Package 'sccomp'

November 13, 2024

Description A robust and outlier-aware method for testing differential tissue composition from singlecell data. This model can infer changes in tissue composition and heterogeneity, and can produce realistic data simulations based on any existing dataset. This model can also transfer knowledge from a large set of integrated datasets to increase accuracy further. License GPL-3 URL https://github.com/MangiolaLaboratory/sccomp BugReports https://github.com/MangiolaLaboratory/sccomp/issues **Encoding** UTF-8 **Roxygen** list(markdown = TRUE) RoxygenNote 7.3.2 **Depends** R (>= 4.2.0) **Imports** instantiate (>= 0.2.3), callr, fs, stats, SeuratObject, SingleCellExperiment, parallel, dplyr, tidyr, purrr, magrittr, rlang, tibble, boot, lifecycle, stats, tidyselect, utils, ggplot2, ggrepel, patchwork, forcats, readr, scales, stringr, glue, withr, digest **Suggests** knitr, rmarkdown, BiocStyle, testthat (>= 3.0.0), markdown, knitr, loo, prettydoc, tidyseurat, tidySingleCellExperiment, bayesplot, posterior Additional_repositories https://mc-stan.org/r-packages/ SystemRequirements CmdStan (https://mc-stan.org/users/interfaces/cmdstan) biocViews Bayesian, Regression, DifferentialExpression, SingleCell LazvData true VignetteBuilder knitr git_url https://git.bioconductor.org/packages/sccomp git_branch RELEASE_3_20

Title Robust Outlier-aware Estimation of Composition and Heterogeneity

for Single-cell Data

git_last_commit 9d3562d

git_last_commit_date 2024-10-29
Repository Bioconductor 3.20

Version 1.10.0

counts_obj

Date/Publication 2024-11-12

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Contents

	sccomp_predict	
	sccomp_remove_outliers	
	sccomp_remove_unwanted_variation	
	•	
	sccomp_replicate	
	sccomp_test	
	sce_obj	22
	seurat_obj	23
	simulate_data	
Tural out		24
Index		26

Description

A tidy example dataset containing cell counts per cell group (cluster), sample, and phenotype for differential analysis. This dataset represents the counts of cells in various phenotypes and cell groups across multiple samples.

Usage

```
data(counts_obj)
```

Format

A tidy data frame with the following columns:

- sample: Factor, representing the sample identifier.
- type: Factor, indicating the sample type (e.g., benign, cancerous).
- phenotype: Factor, representing the cell phenotype (e.g., B_cell, HSC, etc.).
- count: Integer, representing the number of cells for each cell group within each sample.
- cell_group: Factor, representing the cell group (e.g., BM, B1, Dm, etc.).

get_output_samples 3

Value

A tibble representing cell counts per cluster, with columns for sample, type, phenotype, cell group, and counts.

get_output_samples

Get Output Samples from a Stan Fit Object

Description

This function retrieves the number of output samples from a Stan fit object, supporting different methods (MHC and Variational) based on the available data within the object.

Usage

```
get_output_samples(fit)
```

Arguments

fit

A stanfit object, which is the result of fitting a model via Stan.

Value

The number of output samples used in the Stan model. Returns from MHC if available, otherwise from Variational inference.

Examples

```
# Assuming 'fit' is a stanfit object obtained from running a Stan model
print("samples_count = get_output_samples(fit)")
```

multipanel_theme

multipanel_theme

Description

A custom ggplot2 theme used for creating publication-quality multi-panel plots. This theme modifies the appearance of plots by adjusting text sizes, spacing between panels, and axis formatting, ensuring better readability for complex figures.

Usage

```
data(multipanel_theme)
```

Format

A ggplot2 theme with the following adjustments:

- text: Font size adjustments for plot titles, axis labels, and legend text.
- panel.spacing: Adjusts the spacing between panels in multi-panel plots.
- axis.text: Customises axis text appearance for better readability.

4 plot.sccomp_tbl

Value

A ggplot2 theme object.

```
plot.sccomp_tbl
```

plot

Description

This function plots a summary of the results of the model.

Usage

```
## $3 method for class 'sccomp_tbl'
plot(
    x,
    significance_threshold = 0.05,
    test_composition_above_logit_fold_change = attr(.data,
        "test_composition_above_logit_fold_change"),
    ...
)
```

Arguments

х

A tibble including a cell_group name column | sample name column | read counts column | factor columns | Pvalue column | a significance column

significance_threshold

Numeric value specifying the significance threshold for highlighting differences. Default is 0.025.

test_composition_above_logit_fold_change

A positive integer. It is the effect threshold used for the hypothesis test. A value of 0.2 correspond to a change in cell proportion of 10% for a cell type with baseline proportion of 50%. That is, a cell type goes from 45% to 50%. When the baseline proportion is closer to 0 or 1 this effect thrshold has consistent value in the logit uncontrained scale.

... For internal use

Value

A ggplot

Examples

```
message("Use the following example after having installed install.packages(\verb|`"cmdstanr\|", repos = c(\verb|`"https://sizetalled.packages(|"cmdstanr\|", repos = c(|"https://sizetalled.packages(|"cmdstanr\|", repos = c(|"https://sizetalled.packages(|"https://sizetalled.packages(|"https://sizetalled.packages(|"https://sizetalled.packages(|"https://sizetalled.packages(|"https://sizetalled.packages(|"https://sizetalled.packages(|"https://sizet
```

```
if (instantiate::stan_cmdstan_exists()) {
  data("counts_obj")

  estimate = sccomp_estimate(
    counts_obj,
    ~ type, ~1, sample, cell_group, count,
```

plot_1D_intervals 5

```
cores = 1
)

# estimate |> plot()
}
```

plot_1D_intervals

Plot 1D Intervals for Cell-group Effects

Description

This function creates a series of 1D interval plots for cell-group effects, highlighting significant differences based on a given significance threshold.

Usage

```
plot_1D_intervals(
    .data,
    significance_threshold = 0.05,
    test_composition_above_logit_fold_change = attr(.data,
        "test_composition_above_logit_fold_change")
)
```

Arguments

. data Data frame containing the main data.

significance_threshold

Numeric value specifying the significance threshold for highlighting differences. Default is 0.025.

 $test_composition_above_logit_fold_change$

A positive integer. It is the effect threshold used for the hypothesis test. A value of 0.2 correspond to a change in cell proportion of 10% for a cell type with baseline proportion of 50%. That is, a cell type goes from 45% to 50%. When the baseline proportion is closer to 0 or 1 this effect thrshold has consistent value in the logit uncontrained scale.

Value

A combined plot of 1D interval plots.

Examples

```
# Example usage:
# plot_1D_intervals(.data, "cell_group", 0.025, theme_minimal())
```

6 plot_boxplot

plot_2D_intervals

Plot 2D Intervals for Mean-Variance Association

Description

This function creates a 2D interval plot for mean-variance association, highlighting significant differences based on a given significance threshold.

Usage

```
plot_2D_intervals(
    .data,
    significance_threshold = 0.05,
    test_composition_above_logit_fold_change = attr(.data,
        "test_composition_above_logit_fold_change")
)
```

Arguments

. data Data frame containing the main data.

significance_threshold

Numeric value specifying the significance threshold for highlighting differences. Default is 0.025.

test_composition_above_logit_fold_change

A positive integer. It is the effect threshold used for the hypothesis test. A value of 0.2 correspond to a change in cell proportion of 10% for a cell type with baseline proportion of 50%. That is, a cell type goes from 45% to 50%. When the baseline proportion is closer to 0 or 1 this effect thrshold has consistent value in the logit uncontrained scale.

Value

A ggplot object representing the 2D interval plot.

Examples

```
# Example usage:
# plot_2D_intervals(.data, "cell_group", theme_minimal(), 0.025)
```

plot_boxplot

Plot Boxplot of Cell-group Proportion

Description

This function creates a boxplot of cell-group proportions, optionally highlighting significant differences based on a given significance threshold.

plot_scatterplot 7

Usage

```
plot_boxplot(
    .data,
    data_proportion,
    factor_of_interest,
    .cell_group,
    .sample,
    significance_threshold = 0.05,
    my_theme
)
```

Arguments

data Data frame containing the main data. data_proportion

Data frame containing proportions of cell groups.

factor_of_interest

A factor indicating the biological condition of interest.

 $. \verb|cell_group| & The cell group to be analysed.$

. sample The sample identifier.

significance_threshold

Numeric value specifying the significance threshold for highlighting differences.

Default is 0.025.

my_theme A ggplot2 theme object to be applied to the plot.

Value

A ggplot object representing the boxplot.

Examples

```
# Example usage:
# plot_boxplot(.data, data_proportion, "condition", "cell_group", "sample", 0.025, theme_minimal())
```

plot_scatterplot

Plot Scatterplot of Cell-group Proportion

Description

This function creates a scatterplot of cell-group proportions, optionally highlighting significant differences based on a given significance threshold.

```
plot_scatterplot(
   .data,
   data_proportion,
   factor_of_interest,
   .cell_group,
   .sample,
```

8 sccomp_boxplot

```
significance_threshold = 0.05,
my_theme
)
```

Arguments

. data Data frame containing the main data.

data_proportion

Data frame containing proportions of cell groups.

factor_of_interest

A factor indicating the biological condition of interest.

. sample The sample identifier.

significance_threshold

Numeric value specifying the significance threshold for highlighting differences.

Default is 0.025.

my_theme A ggplot2 theme object to be applied to the plot.

Value

A ggplot object representing the scatterplot.

Examples

```
# Example usage:
# plot_scatterplot(.data, data_proportion, "condition", "cell_group", "sample", 0.025, theme_minimal())
```

sccomp_boxplot

sccomp_boxplot

Description

This function plots a boxplot of the results of the model.

```
sccomp_boxplot(
   .data,
   factor,
   significance_threshold = 0.05,
   test_composition_above_logit_fold_change = attr(.data,
        "test_composition_above_logit_fold_change")
)
```

Arguments

.data A tibble including a cell_group name column | sample name column | read

counts column | factor columns | Pvalue column | a significance column

factor A character string for a factor of interest included in the model

 $significance_threshold$

A real. FDR threshold for labelling significant cell-groups.

test_composition_above_logit_fold_change

A positive integer. It is the effect threshold used for the hypothesis test. A value of 0.2 correspond to a change in cell proportion of 10% for a cell type with baseline proportion of 50%. That is, a cell type goes from 45% to 50%. When the baseline proportion is closer to 0 or 1 this effect thrshold has consistent value in the logit uncontrained scale.

Value

A ggplot

Examples

 $message("Use the following example after having installed install.packages(\"cmdstanr\", repos = c(\"https://solution.packages(\"cmdstanr\", repos = c(\"https://solution.packages(\$

```
if (instantiate::stan_cmdstan_exists()) {
   data("counts_obj")

   estimate = sccomp_estimate(
     counts_obj,
     ~ type, ~1, sample, cell_group, count,
     cores = 1
   ) |>
   sccomp_test()

# estimate |> sccomp_boxplot()
}
```

sccomp_calculate_residuals

Calculate Residuals Between Observed and Predicted Proportions

Description

sccomp_calculate_residuals computes the residuals between observed cell group proportions and the predicted proportions from a fitted sccomp model. This function is useful for assessing model fit and identifying cell groups or samples where the model may not adequately capture the observed data. The residuals are calculated as the difference between the observed proportions and the predicted mean proportions from the model.

```
sccomp_calculate_residuals(.data)
```

Arguments

.data

A tibble of class sccomp_tbl, which is the result of sccomp_estimate(). This tibble contains the fitted model and associated data necessary for calculating residuals.

Details

The function performs the following steps:

- 1. Extracts the predicted mean proportions for each cell group and sample using sccomp_predict().
- 2. Calculates the observed proportions from the original count data.
- 3. Computes residuals by subtracting the predicted proportions from the observed proportions.
- 4. Returns a tibble containing the sample, cell group, residuals, and exposure (total counts per sample).

Value

A tibble (tbl) with the following columns:

- sample A character column representing the sample identifiers.
- **cell_group** A character column representing the cell group identifiers.
- **residuals** A numeric column representing the residuals, calculated as the difference between observed and predicted proportions.
- **exposure** A numeric column representing the total counts (sum of counts across cell groups) for each sample.

Examples

```
if (instantiate::stan_cmdstan_exists() && .Platform$OS.type == "unix") {
# Load example data
data("counts_obj")
# Fit the sccomp model
estimates <- sccomp_estimate(</pre>
  counts_obj,
  formula_composition = ~ type,
  formula_variability = ~1,
  .sample = sample,
  .cell_group = cell_group,
  .count = count,
  approximate_posterior_inference = "all",
  cores = 1
# Calculate residuals
residuals <- sccomp_calculate_residuals(estimates)</pre>
# View the residuals
print(residuals)
}
```

sccomp_estimate 11

sccomp_estimate

Main Function for SCCOMP Estimate

Description

The sccomp_estimate function performs linear modeling on a table of cell counts or proportions, which includes a cell-group identifier, sample identifier, abundance (counts or proportions), and factors (continuous or discrete). The user can define a linear model using an R formula, where the first factor is the factor of interest. Alternatively, sccomp accepts single-cell data containers (e.g., Seurat, SingleCellExperiment, cell metadata, or group-size) and derives the count data from cell metadata.

Usage

```
sccomp_estimate(
  .data,
  formula\_composition = ~1,
  formula_variability = ~1,
  .sample,
  .cell_group,
  .abundance = NULL,
  cores = detectCores(),
 bimodal_mean_variability_association = FALSE,
 percent_false_positive = 5,
  inference_method = "pathfinder",
 prior_mean = list(intercept = c(0, 1), coefficients = c(0, 1)),
 prior_overdispersion_mean_association = list(intercept = c(5, 2), slope = c(0, 0.6),
    standard_deviation = c(10, 20)),
  .sample_cell_group_pairs_to_exclude = NULL,
 output_directory = "sccomp_draws_files",
  verbose = TRUE,
  enable_loo = FALSE,
  noise_model = "multi_beta_binomial",
  exclude_priors = FALSE,
 use_data = TRUE,
 mcmc\_seed = sample(1e+05, 1),
 max_sampling_iterations = 20000,
 pass_fit = TRUE,
  . . . ,
  .count = NULL,
 approximate_posterior_inference = NULL,
  variational_inference = NULL
)
```

Arguments

.data A tibble including cell_group name column, sample name column, abundance column (counts or proportions), and factor columns.

formula_composition

A formula describing the model for differential abundance.

12 sccomp_estimate

formula_variability

A formula describing the model for differential variability.

A column name as a symbol for the sample identifier. .sample A column name as a symbol for the cell-group identifier. .cell_group

A column name as a symbol for the cell-group abundance, which can be counts .abundance

(>0) or proportions (between 0 and 1, summing to 1 across .cell_group).

cores Number of cores to use for parallel calculations.

bimodal_mean_variability_association

Logical, whether to model mean-variability as bimodal.

percent_false_positive

A real number between 0 and 100 for outlier identification.

inference_method

Character string specifying the inference method to use ('pathfinder', 'hmc', or 'variational').

prior_mean

A list specifying prior knowledge about the mean distribution, including inter-

cept and coefficients.

prior_overdispersion_mean_association

A list specifying prior knowledge about mean/variability association.

.sample_cell_group_pairs_to_exclude

A column name indicating sample/cell-group pairs to exclude.

output_directory

A character string specifying the output directory for Stan draws.

verbose Logical, whether to print progression details.

enable_loo Logical, whether to enable model comparison using the LOO package. noise_model A character string specifying the noise model (e.g., 'multi_beta_binomial').

exclude_priors Logical, whether to run a prior-free model. use_data Logical, whether to run the model data-free. mcmc_seed An integer seed for MCMC reproducibility.

max_sampling_iterations

Integer to limit the maximum number of iterations for large datasets.

Additional arguments passed to the cmdstanr::sample function.

pass_fit Logical, whether to include the Stan fit as an attribute in the output.

DEPRECATED. Use . abundance instead. .count

approximate_posterior_inference

DEPRECATED. Use inference_method instead.

variational_inference

DEPRECATED. Use inference_method instead.

Value

A tibble (tbl) with the following columns:

- cell_group The cell groups being tested.
- parameter The parameter being estimated from the design matrix described by the input formula_composition and formula_variability.
- factor The covariate factor in the formula, if applicable (e.g., not present for Intercept or contrasts).

sccomp_predict 13

- c_lower Lower (2.5%) quantile of the posterior distribution for a composition (c) parameter.
- c_effect Mean of the posterior distribution for a composition (c) parameter.
- c_upper Upper (97.5%) quantile of the posterior distribution for a composition (c) parameter.
- c_pH0 Probability of the null hypothesis (no difference) for a composition (c). This is not a p-value.
- c_FDR False-discovery rate of the null hypothesis for a composition (c).
- c_n_eff Effective sample size for a composition (c) parameter.
- c_R_k_hat R statistic for a composition (c) parameter, should be within 0.05 of 1.0.
- v_lower Lower (2.5%) quantile of the posterior distribution for a variability (v) parameter.
- v_effect Mean of the posterior distribution for a variability (v) parameter.
- v_upper Upper (97.5%) quantile of the posterior distribution for a variability (v) parameter.
- v_pH0 Probability of the null hypothesis for a variability (v).
- v_FDR False-discovery rate of the null hypothesis for a variability (v).
- v_n_eff Effective sample size for a variability (v) parameter.
- v_R_k_hat R statistic for a variability (v) parameter.
- count_data Nested input count data.

Examples

message("Use the following example after having installed cmdstanr with install.packages(\"cmdstanr\", repos =

```
if (instantiate::stan_cmdstan_exists()) {
  data("counts_obj")

  estimate <- sccomp_estimate(
    counts_obj,
    ~ type,
    ~1,
    sample,
    cell_group,
    abundance,
    cores = 1
  )
}</pre>
```

sccomp_predict

sccomp_predict

Description

This function replicates counts from a real-world dataset.

14 sccomp_predict

Usage

```
sccomp_predict(
   fit,
   formula_composition = NULL,
   new_data = NULL,
   number_of_draws = 500,
   mcmc_seed = sample(1e+05, 1),
   summary_instead_of_draws = TRUE
)
```

Arguments

fit The result of sccomp_estimate.

formula_composition

A formula. The formula describing the model for differential abundance, for example ~treatment. This formula can be a sub-formula of your estimated model; in this case all other factor will be factored out.

new_data

A sample-wise data frame including the column that represent the factors in your formula. If you want to predict proportions for 10 samples, there should be 10 rows. T

number_of_draws

An integer. How may copies of the data you want to draw from the model joint posterior distribution.

mcmc_seed

An integer. Used for Markov-chain Monte Carlo reproducibility. By default a random number is sampled from 1 to 999999. This itself can be controlled by set.seed()

 $\verb|summary_instead_of_draws| \\$

data("counts_obj")

Return the summary values (i.e. mean and quantiles) of the predicted proportions, or return single draws. Single draws can be helful to better analyse the uncertainty of the prediction.

Value

A tibble (tbl) with the following columns:

- cell_group A character column representing the cell group being tested.
- sample A factor column representing the sample name for which the predictions are made.
- proportion_mean A numeric column representing the predicted mean proportions from the model.
- **proportion_lower** A numeric column representing the lower bound (2.5%) of the 95% credible interval for the predicted proportions.
- **proportion_upper** A numeric column representing the upper bound (97.5%) of the 95% credible interval for the predicted proportions.

Examples

```
message("Use the following example after having installed install.packages(\"cmdstanr\", repos = c(\"https://s

if (instantiate::stan_cmdstan_exists() && .Platform$OS.type == "unix") {
```

```
sccomp_estimate(
  counts_obj,
  ~ type, ~1, sample, cell_group, count,
  cores = 1
) |>
sccomp_predict()
}
```

```
sccomp_proportional_fold_change
```

Calculate Proportional Fold Change for sccomp Data

Description

This function calculates the proportional fold change for single-cell composition data from sccomp analysis, comparing two conditions.

Usage

```
sccomp_proportional_fold_change(.data, formula_composition, from, to)
```

Arguments

Details

Note! This statistic is just descriptive and should not be used to define significance. Use sc-comp_test() for that. This statistics is just meant to help interpretation. While fold increase in proportion is easier to understand than fold change in logit space, the first is not linear (the same change for rare cell types does not necessarily have the same weight that for abundant cell types), while the latter is linear, and used to infer probabilities.

Value

A tibble with cell groups and their respective proportional fold change.

Examples

```
## Not run:
# Example usage
result <- sccomp_proportional_fold_change(sccomp_data, formula_composition, "healthy", "cancer")
## End(Not run)</pre>
```

```
sccomp_remove_outliers
```

sccomp_remove_outliers main

Description

The sccomp_remove_outliers function takes as input a table of cell counts with columns for cell-group identifier, sample identifier, integer count, and factors (continuous or discrete). The user can define a linear model using an input R formula, where the first factor is the factor of interest. Alternatively, sccomp accepts single-cell data containers (e.g., Seurat, SingleCellExperiment, cell metadata, or group-size) and derives the count data from cell metadata.

Usage

```
sccomp_remove_outliers(
    .estimate,
    percent_false_positive = 5,
    cores = detectCores(),
    inference_method = "pathfinder",
    output_directory = "sccomp_draws_files",
    verbose = TRUE,
    mcmc_seed = sample(1e+05, 1),
    max_sampling_iterations = 20000,
    enable_loo = FALSE,
    approximate_posterior_inference = NULL,
    variational_inference = NULL,
    ...
)
```

Arguments

.estimate

A tibble including a cell_group name column, sample name column, read counts column (optional depending on the input class), and factor columns.

percent_false_positive

A real number between 0 and 100 (not inclusive), used to identify outliers with a specific false positive rate.

cores

Integer, the number of cores to be used for parallel calculations.

inference_method

Character string specifying the inference method to use ('pathfinder', 'hmc', or 'variational').

output_directory

A character string specifying the output directory for Stan draws.

verbose

Logical, whether to print progression details.

mcmc_seed

Integer, used for Markov-chain Monte Carlo reproducibility. By default, a random number is sampled from 1 to 999999.

max_sampling_iterations

Integer, limits the maximum number of iterations in case a large dataset is used, to limit computation time.

enable_loo Logical, whether to enable model comparison using the R package LOO. This is useful for comparing fits between models, similar to ANOVA.

approximate_posterior_inference

DEPRECATED, use the variational_inference argument.

variational_inference

Logical, whether to use variational Bayes for posterior inference. It is faster and convenient. Setting this argument to FALSE runs full Bayesian (Hamiltonian Monte Carlo) inference, which is slower but the gold standard.

... Additional arguments passed to the cmdstanr::sample function.

Value

A tibble (tbl), with the following columns:

- cell_group The cell groups being tested.
- parameter The parameter being estimated from the design matrix described by the input formula_composition and formula_variability.
- factor The covariate factor in the formula, if applicable (e.g., not present for Intercept or contrasts).
- c lower Lower (2.5%) quantile of the posterior distribution for a composition (c) parameter.
- c_effect Mean of the posterior distribution for a composition (c) parameter.
- c upper Upper (97.5%) quantile of the posterior distribution for a composition (c) parameter.
- c_n_eff Effective sample size, the number of independent draws in the sample. The higher, the better.
- c_R_k_hat R statistic, a measure of chain equilibrium, should be within 0.05 of 1.0.
- v_lower Lower (2.5%) quantile of the posterior distribution for a variability (v) parameter.
- v_effect Mean of the posterior distribution for a variability (v) parameter.
- v_upper Upper (97.5%) quantile of the posterior distribution for a variability (v) parameter.
- v_n_eff Effective sample size for a variability (v) parameter.
- v_R_k_hat R statistic for a variability (v) parameter, a measure of chain equilibrium.
- count_data Nested input count data.

Examples

message("Use the following example after having installed install.packages(\"cmdstanr\", repos = c(\"https://s

}

```
sccomp\_remove\_unwanted\_variation \\ sccomp\_remove\_unwanted\_variation
```

Description

This function uses the model to remove unwanted variation from a dataset using the estimates of the model. For example, if you fit your data with the formula ~ factor_1 + factor_2 and use the formula ~ factor_1 to remove unwanted variation, the factor_2 effect will be factored out.

Usage

```
sccomp_remove_unwanted_variation(
   .data,
   formula_composition_keep = NULL,
   formula_composition = NULL,
   formula_variability = NULL,
   cores = detectCores()
)
```

Arguments

.data A tibble. The result of sccomp_estimate.

formula_composition_keep

A formula. The formula describing the model for differential abundance, for example ~type. In this case, only the effect of the type factor will be preserved, while all other factors will be factored out.

formula_composition

DEPRECATED. Use formula_composition_keep instead.

formula_variability

DEPRECATED. Use formula_variability_keep instead.

cores

Integer, the number of cores to be used for parallel calculations.

Value

A tibble (tbl) with the following columns:

- sample A character column representing the sample name for which data was adjusted.
- cell_group A character column representing the cell group being tested.
- adjusted_proportion A numeric column representing the adjusted proportion after removing unwanted variation.
- adjusted_counts A numeric column representing the adjusted counts after removing unwanted variation.
- **logit_residuals** A numeric column representing the logit residuals calculated after adjustment.

sccomp_replicate 19

Examples

 $message ("Use the following example after having installed cmdstanr with install.packages (\verb|`"cmdstanr|", repos = 1.00 to 1$

```
if (instantiate::stan_cmdstan_exists()) {
   data("counts_obj")

   estimates = sccomp_estimate(
      counts_obj,
      ~ type, ~1, sample, cell_group, count,
      cores = 1
   ) |>
      sccomp_remove_unwanted_variation()
}
```

sccomp_replicate

sccomp_replicate

Description

This function replicates counts from a real-world dataset.

Usage

```
sccomp_replicate(
   fit,
   formula_composition = NULL,
   formula_variability = NULL,
   number_of_draws = 1,
   mcmc_seed = sample(1e+05, 1)
)
```

Arguments

fi+

The result of sccomp estimate.

formula_composition

A formula. The formula describing the model for differential abundance, for example ~treatment. This formula can be a sub-formula of your estimated model; in this case all other factor will be factored out.

formula_variability

A formula. The formula describing the model for differential variability, for example ~treatment. In most cases, if differentially variability is of interest, the formula should only include the factor of interest as a large amount of data is needed to define variability depending to each factors. This formula can be a sub-formula of your estimated model; in this case all other factor will be factored out.

number_of_draws

An integer. How may copies of the data you want to draw from the model joint posterior distribution.

20 sccomp_test

mcmc_seed

An integer. Used for Markov-chain Monte Carlo reproducibility. By default a random number is sampled from 1 to 999999. This itself can be controlled by set.seed()

Value

A tibble tbl with cell_group-wise statistics

A tibble (tbl), with the following columns:

- cell_group A character column representing the cell group being tested.
- sample A factor column representing the sample name from which data was generated.
- **generated_proportions** A numeric column representing the proportions generated from the model.
- generated_counts An integer column representing the counts generated from the model.
- **replicate** An integer column representing the replicate number, where each row corresponds to a different replicate of the data.

Examples

```
\label{lem:message} \textit{``Use the following example after having installed install.packages(\'`cmdstanr'', repos = c(\''https://s'))} \\
```

```
if (instantiate::stan_cmdstan_exists() && .Platform$0S.type == "unix") {
  data("counts_obj")

sccomp_estimate(
  counts_obj,
  ~ type, ~1, sample, cell_group, count,
  cores = 1
  ) |>
  sccomp_replicate()
}
```

sccomp_test

 $sccomp_test$

Description

This function test contrasts from a sccomp result.

```
sccomp_test(
   .data,
   contrasts = NULL,
   percent_false_positive = 5,
   test_composition_above_logit_fold_change = 0.1,
   pass_fit = TRUE
)
```

21 sccomp_test

Arguments

A tibble. The result of sccomp_estimate. .data

A vector of character strings. For example if your formula is ~ 0 + treatment contrasts

and the factor treatment has values yes and no, your contrast could be "con-

strasts = c(treatmentyes - treatmentno)".

percent_false_positive

A real between 0 and 100 non included. This used to identify outliers with a specific false positive rate.

test_composition_above_logit_fold_change

A positive integer. It is the effect threshold used for the hypothesis test. A value of 0.2 correspond to a change in cell proportion of 10% for a cell type with baseline proportion of 50%. That is, a cell type goes from 45% to 50%. When the baseline proportion is closer to 0 or 1 this effect thrshold has consistent value

in the logit uncontrained scale.

A boolean. Whether to pass the Stan fit as attribute in the output. Because the pass_fit Stan fit can be very large, setting this to FALSE can be used to lower the memory

imprint to save the output.

Value

A tibble (tbl), with the following columns:

- cell_group The cell groups being tested.
- parameter The parameter being estimated from the design matrix described by the input formula_composition and formula_variability.
- · factor The covariate factor in the formula, if applicable (e.g., not present for Intercept or contrasts).
- c_lower Lower (2.5%) quantile of the posterior distribution for a composition (c) parameter.
- c_effect Mean of the posterior distribution for a composition (c) parameter.
- c_upper Upper (97.5%) quantile of the posterior distribution for a composition (c) parameter.
- c_pHO Probability of the c_effect being smaller or bigger than the test_composition_above_logit_fold_change argument.

c_FDR - False discovery rate of the c_effect being smaller or bigger than the test_composition_above_logit_fo

- argument. False discovery rate for Bayesian models is calculated differently from frequentists models, as detailed in Mangiola et al, PNAS 2023.
- c_n_eff Effective sample size, the number of independent draws in the sample. The higher, the better.
- c_R_k_hat R statistic, a measure of chain equilibrium, should be within 0.05 of 1.0.
- v_lower Lower (2.5%) quantile of the posterior distribution for a variability (v) parameter.
- v_effect Mean of the posterior distribution for a variability (v) parameter.
- v_upper Upper (97.5%) quantile of the posterior distribution for a variability (v) parameter.
- v_pH0 Probability of the v_effect being smaller or bigger than the test_composition_above_logit_fold_chan argument.
- v_FDR False discovery rate of the v_effect being smaller or bigger than the test_composition_above_logit_fo argument. False discovery rate for Bayesian models is calculated differently from frequentists models, as detailed in Mangiola et al, PNAS 2023.
- v_n_eff Effective sample size for a variability (v) parameter.

22 sce_obj

- v_R_k_hat R statistic for a variability (v) parameter, a measure of chain equilibrium.
- count_data Nested input count data.

#'

Examples

 $message ("Use the following example after having installed install.packages (\verb|`"cmdstanr\|", repos = c(\verb|`"https://stanr\|", repos = c(\|"https://stanr\|", repos = c(\|$

```
if (instantiate::stan_cmdstan_exists()) {
   data("counts_obj")

   estimates = sccomp_estimate(
      counts_obj,
      ~ 0 + type, ~1, sample, cell_group, count,
      cores = 1
   ) |>
   sccomp_test("typecancer - typebenign")
}
```

sce_obj

sce_obj

Description

Example SingleCellExperiment object containing gene expression data for 106,297 cells across two assays: counts and logcounts. The object includes metadata and assay data for RNA expression, which can be used directly in differential analysis functions like sccomp_glm.

Usage

```
data(sce_obj)
```

Format

A SingleCellExperiment object with the following structure:

- assays: Two assays: counts (raw RNA counts) and logcounts (log-transformed counts).
- rowData: No additional row-level metadata is present.
- **colData**: Metadata for each cell, including six fields: sample, type, nFeature_RNA, ident, and others.
- dim: 1 feature and 106,297 cells.
- colnames: Cell identifiers for all 106,297 cells.

Value

 $A \ {\tt SingleCellExperiment} \ object \ containing \ single-cell \ RNA \ expression \ data.$

seurat_obj 23

seurat_obj

seurat_obj

Description

Example Seurat object containing gene expression data for 106,297 cells across a single assay. The object includes RNA counts and data layers, but no variable features are defined. This dataset can be directly used with functions like sccomp_glm for differential abundance analysis.

Usage

```
data(seurat_obj)
```

Format

A Seurat object with the following structure:

- assays: Contains gene expression data. The active assay is RNA, with 1 feature and no variable features.
- layers: Two layers: counts and data, representing raw and processed RNA expression values, respectively.
- samples: 106,297 samples (cells) within the RNA assay.

Value

A Seurat object containing single-cell RNA expression data.

simulate_data

simulate_data

Description

This function simulates counts from a linear model.

```
simulate_data(
    .data,
    .estimate_object,
    formula_composition,
    formula_variability = NULL,
    .sample = NULL,
    .cell_group = NULL,
    .coefficients = NULL,
    variability_multiplier = 5,
    number_of_draws = 1,
    mcmc_seed = sample(1e+05, 1),
    cores = detectCores()
)
```

24 simulate_data

Arguments

.data

A tibble including a cell_group name column | sample name column | read counts column | factor columns | Pvalue column | a significance column

.estimate_object

The result of sccomp_estimate execution. This is used for sampling from real-data properties.

formula_composition

A formula. The sample formula used to perform the differential cell_group abundance analysis

formula_variability

A formula. The formula describing the model for differential variability, for example ~treatment. In most cases, if differentially variability is of interest, the formula should only include the factor of interest as a large amount of data is needed to define variability depending to each factors.

. sample A column name as symbol. The sample identifier

.cell_group A column name as symbol. The cell_group identifier

.coefficients The column names for coefficients, for example, c(b_0, b_1)

variability_multiplier

A real scalar. This can be used for artificially increasing the variability of the simulation for benchmarking purposes.

number_of_draws

An integer. How may copies of the data you want to draw from the model joint posterior distribution.

mcmc_seed

An integer. Used for Markov-chain Monte Carlo reproducibility. By default a random number is sampled from 1 to 999999. This itself can be controlled by set.seed()#' @param cores Integer, the number of cores to be used for parallel calculations.

cores

Integer, the number of cores to be used for parallel calculations.

Value

A tibble (tbl) with the following columns:

- sample A character column representing the sample name.
- type A factor column representing the type of the sample.
- phenotype A factor column representing the phenotype in the data.
- **count** An integer column representing the original cell counts.
- **cell_group** A character column representing the cell group identifier.
- **b_0** A numeric column representing the first coefficient used for simulation.
- **b_1** A numeric column representing the second coefficient used for simulation.
- **generated_proportions** A numeric column representing the generated proportions from the simulation.
- **generated_counts** An integer column representing the generated cell counts from the simulation.
- **replicate** An integer column representing the replicate number for each draw from the posterior distribution.

simulate_data 25

Examples

 $message ("Use the following example after having installed install.packages (\verb|`"cmdstanr\|", repos = c(\verb|`"https://stanr\|", repos = c(\|"https://stanr\|", repos =$

```
if (instantiate::stan_cmdstan_exists()) {
   data("counts_obj")
   library(dplyr)

   estimate = sccomp_estimate(
      counts_obj,
      ~ type, ~1, sample, cell_group, count,
      cores = 1
   )

# Set coefficients for cell_groups. In this case all coefficients are 0 for simplicity.
   counts_obj = counts_obj |> mutate(b_0 = 0, b_1 = 0)

# Simulate data
   simulate_data(counts_obj, estimate, ~type, ~1, sample, cell_group, c(b_0, b_1))
}
```

Index

```
* datasets
    counts_obj, 2
    multipanel_theme, 3
    sce_obj, 22
    seurat_obj, 23
counts_obj, 2
{\tt get\_output\_samples}, \\ 3
multipanel\_theme, 3
plot.sccomp_tbl, 4
plot_1D_intervals, 5
plot_2D_intervals, 6
plot_boxplot, 6
\verb|plot_scatterplot|, 7
sccomp\_boxplot, 8
sccomp_calculate_residuals, 9
sccomp_estimate, 11
sccomp_predict, 13
sccomp_proportional_fold_change, 15
sccomp\_remove\_outliers, 16
sccomp_remove_unwanted_variation, 18
sccomp_replicate, 19
sccomp\_test, 20
sce_obj, 22
seurat_obj, 23
simulate\_data, 23
```