

Package ‘SPICEY’

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Title Calculates cell type specificity from single cell data

Version 0.99.3

Description SPICEY (SPecificity Index for Coding and Epigenetic activitY) is an R package designed to quantify cell-type specificity in single-cell transcriptomic and epigenomic data, particularly scRNA-seq and scATAC-seq. It introduces two complementary indices: the Gene Expression Tissue Specificity Index (GETSI) and the Regulatory Element Tissue Specificity Index (RETSI), both based on entropy to provide continuous, interpretable measures of specificity. By integrating gene expression and chromatin accessibility, SPICEY enables standardized analysis of cell-type-specific regulatory programs across diverse tissues and conditions.

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`.add_tss_annotation` *Add TSS annotation to peaks*

Description

Identifies transcription start sites (TSS) overlapping with input peaks and adds logical and gene ID columns to an annotation table.

Usage

```
.add_tss_annotation(
  annotation,
  peaks,
  txdb,
  annot_db,
  protein_coding_only,
  verbose
)
```

Arguments

annotation	A <code>data.frame</code> or <code>GRanges</code> containing peak annotations, including a <code>region_id</code> column.
peaks	A <code>GRanges</code> object containing the original peak set.
txdb	<code>TxDb</code> object for genome annotation (required if annotation requested).
annot_db	<code>AnnotationDbi</code> object for gene ID mapping (required if annotation requested).
protein_coding_only	Logical; restrict to protein-coding genes (default TRUE).
verbose	Logical; print messages (default TRUE).

Value

A `data.frame` or `GRanges` (depending on input) with added columns:

in_TSS Logical, TRUE if the peak overlaps a TSS.

TSS_gene Gene symbol of the overlapping TSS, if any.

.parse_input_diff *Parses input data of various types (e.g., named lists of `GRanges` or `data.frame`, or a `GRangesList`) into a single tidy `data.frame`, with a `cell_type` column.*

Description

Parses input data of various types (e.g., named lists of `GRanges` or `data.frame`, or a `GRangesList`) into a single tidy `data.frame`, with a `cell_type` column.

Usage

```
.parse_input_diff(input)
```

Arguments

input	An object representing differential results, such as:
	• A named list of <code>GRanges</code> objects.
	• A named list of <code>data.frames</code> .
	• A <code>GRangesList</code> .

Value

A `data.frame` combining all elements, with an added `cell_type` column indicating the source.

.standardize_peaks	<i>Standardizes peak input, ensuring that input peaks is a GRanges object, removes alternate scaffolds (_alt, random, fix, Un), and assigns region IDs as names.</i>
--------------------	--

Description

Standardizes peak input, ensuring that input peaks is a GRanges object, removes alternate scaffolds (_alt, random, fix, Un), and assigns region IDs as names.

Usage

```
.standardize_peaks(peaks)
```

Arguments

peaks	A data.frame with genomic coordinates and a region_id column, or a GRanges object with a region_id column.
-------	--

Value

A GRanges object with:

seqnames, start, end Coordinates of the input regions.

region_id Unique identifier of the region (e.g., chr1-5000-5800).

metadata columns Any additional columns present in the input.

annotate_with_coaccessibility

Annotate peaks with co-accessible genes using Cicero links

Description

Links peaks to genes based on Cicero co-accessibility with promoters or TSSs.

Usage

```
annotate_with_coaccessibility(
  peaks,
  txdb,
  links_df,
  annot_db,
  protein_coding_only = TRUE,
  verbose = TRUE,
  add_tss_annotation = FALSE,
  upstream,
  downstream
)
```

Arguments

peaks	A GRanges or data.frame of peaks with at least the following columns:
	seqnames Chromosome name of the regulatory region (e.g., "chr1"). Only for data.frames.
	start Start coordinate of the peak. Only for data.frames.
	end End coordinate of the peak. Only for data.frames.
	region_id Unique identifier of the region (e.g., chr1-5000-5800)
txdb	TxDb object for genome annotation (required if annotation requested).
links_df	A data.frame with Cicero links. Must contain columns: Peak1, Peak2, and coaccess.
annot_dbi	AnnotationDbi object for gene ID mapping (required if annotation requested).
protein_coding_only	Logical; restrict to protein-coding genes (default TRUE).
verbose	Logical; print messages (default TRUE).
add_tss_annotation	Logical; annotate regulatory elements overlapping TSS (default FALSE). If TRUE, use +/- 1bp TSS.
upstream	Single integer value indicating the number of bases upstream from the TSS (transcription start sites) (default "2000kb").
downstream	Single integer values indicating the number of bases downstream from the TSS (transcription start sites) (default "2000kb").

Value

A data.frame with the original metadata columns from peaks, along with an added gene_id column containing the symbol of the co-accessible gene. Peaks with no gene annotation will have NA in the gene_id field.

Examples

```
library(TxDb.Hsapiens.UCSC.hg38.knownGene)
library(org.Hs.eg.db)

data(atac)
data(cicero_links)

peaks <- unique(unlist(atac)[, c("region_id")])
annotation_coacc <- annotate_with_coaccessibility(
  peaks = peaks,
  txdb = TxDb.Hsapiens.UCSC.hg38.knownGene,
  links_df = cicero_links,
  annot_dbi = org.Hs.eg.db,
  protein_coding_only = TRUE,
  verbose = TRUE,
  add_tss_annotation = FALSE,
  upstream = 2000,
  downstream = 2000
)
```

`annotate_with_nearest` *Annotates regulatory elements (e.g., ATAC-seq peaks) to the nearest gene*

Description

based on distance to the transcription start site (TSS), using a TxDb reference and gene annotations from org.*.db packages.

Usage

```
annotate_with_nearest(
  peaks,
  txdb,
  annot_db,
  protein_coding_only = TRUE,
  verbose = TRUE,
  add_tss_annotation = FALSE,
  upstream,
  downstream
)
```

Arguments

<code>peaks</code>	A GRanges or data.frame of peaks with at least the following columns:
	<code>seqnames</code> Chromosome name of the regulatory region (e.g., "chr1"). Only for data.frames.
	<code>start</code> Start coordinate of the peak. Only for data.frames.
	<code>end</code> End coordinate of the peak. Only for data.frames.
	<code>region_id</code> Unique identifier of the region (e.g., chr1-5000-5800)
<code>txdb</code>	TxDb object for genome annotation (required if annotation requested).
<code>annot_db</code>	AnnotationDbi object for gene ID mapping (required if annotation requested).
<code>protein_coding_only</code>	Logical; restrict to protein-coding genes (default TRUE).
<code>verbose</code>	Logical; print messages (default TRUE).
<code>add_tss_annotation</code>	Logical; annotate regulatory elements overlapping TSS (default FALSE). If TRUE, use +/- 1bp TSS.
<code>upstream</code>	Single integer value indicating the number of bases upstream from the TSS (transcription start sites) (default "2000kb").
<code>downstream</code>	Single integer values indicating the number of bases downstream from the TSS (transcription start sites) (default "2000kb").

Value

A data.frame of peaks annotated to its nearest gene, with columns:

<code>distanceToTSS</code>	Distance to the nearest TSS
<code>gene_id</code>	Identifier of the gene. Must be an official gene symbol (e.g., GAPDH)
<code>annotation</code>	"Promoter" or "Distal" based on distance

Examples

```
library(dplyr)
library(TxDb.Hsapiens.UCSC.hg38.knownGene)
library(org.Hs.eg.db)

data(atac)
peaks <- unique(unlist(atac)[, c("region_id")])

annotation_near <- annotate_with_nearest(
  peaks = peaks,
  txdb = TxDb.Hsapiens.UCSC.hg38.knownGene,
  annot_db = org.Hs.eg.db,
  protein_coding_only = TRUE,
  verbose = TRUE,
  add_tss_annotation = FALSE,
  upstream = 2000,
  downstream = 2000
)
```

atac

Example single-cell ATAC-seq differential accessibility data

Description

A toy example dataset representing single-cell ATAC-seq differential accessibility results. Each row corresponds to a chromatin accessibility peak tested for differential accessibility across one or more cell types. The dataset is formatted as a list of GRanges objects or data frames (convertible to GRanges), where each element represents differential accessibility statistics for a specific cell type. Multiple rows may exist per peak, each representing results in a different cell type.

Usage

```
data(atac)
```

Format

A data frame or GRanges-like object with the following required columns:

- region_id** Unique identifier of the region (e.g., chr1-5000-5800).
- avg_log2FC** Average log2 fold-change of accessibility for the peak in the specific cell type
- p_val_adj** Adjusted p-value (e.g., FDR-corrected)
- cell_type** Cell type or cluster label associated with each measurement (e.g., Acinar)

Source

Precomputed using `FindMarkers()` (Wilcoxon test, via Presto if available) on control samples from the Human Pancreas Analysis Program (HPAP), using paired snATAC-seq and snRNA-seq data from three non-diabetic human donors.

cicero_links	<i>Example Cicero co-accessibility links</i>
--------------	--

Description

A toy example dataset of co-accessibility links inferred from single-cell ATAC-seq data using tools such as Cicero or Signac's LinkPeaks(). These links support integrative analysis by associating regulatory elements with putative target genes. The peaks referenced here must exactly match those in the ATAC-seq differential accessibility dataset.

Usage

```
data(cicero_links)
```

Format

A data frame with the following columns:

- Peak1** Genomic coordinate or peak identifier for the first peak in the pair (e.g., chr1-110209621-110211746)
- Peak2** Genomic coordinate or peak identifier for the second peak in the pair (e.g., chr1-110209621-110211746)
- coaccess** Co-accessibility score or correlation value quantifying the linkage

Source

Cicero co-accessibility links were computed from UMAP-reduced snATAC-seq data (HPAP, control donors) using `run_cicero()` with chromosome sizes from hg38. Input data matched the peaks in the provided ATAC dataset.

compute_spicey_index	<i>Compute cell type specificity scores from single-cell RNA and/or ATAC data</i>
----------------------	---

Description

Computes:

- GETSI (Gene Expression Tissue Specificity Index) from single-cell RNA-seq differential expression data.
- RETSI (Regulatory Element Tissue Specificity Index) from single-cell ATAC-seq differential accessibility data.

Either RNA or ATAC input must be provided.

Usage

```
compute_spicey_index(diff = NULL, id = NULL)
```

Arguments

diff	Single data.frame with differential results, with required columns:
id	Identifier for either genes or regions. Gene IDs must be official gene symbols, while region IDs should follow the usual genomic coordinate format such as chr-start-end or chr:start-end. The name of this column must match the id argument.
avg_log2FC	Average log2 fold-change for the gene in that cell type.
p_val_adj	Adjusted p-value (e.g., FDR-corrected)
cell_type	Cell type or cluster label (e.g., Acinar)
id	A character string specifying the name of the column in diff that contains the unique identifiers for features (e.g., genes or regions). Must match a column name in diff , such as "gene_id" or "region_id".

Value

A data frame with specificity scores for each feature:

- score (GETSI or RETSI)** Specificity score (weighted log2FC).
- norm_entropy** Shannon entropy-based specificity measure.

entropy_index

Calculate normalized Shannon-entropy of specificity scores

Description

Computes the normalized entropy of specificity scores (RETSI or GETSI) across cell types. Entropy quantifies how evenly a feature's activity is distributed among cell types, and if normalized, yields scores from 0 to 1, where values close to 1 indicate widespread distribution across cell types, and values near 0 denote dominating distribution towards one cell type.

Usage

```
entropy_index(spec_df, group_col)
```

Arguments

spec_df	A data.frame containing the computed specificity scores containing at least the following columns:
cell_type	Cell type or cluster label.
score	Specificity score for each feature in each cell type.
[group_col]	Column containing the feature identifier (e.g., gene_id or region) The name of this column must match the value passed to the group_col argument
group_col	A string specifying the name of the column in da that identifies each feature, such as gene_id for genes or region for ATAC peaks.

Value

A data.frame with one row per feature, containing:

group_col Feature identifier.

entropy Raw Shannon entropy computed from specificity scores.

norm_entropy Normalized Shannon entropy score ($1 - \exp(-\text{entropy})$) bounded between 0 and 1, where lower values indicate higher specificity.

extract_gene_peak_annotations

Overlap peaks with gene promoters to obtain gene annotations

Description

Identifies overlaps between a set of peaks and promoter regions, optionally restricted to protein-coding genes.

Usage

```
extract_gene_peak_annotations(
  peaks,
  txdb,
  annot_db,
  protein_coding_only = TRUE,
  upstream,
  downstream,
  verbose = FALSE
)
```

Arguments

peaks	A GRanges or data.frame of peaks with at least the following columns:
	seqnames Chromosome name of the regulatory region (e.g., "chr1"). Only for data.frames.
	start Start coordinate of the peak. Only for data.frames.
	end End coordinate of the peak. Only for data.frames.
	region_id Unique identifier of the region (e.g., chr1-5000-5800)
txdb	TxDb object for genome annotation (required if annotation requested).
annot_db	AnnotationDbi object for gene ID mapping (required if annotation requested).
protein_coding_only	Logical; restrict to protein-coding genes (default TRUE).
upstream	Single integer value indicating the number of bases upstream from the TSS (transcription start sites) (default "2000kb").
downstream	Single integer values indicating the number of bases downstream from the TSS (transcription start sites) (default "2000kb").
verbose	Logical; print messages (default TRUE).

Value

A GRanges object with:

seqnames, start, end Coordinates of the peak that overlaps with a gene promoter.

region_id Unique identifier of the region (e.g., chr1-5000-5800).

gene_id Identifier of the gene. Must be an official gene symbol (e.g., GAPDH)

get_promoters

Extract promoter regions annotated gene symbols from a TxDb and AnnotationDbi object

Description

Extract promoter regions annotated gene symbols from a TxDb and AnnotationDbi object

Usage

```
get_promoters(
  txdb,
  annot_db,
  upstream,
  downstream,
  protein_coding_only = TRUE
)
```

Arguments

txdb	TxDb object for genome annotation (required if annotation requested).
annot_db	AnnotationDbi object for gene ID mapping (required if annotation requested).
upstream	Single integer value indicating the number of bases upstream from the TSS (transcription start sites) (default "2000kb").
downstream	Single integer values indicating the number of bases downstream from the TSS (transcription start sites) (default "2000kb").
protein_coding_only	Logical; restrict to protein-coding genes (default TRUE).

Value

A GRanges object with the chromosomes, start and end positions of defined species promoter regions together with the official gene symbol stored in the gene_id metadata column.

<code>link_spicey</code>	<i>Link RETSI regions to GETSI scores using gene-based association methods</i>
--------------------------	--

Description

This function connects regulatory regions scored with RETSI

Usage

```
link_spicey(retsi = NULL, getsi = NULL, annotation = NULL)
```

Arguments

<code>retsi</code>	A data.frame containing RETSI scores for chromatin accessibility regions, as returned by <code>compute_spicey_index()</code> using single-cell ATAC-seq differential accessibility data. Must include at least the following columns: region_id Unique identifier of the region (e.g., chr1-5000-5800). cell_type Cell type or cluster label (e.g., Acinar) RETSI RETSI value: cell-type specificity score norm_entropy Normalized Shannon entropy of RETSI
<code>getsi</code>	A data.frame containing GETSI scores for genes, as returned by <code>compute_spicey_index()</code> using single-cell RNA-seq differential expression data. Must include at least the following columns: gene_id Identifier of the gene. Must be an official gene symbol (e.g., GAPDH). cell_type Cell type or cluster label (e.g., Acinar) GETSI GETSI value: cell-type specificity score norm_entropy Normalized Shannon entropy of GETSI
<code>annotation</code>	(Optional). A data.frame linking <code>gene_id</code> to <code>region_id</code> . They should have the same names provided in the respective parameters. This can be provided by the user or generated using the functions: <code>annotate_with_nearest</code> or <code>annotate_with_coaccessibility</code> . It should contain at least the following columns: region_id Unique identifier of the region (e.g., chr1-5000-5800). The name of this column should match the <code>region_id</code> argument. cell_type Cell type or cluster label. (e.g., Acinar) gene_id Identifier of the gene. Must be an official gene symbol (e.g., GAPDH). The name of this column should match the <code>gene_id</code> argument.

Value

A data.frame where each row represents a regulatory element–gene pair linked within a given cell type. The output includes:

region_id	Unique identifier of the region (e.g., chr1-5000-5800)
gene_id	Identifier of the gene. Must be an official gene symbol (e.g., GAPDH).
cell_type	Cell type or cluster in which the association is observed (e.g., Acinar)
RETSI	RETSI score: regulatory element specificity in this cell type.
RETSI_entropy	Normalized shannon-entropy of RETSI (lower = more specific).

GETSI GETSI score: gene expression specificity in this cell type.

GETSI_entropy Normalized shannon-entropy of GETSI (lower = more specific).

... Any additional columns from the original `retsi` and `getsi` inputs, suffixed with `_ATAC` and `_RNA` respectively (e.g., `avg_log2FC_ATAC`, `p_val_RNA`).

`plot_heatmap`

Plot a z-scored gene-by-cell-type heatmap

Description

Generates a heatmap using `ggplot2` to visualize expression or accessibility z-scores for genes across different cell types. Genes are ordered by their highest-scoring cell type, and then by maximum z-score within that group.

Usage

```
plot_heatmap(df_z, title_text, fill_label)
```

Arguments

<code>df_z</code>	A data frame with z-scored values. Must contain:
<code>gene_id</code>	Identifier of the gene. Must be an official gene symbol (e.g., <code>GAPDH</code>).
<code>cell_type</code>	Cell type or cluster label (e.g., <code>Acinar</code>)
<code>z_score</code>	Numeric. Z-scored values of the specificity score (e.g., <code>RETSI_z</code> , <code>GETSI_z</code>)
<code>title_text</code>	Character. Title of the heatmap.
<code>fill_label</code>	Character. Legend label for the color scale.

Value

A `ggplot2` object representing the heatmap.

See Also

[prepare_heatmap_data](#), [spicey_heatmap](#)

`prepare_heatmap_data` *Prepare data for SPICEY heatmap*

Description

Filters and processes a gene–cell-type matrix for heatmap visualization. Computes z-scores for a specified score column, selects the top `n` genes per cell type, and returns a summary matrix suitable for plotting.

Usage

```
prepare_heatmap_data(df, score_col, top_n)
```

Arguments

<code>df</code>	A data frame with at least: <code>gene_id</code> , <code>cell_type</code> , and a score column.
<code>score_col</code>	Character. Name of the score column to z-score and rank.
<code>top_n</code>	Integer. Number of top-ranked genes per cell type.

Value

A data frame with: `gene_id`, `cell_type`, and `z_score`.

See Also

[plot_heatmap](#), [spicey_heatmap](#)

`rna`

Example single-cell RNA-seq differential expression data

Description

A toy example dataset representing single-cell RNA-seq differential expression results. Each row corresponds to a gene tested across one or more cell types. The dataset is formatted as a list of GRanges objects or data frames (convertible to GRanges), where each element contains differential expression statistics for a specific cell type. Multiple rows may exist per gene, each representing results in a different cell type.

Usage

```
data(rna)
```

Format

A data frame or GRanges-like object with the following required columns:

gene_id Identifier of the gene. Must be an official gene symbol (e.g., GAPDH)

avg_log2FC Average log2 fold-change of expression for the gene in the specific cell type

p_val_adj Adjusted p-value (e.g., FDR-corrected)

cell_type Cell type or cluster label associated with each measurement (e.g., Acinar)

Source

Precomputed using `FindMarkers()` (Wilcoxon test, via Presto if available) on control samples from the Human Pancreas Analysis Program (HPAP), using paired snATAC-seq and scRNA-seq data from three non-diabetic human donors.

<code>specificity_index</code>	<i>Calculate specificity scores for grouped features</i>
--------------------------------	--

Description

This function computes a specificity index for different features (e.g., genes or regions) based on differential expression/accessibility data. It rescales fold-change values and weights them by significance to quantify how specific a feature's activity is to a particular cell type.

Usage

```
specificity_index(da, group_col)
```

Arguments

<code>da</code>	A data.frame containing differential results with at least the following columns: avg_log2FC Average log2 fold-change of the feature (gene or region). cell_type Cell type or cluster label. (e.g., Acinar) [group_col] Column containing the feature identifier (e.g., gene_id or region) The name of this column must match the value passed to the group_col argument
<code>group_col</code>	A string specifying the name of the column in da that identifies each feature, such as gene_id for genes or region for ATAC peaks.

Value

A data.frame identical to the input but with additional columns:

avg_FC	Fold-change converted from log2 scale.
max_FC	Maximum fold-change observed within each feature group.
weight	Normalized significance weight derived from adjusted p-values.
norm_FC	Fold-change normalized by maximum fold-change in the group.
score	Specificity score computed as the product of normalized fold-change significantly weighted.

<code>SPICEY</code>	<i>SPICEY: Tissue specificity analysis for single-cell data</i> The SPICEY package provides a user-friendly pipeline for quantifying and visualizing tissue specificity from single-cell ATAC-seq and/or single cell RNA-seq datasets, typically processed with tools such as Seurat or Signac. The core outputs of SPICEY are two tissue specific metrics, combined with entropy-based measures.
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SPICEY: Tissue specificity analysis for single-cell data The SPICEY package provides a user-friendly pipeline for quantifying and visualizing tissue specificity from single-cell ATAC-seq and/or single cell RNA-seq datasets, typically processed with tools such as Seurat or Signac. The core outputs of SPICEY are two tissue specific metrics, combined with entropy-based measures.

Description

Computes tissue-specificity scores from differential accessibility (RETSI) and/or gene expression (GETSI) data obtained from single cell experiments. Supports:

- RETSI calculation from differential accessibility data in different cell types/clusters (scATAC-seq).
- GETSI calculation from differential expression data in different cell types/clusters (scRNA-seq).
- Optional integration of RETSI and GETSI scores by linking gene associations (see [annotate_with_nearest](#) or [annotate_with_coaccessibility](#))

Usage

```
SPICEY(atac = NULL, rna = NULL, annotation = NULL, verbose = TRUE)
```

Arguments

atac	Either a single <code>data.frame</code> or a named list of <code>data.frames</code> or <code>GRanges</code> where each element corresponds to a cell type. It should contain differential chromatin accessibility results with required columns: region_id Unique identifier of the region (e.g., chr1-5000-5800). avg_log2FC Average log2 fold-change for accessibility in that cell type. p_val_adj Adjusted p-value (e.g., FDR-corrected). cell_type Cell type or cluster label. Only necessary when input is a single <code>data.frame</code> . If input is a list, it will be generated from list names. Note that the same region may appear multiple times across cell types.
rna	Either a single <code>data.frame</code> or a named list of <code>data.frames</code> or <code>GRanges</code> where each element corresponds to a cell type. It should contain differential expression results, with required columns: gene_id Identifier of the gene. Must be an official gene symbol (e.g., GAPDH). The name of this column should match the <code>gene_id</code> argument. avg_log2FC Average log2 fold-change for the gene in that cell type. p_val_adj Adjusted p-value (e.g., FDR-corrected). cell_type Cell type or cluster label. Only necessary when input is a single <code>data.frame</code> . If input is a list, it will be generated from list names Note that the same gene may appear multiple times across cell types.
annotation	(Optional). A <code>data.frame</code> linking <code>gene_id</code> to <code>region_id</code> . They should have the same names provided in the respective parameters. This can be provided by the user or generated using the functions: annotate_with_nearest or annotate_with_coaccessibility . It should contain at least the following columns: region_id Unique identifier of the region (e.g., chr1-5000-5800). The name of this column should match the <code>region_id</code> argument. cell_type Cell type or cluster label. (e.g., Acinar) gene_id Identifier of the gene. Must be an official gene symbol (e.g., GAPDH). The name of this column should match the <code>gene_id</code> argument.
verbose	Logical; print messages (default TRUE).

Value

Depending on inputs, returns RETSI and/or GETSI data frames, optionally linked and annotated.

Examples

```

data(rna)
data(atac)

# Calculate RETSI only
retsi <- SPICEY(atac = atac)

# Calculate GETSI only
getsi <- SPICEY(rna = rna)

# Calculate both
both <- SPICEY(
  rna = rna,
  atac = atac
)

# Integrate RETSI and GETSI with nearest gene
library(TxDb.Hsapiens.UCSC.hg38.knownGene)
library(org.Hs.eg.db)
peaks <- unique(unlist(atac)[, c("region_id")])

annotation_near <- annotate_with_nearest(
  peaks = peaks,
  txdb = TxDb.Hsapiens.UCSC.hg38.knownGene,
  annot_dbi = org.Hs.eg.db,
  protein_coding_only = TRUE,
  verbose = TRUE,
  add_tss_annotation = FALSE,
  upstream = 2000,
  downstream = 2000
)
spicey_near <- SPICEY(
  rna = rna,
  atac = atac,
  annotation = annotation_near
)

# Integrate RETSI and GETSI with coaccessibility
data(cicero_links)

annotation_coacc <- annotate_with_coaccessibility(
  peaks = peaks,
  txdb = TxDb.Hsapiens.UCSC.hg38.knownGene,
  links_df = cicero_links,
  annot_dbi = org.Hs.eg.db,
  protein_coding_only = TRUE,
  verbose = TRUE,
  add_tss_annotation = FALSE,
  upstream = 2000,
  downstream = 2000
)
spicey_coacc <- SPICEY(
  rna = rna,
  atac = atac,
  annotation = annotation_coacc
)

```

`spicey_heatmap`*SPICEY heatmap for gene specificity across cell types*

Description

Visualizes gene-level specificity scores (RETSI and/or GETSI) across cell types using a z-scored heatmap representation. Depending on the chosen mode, the function can display either RETSI or GETSI scores independently, or compute and visualize a combined SPICEY score (mean z-score of RETSI and GETSI). If `spicey_measure = "SPICEY"` and `combined_zscore = TRUE`, RETSI and GETSI scores are scaled, averaged, and shown in a unified heatmap. Otherwise, separate heatmaps are produced for RETSI and GETSI, respectively.

Usage

```
spicey_heatmap(
  df,
  top_n = 5,
  spicey_measure = c("RETSI", "GETSI", "SPICEY"),
  combined_zscore = FALSE
)
```

Arguments

<code>df</code>	A data frame with at least the following columns:
<code>gene_id</code>	Identifier of the gene. Must be an official gene symbol (e.g., GAPDH). If you only have ATAC data, link to nearest gene (annotate_with_nearest) or using coaccessibility (annotate_with_coaccessibility).
<code>cell_type</code>	Cell type or cluster label (e.g., Acinar)
<code>RETSI</code>	Numeric. RETSI specificity scores (optional unless used).
<code>GETSI</code>	Numeric. GETSI specificity scores (optional unless used).
<code>top_n</code>	Integer. Number of top-ranked genes to include per cell type (default "5")
<code>spicey_measure</code>	Character. Score type to visualize. Must be one of the following: "RETSI" Only RETSI will be plotted. "GETSI" Only GETSI will be plotted. "SPICEY" Both RETSI and GETSI are used (requires both columns)
<code>combined_zscore</code>	Logical. Only relevant if <code>spicey_measure = "SPICEY"</code> . If TRUE, a single heatmap of mean RETSI/GETSI z-score is generated. If FALSE, two heatmaps are produced side by side (RETSI and GETSI).

Value

A ggplot2 object, or a patchwork layout if two heatmaps are returned.

See Also

[SPICEY](#), [prepare_heatmap_data](#), [plot_heatmap](#)

Examples

```
library(TxDb.Hsapiens.UCSC.hg38.knownGene)
library(org.Hs.eg.db)

data(rna)
data(atac)
data(cicero_links)

# Obtain annotation with coaccessibility
peaks <- unique(unlist(atac)[, c("region_id")])
annotation_coacc <- annotate_with_coaccessibility(
  peaks = peaks,
  txdb = TxDb.Hsapiens.UCSC.hg38.knownGene,
  links_df = cicero_links,
  annot_db = org.Hs.eg.db,
  protein_coding_only = TRUE,
  verbose = TRUE,
  add_tss_annotation = FALSE,
  upstream = 2000,
  downstream = 2000
)

# Obtain linked SPICEY measures
spicey_coacc <- SPICEY(
  rna = rna,
  atac = atac,
  annotation = annotation_coacc
)

# Make plots
rets1i <- spicey_coacc$RETSI |> dplyr::left_join(annotation_coacc, by = c("region_id"))
spicey_heatmap(rets1i, spicey_measure = "RETSI")

spicey_heatmap(spicey_coacc$GETSI, spicey_measure = "GETSI")

spicey_heatmap(spicey_coacc$linked, spicey_measure = "SPICEY", combined_zscore = FALSE)
spicey_heatmap(spicey_coacc$linked, spicey_measure = "SPICEY", combined_zscore = TRUE)
```

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