## Package 'Isosceles'

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```
Title Isoform Single-Cell and Long-read Expression Suite
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Description Transcript detection and quantification from long reads.
Depends R (>= 4.2.0),
      SingleCellExperiment (>= 1.18.0)
Imports utils (>= 4.2.0),
      methods (>= 4.2.0),
      stats (>= 4.2.0),
      rlang (>= 1.0.4),
      assertthat (>= 0.2.1),
      magrittr (>= 2.0.3),
      tibble (>= 3.1.7),
      tidyselect (\geq 1.1.2),
      dplyr (>= 1.0.9),
      tidyr (>= 1.2.0),
      glue (>= 1.6.2),
      digest (>= 0.6.29),
      Rcpp (>= 1.0.9),
      Matrix (>= 1.4-1),
      BiocParallel (>= 1.30.3),
      BiocNeighbors (>= 1.14.0),
      S4Vectors (>= 0.34.0),
      BiocGenerics (>= 0.42.0),
      Biostrings (\geq 2.64.0),
      BSgenome (>= 1.64.0),
      GenomeInfoDb (>= 1.32.2),
      IRanges (>= 2.30.0),
      GenomicRanges (>= 1.48.0),
      Rsamtools (>= 2.12.0),
      GenomicAlignments (>= 1.32.1),
      rtracklayer (>= 1.56.1),
      GenomicFeatures (>= 1.48.3),
      SummarizedExperiment (>= 1.26.1),
      igraph (>= 1.3.4),
      scuttle (>= 1.6.2),
```

```
fastmatch (>= 1.1-3),
     pheatmap (>= 1.0.12),
     ggbio (>= 1.44.1),
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Isosceles-package

Isosceles: Isoform Single-Cell and Long-read Expression Suite

## Description

Transcript detection and quantification from long reads

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export\_gtf

Data export to a GTF file

## Description

Export transcripts from a SummarizedExperiment to a GTF file

## Usage

```
export_gtf(se, file)
```

## **Arguments**

se A transcript-level SummarizedExperiment object returned by the prepare\_transcript\_se

function

file A string specifying the output file path

### Value

Nothing is returned

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```
extract_read_structures
```

Read structure extraction from BAM files

## **Description**

Extract non-redundant read structures from one or multiple BAM files

## Usage

```
extract_read_structures(bam_files, chunk_size = 1e+06, ncpu = 1)
```

#### **Arguments**

bam\_files A character vector containing BAM file paths

chunk\_size An integer scalar specifying the chunk size for reading the BAM files

ncpu An integer scalar specifying the number of cores to use for multicore paralleliza-

tion

#### Value

A data frame containing non-redundant read structure data obtained from the BAM files

 $merge\_sc\_neighbors \qquad \textit{Merging neighboring cell TCC values in scRNA-Seq data}$ 

## **Description**

Prepares a TCC SummarizedExperiment object where count values from the nearest k neighbors are added to the count values of each cell

## Usage

```
merge_sc_neighbors(se_tcc, pca_mat, k = 10, use_annoy = FALSE, ncpu = 1)
```

## **Arguments**

se_tcc	A TCC SummarizedExperiment	object returned by t	he prepare_tcc_se func-
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tion

pca\_mat A matrix containing PCA coordinates of each cell

k An integer scalar specifying the number of nearest neighbors to use

use\_annoy A logical scalar indicating whether to use the Annoy algorithm for approximate

nearest neighbor identification (recommended for big datasets)

ncpu An integer scalar specifying the number of cores to use for multicore paralleliza-

tion

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## Value

A SummarizedExperiment object containing merged TCC data

## Description

Creates a heatmap of PSI (Percent Spliced In) values for the regions of a given gene across samples or cells

## Usage

```
plot_psi_heatmap(
    se_psi,
    gene_id,
    heatmap_colors = viridis::cividis(100),
    region_colors = NULL,
    ...
)
```

## Arguments

```
se_psi A PSI SummarizedExperiment object returned by the prepare_psi_se function
gene_id A string containing the identifier of the gene to plot
heatmap_colors A character vector containing the color palette used in the heatmap
region_colors A named character vector of colors for the region type annotations
... Additional parameters for the plot, passed to the pheatmap function
```

#### Value

A plot object

## **Description**

Creates a plot showing PSI regions and transcript structures for the given gene. Individual transcript structures are colored by their relative expression, calculated from the overall TPM values and expressed in percentages. For better visualization, introns can be shrinked using the max\_intron\_length argument

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#### Usage

```
plot_psi_regions(
    se_psi,
    se_transcript,
    gene_id,
    max_transcripts = Inf,
    max_intron_length = NULL,
    region_colors = NULL
)
```

#### **Arguments**

se\_psi A PSI SummarizedExperiment object returned by the prepare\_psi\_se function

se\_transcript A transcript-level SummarizedExperiment object returned by the prepare\_transcript\_se

function

gene\_id A string containing the identifier of the gene to plot

 ${\tt max\_transcripts}$ 

An integer scalar specifying the maximum number of transcripts with the highest

relative expression to plot

max\_intron\_length

An integer scalar specifying the maximum intron length after shrinking. If set

to NULL, no shrinking is performed

region\_colors A named character vector of colors for the PSI region types

#### Value

A plot object

prepare\_gene\_se

Prepare a gene-level SummarizedExperiment object

## Description

Prepares a gene-level SummarizedExperiment from TCC data

#### Usage

```
prepare_gene_se(se_tcc)
```

#### **Arguments**

se\_tcc

A TCC SummarizedExperiment object returned by a function from the Isosceles-package

#### Value

A SummarizedExperiment object containing gene annotation and quantification data

prepare\_pseudobulk\_se

prepare\_pseudobulk\_se Prepare a pseudobulk TCC SummarizedExperiment object

## **Description**

Prepares a pseudobulk TCC SummarizedExperiment from TCC data and given cell labels

#### Usage

```
prepare_pseudobulk_se(se_tcc, cell_labels)
```

## Arguments

se\_tcc A TCC SummarizedExperiment object returned by the prepare\_tcc\_se func-

tion

cell\_labels A vector or a factor containing cell labels acting as a grouping variable

#### Value

A pseudobulk SummarizedExperiment object containing TCC annotation and quantification data

prepare\_psi\_se

Prepare a PSI SummarizedExperiment object

## **Description**

Prepares a PSI (Percent Spliced In) SummarizedExperiment object for the given transcript-level SummarizedExperiment object. PSI values are calculated for the following types of regions:

- TSS transcription start sites
- TES transcription end sites
- CE core exonic regions
- RI retained intronic regions
- A5 5' alternative exonic regions
- A3 3' alternative exonic regions

TSS and TES positions are calculated based on transcripts' binned start and end coordinates extracted from their identifiers

## Usage

```
prepare_psi_se(se, ncpu = 1)
```

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#### **Arguments**

se A transcript-level SummarizedExperiment object returned by the prepare\_transcript\_se

function

ncpu An integer scalar specifying the number of cores to use for multicore paralleliza-

tion

#### Value

A SummarizedExperiment object containing PSI annotation and quantification data

prepare\_tcc\_se

Prepare a TCC SummarizedExperiment object

## **Description**

Prepares a TCC (Transcript Compatibility Counts) SummarizedExperiment object for the given BAM files and transcript set

## Usage

```
prepare_tcc_se(
  bam_files,
  transcript_data,
  run_mode = "strict",
  min_read_count = 1,
  min_relative_expression = 0.1,
  extend_spliced_transcripts = 100,
  is_single_cell = FALSE,
  barcode_tag = "BC",
  chunk_size = 1e+06,
  ncpu = 1
)
```

## **Arguments**

bam\_files A named character vector containing BAM file paths

transcript\_data

A named list containing transcript data returned by the prepare\_transcripts

function

run\_mode A string specifying the mode for choosing the transcript set ('strict', 'de novo strict',

'de\_novo\_loose' or 'de\_novo\_full')

min\_read\_count An integer scalar specifying the read count threshold for transcripts extracted

from the BAM files

min\_relative\_expression

A numeric scalar specifying the relative expression threshold for transcripts ex-

tracted from the BAM files

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```
extend_spliced_transcripts
```

An integer scalar specifying the number of base pairs by which transcript starts

and ends are extended for spliced read compatibility search

is\_single\_cell A logical scalar specifying if the BAM files contain single cell data

barcode\_tag A string specifying the name of the BAM file tag containing cell barcodes chunk\_size An integer scalar specifying the chunk size for reading the BAM files

ncpu An integer scalar specifying the number of cores to use for multicore paralleliza-

tion

#### Value

A SummarizedExperiment object containing TCC annotation and quantification data

```
prepare_transcripts Transcript data preparation
```

## **Description**

Prepare transcript data (reference and extracted from the BAM files) for further analysis

## Usage

```
prepare_transcripts(
  gtf_file,
  genome_fasta_file,
  bam_parsed,
  min_intron_length = 30,
  known_intron_motifs = c("GT-AG"),
  rescue_annotated_introns = FALSE,
  known_intron_granges = NULL,
  min_bam_splice_read_count = 2,
  min_bam_splice_fraction = 0.1,
  bin_size = 50
)
```

#### **Arguments**

```
gtf_file A string containing a GTF file path genome_fasta_file
```

A string containing a genome FASTA file path

bam\_parsed A data frame containing non-redundant read structure data returned by the extract\_read\_structures

function. If NULL, only reference transcripts are used

min\_intron\_length

An integer scalar specifying the minimal length of introns to assign strand to

known\_intron\_motifs

A character vector specifying the known intron motifs

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rescue\_annotated\_introns

A logical scalar specifying if introns found in genome annotations should be kept even if they don't have known intron motifs

known\_intron\_granges

A GRanges object storing known intron positions (e.g. from short read data) used for transcript classification. If set to NULL, only introns from reference annotations are used

min\_bam\_splice\_read\_count

An integer scalar specifying the read count threshold for splice sites confirmed by aligned reads

min\_bam\_splice\_fraction

A numeric scalar specifying the minimum connectivity fraction to a known splice site for splice sites confirmed by aligned reads

bin\_size An integer scalar specifying the bin size for transcript start and end position binning

#### Value

A named list containing following elements:

tx\_df a data frame storing extracted transcript data

tx\_granges a GRanges object storing genomic positions of extracted transcript

tx\_exon\_granges\_list a GRangesList object storing exon genomic positions of extracted transcript

tx\_intron\_granges\_list a GRangesList object storing intron genomic positions of extracted transcript

prepare\_transcript\_se Prepare a transcript-level SummarizedExperiment object

## Description

Prepares a transcript-level SummarizedExperiment from TCC data using the EM algorithm

#### Usage

```
prepare_transcript_se(
    se_tcc,
    em.maxiter = 250,
    em.conv = 0.01,
    use_length_normalization = TRUE,
    ncpu = 1
)
```

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## **Arguments**

se\_tcc A TCC SummarizedExperiment object returned by a function from the Isosceles-package

em.maxiter An integer scalar specifying the maximum number of EM iterations

em. conv A numeric scalar specifying the EM convergence threshold

 $use\_length\_normalization$ 

A logical scalar specifying if normalization using effective transcript lengths

should be used during EM

ncpu An integer scalar specifying the number of cores to use for multicore paralleliza-

tion

#### Value

A SummarizedExperiment object containing transcript annotation and quantification data

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