

Package ‘tidyHeatmap’

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Type Package

Title A Tidy Implementation of Heatmap

Version 1.8.1

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Description This is a tidy implementation for heatmap. At the moment it is based on the (great) package 'ComplexHeatmap'. The goal of this package is to interface a tidy data frame with this powerful tool. Some of the advantages are: Row and/or columns colour annotations are easy to integrate just specifying one parameter (column names). Custom grouping of rows is easy to specify providing a grouped tbl. For example: df %>% group_by(...). Labels size adjusted by row and column total number. Default use of Brewer and Viridis palettes.

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URL <https://www.r-project.org>,
<https://github.com/stemangiola/tidyHeatmap>

BugReports <https://github.com/stemangiola/tidyHeatmap>

Depends R (>= 3.6)

Imports methods,

```
stats,  
utils,  
dplyr (>= 0.8.5),  
magrittr (>= 1.5),  
tidyR (>= 1.0.3),  
rlang (>= 0.4.5),  
purrr (>= 0.3.3),  
tibble,  
ComplexHeatmap (>= 2.2.0),  
viridis (>= 0.5.1),  
circlize (>= 0.4.8),  
RColorBrewer (>= 1.1),  
grid,  
grDevices,  
lifecycle (>= 0.2.0),  
dendextend,  
patchwork
```

Suggests spelling,
 testthat,
 vdiffr,
 BiocManager,
 knitr,
 rmarkdown,
 qpdf,
 covr,
 roxygen2,
 forcats,
 ggplot2

VignetteBuilder knitr

RdMacros lifecycle

Biarch true

biocViews AssayDomain, Infrastructure

Encoding UTF-8

LazyData true

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add_annotation	add_annotation
----------------	----------------

Description

add_annotation() takes a `tbl` object and easily produces a ComplexHeatmap plot, with integration with `tibble` and `dplyr` frameworks.

Usage

```
add_annotation(
  my_input_heatmap,
  annotation,
  type = rep("tile", length(quo_names(annotation))),
  palette_discrete = list(),
  palette_continuous = list(),
  size = NULL,
  ...
)
```

Arguments

my_input_heatmap	A ‘InputHeatmap‘ formatted as <SAMPLE> <TRANSCRIPT> <COUNT> <...>
annotation	Vector of quotes
type	A character vector of the set c("tile", "point", "bar", "line")
palette_discrete	A list of character vectors. This is the list of palettes that will be used for horizontal and vertical discrete annotations. The discrete classification of annotations depends on the column type of your input tibble (e.g., character and factor).

palette_continuous

A list of character vectors. This is the list of palettes that will be used for horizontal and vertical continuous annotations. The continuous classification of annotations depends on the column type of your input tibble (e.g., integer, numerical, double).

size

A grid::unit object, e.g. unit(2, "cm"). This is the height or width of the annotation depending on the orientation.

...

The arguments that will be passed to top_annotation or left_annotation of the ComplexHeatmap container

Details

To be added.

Value

A ‘ComplexHeatmap’ object

add_attr

Add attribute to abject

Description

Add attribute to abject

Usage

```
add_attr(var, attribute, name)
```

Arguments

var A tibble

attribute An object

name A character name of the attribute

Value

A tibble with an additional attribute

add_bar	<i>Adds a bar annotation layer to a ‘InputHeatmap’, that on evaluation creates a ‘ComplexHeatmap’</i>
---------	---

Description

add_bar() from a ‘InputHeatmap’ object, adds a bar annotation layer.

Usage

```
add_bar(.data, .column, palette = NULL, size = NULL, ...)
## S4 method for signature 'InputHeatmap'
add_bar(.data, .column, palette = NULL, size = NULL, ...)
```

Arguments

.data	A ‘tbl_df‘ formatted as <ELEMENT> <FEATURE> <VALUE> <...>
.column	Vector of quotes
palette	A character vector of colors, or a function such as colorRamp2 (see examples).
size	A grid::unit object, e.g. unit(2, "cm"). This is the height or width of the annotation depending on the orientation.
...	The arguments that will be passed to top_annotation or left_annotation of the ComplexHeatmap container

Details

[Maturing]

It uses ‘ComplexHeatmap’ as visualisation tool.

Value

A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% add_bar(inflexion)
```

add_class	<i>Add class to abject</i>
-----------	----------------------------

Description

Add class to abject

Usage

```
add_class(var, name)
```

Arguments

var	A tibble
name	A character name of the attribute

Value

A tibble with an additional attribute

add_line	<i>Adds a line annotation layer to a 'InputHeatmap', that on evaluation creates a 'ComplexHeatmap'</i>
----------	--

Description

`add_line()` from a 'InputHeatmap' object, adds a line annotation layer.

Usage

```
add_line(.data, .column, palette = NULL, size = NULL, ...)
## S4 method for signature 'InputHeatmap'
add_line(.data, .column, palette = NULL, size = NULL, ...)
```

Arguments

.data	A 'tbl_df' formatted as <ELEMENT> <FEATURE> <VALUE> <...>
.column	Vector of quotes
palette	A character vector of colors, or a function such as colorRamp2 (see examples).
size	A grid::unit object, e.g. unit(2, "cm"). This is the height or width of the annotation depending on the orientation.
...	The arguments that will be passed to top_annotation or left_annotation of the ComplexHeatmap container

Details

[Maturing]

It uses 'ComplexHeatmap' as visualisation tool.

Value

A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% add_line(inflexion)
```

add_point

Adds a point annotation layer to a ‘InputHeatmap‘, that on evaluation creates a ‘ComplexHeatmap‘

Description

add_point() from a ‘InputHeatmap‘ object, adds a point annotation layer.

Usage

```
add_point(.data, .column, palette = NULL, size = NULL, ...)
## S4 method for signature 'InputHeatmap'
add_point(.data, .column, palette = NULL, size = NULL, ...)
```

Arguments

.data	A ‘tbl_df‘ formatted as <ELEMENT> <FEATURE> <VALUE> <...>
.column	Vector of quotes
palette	A character vector of colors, or a function such as colorRamp2 (see examples).
size	A grid::unit object, e.g. unit(2, "cm"). This is the height or width of the annotation depending on the orientation.
...	The arguments that will be passed to top_annotation or left_annotation of the ComplexHeatmap container

Details**[Maturing]**

It uses ‘ComplexHeatmap‘ as visualisation tool.

Value

A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% add_point(inflexion)
```

add_tile

Adds a tile annotation layer to a ‘InputHeatmap‘, that on evaluation creates a ‘ComplexHeatmap‘

Description

`add_tile()` from a ‘InputHeatmap‘ object, adds a tile annotation layer.

Usage

```
add_tile(.data, .column, palette = NULL, size = NULL, ...)
## S4 method for signature 'InputHeatmap'
add_tile(.data, .column, palette = NULL, size = NULL, ...)
```

Arguments

.data	A ‘tbl_df‘ formatted as <ELEMENT> <FEATURE> <VALUE> <...>
.column	Vector of quotes
palette	A character vector of colors, or a function such as <code>colorRamp2</code> (see examples).
size	A <code>grid::unit</code> object, e.g. <code>unit(2, "cm")</code> . This is the height or width of the annotation depending on the orientation.
...	The arguments that will be passed to <code>top_annotation</code> or <code>left_annotation</code> of the <code>ComplexHeatmap</code> container

Details**[Maturing]**

It uses ‘ComplexHeatmap‘ as visualisation tool.

Value

A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)

hm %>% add_tile(CAPRA_TOTAL)

hm %>% add_tile(inflection, palette = circlize::colorRamp2(c(0, 3, 10), c("white", "green")))
```

annot_to_list *annot_to_list*

Description

annot_to_list

Usage

```
annot_to_list(.data)
```

Arguments

.data A data frame

Value

A list

`as_ComplexHeatmap` *Creates a ‘ComplexHeatmap‘ object for less standard plot manipulation (e.g. changing legend position)*

Description

`as_ComplexHeatmap()` takes a ‘InputHeatmap‘ object and produces a ‘Heatmap‘ object

Usage

```
as_ComplexHeatmap(tidyHeatmap)

## S4 method for signature 'InputHeatmap'
as_ComplexHeatmap(tidyHeatmap)
```

Arguments

`tidyHeatmap` A ‘InputHeatmap‘ object from `tidyHeatmap::heatmap()` call

Details

[Maturing]

Value

A ‘ComplexHeatmap‘

Examples

```
tidyHeatmap::N52 |>
  tidyHeatmap::heatmap(
    .row = symbol_ct,
    .column = UBR,
    .value = `read count normalised log`,
  ) |>
  as_ComplexHeatmap()
```

`as_matrix`

Get matrix from tibble

Description

Get matrix from tibble

Usage

```
as_matrix(tbl, rownames = NULL, do_check = TRUE)
```

Arguments

tbl	A tibble
rownames	A character string of the rownames
do_check	A boolean

Value

A matrix

check_if_counts_is_na

Check whether there are NA counts

Description

Check whether there are NA counts

Usage

```
check_if_counts_is_na(.data, .abundance)
```

Arguments

.data	A tibble of read counts
.abundance	A character name of the read count column

Value

A tbl

check_if_duplicated_genes

Check whether there are duplicated genes/transcripts

Description

Check whether there are duplicated genes/transcripts

Usage

```
check_if_duplicated_genes(.data, .sample, .transcript, .abundance)
```

Arguments

.data	A tibble of read counts
.sample	A character name of the sample column
.transcript	A character name of the transcript/gene column
.abundance	A character name of the read count column

Value

A tbl

`check_if_wrong_input`

Check whether there are NA counts

Description

Check whether there are NA counts

Usage

```
check_if_wrong_input(.data, list_input, expected_type)
```

Arguments

<code>.data</code>	A tibble of read counts
<code>list_input</code>	A list
<code>expected_type</code>	A character string

Value

A `tbl`

`drop_class`

Remove class to abject

Description

Remove class to abject

Usage

```
drop_class(var, name)
```

Arguments

<code>var</code>	A tibble
<code>name</code>	A character name of the class

Value

A tibble with an additional attribute

```
error_if_log_transformed
```

Check whether a numeric vector has been log transformed

Description

Check whether a numeric vector has been log transformed

Usage

```
error_if_log_transformed(x, .abundance)
```

Arguments

x A numeric vector

.abundance A character name of the transcript/gene abundance column

Value

NA

```
get_abundance_norm_if_exists
```

Get column names either from user or from attributes

Description

Get column names either from user or from attributes

Usage

```
get_abundance_norm_if_exists(.data, .abundance)
```

Arguments

.data A tibble

.abundance A character name of the abundance column

Value

A list of column enquo or error

`get_elements` *Get column names either from user or from attributes*

Description

Get column names either from user or from attributes

Usage

```
get_elements(.data, .element, of_samples = TRUE)
```

Arguments

<code>.data</code>	A tibble
<code>.element</code>	A character name of the sample column
<code>of_samples</code>	A boolean

Value

A list of column enquo or error

`get_elements_features`
Get column names either from user or from attributes

Description

Get column names either from user or from attributes

Usage

```
get_elements_features(.data, .element, .feature, of_samples = TRUE)
```

Arguments

<code>.data</code>	A tibble
<code>.element</code>	A character name of the sample column
<code>.feature</code>	A character name of the transcript/gene column
<code>of_samples</code>	A boolean

Value

A list of column enquo or error

```
get_elements_features_abundance
    Get column names either from user or from attributes
```

Description

Get column names either from user or from attributes

Usage

```
get_elements_features_abundance (
  .data,
  .element,
  .feature,
  .abundance,
  of_samples = TRUE
)
```

Arguments

.data	A tibble
.element	A character name of the sample column
.feature	A character name of the transcript/gene column
.abundance	A character name of the read count column
of_samples	A boolean

Value

A list of column enquo or error

```
get_sample_counts  Get column names either from user or from attributes
```

Description

Get column names either from user or from attributes

Usage

```
get_sample_counts(.data, .sample, .abundance)
```

Arguments

.data	A tibble
.sample	A character name of the sample column
.abundance	A character name of the read count column

Value

A list of column enquo or error

```
get_sample_transcript
```

Get column names either from user or from attributes

Description

Get column names either from user or from attributes

Usage

```
get_sample_transcript(.data, .sample, .transcript)
```

Arguments

- .data A tibble
- .sample A character name of the sample column
- .transcript A character name of the transcript/gene column

Value

A list of column enquo or error

```
get_sample_transcript_counts
```

Get column names either from user or from attributes

Description

Get column names either from user or from attributes

Usage

```
get_sample_transcript_counts(.data, .sample, .transcript, .abundance)
```

Arguments

- .data A tibble
- .sample A character name of the sample column
- .transcript A character name of the transcript/gene column
- .abundance A character name of the read count column

Value

A list of column enquo or error

```
get_x_y_annotation_columns
  get_x_y_annotation_columns
```

Description

`get_x_y_annotation_columns`

Usage

```
get_x_y_annotation_columns(.data, .column, .row, .abundance)
```

Arguments

<code>.data</code>	A ‘tbl‘ formatted as <SAMPLE> <TRANSCRIPT> <COUNT> <...>
<code>.column</code>	The name of the column horizontally presented in the heatmap
<code>.row</code>	The name of the column vertically presented in the heatmap
<code>.abundance</code>	The name of the transcript/gene abundance column

Value

A list

<code>heatmap</code>	<i>Creates a ‘InputHeatmap‘ object from ‘tbl_df‘ on evaluation creates a ‘ComplexHeatmap‘</i>
----------------------	---

Description

`heatmap()` takes a `tbl` object and easily produces a `ComplexHeatmap` plot, with integration with `tibble` and `dplyr` frameworks.

Usage

```
heatmap(
  .data,
  .row,
  .column,
  .value,
  transform = NULL,
  scale = "none",
  palette_value = c("#440154FF", "#21908CFF", "#fefada"),
  palette_grouping = list(),
  .scale = NULL,
  ...
)
heatmap_(
  .data,
```

```

.row,
.column,
.value,
transform = NULL,
scale = "none",
palette_value = c("#440154FF", "#21908CFF", "#fefada"),
palette_grouping = list(),
.scale = NULL,
...
)

## S4 method for signature 'tbl'
heatmap(
  .data,
  .row,
  .column,
  .value,
  transform = NULL,
  scale = "none",
  palette_value = c("#440154FF", "#21908CFF", "#fefada"),
  palette_grouping = list(),
  .scale = NULL,
  ...
)

## S4 method for signature 'tbl_df'
heatmap(
  .data,
  .row,
  .column,
  .value,
  transform = NULL,
  scale = "none",
  palette_value = c("#440154FF", "#21908CFF", "#fefada"),
  palette_grouping = list(),
  .scale = NULL,
  ...
)

```

Arguments

.data	A ‘tbl_df’ formatted as <ELEMENT> <FEATURE> <VALUE> <...>
.row	The name of the column vertically presented in the heatmap
.column	The name of the column horizontally presented in the heatmap
.value	The name of the column for the value of the element/feature pair
transform	A function, used to transform .value row-wise (e.g., transform = log1p)
scale	A character string. Possible values are c("none", "row", "column", "both")
palette_value	A character vector This is the palette that will be used as gradient for .value. For example c("red", "white", "blue"). For higher flexibility you can use circlize::colorRamp2(c(-2, -1, 0, 1, 2)), viridis::magma(5))

palette_grouping	A list of character vectors. This is the list of palettes that will be used for grouping. For example list(RColorBrewer::brewer.pal(8, "Accent")) or list(c("#B3E2CD", "#FDCDAC", "#CBD5E8")) or list(c("black", "red"))
.scale	DEPRECATED. please use scale instead \(` with no dot prefix \`).
...	The arguments that will be passed to the Heatmap function of ComplexHeatmap backend

Details

[Maturing]

This function takes a `tbl` as an input and creates a ‘ComplexHeatmap’ plot. The information is stored in a ‘InputHeatmap’ object that is updated along the pipe statement, for example adding annotation layers.

Value

- A ‘InputHeatmap‘ objects that gets evaluated to a ‘ComplexHeatmap‘ object
- A ‘InputHeatmap‘ object
- A ‘InputHeatmap‘ object
- A ‘InputHeatmap‘ object

Examples

```
library(dplyr)

tidyHeatmap::N52 %>%
group_by(`Cell type`) %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`,
)
```

ifelse2_pipe

This is a generalisation of ifelse that accepts an object and return an objects

Description

This is a generalisation of `ifelse` that accepts an object and return an objects

Usage

```
ifelse2_pipe(.x, .p1, .p2, .f1, .f2, .f3 = NULL)
```

Arguments

.x	A tibble
.p1	A boolean
.p2	ELSE IF condition
.f1	A function
.f2	A function
.f3	A function

Value

A tibble

ifelse_pipe

This is a generalisation of ifelse that accepts an object and return an objects

Description

This is a generalisation of ifelse that accepts an object and return an objects

Usage

```
ifelse_pipe(.x, .p, .f1, .f2 = NULL)
```

Arguments

.x	A tibble
.p	A boolean
.f1	A function
.f2	A function

Value

A tibble

input_heatmap	<i>input_heatmap</i>
---------------	----------------------

Description

`input_heatmap()` takes a `tbl` object and easily produces a `ComplexHeatmap` plot, with integration with `tibble` and `dplyr` frameworks.

Usage

```
input_heatmap(
  .data,
  .horizontal,
  .vertical,
  .abundance,
  transform = NULL,
  scale = "none",
  palette_value = c("#440154FF", "#21908CFF", "#fefada"),
  palette_grouping = list(),
  ...
)
```

Arguments

.data	A ‘tbl‘ formatted as <SAMPLE> <TRANSCRIPT> <COUNT> <...>
.horizontal	The name of the column horizontally presented in the heatmap
.vertical	The name of the column vertically presented in the heatmap
.abundance	The name of the transcript/gene abundance column
transform	A function, used to transform .value, for example <code>log1p</code>
scale	A character string. Possible values are <code>c("none", "row", "column", "both")</code>
palette_value	A character vector, or a function for higher customisation (<code>colorRamp2</code>). This is the palette that will be used as gradient for abundance. If <code>palette_value</code> is a vector of hexadecimal colours, it should have 3 values. If you want more customisation, you can pass to <code>palette_value</code> a function, that is derived as for example <code>'colorRamp2(c(-2, 0, 2), palette_value)'</code>
palette_grouping	A list of character vectors. This is the list of palettes that will be used for grouping
...	Further arguments to be passed to <code>ComplexHeatmap::Heatmap</code>

Details

To be added.

Value

A ‘`ComplexHeatmap`‘ object

<code>layer_arrow_down</code>	<i>Adds a layers of symbols above the heatmap tiles to a ‘InputHeatmap’, that on evaluation creates a ‘ComplexHeatmap’</i>
-------------------------------	--

Description

`layer_arrow_down()` from a ‘InputHeatmap‘ object, adds a bar annotation layer.

Usage

```
layer_arrow_down(.data, ...)
## S4 method for signature 'InputHeatmap'
layer_arrow_down(.data, ...)
```

Arguments

<code>.data</code>	A ‘InputHeatmap‘
<code>...</code>	Expressions that return a logical value, and are defined in terms of the variables in <code>.data</code> . If multiple expressions are included, they are combined with the <code>&</code> operator. Only rows for which all conditions evaluate to TRUE are kept.

Details

[Maturing]

It uses ‘ComplexHeatmap‘ as visualisation tool.

Value

A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% layer_arrow_down()
```

layer_arrow_up	<i>Adds a layers of symbols above the heatmap tiles to a ‘InputHeatmap’, that on evaluation creates a ‘ComplexHeatmap’</i>
----------------	--

Description

layer_arrow_up() from a ‘InputHeatmap‘ object, adds a bar annotation layer.

Usage

```
layer_arrow_up(.data, ...)
## S4 method for signature 'InputHeatmap'
layer_arrow_up(.data, ...)
```

Arguments

- .data A ‘InputHeatmap‘
 ... Expressions that return a logical value, and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept.

Details

[Maturing]

It uses ‘ComplexHeatmap‘ as visualisation tool.

Value

- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
 A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% layer_arrow_up()
```

`layer_diamond` *Adds a layers of symbols above the heatmap tiles to a ‘InputHeatmap’, that on evaluation creates a ‘ComplexHeatmap’*

Description

`layer_diamond()` from a ‘InputHeatmap’ object, adds a bar annotation layer.

Usage

```
layer_diamond(.data, ...)
## S4 method for signature 'InputHeatmap'
layer_diamond(.data, ...)
```

Arguments

- | | |
|-------|---|
| .data | A ‘InputHeatmap’ |
| ... | Expressions that return a logical value, and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept. |

Details

[Maturing]

It uses ‘ComplexHeatmap’ as visualisation tool.

Value

- A ‘InputHeatmap’ object that gets evaluated to a ‘ComplexHeatmap’
- A ‘InputHeatmap’ object that gets evaluated to a ‘ComplexHeatmap’

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% layer_diamond()
```

layer_point	<i>Adds a layers of symbols above the heatmap tiles to a ‘InputHeatmap’, that on evaluation creates a ‘ComplexHeatmap’</i>
-------------	--

Description

layer_point() from a ‘InputHeatmap‘ object, adds a bar annotation layer.

Usage

```
layer_point(.data, ...)
## S4 method for signature 'InputHeatmap'
layer_point(.data, ...)
```

Arguments

- .data A ‘InputHeatmap‘
- ... Expressions that return a logical value, and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept.

Details

[Maturing]

It uses ‘ComplexHeatmap‘ as visualisation tool.

Value

- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% layer_point()
```

layer_square	<i>Adds a layers of symbols above the heatmap tiles to a ‘InputHeatmap’, that on evaluation creates a ‘ComplexHeatmap’</i>
--------------	--

Description

layer_square() from a ‘InputHeatmap‘ object, adds a bar annotation layer.

Usage

```
layer_square(.data, ...)
## S4 method for signature 'InputHeatmap'
layer_square(.data, ...)
```

Arguments

- | | |
|-------|---|
| .data | A ‘InputHeatmap‘ |
| ... | Expressions that return a logical value, and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept. |

Details

[Maturing]

It uses ‘ComplexHeatmap‘ as visualisation tool.

Value

- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% layer_square()
```

N52

Example data set N52

Description

Example data set N52

Usage

N52

Format

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 520 rows and 15 columns.

`parse_formula`

formula parser

Description

.formula parser

Usage

`parse_formula(fm)`

Arguments

`fm` a formula

Value

A character vector

`pasilla`

Example data set Pasilla

Description

Example data set Pasilla

Usage

`pasilla`

Format

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 504 rows and 8 columns.

`prepend` *From rlang deprecated*

Description

From rlang deprecated

Usage

```
prepend(x, values, before = 1)
```

Arguments

<code>x</code>	An array
<code>values</code>	An array
<code>before</code>	A boolean

Value

An array

`quo_names` *Convert array of quosure (e.g. c(col_a, col_b)) into character vector*

Description

Convert array of quosure (e.g. c(col_a, col_b)) into character vector

Usage

```
quo_names(v)
```

Arguments

<code>v</code>	A array of quosures (e.g. c(col_a, col_b))
----------------	--

Value

A character vector

<code>save_pdf</code>	<i>Save plot on PDF file</i>
-----------------------	------------------------------

Description

`save_pdf()` takes as input a Heatmap from ComplexHeatmap and save it to PDF file

Usage

```
save_pdf (
  .heatmap,
  filename,
  width = NULL,
  height = NULL,
  units = c("in", "cm", "mm")
)
```

Arguments

<code>.heatmap</code>	A ‘Heatmap’
<code>filename</code>	A character string. The name of the output file/path
<code>width</code>	A ‘double’. Plot width
<code>height</code>	A ‘double’. Plot height
<code>units</code>	A character string. units ("in", "cm", or "mm")

Details

[Maturing]

It simply save an ‘Heatmap’ to a PDF file use `pdf()` function in the back end

Value

NA

Examples

```
library(dplyr)
tidyHeatmap::heatmap(
  dplyr::group_by(tidyHeatmap::pasilla, location, type),
  .column = sample,
  .row = symbol,
  .value = `count normalised adjusted`,
) %>%
  save_pdf(tempfile())
```

`save_pdf, Heatmap-method`
`save_pdf`

Description

`save_pdf`

Usage

```
## S4 method for signature 'Heatmap'
save_pdf(
  .heatmap,
  filename,
  width = NULL,
  height = NULL,
  units = c("in", "cm", "mm")
)
```

Arguments

<code>.heatmap</code>	A ‘Heatmap’
<code>filename</code>	A character string. The name of the output file/path
<code>width</code>	A ‘double’. Plot width
<code>height</code>	A ‘double’. Plot height
<code>units</code>	A character string. units ("in", "cm", or "mm")

`save_pdf, InputHeatmap-method`
`save_pdf`

Description

`save_pdf`

Usage

```
## S4 method for signature 'InputHeatmap'
save_pdf(
  .heatmap,
  filename,
  width = NULL,
  height = NULL,
  units = c("in", "cm", "mm")
)
```

Arguments

.heatmap	A ‘Heatmap’
filename	A character string. The name of the output file/path
width	A ‘double’. Plot width
height	A ‘double’. Plot height
units	A character string ("in", "cm", or "mm")

scale_design *Scale design matrix*

Description

Scale design matrix

Usage

```
scale_design(df, .formula)
```

Arguments

df	A tibble
.formula	a formula

Value

A tibble

scale_robust *Scale counts in a robust way against sd == 0*

Description

Scale counts in a robust way against sd == 0

Usage

```
scale_robust(y)
```

Arguments

y	A numerical array
---	-------------------

Value

A scaled and centred numerical array

```
select_closest_pairs
```

Sub function of remove_redundancy_elements_though_reduced_dimensions

Description

Sub function of `remove_redundancy_elements_though_reduced_dimensions`

Usage

```
select_closest_pairs(df)
```

Arguments

<code>df</code>	A tibble
-----------------	----------

Value

A tibble with pairs to drop

```
split_rows
```

Split the heatmap row-wise depending on the biggest branches in the cladogram.

Description

`split_rows()` from a ‘InputHeatmap’ object, split the row cladogram.

`split_columns()` from a ‘InputHeatmap’ object, split the column cladogram.

Usage

```
split_rows(.data, number_of_groups)

## S4 method for signature 'InputHeatmap'
split_rows(.data, number_of_groups)

split_columns(.data, number_of_groups)

## S4 method for signature 'InputHeatmap'
split_columns(.data, number_of_groups)
```

Arguments

<code>.data</code>	A ‘InputHeatmap’
--------------------	------------------

<code>number_of_groups</code>	
-------------------------------	--

	An integer. The number of groups to split the cladogram into.
--	---

Details

[Maturing]

It uses ‘ComplexHeatmap’ as visualisation tool.

[Maturing]

It uses ‘ComplexHeatmap’ as visualisation tool.

Value

- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘
- A ‘InputHeatmap‘ object that gets evaluated to a ‘ComplexHeatmap‘

Examples

```
library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% split_rows(2)

library(dplyr)

hm =
tidyHeatmap::N52 %>%
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`
)
hm %>% split_columns(2)
```

`wrap_heatmap`

Wrap tidyHeatmap (ComplexHeatmap) in a patchwork-compliant patch

Description

In order to add tidyHeatmap (ComplexHeatmap) element to a patchwork they can be converted to a compliant representation using the ‘`wrap_heatmap()`‘ function. This allows you to position either grobs, ggplot objects, patchwork objects, or even base graphics (if passed as a formula) in either the full area, the full plotting area (anything between and including the axis label), or the panel area (only the actual area where data is drawn).

Usage

```
wrap_heatmap(
  panel = NULL,
  plot = NULL,
  full = NULL,
  clip = TRUE,
  ignore_tag = FALSE
)

## S4 method for signature 'InputHeatmap'
wrap_heatmap(
  panel = NULL,
  plot = NULL,
  full = NULL,
  clip = TRUE,
  ignore_tag = FALSE
)
```

Arguments

<code>panel, plot, full</code>	A grob, ggplot, patchwork, formula, raster, or nativeRaster object to add to the respective area.
<code>clip</code>	Should the grobs be clipped if expanding outside its area
<code>ignore_tag</code>	Should tags be ignored for this patch. This is relevant when using automatic tagging of plots and the content of the patch does not qualify for a tag.

Value

A wrapped_patch object
A wrapped_patch object

Examples

```
tidyHeatmap::N52 |>
tidyHeatmap::heatmap(
  .row = symbol_ct,
  .column = UBR,
  .value = `read count normalised log`,
) |>
wrap_heatmap()
```

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