector specyfying the read length(s) to be included in the analysis. Default is "all" i.e. all read lengths are used.

transcripts

Character string vector listing the name of transcripts to be included in the analysis. Default is NULL i.e. all transcripts are used. Please note: transcripts with either 5' UTR, coding sequence or 3' UTR shorter than utr51, 2*cds1 and utr31, respectively, are automatically discarded.

frequency

Logical value whether to normalize the metaprofile(s) such that the area under the curve(s) is 1. If TRUE and multisamples is set to "average" or "sum", the normalization is performed before combining the signal from multiple samples. Default is FALSE.

utr5l

Positive integer specifying the length (in nucleotides) of the 5' UTR region flanking the start codon to be considered in the analysis. Default is 25.

cdsl

Positive integer specifying the length (in nucleotides) of the CDS regions flanking both the start and stop codon to be considered in the analysis. Default is 50.

utr31

Positive integer specifying the length (in nucleotides) of the 3' UTR region flanking the stop codon to be considered in the analysis. Default is 25.

colour

Character string or character string vector specifying the colour of the metaprofile(s). If sample is a list of multiple elements and multisamples is set to "average" or "sum", a colour for each element of the list is required. If this parameter is not specified the R default palette is employed. Default is NULL.

plot_title

Character string specifying the title of the plot. It can be any string provided by the user or "sample", "transcript" and "length_range" for automatically displaying the name of the sample(s) specified by sample, the number of transcripts and the most frequent read lengths (i.e. associated to 90 the metaprofiles, respectively. A combination of the three strings, dot-separated, can be used for displaying multiple information. For example, specifying "sample.length_range" the title reports both the name of the sample(s) and the read length(s). Default is NULL i.e. no title is displayed.

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Value

A list containing one or more ggplot2 object(s) and the data table with the associated data ("dt").

Examples

```
## data(reads_list)
## data(mm81cdna)
##
## ## compute and add p-site datails
## psite_offset <- psite(reads_list, flanking = 6, extremity = "auto")</pre>
## reads_psite_list <- psite_info(reads_list, psite_offset)</pre>
##
## ## Generate metaprofiles employing all read lengths:
## metaprof_whole <- metaprofile_psite(reads_psite_list, mm81cdna,</pre>
                                         sample = "Samp1")
##
## metaprof_whole[["plot_Samp1"]]
##
## ## Generate metaprofiles employing reads of 27, 28 and 29 nucleotides and
## ## a subset of transcripts (in this example only transcripts with at least
## ## one P-site mapping on the translation initiation site are kept):
## sample_name <- "Samp1"
## sub_reads_psite_list <- reads_psite_list[[sample_name]][psite_from_start == 0]</pre>
## transcript_names <- as.character(sub_reads_psite_list$transcript)</pre>
## metaprof_sub <- metaprofile_psite(reads_psite_list, mm81cdna,</pre>
                                       sample = sample_name,
##
##
                                       length_range = 27:29,
##
                                       transcripts = transcript_names)
```

mm81cdna

Annotation data table

Description

A dataset containing basic information about 25,892 mouse mRNAs (Ensembl v81 transcript annotation).

Usage

mm81cdna

Format

A data table with 25,892 rows and 5 variables:

transcript Name of the transcript (ENST ID and version, dot separated)

l_tr Length of the transcript, in nucleotides

l_utr5 Length of the annotated 5' UTR (if any), in nucleotides

l_cds Length of the annotated CDS (if any), in nucleotides

l_utr3 Length of the annotated 3' UTR (if any), in nucleotides

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psite

Ribosome P-sites position within reads.

Description

This function identifies the exact position of the ribosome P-site within each read, determined by the localisation of its first nucleotide (see Details). It returns a data table containing, for all samples and read lengths: i) the percentage of reads in the whole dataset, ii) the percentage of reads aligning on the start codon (if any); iii) the distance of the P-site from the two extremities of the reads before and after the correction step; iv) the name of the sample. Optionally, this function plots a collection of read length-specific occupancy metaprofiles displaying the P-site offsets computed through the process.

Usage

```
psite(
  data,
  flanking = 6,
  start = TRUE,
  extremity = "auto",
  plot = FALSE,
  plot_dir = NULL,
  plot_format = "png",
  cl = 99,
  txt = FALSE,
  txt_file = NULL
)
```

is TRUE.

Arguments

data	Either list of data tables or GRangesList object from bamtolist, bedtolist, duplicates_filter or length_filter.
flanking	Integer value specifying for the selected reads the minimum number of nucleotides that must flank the reference codon in both directions. Default is 6.
start	Logical value whether to use the translation initiation site as reference codon. Default is TRUE. If FALSE, the second to last codon is used instead.
extremity	Either "5end", "3end" or "auto". It specifies if the correction step should be based on 5' extremities ("5end") or 3' extremities ("3end"). Default is "auto" i.e. the optimal extremity is automatically selected.
plot	Logical value whether to plot the occupancy metaprofiles displaying the P-site offsets computed in both steps of the algorithm. Default is FALSE.
plot_dir	Character string specifying the directory where read length-specific occupancy metaprofiles shuold be stored. If the specified folder doesn't exist, it is automatically created. If NULL (the default), the metaprofiles are stored in a new subfolder of the working directory, called <i>offset_plot</i> . This parameter is considered only if plot is TRUE.
plot_format	Either "png" (the default) or "pdf". This parameter specifies the file format storing the length-specific occupancy metaprofiles. It is considered only if plot
plot plot_dir	based on 5' extremities ("5end") or 3' extremities ("3end"). Default is "auto i.e. the optimal extremity is automatically selected. Logical value whether to plot the occupancy metaprofiles displaying the P-site offsets computed in both steps of the algorithm. Default is FALSE. Character string specifying the directory where read length-specific occupancy metaprofiles shuold be stored. If the specified folder doesn't exist, it is automatically created. If NULL (the default), the metaprofiles are stored in a new subfolder of the working directory, called <i>offset_plot</i> . This parameter is considered only if plot is TRUE. Either "png" (the default) or "pdf". This parameter specifies the file formatical contents and the subspecifies is the file formatical contents.

psite_info 25

cl Integer value in [1,100] specifying a confidence level for generating occupancy metaprofiles for to a sub-range of read lengths i.e. for the cl 99. This parameter is considered only if plot is TRUE.

Logical value whether to write in a txt file reporting the extremity used for the correction step and the best offset for each sample. Similar information are displayed by default in the console. Default is FALSE.

Character string specifying the path, name and extension (e.g. "PATH/NAME.extension") of the plain text file where the extremity used for the correction step and the best offset for each sample shuold be written. If the specified folder doesn't exist, it is automatically created. If NULL (the default), the information are written in "best_offset.txt" and saved in the working directory. This parameter is considered only if txt is TRUE.

Details

txt

txt_file

The P-site offset (PO) is defined as the distance between the extremities of a read and the first nucleotide of the P-site itself. The function processes all samples separately starting from reads mapping on the reference codon (either the start codon or the second to last codon, see start) of any annotated coding sequences. Read lengths-specific POs are inferred in two steps. First, reads mapping on the reference codon are grouped according to their length, each group corresponding to a bin. Reads whose extremities are too close to the reference codon are discarded (see flanking). For each bin temporary 5' and 3' POs are defined as the distances between the first nucleotide of the reference codon and the nucleotide corresponding to the global maximum found in the profiles of the 5' and the 3' end at the left and at the right of the reference codon, respectively. After the identification of the P-site for all reads aligning on the reference codon, the POs corresponding to each length are assigned to each read of the dataset. Second, the most frequent temporary POs associated to the optimal extremity (see extremity) and the predominant bins are exploited as reference values for correcting the temporary POs of smaller bins. Briefly, the correction step defines for each length bin a new PO based on the local maximum, whose distance from the reference codon is the closest to the most frequent temporary POs. For further details please refer to the riboWaltz article (available here).

Value

A data table.

```
data(reads_list)
## Compute the P-site offset automatically selecting the optimal read
## extremity for the correction step and not plotting any metaprofile:
psite(reads_list, flanking = 6, extremity="auto")
```

26 psite_info

Description

This function provides additional reads information according to the position of the P-site identfied by psite. It attaches to each data table in a list four columns reporting i) the P-site position with respect to the 1st nucleotide of the transcript, ii) the P-site position with respect to the start and the stop codon of the annotated coding sequence (if any) and iii) the region of the transcript (5' UTR, CDS, 3' UTR) that includes the P-site. Please note: 1) for transcripts not associated to any annotated CDS the P-site position with respect to the start and the stop codon is set to NA; 2) P-sites of short reads (<20 nts) might be located very close to the 5' or 3' extremity, with no biological meaning and causing potential downstream issues; for these reasons, all read lengths showing this feature will be removed. Optionally, additional columns reporting the three nucleotides covered by the P-site, the A-site and the E-site are attached, based on FASTA files or BSgenome data packages containing the transcript nucleotide sequences.

Usage

```
psite_info(
  data,
  offset,
  site = NULL,
  fastapath = NULL,
  fasta_genome = TRUE,
  refseq_sep = NULL,
  bsgenome = NULL,
  gtfpath = NULL,
  txdb = NULL,
  dataSource = NA,
  organism = NA,
  output_class = "datatable"
)
```

Arguments

data Either list of data tables or GRangesList object from bamtolist, bedtolist,

duplicates_filter or length_filter.

offset Data table from psite.

site Either "psite, "asite", "esite" or a combination of these strings. It specifies if

additional column(s) reporting the three nucleotides covered by the ribosome P-site ("psite"), A-site ("asite") and E-site ("esite") should be added. Note: either

 $\label{thm:continuous} \mbox{fastapath or bsgenome is required for this purpose. Default is NULL.}$

fastapath Character string specifying the FASTA file used in the alignment step, including

its path, name and extension. This file can contain reference nucleotide sequences either of a genome assembly or of all the transcripts (see Details and fasta_genome). Please make sure the sequences derive from the same release of the annotation file used in the create_annotation function. Note: either fastapath or bsgenome is required to generate additional column(s) specified

by site. Default is NULL.

fasta_genome Logical value whether the FASTA file specified by fastapath contains nu-

cleotide sequences of a genome assembly. If TRUE (the default), an annotation object is required (see gtfpath and txdb). FALSE implies the nucleotide

sequences of all the transcripts is provided instead.

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refseq_sep

Character specifying the separator between reference sequences' name and additional information to discard, stored in the headers of the FASTA file specified by fastapath (if any). It might be required for matching the reference sequences' identifiers reported in the input list of data tables. All characters before the first occurrence of the specified separator are kept. Default is NULL i.e. no string splitting is performed.

bsgenome

Character string specifying the BSgenome data package with the genome sequences to be loaded. If not already present in the system, it is automatically installed through the biocLite.R script (check the list of available BSgenome data packages by running the available.genomes function of the BSgenome package). This parameter must be coupled with an annotation object (see gtfpath and txdb). Please make sure the sequences included in the specified BSgenome data pakage are in agreement with the sequences used in the alignment step. Note: either fastapath or bsgenome is required to generate additional column(s) specified by site. Default is NULL.

gtfpath

Character string specifying the location of a GTF file, including its path, name and extension. Please make sure the GTF file and the sequences specified by fastapath or bsgenome derive from the same release. Note that either gtfpath or txdb is required if and only if nucleotide sequences of a genome assembly are provided (see fastapath or bsgenome). Default is NULL.

txdb

Character string specifying the TxDb annotation package to be loaded. If not already present in the system, it is automatically installed through the biocLite.R script (check here the list of available TxDb annotation packages). Please make sure the TxDb annotation package and the sequences specified by fastapath or bsgenome derive from the same release. Note that either gtfpath or txdb is required if and only if nucleotide sequences of a genome assembly are provided (see fastapath or bsgenome). Default is NULL.

dataSource

Optional character string describing the origin of the GTF data file. This parameter is considered only if gtfpath is specified. For more information about this parameter please refer to the description of *dataSource* of the makeTxDbFromGFF function included in the GenomicFeatures package.

organism

Optional character string reporting the genus and species of the organism of the GTF data file. This parameter is considered only if gtfpath is specified. For more information about this parameter please refer to the description of *organism* of the makeTxDbFromGFF function included in the GenomicFeatures package.

output_class

Either "datatable" or "granges". It specifies the format of the output i.e. a list of data tables or a GRangesList object. Default is "datatable".

Details

riboWaltz only works for read alignments based on transcript coordinates. This choice is due to the main purpose of RiboSeq assays to study translational events through the isolation and sequencing of ribosome protected fragments. Most reads from RiboSeq are supposed to map on mRNAs and not on introns and intergenic regions. Nevertheless, BAM based on transcript coordinates can be generated in two ways: i) aligning directly against transcript sequences; ii) aligning against standard chromosome sequences, requiring the outputs to be translated in transcript coordinates. The first option can be easily handled by many aligners (e.g. Bowtie), given a reference FASTA file where each sequence represents a transcript, from the beginning of the 5' UTR to the end of the 3' UTR. The second procedure is based on reference FASTA files where each sequence represents a chromosome, usually coupled with comprehensive gene annotation files (GTF or GFF).

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The STAR aligner, with its option –quantMode TranscriptomeSAM (see Chapter 6 of its manual), is an example of tool providing such a feature.

Value

A list of data tables or a GRangesList object.

Examples

```
data(reads_list)
data(psite_offset)
data(mm81cdna)
reads_psite_list <- psite_info(reads_list, psite_offset)</pre>
```

psite_offset

P-site offsets data table

Description

An example dataset containing length-specific ribosome P-site offsets as returned by psite applied to reads_list.

Usage

```
psite_offset
```

Format

A data table with 31 rows and 9 variables:

length Length of the read, in nucleotides (intended as the width of the reference sequence region covered by the RNA fragment. For further information about this choice please refer to section Details of function bamtolist)

total_percentage Percentage of reads of the considered length in the whole dataset

start_percentage Percentage of reads of the considered length aligning on the start codon (if any)

 $around_start$ A logical value whether at least one read of the considered length aligns on the start codon (T = yes, F = no)

offset_from_5 Temporary P-site offset from the 5' end of the read, in nucleotides (before the correction step)

offset_from_3 Temporary P-site offset from the 3' end of the read, in nucleotides (before the correction step)

corrected_offset_from_5 P-site offset from the 5' end of the read, in nucleotides (after the correction step)

corrected_offset_from_3 P-site offset from the 3' end of the read, in nucleotides (after the correction step)

sample Name of the sample

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reads_list

Reads information data table

Description

An example dataset containing details on reads mapping on the mouse transcriptome, generated from BAM or BED files. A subset of the original dataset is provided, including only reads aligning on the translation initiation site. Please contact the authors for further information.

Usage

reads_list

Format

A list of data tables with 1 object (named Samp 1) of 100,062 rows and 6 variables:

transcript Name of the transcript (ENST ID and version, dot separated)

- end5 Position of the 5' end of the read with respect to the first nucleotide of the transcript, in nucleotides
- end3 Position of the 3' end of the read with respect to the first nucleotide of the transcript, in nucleotides
- cds_start Leftmost position of the annotated CDS with respect to the first nucleotide of the transcript, in nucleotides
- cds_stop Rightmost position of the annotated CDS with respect to the first nucleotide of the transcript, in nucleotides

region_psite

Percentage of P-sites per transcript region.

Description

This function computes the percentage of P-sites falling in the three annotated regions of the transcripts (5' UTR, CDS and 3' UTR) and generates a bar plot of the resulting values.

Usage

```
region_psite(
  data,
  annotation,
  sample = NULL,
  transcripts = NULL,
  label_sample = NULL,
  colour = c("gray70", "gray40", "gray10"))
```

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Arguments

data	Either list of data tables or GRangesList object from psite_info.
annotation	Data table as generated by create_annotation.
sample	Character string vector specifying the name of the sample(s) of interest. Default is NULL i.e. all samples in data are processed.
transcripts	Character string vector listing the name of transcripts to be included in the analysis. Default is NULL i.e. all transcripts are used. Please note: transcripts without annotated 5' UTR, CDS and 3' UTR are automatically discarded.
label_sample	Named character string vector the same length as sample specifying the sample names to be displayed in the plot. Plase be careful to name each element of the vector after the correct corresponding string in sample. No specific order is required. Default is NULL i.e. sample names in sample are used.
colour	Character string vector of three elements specifying the colour for the 5' UTR, CDS and 3' UTR bars, respectively. Default is a grayscale.

Details

Column "RNAs" reports the percentage of region length for the transcripts included in the analysis, based on the cumulative nucleotide length of 5' UTRs, CDSs and 3' UTRs. These values reflect the expected read distribution from a random fragmentation of RNA and can be used as a baseline to verify the expected enrichment of ribosome (P-site) signal in CDSs.

Value

A list containing a ggplot2 object ("plot") and the data table with the associated data ("dt").

Examples

```
## data(reads_list)
## data(mm81cdna)
##
## ## compute and add p-site datails
## psite_offset <- psite(reads_list, flanking = 6, extremity = "auto")
## reads_psite_list <- psite_info(reads_list, psite_offset)
##
## reg_psite <- region_psite(reads_psite_list, mm81cdna, sample = "Samp1")
## reg_psite[["plot"]]</pre>
```

rends_heat

Metaheatmaps of the two extremities of the reads.

Description

This function generates four metaheatmaps displaying the abundance of the 5' and 3' extremity of reads mapping around the start and the stop codon of annotated CDSs, stratified by their length.

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Usage

```
rends_heat(
 data,
  annotation,
  sample,
  transcripts = NULL,
  c1 = 95,
  utr51 = 50.
  cdsl = 50,
 utr31 = 50,
  log_colour = F,
  colour = "black"
```

Arguments

data	Either list of data tables or GRangesList object from bamtolist, bedtolist, length_filter or psite_info.
annotation	Data table as generated by create_annotation.
sample	Character string specifying the name of the sample of interest.
transcripts	Character string vector listing the name of transcripts to be included in the analysis. Default is NIII i.e. all transcripts are used. Please note: transcripts

ysis. Default is NULL i.e. all transcripts are used. Please note: transcripts with either 5' UTR, coding sequence or 3' UTR shorter than utr51, 2*cds1 and utr31, respectively, are automatically discarded.

cl Integer value in [1,100] specifying a confidence level for restricting the plot to a sub-range of read lengths. The new range is associated to the most abundant

populations of reads accounting for the cl of the sample. Default is 95.

utr5l Positive integer specifying the length (in nucleotides) of the 5' UTR region

flanking the start codon to be considered in the analysis. Default is 50.

cdsl Positive integer specifying the length (in nucleotides) of the CDS regions flank-

ing both the start and stop codon to be considered in the analysis. Default is

Positive integer specifying the length (in nucleotides) of the 3' UTR region utr31

flanking the stop codon to be considered in the analysis. Default is 50.

Logical value whether to use a logarithmic colour scale (strongly suggested in log_colour

case of large signal variations). Default is FALSE.

colour Character string specifying the colour of the plot. The colour scheme is as

> follow: tiles corresponding to the lowest signal are always white, tiles corresponding to the highest signal are of the specified colour and the progression between these two colours follows either linear or logarithmic gradients (see

log_colour). Default is "black".

Value

A list containing a ggplot2 object ("plot") and the data table with the associated data ("dt").

```
data(reads_list)
data(mm81cdna)
```

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```
## Generate metaheatmaps for all read lengths:
heatend_whole <- rends_heat(reads_list, mm81cdna, sample = "Samp1", cl = 100)
## Generate metaheatmaps for a sub-range of read lengths shortening the
## flanking regions around the start and stop codon:
heatend_sub95 <- rends_heat(reads_list, mm81cdna, sample = "Samp1", cl = 95,
utr51 = 30, cds1 = 40, utr31 = 30)</pre>
```

rlength_distr

Read length distributions.

Description

This function generates read length distributions, displayed as bar plots. Multiple samples can be handled in several ways.

Usage

```
rlength_distr(
  data,
  sample,
  multisamples = "separated",
  plot_style = "split",
  transcripts = NULL,
  cl = 100,
  colour = NULL
)
```

Arguments

data

Either list of data tables or GRangesList object from bamtolist, bedtolist, length_filter or psite_info.

sample

Either character string or character string vector specifying the name of the sample(s) of interest. A named list of one or more character strings and/or character string vectors can be provided. In this case i) each list element should include the name of the replicate(s) related to the sample of interest and ii) the name assigned to the elements of the list are displayed in the plot. Multiple replicates specified in character string vectors are handled according to multisample.

multisamples

Either "separated", "average" or "sum". It specifies how to handle multiple samples and replicates. If "saparated", one metaprofile for each sample included in sample is returned as an independent ggplot object. If sample is a list, it is unlisted, coerced to character string and handled accordingly. If "average" or "sum" i) one metaprofiles is returned if sample is a character string vector or ii) one metaprofiles is built for each element of sample when it is a list. If "average", the metaprofiles display for each nucleotide the mean signal and the corresponding standard error computed among the replicates. In this case a single ggplot object is returned, where multiple bar plots are organized and displayed according to plot_style. Default is "separated".

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plot_style Either "split", "dodged" or "mirrored". It specifies how to organize and display

multiple bar plots. If "split", the bar plots are placed one next to the other, in independent boxes. If "dodged", all bar plots are included in one box, with each bar side by side with the others. If "mirrored" sample must be a list of exactly two elements and the two bar plots are mirrored along the x axis. Default is

"split".

transcripts Character string vector listing the name of transcripts to be included in the anal-

ysis. Default is NULL i.e. all transcripts are used.

cl Integer value in [1,100] specifying a confidence level for restricting the plot to

a sub-range of read lengths. The new range is associated to the most abundant populations of reads accounting for the cl of the sample. If multiple sample names are provided one range of read lengths is computed, such that at least the

cl represented. Default is 100.

colour Character string or character string vector specifying the colour of the bar plot(s).

If sample is a list of multiple elements and multisamples is set to "average", a colour for each element of the list is required. If this parameter is not specified

the R default palette is employed. Default is NULL.

Value

List containing one or more ggplot2 object(s) and the data table with the associated data ("dt").

```
data(reads_list)
## Generate the length distribution for all read lengths:
lendist_whole <- rlength_distr(reads_list, sample = "Samp1", cl = 100)
lendist_whole[["plot_Samp1"]]

## Generate the length distribution for a sub-range of read lengths:
lendist_sub95 <- rlength_distr(reads_list, sample = "Samp1", cl = 95)
lendist_sub95[["plot_Samp1"]]</pre>
```

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