

Package ‘PopComm’

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Title Population-Level Cell-Cell Communication Analysis Tools

Version 0.1.0.1

Description Facilitates population-level analysis of ligand-receptor (LR) interactions using large-scale single-cell transcriptomic data. Identifies significant LR pairs and quantifies their interactions through correlation-based filtering and projection score computations. Designed for large-sample single-cell studies, the package employs statistical modeling, including linear regression, to investigate LR relationships between cell types. It provides a systematic framework for understanding cell-cell communication, uncovering regulatory interactions and signaling mechanisms. Offers tools for LR pair-level, sample-level, and differential interaction analyses, with comprehensive visualization support to aid biological interpretation. The methodology is described in a manuscript currently under review and will be referenced here once published or publicly available.

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Contents

boxplot_lr_group_comparison	2
circle_plot	4
dotplot_lr_continuous_group	5
dot_plot	6
filtered_lr_eg	7
filter_lr_all	7
filter_lr_single	10
heatmap_sample	12
lr_db	13
lr_linear_model_discrete	14
lr_scores_eg	15
metadata_eg	15
one_step_all	16
one_step_single	18
pca_sample	20
score_lr_all	21
score_lr_single	23

Index

26

boxplot_lr_group_comparison
Boxplot Comparison of Ligand-Receptor Interaction Scores Across Groups

Description

Generates a boxplot comparing LR (ligand-receptor) interaction scores across sample groups. with optional significance testing (t-test or Wilcoxon).

Usage

```
boxplot_lr_group_comparison(
  lr_scores,
  metadata,
  ligand,
  receptor,
  sender,
  receiver,
  group_by,
  score = c("normalized", "raw"),
  test = TRUE,
  paired = FALSE,
  test_method = c("wilcox.test", "t.test"),
  colors = c("#5fa9d1", "#154778"),
  title = NULL
)
```

Arguments

<code>lr_scores</code>	Data frame containing LR interaction scores per sample (data frame).
<code>metadata</code>	Data frame containing sample metadata (data frame).
<code>ligand</code>	Ligand gene name to filter (character).
<code>receptor</code>	Receptor gene name to filter (character).
<code>sender</code>	Sender cell type to filter (character).
<code>receiver</code>	Receiver cell type to filter (character).
<code>group_by</code>	Column name in <code>metadata</code> to group samples (character).
<code>score</code>	Use 'normalized' or 'raw' score (default: "normalized") (character).
<code>test</code>	Whether to add a statistical test annotation (logical, default: TRUE).
<code>paired</code>	Whether to treat the comparison as paired (logical, default: FALSE).
<code>test_method</code>	Statistical test to use: "t.test" or "wilcox.test" (default = "wilcox.test") (character).
<code>colors</code>	Vector of colors for groups (default: c("#5fa9d1", "#154778")).
<code>title</code>	Custom plot title (optional).

Value

A list containing:

- `plot` - ggplot object of the boxplot
- `df` - data frame used for plotting

Examples

```
# Boxplot of LR Score by group
data(lr_scores_eg)
data(metadata_eg)
res <- boxplot_lr_group_comparison(
  lr_scores_eg, metadata_eg,
  ligand = "TAC4", receptor = "TACR1",
  sender = "Perivascular", receiver = "Cardiac",
  group_by = "IFN_type", score = "normalized"
)
print(res$plot)
head(res$df)
```

circle_plot*Plot Circular Ligand-Receptor Interaction Network***Description**

Plots a circular ligand-receptor (LR) interaction network with curved directed edges. Nodes are arranged in a circle, and edge widths and colors represent interaction strengths.

Usage

```
circle_plot(
  filtered_lr,
  edge_width = c("count", "cor"),
  node_colors = NULL,
  show_self_interactions = TRUE,
  cutoff = 0
)
```

Arguments

<code>filtered_lr</code>	A data frame of ligand-receptor pairs from prior analysis (e.g., output of <code>filter_lr_all</code>), containing at least the columns "sender", "receiver", and "cor".
<code>edge_width</code>	Determines edge weights, either "cor" (correlation) or "count" (interaction count) (default: "count").
<code>node_colors</code>	Named vector mapping cell types to colors. Example: <code>c("Cardiac" = "#E41A1C", "Fibroblast" = "#377EB8")</code> . If <code>NULL</code> , uses default palette.
<code>show_self_interactions</code>	Logical indicating whether to display self-interactions (logical, default: <code>TRUE</code>).
<code>cutoff</code>	Minimum edge weight to display (numeric, default: 0).

Value

A recordedplot object representing the network plot.

Examples

```
# Plot Circular Cell-Cell Interaction Network
data(filtered_lr_eg)
p <- circle_plot(filtered_lr_eg, edge_width = "count", show_self_interactions = TRUE)
print(p)
```

```
dotplot_lr_continuous_group
```

Dotplot of Ligand-Receptor Interaction Scores Against Continuous Group Variable

Description

Creates a dotplot (scatter plot) of LR interaction scores against a continuous variable with optional regression line.

Usage

```
dotplot_lr_continuous_group(  
  lr_scores,  
  metadata,  
  ligand,  
  receptor,  
  sender,  
  receiver,  
  group_by,  
  score = c("normalized", "raw"),  
  point_size = 3,  
  point_color = "dodgerblue4",  
  add_regression = TRUE,  
  title = NULL  
)
```

Arguments

lr_scores	Data frame containing LR interaction scores per sample (data frame).
metadata	Data frame containing sample metadata (data frame).
ligand	Ligand gene name to filter (character).
receptor	Receptor gene name to filter (character).
sender	Sender cell type to filter (character).
receiver	Receiver cell type to filter (character).
group_by	Continuous variable column in <code>metadata</code> (e.g., age, severity score) (character).
score	Use 'normalized' or 'raw' score (default: "normalized") (character).
point_size	Size of the points in the plot (numeric, default: 3).
point_color	Color of the points in the plot (default: "dodgerblue4").
add_regression	Whether to add regression line (logical, default: TRUE).
title	Custom plot title (optional).

Value

A list containing:

- plot - ggplot object of the dotplot
- df - data frame used for plotting

Examples

```
# Dotplot of LR Score Against Continuous Group Variable
data(lr_scores_eg)
data(metadata_eg)
res <- dotplot_lr_continuous_group(
  lr_scores_eg, metadata_eg,
  ligand = "TAC4", receptor = "TACR1",
  sender = "Perivascular", receiver = "Cardiac",
  group_by = "IFNscore"
)
print(res$plot)
head(res$df)
```

dot_plot*Create Ligand-Receptor Interaction Dot Plot***Description**

Generates a dot plot to visualize ligand-receptor (LR) interaction. Dot sizes are scaled by the correlation coefficient and dot colors represent -log10(adjust.p). The function supports plotting the top interactions per sender-receiver pair or user-specified ligand-receptor pairs.

Usage

```
dot_plot(
  filtered_lr,
  top_n = 5,
  axis = c("LR-SR", "SR-LR"),
  type_scale = c("size", "radius"),
  selected_LR = NULL
)
```

Arguments

<code>filtered_lr</code>	A data frame containing ligand-receptor interaction data.
<code>top_n</code>	Integer specifying the number of top interactions to select per sender-receiver pair (numeric, default: 5).
<code>axis</code>	Character indicating the configuration of rows and columns in the plot. Options: "LR-SR" (default, rows = ligand-receptor pairs, columns = sender-receiver pairs) or "SR-LR".

type_scale	Character indicating the scaling method for the plot. Options: "size" (default, uses <code>scale_size()</code> for point scaling) or "radius" (uses <code>scale_radius()</code> for point scaling).
selected_LR	Optional character vector of ligand-receptor pair identifiers (e.g., <code>c("TIMP1_CD63", "DSCAM_DCC")</code>). If NULL, the top_n interactions per sender-receiver pair are used.

Value

A ggplot object representing the dot plot.

Examples

```
# Plot LR Interaction Dot Plot
data(filtered_lr_eg)
p <- dot_plot(filtered_lr_eg, axis = "LR-SR", type_scale = "size")
print(p)
```

filtered_lr_eg

Example for filtered_lr

Description

Example for filtered_lr

Usage

`filtered_lr_eg`

Format

An object of class `data.frame` with 5904 rows and 12 columns.

filter_lr_all

Filter and Analyze Ligand-Receptor Pair Correlations (All Cell Types)

Description

Filters ligand-receptor (LR) pairs and analyzes their correlations for all possible cell type pairs, and returns significant LR pairs based on user-defined thresholds.

Usage

```
filter_lr_all(
  rna,
  lr_database = PopComm::lr_db,
  sample_col,
  cell_type_col,
  min_cells = 50,
  min_samples = 10,
  min_cell_ratio = 0.1,
  min_sample_ratio = 0.1,
  cor_method = "spearman",
  adjust_method = "BH",
  min_adjust_p = 0.05,
  min_cor = 0,
  num_cores = 10,
  verbose = TRUE
)
```

Arguments

<code>rna</code>	A Seurat object containing single-cell RNA expression data.
<code>lr_database</code>	A data frame of ligand-receptor pairs with columns "ligand_gene_symbol" and "receptor_gene_symbol".
<code>sample_col</code>	Column name in Seurat metadata indicating sample identifiers (character).
<code>cell_type_col</code>	Column name in Seurat metadata indicating cell type classifications (character).
<code>min_cells</code>	Minimum cells required per sample for both sender and receiver (numeric, default 50).
<code>min_samples</code>	Minimum valid samples required to proceed (numeric, default 10).
<code>min_cell_ratio</code>	Minimum ratio of cells expressing ligand and receptor genes in sender or receiver cells (numeric, default 0.1).
<code>min_sample_ratio</code>	Minimum ratio of samples in which both the ligand and receptor genes must be expressed (numeric, default 0.1).
<code>cor_method</code>	Correlation method: "spearman" (default), "pearson", or "kendall".
<code>adjust_method</code>	P-value adjustment method (default "BH" for Benjamini-Hochberg). Options: "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none".
<code>min_adjust_p</code>	Adjusted p-value threshold for significance (numeric, default 0.05).
<code>min_cor</code>	Minimum correlation coefficient threshold (numeric, default 0). Must be ≥ 0 .
<code>num_cores</code>	Number of CPU cores for parallel processing (numeric, default 10). Automatically capped at (system cores - 1).
<code>verbose</code>	Logical indicating whether to print progress messages (logical, default: TRUE).

Value

A data frame includes LR pairs with sufficient correlation and expression support across samples.

ligand, receptor	Ligand and receptor gene symbols.
cor	Correlation coefficient.
p_val	Raw p-value.
adjust.p	Adjusted p-value.
sender, receiver	Sender and receiver cell types.
slope	Slope of the linear regression model.
intercept	Intercept of the linear regression model.

Rows are ordered by ascending adjust.p and descending cor.

Returns NULL if:

- No cell types are found in the metadata.
- Insufficient samples or cells remain after filtering.
- No ligand-receptor pairs pass the filtering thresholds.

Examples

```
# Long-running example (may take >10s)
seurat_object <- load_example_seurat()
data(lr_db)

# Analyzing ligand-receptor interactions between all cell types
result01a <- filter_lr_all(
  rna = seurat_object,
  lr_database = lr_db,
  sample_col = "sample",
  cell_type_col = "cell.type",
  min_cells = 20,
  min_samples = 10,
  min_adjust_p = 0.5,
  num_cores = 1,
  verbose = TRUE
)

if (!is.null(result01a)) {
  print(head(result01a))
}
```

filter_lr_single	<i>Filter and Analyze Ligand-Receptor Pair Correlations (Specified Sender and Receiver)</i>
------------------	---

Description

Filters ligand-receptor (LR) pairs and analyzes their correlations for specified sender and receiver cell types, and returns significant LR pairs based on user-defined thresholds.

Usage

```
filter_lr_single(
  rna,
  sender,
  receiver,
  lr_database = PopComm::lr_db,
  sample_col,
  cell_type_col,
  min_cells = 50,
  min_samples = 10,
  min_cell_ratio = 0.1,
  min_sample_ratio = 0.1,
  cor_method = "spearman",
  adjust_method = "BH",
  min_adjust_p = 0.05,
  min_cor = 0,
  num_cores = 10,
  verbose = TRUE
)
```

Arguments

<code>rna</code>	A Seurat object containing single-cell RNA expression data.
<code>sender</code>	Cell type designated as the ligand sender (character).
<code>receiver</code>	Cell type designated as the receptor receiver (character).
<code>lr_database</code>	A data frame of ligand-receptor pairs with columns "ligand_gene_symbol" and "receptor_gene_symbol".
<code>sample_col</code>	Column name in Seurat metadata indicating sample identifiers (character).
<code>cell_type_col</code>	Column name in Seurat metadata indicating cell type classifications (character).
<code>min_cells</code>	Minimum cells required per sample for both sender and receiver (numeric, default 50).
<code>min_samples</code>	Minimum valid samples required to proceed (numeric, default 10).
<code>min_cell_ratio</code>	Minimum ratio of cells expressing ligand and receptor genes in sender or receiver cells (numeric, default 0.1).

<code>min_sample_ratio</code>	Minimum ratio of samples in which both the ligand and receptor genes must be expressed (numeric, default 0.1).
<code>cor_method</code>	Correlation method: "spearman" (default), "pearson", or "kendall".
<code>adjust_method</code>	P-value adjustment method (default "BH" for Benjamini-Hochberg). Options: "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none".
<code>min_adjust_p</code>	Adjusted p-value threshold for significance (numeric, default 0.05).
<code>min_cor</code>	Minimum correlation coefficient threshold (numeric, default 0). Must be ≥ 0 .
<code>num_cores</code>	Number of CPU cores for parallel processing (numeric, default 10). Automatically capped at (system cores - 1).
<code>verbose</code>	Logical indicating whether to print progress messages (logical, default: TRUE).

Value

A data frame includes LR pairs with sufficient correlation and expression support across samples.

<code>ligand, receptor</code>	Ligand and receptor gene symbols.
<code>cor</code>	Correlation coefficient.
<code>p_val</code>	Raw p-value.
<code>adjust.p</code>	Adjusted p-value.
<code>sender, receiver</code>	Sender and receiver cell types.
<code>slope</code>	Slope of the linear regression model.
<code>intercept</code>	Intercept of the linear regression model.

Rows are ordered by ascending `adjust.p` and descending `cor`.

Returns NULL if:

- No cell types are found in the metadata.
- Insufficient samples or cells remain after filtering.
- No ligand-receptor pairs pass the filtering thresholds.

Examples

```
# Long-running example (may take >10s)
seurat_object <- load_example_seurat()
data(lr_db)

# Analyzing ligand-receptor interactions: Cardiac -> Perivascular
result01s <- filter_lr_single(
  rna = seurat_object,
  sender = "Cardiac",
  receiver = "Perivascular",
  lr_database = lr_db,
  sample_col = "sample",
  cell_type_col = "cell.type",
```

```

min_cells = 20,
min_samples = 10,
min_adjust_p = 0.5,
num_cores = 1,
verbose = TRUE
)

if (!is.null(result01s)) {
print(head(result01s))
}

```

heatmap_sample*Generate Heatmap of Ligand-Receptor Interaction Scores***Description**

This function generates a heatmap to visualize the ligand-receptor (LR) interaction scores across samples. Rows represent LR pairs and columns represent samples. Optionally, sample metadata can be used to annotate the columns.

Usage

```

heatmap_sample(
  lr_scores,
  metadata,
  score = c("normalized", "raw"),
  selected_sender = NULL,
  selected_receiver = NULL,
  selected_metadata = NULL
)

```

Arguments

- lr_scores** Data frame containing LR interaction scores per sample (data frame).
- metadata** Data frame containing sample metadata (data frame).
- score** Character string indicating which score to use: "normalized" (default) or "raw"
- selected_sender** Specific sender cell type to filter, default is None (use all) (character).
- selected_receiver** Specific receiver cell type to filter, default is None (use all) (character).
- selected_metadata** List of column names in `metadata` to annotate samples (default: None, use all)(character vector).

Value

Heatmap of average LR interaction scores per sample.

Examples

```
# Heatmap of LR Interaction Scores
data(lr_scores_eg)
data(metadata_eg)
p <- heatmap_sample(lr_scores_eg, metadata_eg, score = "normalized", selected_sender = "Cardiac",
                     selected_receiver = "Perivascular", selected_metadata = c("Sex", "Age_group", "IFN_type"))
print(p)
```

lr_db*Ligand-Receptor Pair Database*

Description

A comprehensive database of human ligand-receptor pairs with gene/protein identifiers and supporting evidence from literature. Data imported from `human_lr_pair.txt`.

Usage

`lr_db`

Format

A data frame with 3,398 rows (pairs) and 10 columns:

lr_pair Character. Unique identifier for ligand-receptor pair, formatted as "LIGAND_RECEPTOR" (e.g., "SEMA3F_PLXNA3")
ligand_gene_symbol Character. Official HGNC symbol of the ligand gene (e.g., "SEMA3F")
receptor_gene_symbol Character. Official HGNC symbol of the receptor gene (e.g., "PLXNA3")
ligand_gene_id Integer. Entrez Gene ID of the ligand gene (NCBI identifier)
receptor_gene_id Integer. Entrez Gene ID of the receptor gene (NCBI identifier)
ligand_ensembl_protein_id Character. Ensembl protein ID of the ligand (e.g., "ENSP00000002829")
receptor_ensembl_protein_id Character. Ensembl protein ID of the receptor (e.g., "ENSP00000358696")
ligand_ensembl_gene_id Character. Ensembl gene ID of the ligand (e.g., "ENSG00000001617")
receptor_ensembl_gene_id Character. Ensembl gene ID of the receptor (e.g., "ENSG00000130827")
evidence Character. PubMed IDs (PMIDs) supporting the interaction, comma-separated (e.g., "15721238")

Source

Source from CellTalkDB (PMID: 33147626).

lr_linear_model_discrete

Compare Ligand-Receptor Interaction Scores with Group Variable using Linear Regression

Description

Perform linear regression analysis to compare ligand-receptor (LR) interaction scores across groups, handling both continuous and binary group variables (ident1 vs ident2 or all others).

Usage

```
lr_linear_model_discrete(
  lr_scores,
  metadata,
  group_variable,
  ident1,
  ident2 = NULL,
  covariates = NULL,
  fdr_threshold = 0.05
)
```

Arguments

lr_scores	Data frame containing LR interaction scores per sample (data frame).
metadata	Data frame containing sample metadata (data frame).
group_variable	Column name in metadata to compare groups (categorical or continuous) (character).
ident1	If categorical, group to compare (coded as 1) (character).
ident2	Reference group or list of groups (coded as 0). If None, uses all others (character).
covariates	Optional list of covariate column names (character vector).
fdr_threshold	Significance cutoff for adjusted p-values (numeric, default: 0.05).

Value

Data frame with ligand, receptor, sender, receiver, coef (coefficient, logFC), p-values, and adjusted p-values.

Examples

```
# Long-running example (may take >10s)
data(lr_scores_eg)
data(metadata_eg)

res <- lr_linear_model_discrete(
```

```
lr_scores_eg, metadata_eg,  
group_variable = "IFN_type",  
indent1 = "high",  
covariates = c("Age_group", "Sex")  
)  
head(res)
```

lr_scores_eg

Example for lr_scores

Description

Example for lr_scores

Usage

lr_scores_eg

Format

An object of class `data.frame` with 377006 rows and 15 columns.

metadata_eg

Example for metadata

Description

Example for metadata

Usage

metadata_eg

Format

An object of class `data.frame` with 163 rows and 9 columns.

one_step_all	<i>Analyze Ligand-Receptor Pair Correlations and Projection Scores (Across All Cell Types)</i>
---------------------	--

Description

Performs integrated analysis of ligand-receptor (LR) pairs through two consecutive phases:

1. Filters LR pairs and analyzes correlations across all cell types.
2. Calculates projection scores based on regression models for valid pairs. Returns comprehensive results combining statistical metrics.

Usage

```
one_step_all(
  rna,
  lr_database,
  sample_col,
  cell_type_col,
  min_cells = 50,
  min_samples = 10,
  min_cell_ratio = 0.1,
  min_sample_ratio = 0.1,
  cor_method = "spearman",
  adjust_method = "BH",
  min_adjust_p = 0.05,
  min_cor = 0,
  num_cores = 10,
  verbose = TRUE
)
```

Arguments

rna	A Seurat object containing single-cell RNA expression data.
lr_database	A data frame of ligand-receptor pairs with columns "ligand_gene_symbol" and "receptor_gene_symbol".
sample_col	Column name in Seurat metadata indicating sample identifiers (character).
cell_type_col	Column name in Seurat metadata indicating cell type classifications (character).
min_cells	Minimum cells required per sample for both sender and receiver (numeric, default 50).
min_samples	Minimum valid samples required to proceed (numeric, default 10).
min_cell_ratio	Minimum ratio of cells expressing ligand and receptor genes in sender or receiver cells (numeric, default 0.1).
min_sample_ratio	Minimum ratio of samples in which both the ligand and receptor genes must be expressed (numeric, default 0.1).

<code>cor_method</code>	Correlation method: "spearman" (default), "pearson", or "kendall".
<code>adjust_method</code>	P-value adjustment method (default "BH" for Benjamini-Hochberg). Options: "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none".
<code>min_adjust_p</code>	Adjusted p-value threshold for significance (numeric, default 0.05).
<code>min_cor</code>	Minimum correlation coefficient threshold (numeric, default 0). Must be ≥ 0 .
<code>num_cores</code>	Number of CPU cores for parallel processing (numeric, default 10). Automatically capped at (system cores - 1).
<code>verbose</code>	Logical indicating whether to print progress messages (logical, default: TRUE).

Value

Two data frames with columns:

<code>ligand, receptor</code>	Ligand and receptor gene symbols (res1/res2).
<code>cor</code>	Correlation coefficient (res1/res2).
<code>p_val</code>	Raw p-value (res1/res2).
<code>adjust.p</code>	Adjusted p-value (res1/res2).
<code>sender, receiver</code>	Sender and receiver cell types (res1/res2).
<code>slope</code>	Slope of the linear regression model (res1/res2).
<code>intercept</code>	Intercept of the linear regression model (res1/res2).
<code>sample</code>	Sample identifier (res2).
<code>score</code>	Projection score (raw co-expression intensity) (res2).
<code>normalized_score</code>	Normalized score scaled between 0-1 (res2).

Returns NULL if:

- No cell types are found in the metadata.
- Insufficient samples or cells remain after filtering.
- No ligand-receptor pairs pass the filtering thresholds.
- One or both of the specified sender and receiver cell types are missing in the data.
- Fewer than two valid samples remain after filtering based on minimum cell number per sample.

Examples

```
# Long-running example (may take >10s)
seurat_object <- load_example_seurat()
data(lr_db)

# Integrated analysis across all cell types
res_all <- one_step_all(
  rna = seurat_object,
  lr_database = lr_db,
```

```

sample_col = "sample",
cell_type_col = "cell.type",
min_cells = 20,
min_samples = 10,
min_adjust_p = 0.5,
num_cores = 1,
verbose = TRUE
)

if (!is.null(res_all)) {
  print(head(res_all$res1))
  print(head(res_all$res2))
}

```

one_step_single

*Analyze Ligand-Receptor Pair Correlations and Projection Scores
(Specified Sender and Receiver)*

Description

Performs integrated analysis of ligand-receptor (LR) pairs through two consecutive phases:

1. Filters LR pairs and analyzes correlations between specified cell types.
2. Calculates projection scores based on regression models for valid pairs. Returns comprehensive results combining statistical metrics.

Usage

```

one_step_single(
  rna,
  sender,
  receiver,
  lr_database = PopComm::lr_db,
  sample_col,
  cell_type_col,
  min_cells = 50,
  min_samples = 10,
  min_cell_ratio = 0.1,
  min_sample_ratio = 0.1,
  cor_method = "spearman",
  adjust_method = "BH",
  min_adjust_p = 0.05,
  min_cor = 0,
  num_cores = 10,
  verbose = TRUE
)

```

Arguments

<code>rna</code>	A Seurat object containing single-cell RNA expression data.
<code>sender</code>	Cell type designated as the ligand sender (character).
<code>receiver</code>	Cell type designated as the receptor receiver (character).
<code>lr_database</code>	A data frame of ligand-receptor pairs with columns "ligand_gene_symbol" and "receptor_gene_symbol".
<code>sample_col</code>	Column name in Seurat metadata indicating sample identifiers (character).
<code>cell_type_col</code>	Column name in Seurat metadata indicating cell type classifications (character).
<code>min_cells</code>	Minimum cells required per sample for both sender and receiver (numeric, default 50).
<code>min_samples</code>	Minimum valid samples required to proceed (numeric, default 10).
<code>min_cell_ratio</code>	Minimum ratio of cells expressing ligand and receptor genes in sender or receiver cells (numeric, default 0.1).
<code>min_sample_ratio</code>	Minimum ratio of samples in which both the ligand and receptor genes must be expressed (numeric, default 0.1).
<code>cor_method</code>	Correlation method: "spearman" (default), "pearson", or "kendall".
<code>adjust_method</code>	P-value adjustment method (default "BH" for Benjamini-Hochberg). Options: "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none".
<code>min_adjust_p</code>	Adjusted p-value threshold for significance (numeric, default 0.05).
<code>min_cor</code>	Minimum correlation coefficient threshold (numeric, default 0). Must be ≥ 0 .
<code>num_cores</code>	Number of CPU cores for parallel processing (numeric, default 10). Automatically capped at (system cores - 1).
<code>verbose</code>	Logical indicating whether to print progress messages (logical, default: TRUE).

Value

Two data frames with columns:

<code>ligand, receptor</code>	Ligand and receptor gene symbols (res1/res2).
<code>cor</code>	Correlation coefficient (res1/res2).
<code>p_val</code>	Raw p-value (res1/res2).
<code>adjust.p</code>	Adjusted p-value (res1/res2).
<code>sender, receiver</code>	Sender and receiver cell types (res1/res2).
<code>slope</code>	Slope of the linear regression model (res1/res2).
<code>intercept</code>	Intercept of the linear regression model (res1/res2).
<code>sample</code>	Sample identifier (res2).
<code>score</code>	Projection score (raw co-expression intensity) (res2).
<code>normalized_score</code>	Normalized score scaled between 0-1 (res2).

Returns NULL if:

- No cell types are found in the metadata.
- Insufficient samples or cells remain after filtering.
- No ligand-receptor pairs pass the filtering thresholds.
- One or both of the specified sender and receiver cell types are missing in the data.
- Fewer than two valid samples remain after filtering based on minimum cell number per sample.

Examples

```
# Long-running example (may take >10s)
seurat_object <- load_example_seurat()
data(lr_db)

# Integrated analysis with Cardiac -> Perivascular
res_single <- one_step_single(
  rna = seurat_object,
  sender = "Cardiac",
  receiver = "Perivascular",
  lr_database = lr_db,
  sample_col = "sample",
  cell_type_col = "cell.type",
  min_cells = 20,
  min_samples = 10,
  min_adjust_p = 0.5,
  num_cores = 1,
  verbose = TRUE
)
if (!is.null(res_single)) {
  print(head(res_single$res1))
  print(head(res_single$res2))
}
```

Description

This function performs principal component analysis (PCA) on ligand-receptor (LR) interaction scores across samples, and generates a scatter plot of the first two principal components. Optionally, sample metadata can be used to color the points.

Usage

```
pca_sample(
  lr_scores,
  metadata,
  selected_sender = NULL,
  selected_receiver = NULL,
  color_by = NULL,
  n_components = 2
)
```

Arguments

lr_scores	Data frame containing LR interaction scores per sample (data frame).
metadata	Data frame containing sample metadata (data frame).
selected_sender	Specific sender cell type to filter, default is None (use all) (character).
selected_receiver	Specific receiver cell type to filter, default is None (use all) (character).
color_by	metadata column name to color points in PCA plot (character).
n_components	Number of principal components to extract (numeric, default: 2).

Value

A list with two elements: the first is a ggplot2 PCA scatter plot and the second is the PCA results data frame.

Examples

```
# PCA of LR Interaction Scores
data(lr_scores_eg)
data(metadata_eg)
res <- pca_sample(lr_scores_eg, metadata_eg, selected_sender = "Cardiac",
  selected_receiver = "Perivascular", color_by = "IFN_type")

print(res$plot)
head(res$df)
```

Description

This function calculates the ligand-receptor (LR) projection scores between all combinations of sender and receiver cell types. The projection score is computed based on linear regression models, measuring the normalized distance of each sample's LR expression from the origin of the regression line.

Usage

```
score_lr_all(
  rna,
  filtered_lr,
  sample_col,
  cell_type_col,
  min_cells = 50,
  num_cores = 10,
  verbose = TRUE
)
```

Arguments

<code>rna</code>	A Seurat object containing single-cell RNA expression data.
<code>filtered_lr</code>	A data frame of ligand-receptor pairs from prior analysis (e.g., output of <code>filter_lr_single</code>). Must contain an "lr" column with pair identifiers in "Ligand_Receptor" format.
<code>sample_col</code>	Column name in Seurat metadata indicating sample identifiers (character).
<code>cell_type_col</code>	Column name in Seurat metadata indicating cell type classifications (character).
<code>min_cells</code>	Minimum cells required per sample for both sender and receiver (numeric, default 50).
<code>num_cores</code>	Number of CPU cores for parallel processing (numeric, default 10). Automatically capped at (system cores - 1).
<code>verbose</code>	Logical indicating whether to print progress messages (logical, default: TRUE).

Value

A data frame with projection scores per sample and LR pair. Columns:

All input from <code>filtered_lr</code>	
	Original columns provided by the user in <code>filtered_lr</code> .
<code>sample</code>	Sample identifier.
<code>score</code>	Projection score (raw co-expression intensity).
<code>normalized_score</code>	Normalized score scaled between 0-1.

Rows are ordered by `filtered_lr` columns and descending `score`.

Returns NULL if:

- No cell types are found in the metadata.
- One or both of the specified sender and receiver cell types are missing in the data.
- Fewer than two valid samples remain after filtering based on minimum cell number per sample.

Examples

```
# Long-running example (may take >10s)
seurat_object <- load_example_seurat()
data(lr_db)

# Analyzing ligand-receptor interactions between all cell types
result01a <- filter_lr_all(
  rna = seurat_object,
  lr_database = lr_db,
  sample_col = "sample",
  cell_type_col = "cell.type",
  min_cells = 20,
  min_samples = 10,
  min_adjust_p = 0.5,
  num_cores = 1,
  verbose = TRUE
)

# Analyzing ligand-receptor projection scores between all cell types
result02a <- score_lr_all(
  rna = seurat_object,
  filtered_lr = result01a,
  sample_col = "sample",
  cell_type_col = "cell.type",
  min_cells = 20,
  num_cores = 1,
  verbose = TRUE
)

if (!is.null(result02a)) {
  print(head(result02a))
}
```

score_lr_single

Analyze Ligand-Receptor Projection Scores (Specified Sender and Receiver)

Description

This function calculates the projection scores for ligand-receptor (LR) pairs between specified sender and receiver cell types. The projection score is computed based on linear regression models, measuring the normalized distance of each sample's LR expression from the origin of the regression line.

Usage

```
score_lr_single(
  rna,
```

```

    sender,
    receiver,
    filtered_lr,
    sample_col,
    cell_type_col,
    min_cells = 50,
    num_cores = 10,
    verbose = TRUE
)

```

Arguments

<code>rna</code>	A Seurat object containing single-cell RNA expression data.
<code>sender</code>	Cell type designated as the ligand sender (character).
<code>receiver</code>	Cell type designated as the receptor receiver (character).
<code>filtered_lr</code>	A data frame of filtered ligand-receptor pairs from prior analysis (e.g., output of <code>filter_lr_single</code>). Must contain an "lr" column with pair identifiers in "Ligand_Receptor" format.
<code>sample_col</code>	Column name in Seurat metadata indicating sample identifiers (character).
<code>cell_type_col</code>	Column name in Seurat metadata indicating cell type classifications (character).
<code>min_cells</code>	Minimum cells required per sample for both sender and receiver (numeric, default 50).
<code>num_cores</code>	Number of CPU cores for parallel processing (numeric, default 10). Automatically capped at (system cores - 1).
<code>verbose</code>	Logical indicating whether to print progress messages (logical, default: TRUE).

Value

A data frame with projection scores per sample and LR pair. Columns:

All input from <code>filtered_lr</code>	Original columns provided by the user in <code>filtered_lr</code> .
<code>sample</code>	Sample identifier.
<code>score</code>	Projection score (raw co-expression intensity).
<code>normalized_score</code>	Normalized score scaled between 0-1.

Rows are ordered by `filtered_lr` columns and descending `score`.

Returns NULL if:

- No cell types are found in the metadata.
- One or both of the specified sender and receiver cell types are missing in the data.
- Fewer than two valid samples remain after filtering based on minimum cell number per sample.

Examples

```
# Long-running example (may take >10s)
seurat_object <- load_example_seurat()
data(lr_db)

# Analyzing ligand-receptor interactions: Cardiac -> Perivascular
result01s <- filter_lr_single(
  rna = seurat_object,
  sender = "Cardiac",
  receiver = "Perivascular",
  lr_database = lr_db,
  sample_col = "sample",
  cell_type_col = "cell.type",
  min_cells = 20,
  min_samples = 10,
  min_adjust_p = 0.5,
  num_cores = 1,
  verbose = TRUE
)

# Analyzing ligand-receptor projection scores: Cardiac -> Perivascular
result02s <- score_lr_single(
  rna = seurat_object,
  sender = "Cardiac",
  receiver = "Perivascular",
  filtered_lr = result01s,
  sample_col = "sample",
  cell_type_col = "cell.type",
  min_cells = 20,
  num_cores = 1,
  verbose = TRUE
)

if (!is.null(result02s)) {
  print(head(result02s))
}
```

Index

- * **datasets**
 - filtered_lr_eg, [7](#)
 - lr_db, [13](#)
 - lr_scores_eg, [15](#)
 - metadata_eg, [15](#)
- boxplot_lr_group_comparison, [2](#)
- circle_plot, [4](#)
- dot_plot, [6](#)
- dotplot_lr_continuous_group, [5](#)
- filter_lr_all, [7](#)
- filter_lr_single, [10](#)
- filtered_lr_eg, [7](#)
- heatmap_sample, [12](#)
- lr_db, [13](#)
- lr_linear_model_discrete, [14](#)
- lr_scores_eg, [15](#)
- metadata_eg, [15](#)
- one_step_all, [16](#)
- one_step_single, [18](#)
- pca_sample, [20](#)
- score_lr_all, [21](#)
- score_lr_single, [23](#)