

Package ‘romic’

October 14, 2022

Type Package

Title R for High-Dimensional Omic Data

Version 1.0.0

Maintainer Sean Hackett <sean@calicolabs.com>

Description Represents high-dimensional data as tables of features, samples and measurements, and a design list for tracking the meaning of individual variables. Using this format, filtering, normalization, and other transformations of a dataset can be carried out in a flexible manner. 'romic' takes advantage of these transformations to create interactive 'shiny' apps for exploratory data analysis such as an interactive heatmap.

Depends R (>= 3.2.3)

Imports checkmate, dplyr, ggplot2, glue, purrr, readr, reshape2, rlang, shiny (>= 1.5.0), stringr, tibble, tidyr (>= 1.0.0)

Suggests knitr, lazyeval, plotly, rmarkdown, usethis, testthat

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Encoding UTF-8

LazyData true

RoxygenNote 7.1.1

VignetteBuilder knitr

NeedsCompilation no

Author Sean Hackett [aut, cre] (<<https://orcid.org/0000-0002-9553-4341>>), Calico Life Sciences LLC [cph, fnd]

Repository CRAN

Date/Publication 2021-07-20 09:00:02 UTC

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`add_pca_loadings` *Add PCA Loadings*

Description

Add Principal Components Analysis Loadings to a tidy or triple omics dataset.

Usage

```
add_pca_loadings(  
  tomic,  
  value_var = NULL,  
  center_rows = TRUE,  
  npcs = NULL,  
  missing_val_method = "drop_samples"  
)
```

Arguments

<code>tomic</code>	Either a <code>tidy_omic</code> or <code>triple_omic</code> object
<code>value_var</code>	An abundance value to use with <code>hclust</code>
<code>center_rows</code>	center rows before performing PCA
<code>npcs</code>	number of principal component loadings to add to samples (default is number of samples)
<code>missing_val_method</code>	Approach to remove missing values: drop_features Drop features with missing values drop_samples Drop samples which are missing all features, then drop features impute Impute missing values

Value

A `tomic` object with principal components added to samples.

Examples

```
add_pca_loadings(brauer_2008_triple, npcs = 5)
```

app_flow	<i>Flow</i>
----------	-------------

Description

Using shiny comb through datasets by iterating between plotting steps, and lassoing steps to select points of interest.

Usage

```
app_flow(tomic)
```

Arguments

tomic Either a tidy_omic or triple_omic object

Value

A shiny app

Examples

```
if (interactive()) {
  # library(reactlog)
  # reactlog_enable()
  app_flow(brauer_2008_triple)
  # shiny::reactlogShow()
}
```

app_heatmap	<i>Interactive Heatmap</i>
-------------	----------------------------

Description

Generate a shiny interactive heatmap that allows for on demand filtering, ordering and faceting by variables of interest.

Usage

```
app_heatmap(tomic)
```

Arguments

tomic Either a tidy_omic or triple_omic object

Value

A shiny app

Examples

```
if (interactive()) {  
  app_heatmap(brauer_2008_tidy)  
}
```

app_pcs

PC Plot

Description

Generate a Shiny interactive scatter plot which allows visualization of features, measurements, and samples (with principal components added).

Usage

```
app_pcs(tomic)
```

Arguments

tomic Either a tidy_omic or triple_omic object

Value

A shiny app

Examples

```
if (interactive()) {  
  app_pcs(brauer_2008_tidy)  
}
```

`brauer_2008`*Brauer 2008*

Description

An RNA expression (microarray) dataset looking at how yeast gene expression changes as nutrient sources and nutrient richness changes.

`brauer_2008` formatted as a `tidy_omic` object

`brauer_2008` formatted as a `triple_omic` object

Usage

```
brauer_2008
```

```
brauer_2008_tidy
```

```
brauer_2008_triple
```

Format

A tibble with 18,000 rows and 8 columns:

name Common gene name

BP Gene ontology biological process of the gene

MF Gene ontology molecular function of the gene

sample Sample name

nutrient Which nutrient limits growth (Glucose, Nitrogen, Phosphorous, Sulfur, Uracil, Leucine

DR Dilution rate of the culture - basically how fast the cells are growing

expression Expression level of the gene, log2 observation relative to a replicate of G0.3

An object of class `tidy_omic` (inherits from `tomic`, `general`) of length 2.

An object of class `triple_omic` (inherits from `tomic`, `general`) of length 4.

Details

This version of the dataset contains only 500 genes randomly selected from the ~6K genes in the complete dataset.

Source

<https://pubmed.ncbi.nlm.nih.gov/17959824/>

center_tomic	<i>Center T* Omic</i>
--------------	-----------------------

Description

Center each measurement by subtracting the mean.

Usage

```
center_tomic(tomic, measurement_vars = "all")
```

Arguments

tomic	Either a tidy_omic or triple_omic object
measurement_vars	measurement variables to center

Value

A tomic object where one or more measurements have been centered on a feature-by-feature basis.

Examples

```
center_tomic(brauer_2008_tidy)
```

check_tidy_omic	<i>Check Tidy Omic</i>
-----------------	------------------------

Description

Check a tidy_omic dataset for consistency between the data and design and validate that the dataset follows the tidy_omic/tomic specification.

Usage

```
check_tidy_omic(tidy_omic, fast_check = TRUE)
```

Arguments

tidy_omic	an object of class tidy_omic produced by create_tidy_omic
fast_check	if TRUE then skip some checks which are slow and that are generally only needed when a tomic object is first created.

Value

Error and warning messages are printed and the input tidy_omic object is returned

check_tomic	<i>Check T*Omic</i>
-------------	---------------------

Description

Check a tidy or triple 'omic object for common pathologies, such as a mismatch between data and schema and non-uniqueness of primary keys.

Usage

```
check_tomic(tomic, fast_check = TRUE)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
fast_check	if TRUE then skip some checks which are slow and that are generally only needed when a tomic object is first created.

Value

0 invisibly

Examples

```
check_tomic(brauer_2008_triple)
```

check_triple_omic	<i>Check Triple Omic</i>
-------------------	--------------------------

Description

Check a triple omic dataset for consistency between the data and design and validate that the dataset follows the triple_omic/tomic specification.

Usage

```
check_triple_omic(triple_omic, fast_check = TRUE)
```

Arguments

triple_omic	an object of class triple_omic produced by create_triple_omic
fast_check	if TRUE then skip some checks which are slow and that are generally only needed when a tomic object is first created.

Value

Error and warning messages are printed and the input tidy_omic object is returned

```
convert_wide_to_tidy_omic
```

Convert Wide to Tidy Omic

Description

Convert a wide dataset of species' abundances (gene product, metabolites, lipids, ...) into a triple_omic dataset (one observation per row)

Usage

```
convert_wide_to_tidy_omic(  
  wide_df,  
  feature_pk,  
  feature_vars = NULL,  
  sample_var = "sample",  
  measurement_var = "abundance",  
  omic_type_tag = "general"  
)
```

Arguments

wide_df	a data.frame (or tibble) containing 1+ columns of feature attributes and many columns of samples
feature_pk	A unique identifier for features
feature_vars	a character vector of additional feature-level variables (or NULL if there are no additional variables)
sample_var	variable name to use for samples
measurement_var	variable name to use for measurements
omic_type_tag	an optional subtype of omic data: metabolomics, lipidomics, proteomics, genomics, general

Value

A tidy_omic object as produced by create_tidy_omic.

Examples

```
library(dplyr)  
  
wide_measurements <- brauer_2008_triple[["measurements"]] %>%  
  tidyr::spread(sample, expression)  
  
wide_df <- brauer_2008_triple[["features"]] %>%
```

```

left_join(wide_measurements, by = "name")

convert_wide_to_tidy_omic(wide_df,
  feature_pk = "name",
  feature_vars = c("BP", "MF", "systematic_name")
)

```

create_tidy_omic *Create Tidy Omic*

Description

A tidy omics object contains a formatted dataset and a summary of the experimental design.

Usage

```

create_tidy_omic(
  df,
  feature_pk,
  feature_vars = NULL,
  sample_pk,
  sample_vars = NULL,
  omic_type_tag = "general"
)

```

Arguments

df	a data.frame (or tibble) containing some combination of feature, sample and observation-level variables
feature_pk	A unique identifier for features
feature_vars	a character vector of additional feature-level variables (or NULL if there are no additional variables)
sample_pk	A unique identifier for samples
sample_vars	a character vector of additional sample-level variables (or NULL if there are no additional variables)
omic_type_tag	an optional subtype of omic data: metabolomics, lipidomics, proteomics, genomics, general

Value

An S3 tidy_omic/tomic object built on a list:

data A tibble with one row per measurement (i.e., features x samples)

design A list which organized the dataset's meta-data:

feature_pk variable specifying a unique feature

sample_pk variable specifying a unique sample

features tibble of feature attributes
samples tibble of sample attributes
measurements tibble of measurement attributes

Examples

```
library(dplyr)

measurement_df <- tidyr::expand_grid(
  feature_id = 1:10,
  sample_id = LETTERS[1:5]
) %>%
  dplyr::mutate(value = rnorm(n()))

feature_df <- tibble(
  feature_id = 1:10,
  feature_group = rep(c("a", "b"), each = 5)
)
sample_df <- tibble(
  sample_id = LETTERS[1:5],
  sample_group = c("a", "a", "b", "b", "b")
)

triple_omic <- create_triple_omic(
  measurement_df, feature_df, sample_df,
  "feature_id", "sample_id"
)
raw_tidy_omic <- triple_to_tidy(triple_omic)$data

create_tidy_omic(raw_tidy_omic,
  feature_pk = "feature_id",
  feature_vars = "feature_group", sample_pk = "sample_id",
  sample_vars = "sample_group"
)
```

create_triple_omic *Create Triple Omic*

Description

A triple omics class contains three data.frames, one for features, one for samples, and one for abundances. This is a good format when there is a large amount of meta data associated with features or samples.

Usage

```
create_triple_omic(
  measurement_df,
```

```

feature_df = NULL,
sample_df = NULL,
feature_pk,
sample_pk,
omic_type_tag = "general"
)

```

Arguments

measurement_df A data.frame (or tibble) of measurements - one row for each combination of feature and sample

feature_df A data.frame (or tibble) of features - one row per feature

sample_df A data.frame (or tibble) of samples - one row per sample

feature_pk A unique identifier for features

sample_pk A unique identifier for samples

omic_type_tag an optional subtype of omic data: metabolomics, lipidomics, proteomics, genomics, general

Details

for now primary keys are unique (rather than allowing for a multi-index)

Value

An S3 triple_omic/tomic object built on a list:

features A tibble of feature meta-data (one row per feature)

samples A tibble of sample meta-data (one row per sample)

measurements A tibble with one row per measurement (i.e., features x samples)

design A list which organized the dataset's meta-data:

feature_pk variable specifying a unique feature

sample_pk variable specifying a unique sample

features tibble of feature attributes

samples tibble of sample attributes

measurements tibble of measurement attributes

Examples

```

library(dplyr)

measurement_df <- tidyr::expand_grid(
  feature_id = 1:10,
  sample_id = LETTERS[1:5]
) %>%
  dplyr::mutate(value = rnorm(n()))

```

```
feature_df <- tibble(
  feature_id = 1:10,
  feature_group = rep(c("a", "b"), each = 5)
)
sample_df <- tibble(
  sample_id = LETTERS[1:5],
  sample_group = c("a", "a", "b", "b", "b")
)

triple_omic <- create_triple_omic(
  measurement_df, feature_df, sample_df,
  "feature_id", "sample_id"
)
```

export_tomic_as_tidy *Export T*Omic in Tidy Format*

Description

Export a data table including all fields from features, samples and measurements.

Usage

```
export_tomic_as_tidy(tomic, dir_path, name_preamble)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
dir_path	path to save outputs
name_preamble	start of output file name

Value

Export one table which is one row per peak, which includes all feature and sample attributes.

Examples

```
if (interactive()) {
  export_tomic_as_tidy(brauer_2008_triple, "/tmp", "brauer")
}
```

`export_tomic_as_triple`*Export T*Omic as Triple*

Description

Export features, samples and measurements tables

Usage

```
export_tomic_as_triple(tomic, dir_path, name_preamble)
```

Arguments

<code>tomic</code>	Either a <code>tidy_omic</code> or <code>triple_omic</code> object
<code>dir_path</code>	path to save outputs
<code>name_preamble</code>	start of output file name

Value

Export three tables:

- features: one row per features measured (i.e., a metabolite)
- sample: one row per sample
- measurements: one row per measurement (i.e., one metabolite in one sample)

Examples

```
if (interactive()) {  
  export_tomic_as_triple(brauer_2008_triple, "/tmp", "brauer")  
}
```

`export_tomic_as_wide` *Export T*Omic as Wide Data*

Description

abundances form a matrix with metabolites as rows and samples as columns. Use transpose to treat samples as rows filename

Usage

```
export_tomic_as_wide(
  tomic,
  dir_path,
  name_preamble,
  value_var = NULL,
  transpose = FALSE
)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
dir_path	path to save outputs
name_preamble	start of output file name
value_var	measurement variable to use for the matrix
transpose	if TRUE then samples will be stored as rows

Value

Export one table which contains metabolites as rows and samples as columns.

Examples

```
if (interactive()) {
  export_tomic_as_wide(brauer_2008_triple, "/tmp", "brauer")
}
```

 filterInput

Filter Input

Description

UI components for the filter module.

Usage

```
filterInput(id, filter_table)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
filter_table	table to filter

Value

A shiny UI

filterServer	<i>Filter Server</i>
--------------	----------------------

Description

Server components for the filter module.

Usage

```
filterServer(id, tidy_omic, filter_table)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
tidy_omic	an object of class tidy_omic produced by create_tidy_omic
filter_table	table to filter

Value

A tidy_omic with some features and/or samples filtered.

filter_tomic	<i>Filter T* Omics</i>
--------------	------------------------

Description

Filter a tidy or triple omic to entries of interest.

Usage

```
filter_tomic(
  tomic,
  filter_type,
  filter_table,
  filter_value,
  filter_variable = NULL
)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
filter_type	category filter filter_variable to categories specified in filter_value range filter filter_variable to using the range (i.e., lower and upper limit) provided in filter_value apply a quosure as a filter_value to a table of interest
filter_table	table where the filter should be applied
filter_value	values to filter based on
filter_variable	variable to apply the filter to

Value

A tomic object where a subset of features, samples or measurmenets have been filtered.

Examples

```
filter_tomic(
  brauer_2008_triple,
  filter_type = "category",
  filter_table = "features",
  filter_variable = "BP",
  filter_value = c("biological process unknown", "vacuolar acidification")
)

filter_tomic(
  brauer_2008_triple,
  filter_type = "category",
  filter_table = "samples",
  filter_variable = "DR",
  filter_value = 0.05
)

filter_tomic(
  brauer_2008_tidy,
  filter_type = "range",
  filter_table = "samples",
  filter_variable = "DR",
  filter_value = c(0, 0.2)
)

filter_tomic(
  brauer_2008_triple,
  filter_type = "quo",
  filter_table = "features",
  filter_value = rlang::quo(BP == "biological process unknown")
)
```

```
format_names_for_plotting
```

Format Names for Plotting

Description

Wrap long names over multiple lines so that they will look better on plots.

Usage

```
format_names_for_plotting(chars, width = 40, truncate_at = 80)
```

Arguments

chars	a character vector (or a variable that can be converted to one)
width	positive integer giving target line width in characters. A width less than or equal to 1 will put each word on its own line.
truncate_at	max character length

Value

a reformatted character vector of the same length as the input.

Examples

```
chars <- "Lorem ipsum dolor sit amet, consectetur adipiscing elit. Integer  
ac arcu semper erat porttitor egestas. Etiam sagittis, sapien at mattis."
```

```
format_names_for_plotting(chars)
```

```
get_design_tbl
```

Get Design Table

Description

Get a tabular summary of all variables.

Usage

```
get_design_tbl(tomic)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
-------	--

Value

a tibble reflecting the tomic object's design.

Examples

```
get_design_tbl(brauer_2008_triple)
```

ggBivOutput	<i>ggBivariate Output</i>
-------------	---------------------------

Description

UI components for the ggBivariate module.

Usage

```
ggBivOutput(id, return_brushed_points = FALSE)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
return_brushed_points	Return values selected on the plot

Value

A shiny UI

ggBivServer	<i>ggBivariate Server</i>
-------------	---------------------------

Description

Server components for the ggBivariate module.

Usage

```
ggBivServer(id, tomic, plot_table, return_brushed_points = FALSE)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
tomic	Either a tidy_omic or triple_omic object
plot_table	table containing the data to be plotted
return_brushed_points	Return values selected on the plot

Value

a `tomic_table` if `return_brushed_points` is `TRUE`, and 0 otherwise

<code>ggplotOutput</code>	<i>ggplot Output</i>
---------------------------	----------------------

Description

UI components for the `ggplot` module.

Usage

```
ggplotOutput(id)
```

Arguments

`id` An ID string that corresponds with the ID used to call the module's UI function.

Value

A shiny UI

<code>ggplotServer</code>	<i>ggplot Server</i>
---------------------------	----------------------

Description

Server components for the `ggplot` module.

Usage

```
ggplotServer(id, tomic, return_brushed_points = FALSE)
```

Arguments

`id` An ID string that corresponds with the ID used to call the module's UI function.
`tomic` Either a `tidy_omic` or `triple_omic` object
`return_brushed_points`
 Return values selected on the plot

Value

a tibble of selected observations if `return_brushed_points` is `TRUE`. Otherwise, returns `NULL`.

ggUnivOutput	<i>ggUnivariate Output</i>
--------------	----------------------------

Description

UI components for the ggUnivariate module.

Usage

```
ggUnivOutput(id, return_brushed_points = FALSE)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
return_brushed_points	Return values selected on the plot

Value

A shiny UI

ggUnivServer	<i>ggUnivariate Server</i>
--------------	----------------------------

Description

Server components for the ggUnivariate module

Usage

```
ggUnivServer(id, tomic, plot_table, return_brushed_points = FALSE)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
tomic	Either a tidy_omic or triple_omic object
plot_table	table containing the data to be plotted
return_brushed_points	Return values selected on the plot

Value

a tomic_table if return_brushed_points is TRUE, and 0 otherwise.

hclust_order	<i>Hierarchical clustering order</i>
--------------	--------------------------------------

Description

Format and hierarchically cluster a data.frame. If hclust could not normally be produced (usually because no samples are in common for a feature) pad the matrix with zeros and still calculate the distance

Usage

```
hclust_order(
  df,
  feature_pk,
  sample_pk,
  value_var,
  cluster_dim,
  distance_measure = "dist",
  hclust_method = "ward.D2"
)
```

Arguments

df	data.frame to cluster
feature_pk	variable uniquely defining a row
sample_pk	variable uniquely defining a sample
value_var	An abundance value to use with hclust
cluster_dim	rows, columns, or both
distance_measure	variable to use for computing dis-similarity
	corr pearson correlation
	dist euclidean distance
hclust_method	method from stats::hclust to use for clustering

Value

a list containing a hierarchically clustered set of rows and/or columns

Examples

```
library(dplyr)

df <- tidyr::crossing(letters = LETTERS, numbers = 1:10) %>%
  mutate(noise = rnorm(n()))
hclust_order(df, "letters", "numbers", "noise", "rows")
```

`infer_tomic_table_type`*Infer Tomic Table Type*

Description

From a `tomic_table`, choose whether it reflects features, samples or measurements

Usage

```
infer_tomic_table_type(tomic, tomic_table)
```

Arguments

<code>tomic</code>	Either a <code>tidy_omic</code> or <code>triple_omic</code> object
<code>tomic_table</code>	A table taken from a tidy (i.e., augmented measurements) or triple omic dataset

Value

features, samples or measurements

`lassoInput`*Lasso Input*

Description

UI components for the lasso module.

Usage

```
lassoInput(id)
```

Arguments

<code>id</code>	An ID string that corresponds with the ID used to call the module's UI function.
-----------------	--

Value

A shiny UI

lassoServer	<i>Lasso Server</i>
-------------	---------------------

Description

Take a subset of entries from a topic table (generally selected using the lasso function) and then either filter a topic object to these entries or tag the entries of interest with a user-specified variable.

Usage

```
lassoServer(id, topic, topic_table)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
topic	Either a tidy_omic or triple_omic object
topic_table	A table taken from a tidy (i.e., augmented measurements) or triple omic dataset

Value

A topic object amended based on the lasso selection.

organizeInput	<i>Organize Input</i>
---------------	-----------------------

Description

UI components for the organize input module.

Usage

```
organizeInput(id)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
----	--

Value

A shiny UI

organizeServer	<i>Organize Servers</i>
----------------	-------------------------

Description

Server components for the organize input module.

Usage

```
organizeServer(id, tidy_omic, feature_vars, sample_vars, value_var)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
tidy_omic	an object of class tidy_omic produced by create_tidy_omic
feature_vars	variables available for arranging features
sample_vars	variables available for arrange samples
value_var	An abundance value to use with hclust

Value

A tomic with sorted features and/or samples.

plotsaverInput	<i>Plot Saver Input</i>
----------------	-------------------------

Description

UI components for the plot saver module.

Usage

```
plotsaverInput(id)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
----	--

Value

a shiny UI

plotsaverServer *Plot Saver Server*

Description

Server components for the plot saver module.

Usage

```
plotsaverServer(id, grob)
```

Arguments

id	An ID string that corresponds with the ID used to call the module's UI function.
grob	a ggplot2 plot

Value

None

plot_bivariate *Bivariate Plot*

Description

Create a scatter or boxplot from a tomic dataset.

Usage

```
plot_bivariate(tomic_table, x_var, y_var, color_var = NULL)
```

Arguments

tomic_table	A table taken from a tidy (i.e., augmented measurements) or triple omic dataset
x_var	x-axis variable
y_var	y-axis variable
color_var	coloring variable (NULL to suppress coloring)

Value

a ggplot2 grob

Examples

```
library(dplyr)

brauer_augmented <- brauer_2008_tidy %>%
  add_pca_loadings(npcs = 5) %>%
  tomic_to("triple_omic")

tomic_table <- brauer_augmented$samples
plot_bivariate(tomic_table, "PC1", "PC2", "nutrient")
plot_bivariate(tomic_table, "PC1", "PC2", NULL)
plot_bivariate(tomic_table, "nutrient", "PC2", "nutrient")
```

plot_heatmap	<i>Plot Heatmap</i>
--------------	---------------------

Description

Generate a heatmap visualization of a features x samples matrix of measurements.

Usage

```
plot_heatmap(
  tomic,
  feature_var = NULL,
  sample_var = NULL,
  value_var = NULL,
  cluster_dim = "both",
  distance_measure = "dist",
  hclust_method = "ward.D2",
  change_threshold = Inf,
  plot_type = "grob"
)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
feature_var	variable from "features" to use as a unique feature label.
sample_var	variable from "samples" to use as a unique sample label.
value_var	which variable in "measurements" to use for quantification.
cluster_dim	rows, columns, or both
distance_measure	variable to use for computing dis-similarity
	corr pearson correlation
	dist euclidean distance
hclust_method	method from stats::hclust to use for clustering

change_threshold values with a more extreme absolute change will be thresholded to this value.
 plot_type plotly (for interactivity) or grob (for a static ggplot)

Value

a ggplot2 grob

Examples

```
library(dplyr)

tomic <- brauer_2008_triple %>%
  filter_tomic(
    filter_type = "category",
    filter_table = "features",
    filter_variable = "BP",
    filter_value = c(
      "protein biosynthesis",
      "rRNA processing", "response to stress"
    )
  )

plot_heatmap(
  tomic = tomic,
  value_var = "expression",
  change_threshold = 5,
  cluster_dim = "rows",
  plot_type = "grob",
  distance_measure = "corr",
  hclust_method = "complete"
)
```

plot_univariate *Univariate Plot*

Description

Create a histogram from a tomic dataset.

Usage

```
plot_univariate(tomic_table, x_var, color_var = NULL)
```

Arguments

tomic_table A table taken from a tidy (i.e., augmented measurements) or triple omic dataset
 x_var x-axis variable
 color_var coloring variable (NULL to suppress coloring)

Value

A ggplot2 grob

Examples

```
library(dplyr)

brauer_augmented <- brauer_2008_tidy %>%
  add_pca_loadings(npcs = 5) %>%
  tomic_to("triple_omic")

plot_univariate(brauer_augmented$samples, "PC1", "nutrient")
plot_univariate(brauer_augmented$measurements, "expression", NULL)
```

prepare_example_datasets

Prepare Example Datasets

Description

Format example datasets and add them to the package.

Usage

```
prepare_example_datasets(seed = 1234)
```

Arguments

seed a seed value used to reproducibly sample random genes.

Value

None; used for side-effects.

reconcile_triple_omic *Reconcile Triple Omic*

Description

If some samples, feature or measurements have been dropped; update other tables.

Usage

```
reconcile_triple_omic(triple_omic)
```

Arguments

triple_omic an object of class triple_omic produced by [create_triple_omic](#)

Value

a triple_omic object

remove_missing_values *Remove Missing Values*

Description

Account for missing values by dropping features, samples or using imputation.

Usage

```
remove_missing_values(  
  tomic,  
  value_var = NULL,  
  missing_val_method = "drop_samples"  
)
```

Arguments

tomic Either a tidy_omic or triple_omic object

value_var An abundance value to use with hclust

missing_val_method

 Approach to remove missing values:

drop_features Drop features with missing values

drop_samples Drop samples which are missing all features, then drop features

impute Impute missing values

Value

A tomic object where missing values have been accounted for.

Examples

```
remove_missing_values(brauer_2008_triple)
```

romic	<i>romic package</i>
-------	----------------------

Description

'romic' represents high-dimensional data as tables of features, samples and measurements, and a design list for tracking the meaning of individual variables. Using this format, filtering, normalization, and other transformations of a dataset can be carried out in a flexible manner. 'romic' takes advantage of these transformations to create interactive shiny apps for exploratory data analysis such as an interactive heatmap.

shiny_filter_test	<i>Shiny Filter Test</i>
-------------------	--------------------------

Description

Tests the shiny filter module as a stand-alone application.

Usage

```
shiny_filter_test(tidy_omic, filter_table = "features")
```

Arguments

tidy_omic	an object of class tidy_omic produced by create_tidy_omic
filter_table	table to filter

Value

A shiny app

Examples

```
if (interactive()) {  
  shiny_filter_test(brauer_2008_tidy)  
}
```

shiny_ggbiv_test *Shiny ggBivariate Test*

Description

Test the shiny ggBivariate module as a stand-alone application.

Usage

```
shiny_ggbiv_test(tomic, plot_table = "samples")
```

Arguments

tomic Either a tidy_omic or triple_omic object
plot_table table containing the data to be plotted

Value

a shiny app

Examples

```
if (interactive()) {  
  shiny_ggbiv_test(  
    add_pca_loadings(brauer_2008_triple, npcs = 5),  
    plot_table = "samples"  
  )  
  shiny_ggbiv_test(  
    brauer_2008_triple,  
    plot_table = "measurements"  
  )  
}
```

shiny_ggplot_test *Shiny ggplot Test*

Description

Test the shiny ggplot module as a stand-alone application.

Usage

```
shiny_ggplot_test(tomic)
```

Arguments

tomic Either a tidy_omic or triple_omic object

Value

A shiny app

Examples

```
if (interactive()) {  
  shiny_ggplot_test(add_pca_loadings(brauer_2008_triple, npcs = 5))  
  shiny_ggplot_test(brauer_2008_triple)  
}
```

shiny_gguniv_test *Shiny ggUnivariate Test*

Description

Test the shiny ggUnivariate module as a stand-alone application.

Usage

```
shiny_gguniv_test(tomic, plot_table = "samples")
```

Arguments

tomic Either a tidy_omic or triple_omic object
plot_table table containing the data to be plotted

Value

A shiny app

Examples

```
if (interactive()) {  
  shiny_gguniv_test(  
    add_pca_loadings(brauer_2008_triple, npcs = 5),  
    plot_table = "samples"  
  )  
  shiny_gguniv_test(brauer_2008_triple, plot_table = "measurements")  
  shiny_gguniv_test(brauer_2008_triple, plot_table = "features")  
}
```

shiny_lasso_test *Shiny Lasso Test*

Description

Tests the shiny lasso module as a stand-alone application.

Usage

```
shiny_lasso_test(tomic, tomic_table)
```

Arguments

tomic Either a tidy_omic or triple_omic object
tomic_table A table taken from a tidy (i.e., augmented measurements) or triple omic dataset

Value

A shiny app

Examples

```
if (interactive()) {
  tomic <- brauer_2008_triple
  tomic_table <- tomic[["samples"]] %>% dplyr::filter(nutrient == "G")
  shiny_lasso_test(tomic, tomic_table)
}
```

shiny_lasso_test_reactval
Shiny Lasso Test w/ Reactive Values

Description

Tests the shiny lasso module as a stand-alone application when the tomic is a reactiveVal.

Usage

```
shiny_lasso_test_reactval(tomic, tomic_table)
```

Arguments

tomic Either a tidy_omic or triple_omic object
tomic_table A table taken from a tidy (i.e., augmented measurements) or triple omic dataset

Value

A shiny app

Examples

```
if (interactive()) {
  tomic <- brauer_2008_triple
  tomic_table <- tomic[["samples"]] %>% dplyr::filter(nutrient == "G")
  shiny_lasso_test_reactval(tomic, tomic_table)

  tomic_table <- tomic[["measurements"]] %>% dplyr::filter(expression < -3)
  shiny_lasso_test_reactval(tomic, tomic_table)
}
```

shiny_organize_test *Shiny Organize Test*

Description

Tests the shiny organization module as stand-alone application.

Usage

```
shiny_organize_test(tidy_omic, feature_vars, sample_vars, value_var)
```

Arguments

tidy_omic	an object of class tidy_omic produced by create_tidy_omic
feature_vars	variables available for arranging features
sample_vars	variables available for arrange samples
value_var	An abundance value to use with hclust

Value

a shiny app

Examples

```
if (interactive()) {
  shiny_organize_test(
    brauer_2008_tidy,
    feature_vars = c("BP", "MF"),
    sample_vars = c("sample", "nutrient", "DR"),
    value_var = "expression"
  )
}
```

shiny_plotsaver_test *Shiny Plot Saver Test*

Description

Test the shiny plotsaver module as a stand-alone application.

Usage

```
shiny_plotsaver_test()
```

Value

a shiny app

Examples

```
if (interactive()) {  
  shiny_plotsaver_test()  
}
```

shiny_sort_test *Shiny Sort Test*

Description

Test the shiny sorting module as a stand-alone app.

Usage

```
shiny_sort_test(triple_omic, valid_sort_vars, value_var)
```

Arguments

triple_omic an object of class triple_omic produced by [create_triple_omic](#)
valid_sort_vars variables available for categorical arranging
value_var An abundance value to use with hclust

Value

a shiny app

Examples

```

if (interactive()) {
  shiny_sort_test(brauer_2008_triple,
    valid_sort_vars = c("sample", "nutrient", "DR"),
    value_var = "expression"
  )
}

```

`sortInput`*Sort Input*

Description

UI components for the sort module.

Usage

```
sortInput(id, sort_table)
```

Arguments

<code>id</code>	An ID string that corresponds with the ID used to call the module's UI function.
<code>sort_table</code>	table to sort

Value

A shiny UI

`sortServer`*Sort Server*

Description

Server components for the sort module.

Usage

```
sortServer(id, tomic, sort_table, valid_sort_vars = NULL, value_var = NULL)
```

Arguments

<code>id</code>	An ID string that corresponds with the ID used to call the module's UI function.
<code>tomic</code>	Either a <code>tidy_omic</code> or <code>triple_omic</code> object
<code>sort_table</code>	samples or features
<code>valid_sort_vars</code>	variables available for categorical arranging
<code>value_var</code>	An abundance value to use with <code>hclust</code>

Value

A sorted tomic object.

sort_tomic

Sort Triple Omic

Description

Sort a dataset's features or samples

Usage

```
sort_tomic(
  tomic,
  sort_type,
  sort_table,
  sort_variables = NULL,
  value_var = NULL
)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
sort_type	hclust Arrange samples by hierarchical clustering of a provided value_var arrange Arrange samples by the factor or alphanumeric ordering of a set of sort_variables
sort_table	samples or features
sort_variables	A set of attributes in sort_table to sort with in arrange.
value_var	An abundance value to use with hclust

Details

sort_tomic supports the reordering of features or samples using either hierarchical clustering or based on the levels of other variables. Sorting occurs by turning either the feature or sample primary key into a factor whose levels reflect the sort.

Value

A tomic object where feature or sample primary keys have been turned into a factor reflecting how they are sorted.

Examples

```
library(dplyr)

sort_tomic(brauer_2008_triple,
  sort_type = "arrange", sort_table = "samples",
  sort_variables = c("nutrient", "DR")
) %>%
  sort_tomic(
    sort_type = "hclust",
    sort_table = "features",
    value_var = "expression"
  )
```

sort_triple_arrange *Sort Triple Arrange*

Description

Sort a triple_omic object based on the values of one or more variables.

Usage

```
sort_triple_arrange(triple_omic, sort_table, sort_variables)
```

Arguments

triple_omic an object of class triple_omic produced by [create_triple_omic](#)
sort_table samples or features
sort_variables A set of attributes in sort_table to sort with in arrange.

Value

A triple_omic with sorted features or samples.

sort_triple_hclust *Sort Triple Hclust*

Description

Sort a triple_omic object using hierarchical clustering

Usage

```
sort_triple_hclust(triple_omic, sort_table, value_var)
```

Arguments

triple_omic an object of class triple_omic produced by [create_triple_omic](#)
sort_table samples or features
value_var An abundance value to use with hclust

Value

A triple_omic with clustered features or samples.

tidy_to_triple	<i>Tidy omic to triple omic</i>
----------------	---------------------------------

Description

Convert a tidy_omic object into a triple_omic object.

Usage

```
tidy_to_triple(tidy_omic)
```

Arguments

tidy_omic an object of class tidy_omic produced by [create_tidy_omic](#)

Details

The data table will be converted into features, samples, and measurements tables using the design to determine which variables belong in each table. The design will be preserved as-is.

Value

A triple_omic object as created by [create_triple_omic](#)

Examples

```
tidy_to_triple(brauer_2008_tidy)
```

tomic_sort_status	<i>T* Omic Sort Status</i>
-------------------	----------------------------

Description

Determine whether features &/or samples have been sorted and stored as ordered_featureId and ordered_sampleId.

Usage

```
tomic_sort_status(tomic)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
-------	--

Value

length 1 character string indicating whether the tomic is sorted.

Examples

```
tomic_sort_status(brauer_2008_tidy)
```

tomic_to	<i>T* Omic To</i>
----------	-------------------

Description

Takes in any romic representation of a dataset and returns a specific representation.

Usage

```
tomic_to(tomic, to_class)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
to_class	The class to return, either tidy_omic or triple_omic

Value

tomic transformed to to_class class (or un-transformed if it started that way).

Examples

```
tomic_to(brauer_2008_tidy, "triple_omic")
```

triple_to_tidy	<i>Triple Omic to Tidy Omic</i>
----------------	---------------------------------

Description

Convert a `triple_omic` object into a `tidy_omic` object.

Usage

```
triple_to_tidy(triple_omic)
```

Arguments

`triple_omic` an object of class `triple_omic` produced by [create_triple_omic](#)

Details

Features, samples and measurements will be merged into a single data table, and the design will be preserved as-is.

Value

A `tidy_omic` object as created by [create_tidy_omic](#).

Examples

```
library(dplyr)

measurement_df <- tidyr::expand_grid(
  feature_id = 1:10,
  sample_id = LETTERS[1:5]
) %>%
  dplyr::mutate(value = rnorm(n()))

feature_df <- tibble(
  feature_id = 1:10,
  feature_group = rep(c("a", "b"), each = 5)
)
sample_df <- tibble(
  sample_id = LETTERS[1:5],
  sample_group = c("a", "a", "b", "b", "b")
)

triple_omic <- create_triple_omic(
  measurement_df, feature_df, sample_df,
  "feature_id", "sample_id"
)
triple_to_tidy(triple_omic)
```

try_brushedPoints	<i>Try brushedPoints</i>
-------------------	--------------------------

Description

This function wraps brushedPoints in a try statement to catch cases where the brushing is out-of-sync with the df that is selected.

Usage

```
try_brushedPoints(...)
```

Arguments

... args to pass to [brushedPoints](#)

Value

a df of brushed points

update_tidy_omic	<i>Update Tidy Omic</i>
------------------	-------------------------

Description

Update a Tidy 'Omics data and schema to reflect newly added fields.

Usage

```
update_tidy_omic(tidy_omic, updated_tidy_data, new_variable_tables = c())
```

Arguments

tidy_omic an object of class tidy_omic produced by [create_tidy_omic](#)
updated_tidy_data a tibble of data to use to update tidy_omic.
new_variable_tables a named character vector of newly added variables in updated_tidy_data (names) and the table features, samples, measurements they apply to (values).

Value

a tidy_omic object with an updated schema and/or data.

Examples

```
library(dplyr)

tidy_omic <- brauer_2008_tidy
updated_tidy_data <- tidy_omic$data %>%
  mutate(new_sample_var = "foo") %>%
  select(-DR)
new_variable_tables <- c("new_sample_var" = "samples")
```

update_tomic	<i>Update T* Omic</i>
--------------	-----------------------

Description

Provide an updated features, samples or measurements table to a tomic.

Usage

```
update_tomic(tomic, tomic_table)
```

Arguments

tomic	Either a tidy_omic or triple_omic object
tomic_table	A table taken from a tidy (i.e., augmented measurements) or triple omic dataset

Value

A tomic object with updated features, samples or measurements.

Examples

```
library(dplyr)
updated_features <- brauer_2008_triple$features %>%
  dplyr::filter(BP == "biological process unknown") %>%
  dplyr::mutate(chromosome = purrr::map_int(systematic_name, function(x) {
    which(LETTERS == stringr::str_match(x, "Y([A-Z])")[2])
  }))

update_tomic(brauer_2008_triple, updated_features)
```

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