

Package: SCpubr (via r-universe)

August 21, 2024

Type Package

Title Generate Publication Ready Visualizations of Single Cell Transcriptomics Data

Version 2.0.2.9000

Description A system that provides a streamlined way of generating publication ready plots for known Single-Cell transcriptomics data in a “publication ready” format. This is, the goal is to automatically generate plots with the highest quality possible, that can be used right away or with minimal modifications for a research article.

License GPL-3

URL <https://github.com/enblacar/SCpubr/>,
<https://enblacar.github.io/SCpubr-book/>

BugReports <https://github.com/enblacar/SCpubr/issues/>

Depends R (>= 4.0.0)

Suggests assertthat, circlize, cli, cluster, colorspace, ComplexHeatmap, covr, decoupleR, dplyr (>= 1.1.0), enrichplot,forcats, ggalluvial (>= 0.12.5), ggbeeswarm, ggdist, ggExtra, ggh4x, ggplot2 (>= 3.4.0), ggplotify, ggrastr, ggrepel, ggridges, ggsignif, graphics, infercnv, knitr, labeling, magrittr, MASS, Matrix, methods, Nebulosa, org.Hs.eg.db, patchwork, pheatmap, plyr, purrr, qpdf, RColorBrewer, rjags, rlang, rmarkdown, scales, scattermore, Seurat, SeuratObject, sf, stringr, svglite, testthat (>= 3.0.0), tibble, tidyverse, UCell, viridis, waffle, withr, liana (>= 0.1.6)

Remotes saezlab/liana

VignetteBuilder knitr

biocViews Software, SingleCell, Visualization

Config/testthat.edition 3

Encoding UTF-8

LazyData true

LazyDataCompression xz
Rxygen list(markdown = TRUE)
RxygenNote 7.2.1
Repository <https://enblacar.r-universe.dev>
RemoteUrl <https://github.com/enblacar/scpubr>
RemoteRef HEAD
RemoteSha ebc5e868f5f1cd275f541577abeb83a70d5d8f68

Contents

do_AffinityAnalysisPlot	3
do_AlluvialPlot	7
do_BarPlot	10
do_BeeSwarmPlot	14
do_BoxPlot	18
do_CellularStatesPlot	23
do_ChordDiagramPlot	28
do_ColorPalette	30
do_CopyNumberVariantPlot	33
do_CorrelationPlot	37
do_DimPlot	41
do_DotPlot	46
do_EnrichmentHeatmap	50
do_ExpressionHeatmap	55
do_FeaturePlot	58
do_GeyserPlot	64
do_GroupwiseDEPlot	69
do_LigandReceptorPlot	73
do_LoadingsPlot	77
do_MetadataPlot	81
do_NebulosaPlot	83
do_PathwayActivityPlot	87
do_RankedEnrichmentPlot	91
do_RankedExpressionPlot	95
do_RidgePlot	100
do_SCEnrichmentHeatmap	104
do_SCExpressionHeatmap	109
do_TermEnrichmentPlot	113
do_TFActivityPlot	117
do_ViolinPlot	120
do_VolcanoPlot	124
do_WafflePlot	126
human_chr_locations	129
package_report	129
save_Plot	130

do_AffinityAnalysisPlot

Compute affinity of gene sets to cell populations using decoupleR.

Description

Major contributions to this function:

- **Marc Elosua Bayés** for the core concept code and idea.
- **Pau Badia i Mompej** for the network generation.

Usage

```
do_AffinityAnalysisPlot(  
  sample,  
  input_gene_list,  
  subsample = 2500,  
  group.by = NULL,  
  assay = NULL,  
  slot = NULL,  
  statistic = "ulm",  
  number.breaks = 5,  
  use_viridis = FALSE,  
  viridis.palette = "G",  
  viridis.direction = -1,  
  sequential.palette = "YlGnBu",  
  sequential.direction = 1,  
  diverging.palette = "RdBu",  
  diverging.direction = -1,  
  enforce_symmetry = TRUE,  
  legend.position = "bottom",  
  legend.width = 1,  
  legend.length = 20,  
  legend.framewidth = 0.5,  
  legend.tickwidth = 0.5,  
  legend.framecolor = "grey50",  
  legend.tickcolor = "white",  
  legend.type = "colorbar",  
  na.value = "grey75",  
  font.size = 14,  
  font.type = "sans",  
  axis.text.x.angle = 45,  
  flip = FALSE,  
  colors.use = NULL,  
  min.cutoff = NA,  
  max.cutoff = NA,
```

```

verbose = TRUE,
return_object = FALSE,
grid.color = "white",
border.color = "black",
flavor = "Seurat",
nbin = 24,
ctrl = 100,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
input_gene_list	named_list Named list of lists of genes to be used as input.
subsample	numeric Number of cells to subset for the analysis. NA will use all. Cells are selected at random.
group.by	character Metadata variable to group the output by. Has to be a character or factor column.
assay	character Assay to use. Defaults to the current assay.
slot	character Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
statistic	character DecoupleR statistic to use for the analysis. values in the Idents of the Seurat object are reported, assessing how specific a given gene set is for a given cell population compared to other gene sets of equal expression.
number.breaks	numeric Controls the number of breaks in continuous color scales of ggplot2-based plots.
use_viridis	logical Whether to use viridis color scales.
viridis.palette	character A capital letter from A to H or the scale name as in scale_fill_viridis .
viridis.direction	numeric Either 1 or -1. Controls how the gradient of viridis scale is formed.
sequential.palette	character Type of sequential color palette to use. Out of the sequential palettes defined in brewer.pal .
sequential.direction	numeric Direction of the sequential color scale. Either 1 or -1.
diverging.palette	character Type of symmetrical color palette to use. Out of the diverging palettes defined in brewer.pal .

diverging.direction
`numeric` | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.

enforce_symmetry
`logical` | Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.

legend.position
`character` | Position of the legend in the plot. One of:

- top: Top of the figure.
- bottom: Bottom of the figure.
- left: Left of the figure.
- right: Right of the figure.
- none: No legend is displayed.

legend.length, legend.width
`numeric` | Length and width of the legend. Will adjust automatically depending on legend side.

legend.framewidth, legend.tickwidth
`numeric` | Width of the lines of the box in the legend.

legend.framecolor
`character` | Color of the lines of the box in the legend.

legend.tickcolor
`character` | Color of the ticks of the box in the legend.

legend.type
`character` | Type of legend to display. One of:

- normal: Default legend displayed by `ggplot2`.
- colorbar: Redefined colorbar legend, using `guide_colorbar`.

na.value
`character` | Color value for NA.

font.size
`numeric` | Overall font size of the plot. All plot elements will have a size relationship with this font size.

font.type
`character` | Base font family for the plot. One of:

- mono: Mono spaced font.
- serif: Serif font family.
- sans: Default font family.

axis.text.x.angle
`numeric` | Degree to rotate the X labels. One of: 0, 45, 90.

flip
`logical` | Whether to invert the axis of the displayed plot.

colors.use
`named_vector` | Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of group.by. If group.by is not provided, defaults to the unique values of `Idents`. If not provided, a color scale will be set by default.

min.cutoff, max.cutoff
`numeric` | Set the min/max ends of the color scale. Any cell/group with a value lower than min.cutoff will turn into min.cutoff and any cell with a value higher than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.

verbose	<code>logical</code> Whether to show extra comments, warnings,etc.
return_object	<code>logical</code> Returns the Seurat object with the modifications performed in the function. Normally, this contains a new assay with the data that can then be used for any other visualization desired.
grid.color	<code>character</code> Color of the grid in the plot. In heatmaps, color of the border of the cells.
border.color	<code>character</code> Color for the border of the heatmap body.
flavor	<code>character</code> One of: Seurat, UCell. Compute the enrichment scores using <code>AddModuleScore</code> or <code>AddModuleScore_UCell</code> .
nbin	<code>numeric</code> Number of bins to use in <code>AddModuleScore</code> .
ctrl	<code>numeric</code> Number of genes in the control set to use in <code>AddModuleScore</code> .
plot.title.face, axis.title.face, plot.subtitle.face, axis.text.face, plot.caption.face, legend.title.face, legend.text.face	<code>character</code> Controls the style of the font for the corresponding theme element. One of: <ul style="list-style-type: none">• plain: For normal text.• italic: For text in italic.• bold: For text in bold.• bold.italic: For text both in italic and bold.

Value

A list containing different plots.

Examples

```
p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

do_AlluvialPlot *Generate Alluvial plots.*

Description

This function is based on the **ggalluvial** package. It allows you to generate alluvial plots from a given Seurat object.

Usage

```
do_AlluvialPlot(
  sample,
  first_group,
  last_group,
  middle_groups = NULL,
  colors.use = NULL,
  plot.title = NULL,
  plot.subtitle = NULL,
  plot.caption = NULL,
  font.size = 14,
  font.type = "sans",
  xlab = NULL,
  ylab = "Number of cells",
  repel = FALSE,
  fill.by = last_group,
  use_labels = FALSE,
  stratum.color = "black",
  stratum.fill = "white",
  stratum.width = 1/3,
  stratum.fill.conditional = FALSE,
  use_geom_flow = FALSE,
  alluvium.color = "white",
  flow.color = "white",
  flip = FALSE,
  label.color = "black",
  curve_type = "sigmoid",
  use_viridis = FALSE,
  viridis.palette = "G",
  viridis.direction = -1,
```

```

sequential.palette = "YlGnBu",
sequential.direction = 1,
plot.grid = FALSE,
grid.color = "grey75",
grid.type = "dashed",
na.value = "white",
legend.position = "right",
legend.title = NULL,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
first_group	character Categorical metadata variable. First group of nodes of the alluvial plot.
last_group	character Categorical metadata variable. Last group of nodes of the alluvial plot.
middle_groups	character Categorical metadata variable. Vector of groups of nodes of the alluvial plot.
colors.use	character Named list of colors corresponding to the unique values in fill.by (which defaults to last_group).
plot.title, plot.subtitle, plot.caption	character Title, subtitle or caption to use in the plot.
font.size	numeric Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	character Base font family for the plot. One of: <ul style="list-style-type: none"> • mono: Mono spaced font. • serif: Serif font family. • sans: Default font family.
xlab, ylab	character Titles for the X and Y axis.
repel	logical Whether to repel the text labels.
fill.by	character One of first_group, middle_groups (one of the values, if multiple mid_groups) or last_group. These values will be used to color the alluvium/flow.
use_labels	logical Whether to use labels instead of text for the stratum.
stratum.color, alluvium.color, flow.color	character Color for the border of the alluvium (and flow) and stratum.
stratum.fill	character Color to fill the stratum.

```
stratum.width    logical | Width of the stratum.  
stratum.fill.conditional  
                      logical | Whether to fill the stratum with the same colors as the alluvium/flow.  
use_geom_flow   logical | Whether to use geom_flow instead of geom_alluvium. Visual results  
                  might differ.  
flip            logical | Whether to invert the axis of the displayed plot.  
label.color     character | Color for the text labels.  
curve_type      character | Type of curve used in geom_alluvium. One of:  
                  • linear.  
                  • cubic.  
                  • quintic.  
                  • sine.  
                  • arctangent.  
                  • sigmoid.  
                  • xspline.  
use_viridis     logical | Whether to use viridis color scales.  
viridis.palette character | A capital letter from A to H or the scale name as in scale_fill_viridis.  
viridis.direction  
                     numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.  
sequential.palette  
                     character | Type of sequential color palette to use. Out of the sequential  
                           palettes defined in brewer.pal.  
sequential.direction  
                     numeric | Direction of the sequential color scale. Either 1 or -1.  
plot.grid       logical | Whether to plot grid lines.  
grid.color      character | Color of the grid in the plot. In heatmaps, color of the border of the  
                  cells.  
grid.type       character | One of the possible linetype options:  
                  • blank.  
                  • solid.  
                  • dashed.  
                  • dotted.  
                  • dotdash.  
                  • longdash.  
                  • twodash.  
na.value        character | Color value for NA.  
legend.position character | Position of the legend in the plot. One of:  
                  • top: Top of the figure.  
                  • bottom: Bottom of the figure.  
                  • left: Left of the figure.
```

- **right**: Right of the figure.
- **none**: No legend is displayed.

`legend.title` **character** | Title for the legend.
`plot.title.face`, `plot.subtitle.face`, `plot.caption.face`,
`axis.title.face`, `axis.text.face`, `legend.title.face`, `legend.text.face`
character | Controls the style of the font for the corresponding theme element.
One of:

- **plain**: For normal text.
- **italic**: For text in italic.
- **bold**: For text in bold.
- **bold.italic**: For text both in italic and bold.

Value

A ggplot2 object.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_AlluvialPlot", passive = TRUE)
message(value)
if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Compute basic sankey plot.
  p <- SCpubr:::do_AlluvialPlot(sample = sample,
                                 first_group = "orig.ident",
                                 last_group = "seurat_clusters")

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

Description

Create Bar Plots.

Usage

```
do_BarPlot(  
  sample,  
  group.by,  
  order = FALSE,  
  add.n = FALSE,  
  add.n.face = "bold",  
  add.n.expand = c(0, 1.15),  
  add.n.size = 4,  
  order.by = NULL,  
  split.by = NULL,  
  facet.by = NULL,  
  position = "stack",  
  font.size = 14,  
  font.type = "sans",  
  legend.position = "bottom",  
  legend.title = NULL,  
  legend.ncol = NULL,  
  legend.nrow = NULL,  
  legend.byrow = FALSE,  
  axis.text.x.angle = 45,  
  xlab = NULL,  
  ylab = NULL,  
  colors.use = NULL,  
  flip = FALSE,  
  plot.title = NULL,  
  plot.subtitle = NULL,  
  plot.caption = NULL,  
  plot.grid = FALSE,  
  grid.color = "grey75",  
  grid.type = "dashed",  
  plot.title.face = "bold",  
  plot.subtitle.face = "plain",  
  plot.caption.face = "italic",  
  axis.title.face = "bold",  
  axis.text.face = "plain",  
  legend.title.face = "bold",  
  legend.text.face = "plain",  
  strip.text.face = "bold",  
  return_data = FALSE  
)
```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
group.by	<code>character</code> Metadata column to compute the counts of. Has to be either a character or factor column.
order	<code>logical</code> Whether to order the results in descending order of counts.

add.n	<code>logical</code> Whether to add the total counts on top of each bar.
add.n.face	<code>character</code> Font face of the labels added by add.n.
add.n.expand	<code>numeric</code> Vector of two numerics representing the start and end of the scale. Minimum should be 0 and max should be above 1. This basically expands the Y axis so that the labels fit when <code>flip = TRUE</code> . <ul style="list-style-type: none"> • <code>stack</code>: Set the bars side by side, displaying the total number of counts. Uses <code>position_stack</code>. • <code>fill</code>: Set the bars on top of each other, displaying the proportion of counts from the total that each group represents. Uses <code>position_fill</code>.
add.n.size	<code>numeric</code> Size of the labels
order.by	<code>character</code> When <code>split.by</code> is used, value of <code>group.by</code> to reorder the columns based on its value.
split.by	<code>character</code> Metadata column to split the values of <code>group.by</code> by. If not used, defaults to the active idents.
facet.by	<code>character</code> Metadata column to gather the columns by. This is useful if you have other overarching metadata.
position	<code>character</code> Position function from ggplot2 . Either stack or fill.
font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • <code>mono</code>: Mono spaced font. • <code>serif</code>: Serif font family. • <code>sans</code>: Default font family.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • <code>top</code>: Top of the figure. • <code>bottom</code>: Bottom of the figure. • <code>left</code>: Left of the figure. • <code>right</code>: Right of the figure. • <code>none</code>: No legend is displayed.
legend.title	<code>character</code> Title for the legend.
legend.ncol	<code>numeric</code> Number of columns in the legend.
legend.nrow	<code>numeric</code> Number of rows in the legend.
legend.byrow	<code>logical</code> Whether the legend is filled by row or not.
axis.text.x.angle	<code>numeric</code> Degree to rotate the X labels. One of: 0, 45, 90.
xlab, ylab	<code>character</code> Titles for the X and Y axis.
colors.use	<code>named_vector</code> Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of <code>group.by</code> . If <code>group.by</code> is not provided, defaults to the unique values of <code>Idents</code> . If not provided, a color scale will be set by default.

```

flip           logical | Whether to invert the axis of the displayed plot.
plot.title, plot.subtitle, plot.caption
               character | Title, subtitle or caption to use in the plot.
plot.grid      logical | Whether to plot grid lines.
grid.color     character | Color of the grid in the plot. In heatmaps, color of the border of the
cells.
grid.type      character | One of the possible linetype options:
  • blank.
  • solid.
  • dashed.
  • dotted.
  • dotdash.
  • longdash.
  • twodash.
plot.title.face,       plot.subtitle.face,       plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
               character | Controls the style of the font for the corresponding theme element.
One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.
strip.text.face
               character | Controls the style of the font for the strip text. One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.
return_data    logical | Returns a data.frame with the count and proportions displayed in the
plot.

```

Value

A ggplot2 object containing a Bar plot.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_BarPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))
}

```

```

# Basic bar plot, horizontal.
p1 <- SCpubr::do_BarPlot(sample = sample,
                         group.by = "seurat_clusters",
                         legend.position = "none",
                         plot.title = "Number of cells per cluster")

# Split by a second variable.
sample$modified_orig.ident <- sample(x = c("Sample_A", "Sample_B", "Sample_C"),
                                      size = ncol(sample),
                                      replace = TRUE,
                                      prob = c(0.2, 0.7, 0.1))

p <- SCpubr::do_BarPlot(sample,
                        group.by = "seurat_clusters",
                        split.by = "modified_orig.ident",
                        plot.title = "Number of cells per cluster in each sample",
                        position = "stack")

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_BeeSwarmPlot *BeeSwarm plot.*

Description

BeeSwarm plot.

Usage

```
do_BeeSwarmPlot(
  sample,
  feature_to_rank,
  group.by = NULL,
  assay = NULL,
  reduction = NULL,
  slot = NULL,
  continuous_feature = FALSE,
  order = FALSE,
  colors.use = NULL,
  legend.title = NULL,
  legend.type = "colorbar",
  legend.position = "bottom",
  legend.framewidth = 0.5,
```

```

legend.tickwidth = 0.5,
legend.length = 20,
legend.width = 1,
legend.framecolor = "grey50",
legend.tickcolor = "white",
legend.ncol = NULL,
legend.icon.size = 4,
plot.title = NULL,
plot.subtitle = NULL,
plot.caption = NULL,
xlab = NULL,
ylab = NULL,
font.size = 14,
font.type = "sans",
remove_x_axis = FALSE,
remove_y_axis = FALSE,
flip = FALSE,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = 1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
verbose = TRUE,
raster = FALSE,
raster.dpi = 300,
plot_cell_borders = TRUE,
border.size = 1.5,
border.color = "black",
pt.size = 2,
min.cutoff = NA,
max.cutoff = NA,
na.value = "grey75",
number.breaks = 5,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
feature_to_rank	character Feature for which the cells are going to be ranked. Ideal case is that this feature is stored as a metadata column.

group.by	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
assay	<code>character</code> Assay to use. Defaults to the current assay.
reduction	<code>character</code> Reduction to use. Can be the canonical ones such as "umap", "pca", or any custom ones, such as "diffusion". If you are unsure about which reductions you have, use Seurat::Reductions(sample). Defaults to "umap" if present or to the last computed reduction if the argument is not provided.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
continuous_feature	<code>logical</code> Is the feature to rank and color for continuous? I.e: an enrichment score.
order	<code>logical</code> Whether to reorder the groups based on the median of the ranking.
colors.use	<code>named_vector</code> Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of group.by. If group.by is not provided, defaults to the unique values of <code>Idents</code> . If not provided, a color scale will be set by default.
legend.title	<code>character</code> Title for the legend.
legend.type	<code>character</code> Type of legend to display. One of: <ul style="list-style-type: none"> • <code>normal</code>: Default legend displayed by <code>ggplot2</code>. • <code>colorbar</code>: Redefined colorbar legend, using <code>guide_colorbar</code>.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • <code>top</code>: Top of the figure. • <code>bottom</code>: Bottom of the figure. • <code>left</code>: Left of the figure. • <code>right</code>: Right of the figure. • <code>none</code>: No legend is displayed.
legend.framewidth, legend.tickwidth	<code>numeric</code> Width of the lines of the box in the legend.
legend.length, legend.width	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.
legend.framecolor	<code>character</code> Color of the lines of the box in the legend.
legend.tickcolor	<code>character</code> Color of the ticks of the box in the legend.
legend.ncol	<code>numeric</code> Number of columns in the legend.
legend.icon.size	<code>numeric</code> Size of the icons in legend.
plot.title, plot.subtitle, plot.caption	<code>character</code> Title, subtitle or caption to use in the plot.
xlab, ylab	<code>character</code> Titles for the X and Y axis.

```

font.size      numeric | Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type      character | Base font family for the plot. One of:
                      • mono: Mono spaced font.
                      • serif: Serif font family.
                      • sans: Default font family.
remove_x_axis, remove_y_axis
                      logical | Remove X axis labels and ticks from the plot.
flip            logical | Whether to invert the axis of the displayed plot.
use_viridis    logical | Whether to use viridis color scales.
viridis.palette
                      character | A capital letter from A to H or the scale name as in scale_fill_viridis.
viridis.direction
                      numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.
sequential.palette
                      character | Type of sequential color palette to use. Out of the sequential palettes defined in brewer.pal.
sequential.direction
                      numeric | Direction of the sequential color scale. Either 1 or -1.
verbose          logical | Whether to show extra comments, warnings,etc.
raster           logical | Whether to raster the resulting plot. This is recommendable if plotting a lot of cells.
raster.dpi       numeric | Pixel resolution for rasterized plots. Defaults to 1024. Only activates on Seurat versions higher or equal than 4.1.0.
plot_cell_borders
                      logical | Whether to plot border around cells.
border.size     numeric | Width of the border of the cells.
border.color    character | Color for the border of the heatmap body.
pt.size         numeric | Size of the dots.
min.cutoff, max.cutoff
                      numeric | Set the min/max ends of the color scale. Any cell/group with a value lower than min.cutoff will turn into min.cutoff and any cell with a value higher than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
na.value        character | Color value for NA.
number.breaks   numeric | Controls the number of breaks in continuous color scales of ggplot2-based plots.
plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
                      character | Controls the style of the font for the corresponding theme element.
One of:
                      • plain: For normal text.
                      • italic: For text in italic.
                      • bold: For text in bold.
                      • bold.italic: For text both in italic and bold.

```

Value

A ggplot2 object containing a Bee Swarm plot.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_BeeSwarmPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Basic Bee Swarm plot - categorical coloring.
  # This will color based on the unique values of seurat_clusters.
  p <- SCpubr:::do_BeeSwarmPlot(sample = sample,
                                 feature_to_rank = "PC_1",
                                 group.by = "seurat_clusters",
                                 continuous_feature = FALSE)

  # Basic Bee Swarm plot - continuous coloring.
  # This will color based on the PC_1 values.
  p <- SCpubr:::do_BeeSwarmPlot(sample = sample,
                                 feature_to_rank = "PC_1",
                                 group.by = "seurat_clusters",
                                 continuous_feature = TRUE)
} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

do_BoxPlot

Generate Box Plots.

Description

Generate Box Plots.

Usage

```
do_BoxPlot(
  sample,
  feature,
  group.by = NULL,
  split.by = NULL,
  assay = NULL,
```

```
slot = "data",
font.size = 14,
font.type = "sans",
axis.text.x.angle = 45,
colors.use = NULL,
na.value = "grey75",
plot.title = NULL,
plot.subtitle = NULL,
plot.caption = NULL,
xlab = NULL,
ylab = NULL,
legend.title = NULL,
legend.title.position = "top",
legend.position = "bottom",
legend.ncol = NULL,
legend.nrow = NULL,
legend.byrow = FALSE,
boxplot.line.color = "black",
outlier.color = "black",
outlier.alpha = 0.5,
boxplot.linewidth = 0.5,
boxplot.width = NULL,
plot.grid = TRUE,
grid.color = "grey75",
grid.type = "dashed",
flip = FALSE,
order = FALSE,
use_silhouette = FALSE,
use_test = FALSE,
comparisons = NULL,
test = "wilcox.test",
map_signif_level = c(`***` = 0.001, `**` = 0.01, `*` = 0.05),
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)
```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
feature	<code>character</code> Feature to represent.
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.

<code>split.by</code>	<code>character</code> Secondary metadata variable to further group (split) the output by. Has to be a character or factor column.
<code>assay</code>	<code>character</code> Assay to use. Defaults to the current assay.
<code>slot</code>	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
<code>font.size</code>	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
<code>font.type</code>	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none">• <code>mono</code>: Mono spaced font.• <code>serif</code>: Serif font family.• <code>sans</code>: Default font family.
<code>axis.text.x.angle</code>	<code>numeric</code> Degree to rotate the X labels. One of: 0, 45, 90.
<code>colors.use</code>	<code>named_vector</code> Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of <code>group.by</code> . If <code>group.by</code> is not provided, defaults to the unique values of <code>Idents</code> . If not provided, a color scale will be set by default.
<code>na.value</code>	<code>character</code> Color value for NA.
<code>plot.title, plot.subtitle, plot.caption</code>	<code>character</code> Title, subtitle or caption to use in the plot.
<code>xlab, ylab</code>	<code>character</code> Titles for the X and Y axis.
<code>legend.title</code>	<code>character</code> Title for the legend.
<code>legend.title.position</code>	<code>character</code> Position for the title of the legend. One of: <ul style="list-style-type: none">• <code>top</code>: Top of the legend.• <code>bottom</code>: Bottom of the legend.• <code>left</code>: Left of the legend.• <code>right</code>: Right of the legend.
<code>legend.position</code>	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none">• <code>top</code>: Top of the figure.• <code>bottom</code>: Bottom of the figure.• <code>left</code>: Left of the figure.• <code>right</code>: Right of the figure.• <code>none</code>: No legend is displayed.
<code>legend.ncol</code>	<code>numeric</code> Number of columns in the legend.
<code>legend.nrow</code>	<code>numeric</code> Number of rows in the legend.
<code>legend.byrow</code>	<code>logical</code> Whether the legend is filled by row or not.
<code>boxplot.line.color</code>	<code>character</code> Color of the borders of the boxplots if <code>use_silhouette</code> is FALSE.
<code>outlier.color</code>	<code>character</code> Color of the outlier dots.

outlier.alpha **numeric** | Alpha applied to the outliers.
 boxplot.linewidth **numeric** | Width of the lines in the boxplots. Also controls the lines of the tests applied if use_test is set to true.
 boxplot.width **numeric** | Width of the boxplots.
 plot.grid **logical** | Whether to plot grid lines.
 grid.color **character** | Color of the grid in the plot. In heatmaps, color of the border of the cells.
 grid.type **character** | One of the possible linetype options:
 • blank.
 • solid.
 • dashed.
 • dotted.
 • dotdash.
 • longdash.
 • twodash.
 flip **logical** | Whether to invert the axis of the displayed plot.
 order **logical** | Whether to order the boxplots by average values. Can not be used alongside split.by.
 use_silhouette **logical** | Whether to color the borders of the boxplots instead of the inside area.
 use_test **logical** | Whether to apply a statistical test to a given pair of elements. Can not be used alongside split.by.
 comparisons A list of length-2 vectors. The entries in the vector are either the names of 2 values on the x-axis or the 2 integers that correspond to the index of the columns of interest.
 test the name of the statistical test that is applied to the values of the 2 columns (e.g. t.test, wilcox.test etc.). If you implement a custom test make sure that it returns a list that has an entry called p.value.
 map_signif_level Boolean value, if the p-value are directly written as annotation or asterisks are used instead. Alternatively one can provide a named numeric vector to create custom mappings from p-values to annotation: For example: c("***=0.001, **=0.01, *=0.05). Alternatively, one can provide a function that takes a numeric argument (the p-value) and returns a string.
 plot.title.face, plot.subtitle.face, plot.caption.face,
 axis.title.face, axis.text.face, legend.title.face, legend.text.face
character | Controls the style of the font for the corresponding theme element.
 One of:
 • plain: For normal text.
 • italic: For text in italic.
 • bold: For text in bold.
 • bold.italic: For text both in italic and bold.

Value

A ggplot2 object.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_BoxPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Basic box plot.
  p <- SCpubr:::do_BoxPlot(sample = sample,
                            feature = "nCount_RNA")
  p

  # Use silhouette style.
  p <- SCpubr:::do_BoxPlot(sample = sample,
                            feature = "nCount_RNA",
                            use_silhouette = TRUE)
  p

  # Order by mean values.
  p <- SCpubr:::do_BoxPlot(sample = sample,
                            feature = "nCount_RNA",
                            order = TRUE)
  p

  # Apply second grouping.
  sample$orig.ident <- ifelse(sample$seurat_clusters %in% c("0", "1", "2", "3"), "A", "B")
  p <- SCpubr:::do_BoxPlot(sample = sample,
                            feature = "nCount_RNA",
                            split.by = "orig.ident")
  p

  # Apply statistical tests.
  p <- SCpubr:::do_BoxPlot(sample = sample,
                            feature = "nCount_RNA",
                            group.by = "orig.ident",
                            use_test = TRUE,
                            comparisons = list(c("A", "B")))
  p

} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()``.")
}
```

do_CellularStatesPlot *Cellular States plot.*

Description

This plot aims to show the relationships between distinct enrichment scores. If 3 variables are provided, the relationship is between the Y axis and the dual X axis. If 4 variables are provided, each corner of the plot represents how enriched the cells are in that given list. How to interpret this? In a 3-variable plot, the Y axis just means one variable. The higher the cells are in the Y axis the more enriched they are in that given variable. The X axis is a dual parameter one. Cells falling into each extreme of the axis are highly enriched for either x1 or x2, while cells falling in between are not enriched for any of the two. In a 4-variable plot, each corner shows the enrichment for one of the 4 given features. Cells will tend to locate in either of the four corners, but there will be cases of cells locating mid-way between two given corners (enriched in both features) or in the middle of the plot (not enriched for any).

Usage

```
do_CellularStatesPlot(  
  sample,  
  input_gene_list,  
  x1,  
  y1,  
  x2 = NULL,  
  y2 = NULL,  
  group.by = NULL,  
  colors.use = NULL,  
  legend.position = "bottom",  
  legend.icon.size = 4,  
  legend.ncol = NULL,  
  legend.nrow = NULL,  
  legend.byrow = FALSE,  
  plot.title = NULL,  
  plot.subtitle = NULL,  
  plot.caption = NULL,  
  font.size = 14,  
  font.type = "sans",  
  xlab = NULL,  
  ylab = NULL,  
  axis.ticks = TRUE,  
  axis.text = TRUE,  
  verbose = FALSE,  
  enforce_symmetry = FALSE,  
  plot_marginal_distributions = FALSE,  
  marginal.type = "density",  
  marginal.size = 5,  
  marginal.group = TRUE,
```

```

plot_cell_borders = TRUE,
plot_enrichment_scores = FALSE,
border.size = 2,
border.color = "black",
pt.size = 2,
raster = FALSE,
raster.dpi = 1024,
plot_features = FALSE,
features = NULL,
use_viridis = TRUE,
viridis.palette = "G",
viridis.direction = 1,
sequential.palette = "YlGnBu",
sequential.direction = -1,
nbin = 24,
ctrl = 100,
number.breaks = 5,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
input_gene_list	named_list Named list of lists of genes to be used as input.
x1	character A name of a list from input_gene_list. First feature in the X axis. Will go on the right side of the X axis if y2 is not provided and top-right quadrant if provided.
y1	character A name of a list from input_gene_list. First feature on the Y axis. Will become the Y axis if y2 is not provided and bottom-right quadrant if provided.
x2	character A name of a list from input_gene_list. Second feature on the X axis. Will go on the left side of the X axis if y2 is not provided and top-left quadrant if provided.
y2	character A name of a list from input_gene_list. Second feature on the Y axis. Will become the bottom-left quadrant if provided.
group.by	character Metadata variable to group the output by. Has to be a character or factor column.
colors.use	named_vector Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of group.by. If group.by

is not provided, defaults to the unique values of `Idents`. If not provided, a color scale will be set by default.

<code>legend.position</code>	<code>character</code> Position of the legend in the plot. One of:
	<ul style="list-style-type: none"> • <code>top</code>: Top of the figure. • <code>bottom</code>: Bottom of the figure. • <code>left</code>: Left of the figure. • <code>right</code>: Right of the figure. • <code>none</code>: No legend is displayed.
<code>legend.icon.size</code>	<code>numeric</code> Size of the icons in legend.
<code>legend.ncol</code>	<code>numeric</code> Number of columns in the legend.
<code>legend.nrow</code>	<code>numeric</code> Number of rows in the legend.
<code>legend.byrow</code>	<code>logical</code> Whether the legend is filled by row or not.
<code>plot.title, plot.subtitle, plot.caption</code>	<code>character</code> Title, subtitle or caption to use in the plot.
<code>font.size</code>	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
<code>font.type</code>	<code>character</code> Base font family for the plot. One of:
	<ul style="list-style-type: none"> • <code>mono</code>: Mono spaced font. • <code>serif</code>: Serif font family. • <code>sans</code>: Default font family.
<code>xlab, ylab</code>	<code>character</code> Titles for the X and Y axis.
<code>axis.ticks</code>	<code>logical</code> Whether to show axis ticks.
<code>axis.text</code>	<code>logical</code> Whether to show axis text.
<code>verbose</code>	<code>logical</code> Whether to show extra comments, warnings,etc.
<code>enforce_symmetry</code>	<code>logical</code> Whether to enforce the plot to follow a symmetry (3 variables, the X axis has 0 as center, 4 variables, all axis have the same range and the plot is squared).
<code>plot_marginal_distributions</code>	<code>logical</code> Whether to plot marginal distributions on the figure or not.
<code>marginal.type</code>	<code>character</code> One of:
	<ul style="list-style-type: none"> • <code>density</code>: Compute density plots on the margins. • <code>histogram</code>: Compute histograms on the margins. • <code>boxplot</code>: Compute boxplot on the margins. • <code>violin</code>: Compute violin plots on the margins. • <code>densigram</code>: Compute densigram plots on the margins.
<code>marginal.size</code>	<code>numeric</code> Size ratio between the main and marginal plots. A value of 5 means that the main plot is 5 times bigger than the marginal plots.
<code>marginal.group</code>	<code>logical</code> Whether to group the marginal distribution by <code>group.by</code> or current identities.

```

plot_cell_borders
  logical | Whether to plot border around cells.

plot_enrichment_scores
  logical | Whether to report enrichment scores for the input lists as plots.

border.size    numeric | Width of the border of the cells.

border.color   character | Color for the border of the heatmap body.

pt.size        numeric | Size of the dots.

raster         logical | Whether to raster the resulting plot. This is recommendable if plotting
                        a lot of cells.

raster.dpi     numeric | Pixel resolution for rasterized plots. Defaults to 1024. Only activates
                        on Seurat versions higher or equal than 4.1.0.

plot_features  logical | Whether to also report any other feature onto the primary plot.

features       character | Additional features to plot.

use_viridis    logical | Whether to use viridis color scales.

viridis.palette
  character | A capital letter from A to H or the scale name as in scale_fill_viridis.

viridis.direction
  numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.

sequential.palette
  character | Type of sequential color palette to use. Out of the sequential
              palettes defined in brewer.pal.

sequential.direction
  numeric | Direction of the sequential color scale. Either 1 or -1.

nbin           numeric | Number of bins to use in AddModuleScore.

ctrl            numeric | Number of genes in the control set to use in AddModuleScore.

number.breaks  numeric | Controls the number of breaks in continuous color scales of ggplot2-
                        based plots.

plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
  character | Controls the style of the font for the corresponding theme element.
One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.

```

Details

This plots are based on the following publications:

- Neftel, C. *et al.* An Integrative Model of Cellular States, Plasticity, and Genetics for Glioblastoma. *Cell* 178, 835-849.e21 (2019). doi:[10.1016/j.cell.2019.06.024](https://doi.org/10.1016/j.cell.2019.06.024)
- Tirosh, I., Venteicher, A., Hebert, C. *et al.* Single-cell RNA-seq supports a developmental hierarchy in human oligodendrogloma. *Nature* 539, 309–313 (2016). doi:[10.1038/nature20123](https://doi.org/10.1038/nature20123)

Value

A ggplot2 object containing a butterfly plot.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_CellularStatesPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Define some gene sets to query. It has to be a named list.
  gene_set <- list("A" = rownames(sample)[1:10],
                  "B" = rownames(sample)[11:20],
                  "C" = rownames(sample)[21:30],
                  "D" = rownames(sample)[31:40])

  # Using two variables: A scatter plot X vs Y.
  p <- SCpubr::do_CellularStatesPlot(sample = sample,
                                       input_gene_list = gene_set,
                                       x1 = "A",
                                       y1 = "B",
                                       nbin = 1,
                                       ctrl = 10)
  p

  # Using three variables. Figure from: https://www.nature.com/articles/nature20123.
  p <- SCpubr::do_CellularStatesPlot(sample = sample,
                                       input_gene_list = gene_set,
                                       x1 = "A",
                                       y1 = "B",
                                       x2 = "C",
                                       nbin = 1,
                                       ctrl = 10)
  p

  # Using four variables. Figure from: https://pubmed.ncbi.nlm.nih.gov/31327527/
  p <- SCpubr::do_CellularStatesPlot(sample = sample,
                                       input_gene_list = gene_set,
                                       x1 = "A",
                                       y1 = "C",
                                       x2 = "B",
                                       y2 = "D",
                                       nbin = 1,
                                       ctrl = 10)
  p
} else if (base::isFALSE(value)) {
```

```

  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()``.")
}

```

do_ChordDiagramPlot *Generate a Chord diagram.*

Description

Generate a Chord diagram.

Usage

```

do_ChordDiagramPlot(
  sample = NULL,
  from = NULL,
  to = NULL,
  colors.from = NULL,
  colors.to = NULL,
  big.gap = 10,
  small.gap = 1,
  link.border.color = NA,
  link.border.width = 1,
  highlight_group = NULL,
  alpha.highlight = 25,
  link.sort = NULL,
  link.decreasing = TRUE,
  z_index = FALSE,
  self.link = 1,
  symmetric = FALSE,
  directional = 1,
  direction.type = c("diffHeight", "arrows"),
  link.arr.type = "big.arrow",
  scale = FALSE,
  alignment = "default",
  annotationTrack = c("grid", "axis"),
  padding_labels = 4,
  ...
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
from, to	character Categorical metadata variable to be used as origin and end points of the interactions.

colors.from, colors.to
`named_vector` | Named vector of colors corresponding to the unique values of "from" and "to".

big.gap `numeric` | Space between the groups in "from" and "to".

small.gap `numeric` | Space within the groups.

link.border.color
`character` | Color for the border of the links. NA = no color.

link.border.width
`numeric` | Width of the border line of the links.

highlight_group
`character` | A value from from that will be used to highlight only the links coming from it.

alpha.highlight
`numeric` | A value between 00 (double digits) and 99 to depict the alpha of the highlighted links. No transparency needs "FF"

link.sort pass to `chordDiagramFromMatrix` or `chordDiagramFromDataFrame`

link.decreasing
pass to `chordDiagramFromMatrix` or `chordDiagramFromDataFrame`

z_index `logical` | Whether to bring the bigger links to the top.

self.link `numeric` | Behavior of the links. One of:

- 1: Prevents self linking.
- 2: Allows self linking.

symmetric pass to `chordDiagramFromMatrix`

directional `numeric` | Set the direction of the links. One of:

- 0: Non-directional data.
- 1: Links go from "from" to "to".
- -1: Links go from "to" to "from".
- 2: Links go in both directions.

direction.type `character` | How to display the directions. One of:

- diffHeight: Sets a line at the origin of the group showing to how many groups and in which proportion this group is linked to.
- arrows: Sets the connection as arrows.
- both: Sets up both behaviors. Use as: c("diffHeight", "arrows").

link.arr.type `character` | Sets the appearance of the arrows. One of:

- triangle: Arrow with a triangle tip at the end displayed on top of the link.
- big.arrow: The link itself ends in a triangle shape.

scale `logical` | Whether to put all nodes the same width.

alignment `character` | How to align the diagram. One of:

- default: Allows `circize` to set up the plot as it sees fit.
- horizontal: Sets the break between "from" and "to" groups on the horizontal axis.

- vertical: Sets the break between "from" and "to" groups on the vertical axis.
- annotationTrack
 pass to `chordDiagramFromMatrix` or `chordDiagramFromDataFrame`
- padding_labels `numeric` | Number of extra padding (white spaces) of the labels so that they do not overlap with the scales.
- ...
 For internal use only.

Value

A circlize plot.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_ChordDiagramPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Basic chord diagram.
  sample$assignment <- ifelse(sample$seurat_clusters %in% c("0", "4", "7"), "A", "B")
  sample$assignment[sample$seurat_clusters %in% c("1", "2")] <- "C"
  sample$assignment[sample$seurat_clusters %in% c("10", "5")] <- "D"
  sample$assignment[sample$seurat_clusters %in% c("8", "9")] <- "E"

  p <- SCpubr::do_ChordDiagramPlot(sample = sample,
                                    from = "seurat_clusters",
                                    to = "assignment")

  p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

do_ColorPalette

Generate color scales based on a value.

Description

This function is an adaptation of colortools package. As the package was removed from CRAN on 23-06-2022, this utility function came to existence in order to cover the gap. It is, on its basis, an adaptation of the package into a single function. Original code, developed by Gaston Sanchez, can be found in: <https://github.com/gastonstat/colortools>

Usage

```
do_ColorPalette(
  colors.use,
  n = 12,
  opposite = FALSE,
  adjacent = FALSE,
  triadic = FALSE,
  split_complementary = FALSE,
  tetradic = FALSE,
  square = FALSE,
  complete_output = FALSE,
  plot = FALSE,
  font.size = 14,
  font.type = "sans"
)
```

Arguments

colors.use	<code>character</code> One color upon which generate the color scale. Can be a name or a HEX code.
n	<code>numeric</code> Number of colors to include in the color wheel. Use it when all other options are FALSE, otherwise, it becomes 12.
opposite	<code>logical</code> Return the opposing color to the one provided.
adjacent	<code>logical</code> Return the adjacent colors to the one provided.
triadic	<code>logical</code> Return the triadic combination of colors to the one provided.
split_complementary	<code>logical</code> Return the split complementary combination of colors to the one provided.
tetradic	<code>logical</code> Return the tetradic combination of colors to the one provided.
square	<code>logical</code> Return the square combination of colors to the one provided.
complete_output	<code>logical</code> Runs all the previous options and returns all the outputs as a list that contains all color vectors, all plots and a combined plot with everything.
plot	<code>logical</code> Whether to also return a plot displaying the values instead of a vector with the color.
font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • mono: Mono spaced font. • serif: Serif font family. • sans: Default font family.

Value

A character vector with the desired color scale.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_ColorPalette", passive = TRUE)

if (isTRUE(value)){
  # Generate a color wheel based on a single value.
  colors <- SCpubr:::do_ColorPalette(colors.use = "steelblue")
  p <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                plot = TRUE)

  # Generate a pair of opposite colors based on a given one.
  colors <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                      opposite = TRUE)
  p <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                 opposite = TRUE,
                                 plot = TRUE)

  # Generate a trio of adjacent colors based on a given one.
  colors <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                      adjacent = TRUE)
  p <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                 adjacent = TRUE,
                                 plot = TRUE)

  # Generate a trio of triadic colors based on a given one.
  colors <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                      triadic = TRUE)
  p <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                 triadic = TRUE,
                                 plot = TRUE)

  # Generate a trio of split complementary colors based on a given one.
  colors <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                      split_complementary = TRUE)
  p <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                 split_complementary = TRUE,
                                 plot = TRUE)

  # Generate a group of tetradic colors based on a given one.
  colors <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                      tetradiic = TRUE)
  p <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                 tetradiic = TRUE,
                                 plot = TRUE)

  # Generate a group of square colors based on a given one.
  colors <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                      square = TRUE)
  p <- SCpubr:::do_ColorPalette(colors.use = "steelblue",
                                 square = TRUE,
                                 plot = TRUE)
```

```

# Retrieve the output of all options.
out <- SCpubr::do_ColorPalette(colors.use = "steelblue",
                                complete_output = TRUE)
## Retrieve the colors.
colors <- out$colors
## Retrieve the plots.
plots <- out$plots
## Retrieve a combined plot with all the options.
p <- out$combined_plot

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_CopyNumberVariantPlot*Display CNV scores from inferCNV as Feature Plots.***Description**

Display CNV scores from inferCNV as Feature Plots.

Usage

```

do_CopyNumberVariantPlot(
  sample,
  infercnv_object,
  chromosome_locations,
  group.by = NULL,
  using_metacells = FALSE,
  metacell_mapping = NULL,
  include_chr_arms = FALSE,
  legend.type = "colorbar",
  legend.position = "bottom",
  legend.length = 20,
  legend.width = 1,
  legend.framewidth = 0.5,
  legend.tickwidth = 0.5,
  legend.framecolor = "grey50",
  legend.tickcolor = "white",
  font.size = 14,
  pt.size = 1,
  font.type = "sans",
  axis.text.x.angle = 45,
  enforce_symmetry = TRUE,
  legend.title = NULL,

```

```

na.value = "grey75",
viridis.palette = "G",
viridis.direction = 1,
verbose = FALSE,
min.cutoff = NA,
max.cutoff = NA,
number.breaks = 5,
diverging.palette = "RdBu",
diverging.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = -1,
use_viridis = TRUE,
return_object = FALSE,
grid.color = "white",
border.color = "black",
flip = FALSE,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
infercnv_object	<code>infercnv</code> Output inferCNV object run on the same Seurat object.
chromosome_locations	<code>tibble</code> Tibble containing the chromosome regions to use. Can be obtained using <code>utils::data("human_chr_locations", package = "SCpubr")</code> .
group_by	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
using_metacells	<code>logical</code> Whether inferCNV was run using metacells or not.
metacell_mapping	<code>named_vector</code> Vector or cell - metacell mapping.
include_chr_arms	<code>logical</code> Whether the output heatmap should also include chromosome arms or just whole chromosomes.
legend.type	<code>character</code> Type of legend to display. One of: <ul style="list-style-type: none"> • <code>normal</code>: Default legend displayed by <code>ggplot2</code>. • <code>colorbar</code>: Redefined colorbar legend, using <code>guide_colorbar</code>.
legend.position	<code>character</code> Position of the legend in the plot. One of:

- `top`: Top of the figure.
- `bottom`: Bottom of the figure.
- `left`: Left of the figure.
- `right`: Right of the figure.
- `none`: No legend is displayed.

`legend.length, legend.width`
`numeric` | Length and width of the legend. Will adjust automatically depending on legend side.

`legend.framewidth, legend.tickwidth`
`numeric` | Width of the lines of the box in the legend.

`legend.framecolor`
`character` | Color of the lines of the box in the legend.

`legend.tickcolor`
`character` | Color of the ticks of the box in the legend.

`font.size`
`numeric` | Overall font size of the plot. All plot elements will have a size relationship with this font size.

`pt.size`
`numeric` | Size of the dots.

`font.type`
`character` | Base font family for the plot. One of:

- `mono`: Mono spaced font.
- `serif`: Serif font family.
- `sans`: Default font family.

`axis.text.x.angle`
`numeric` | Degree to rotate the X labels. One of: 0, 45, 90.

`enforce_symmetry`
`logical` | Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.

`legend.title`
`character` | Title for the legend.

`na.value`
`character` | Color value for NA.

`viridis.palette`
`character` | A capital letter from A to H or the scale name as in `scale_fill_viridis`.

`viridis.direction`
`numeric` | Either 1 or -1. Controls how the gradient of viridis scale is formed.

`verbose`
`logical` | Whether to show extra comments, warnings,etc.

`min.cutoff, max.cutoff`
`numeric` | Set the min/max ends of the color scale. Any cell/group with a value lower than `min.cutoff` will turn into `min.cutoff` and any cell with a value higher than `max.cutoff` will turn into `max.cutoff`. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.

`number.breaks`
`numeric` | Controls the number of breaks in continuous color scales of ggplot2-based plots.

`diverging.palette`
`character` | Type of symmetrical color palette to use. Out of the diverging palettes defined in `brewer.pal`.

diverging.direction	<code>numeric</code> Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.
sequential.palette	<code>character</code> Type of sequential color palette to use. Out of the sequential palettes defined in <code>brewer.pal</code> .
sequential.direction	<code>numeric</code> Direction of the sequential color scale. Either 1 or -1.
use_viridis	<code>logical</code> Whether to use viridis color scales.
return_object	<code>logical</code> Returns the Seurat object with the modifications performed in the function. Normally, this contains a new assay with the data that can then be used for any other visualization desired.
grid.color	<code>character</code> Color of the grid in the plot. In heatmaps, color of the border of the cells.
border.color	<code>character</code> Color for the border of the heatmap body.
flip	<code>logical</code> Whether to invert the axis of the displayed plot.
plot.title.face, axis.title.face	<code>character</code> Controls the style of the font for the corresponding theme element.
plot.subtitle.face, axis.text.face	One of:
plot.caption.face, legend.title.face, legend.text.face	<ul style="list-style-type: none">• plain: For normal text.• italic: For text in italic.• bold: For text in bold.• bold.italic: For text both in italic and bold.

Value

A list containing Feature Plots for different chromosome regions and corresponding dot plots by groups..

Examples

```
# Get human chromosome locations.  
chromosome_locations = SCpubr::human_chr_locations  
  
# Compute for all chromosomes.  
p <- SCpubr::do_CopyNumberVariantPlot(sample = sample,  
                                         infercnv_object = infercnv_object,  
                                         using_metacells = FALSE,  
                                         chromosome_locations = chromosome_locations)  
  
} else if (base::isFALSE(value)){  
  message("This function can not be used without its suggested packages.")  
  message("Check out which ones are needed using `SCpubr::state_dependencies()`.")  
}  

```

do_CorrelationPlot *Create correlation matrix heatmaps.*

Description

Create correlation matrix heatmaps.

Usage

```
do_CorrelationPlot(  
  sample = NULL,  
  input_gene_list = NULL,  
  cluster = TRUE,  
  remove.diagonal = TRUE,  
  mode = "hvg",  
  assay = NULL,  
  group.by = NULL,  
  legend.title = "Pearson coef.",  
  enforce_symmetry = ifelse(mode == "hvg", TRUE, FALSE),  
  font.size = 14,  
  font.type = "sans",  
  na.value = "grey75",  
  legend.width = 1,  
  legend.length = 20,  
  legend.framewidth = 0.5,  
  legend.tickwidth = 0.5,  
  legend.framecolor = "grey50",  
  legend.tickcolor = "white",  
  legend.type = "colorbar",  
  legend.position = "bottom",  
  min.cutoff = NA,  
  max.cutoff = NA,
```

```

number.breaks = 5,
plot.title = NULL,
plot.subtitle = NULL,
plot.caption = NULL,
diverging.palette = "RdBu",
diverging.direction = -1,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
axis.text.x.angle = 45,
grid.color = "white",
border.color = "black",
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
input_gene_list	<code>named_list</code> Named list of lists of genes to be used as input.
cluster	<code>logical</code> Whether to cluster the elements in the heatmap or not.
remove.diagonal	<code>logical</code> Whether to convert diagonal to NA. Normally this value would be 1, heavily shifting the color scale.
mode	<code>character</code> Different types of correlation matrices can be computed. Right now, the only possible value is "hvg", standing for Highly Variable Genes. The sample is subset for the HVG and the data is re-scaled. Scale data is used for the correlation.
assay	<code>character</code> Assay to use. Defaults to the current assay.
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
legend.title	<code>character</code> Title for the legend.
enforce_symmetry	<code>logical</code> Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.
font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	<code>character</code> Base font family for the plot. One of:

- `mono`: Mono spaced font.
 - `serif`: Serif font family.
 - `sans`: Default font family.
- `na.value` `character` | Color value for NA.
- `legend.length, legend.width`
- `numeric` | Length and width of the legend. Will adjust automatically depending on legend side.
- `legend.framewidth, legend.tickwidth`
- `numeric` | Width of the lines of the box in the legend.
- `legend.framecolor`
- `character` | Color of the lines of the box in the legend.
- `legend.tickcolor`
- `character` | Color of the ticks of the box in the legend.
- `legend.type` `character` | Type of legend to display. One of:
- `normal`: Default legend displayed by `ggplot2`.
 - `colorbar`: Redefined colorbar legend, using `guide_colorbar`.
- `legend.position`
- `character` | Position of the legend in the plot. One of:
- `top`: Top of the figure.
 - `bottom`: Bottom of the figure.
 - `left`: Left of the figure.
 - `right`: Right of the figure.
 - `none`: No legend is displayed.
- `min.cutoff, max.cutoff`
- `numeric` | Set the min/max ends of the color scale. Any cell/group with a value lower than `min.cutoff` will turn into `min.cutoff` and any cell with a value higher than `max.cutoff` will turn into `max.cutoff`. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
- `number.breaks` `numeric` | Controls the number of breaks in continuous color scales of ggplot2-based plots.
- `plot.title, plot.subtitle, plot.caption`
- `character` | Title, subtitle or caption to use in the plot.
- `diverging.palette`
- `character` | Type of symmetrical color palette to use. Out of the diverging palettes defined in `brewer.pal`.
- `diverging.direction`
- `numeric` | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.
- `use_viridis` `logical` | Whether to use viridis color scales.
- `viridis.palette`
- `character` | A capital letter from A to H or the scale name as in `scale_fill_viridis`.
- `viridis.direction`
- `numeric` | Either 1 or -1. Controls how the gradient of viridis scale is formed.

```

sequential.palette
  character | Type of sequential color palette to use. Out of the sequential
  palettes defined in brewer.pal.
sequential.direction
  numeric | Direction of the sequential color scale. Either 1 or -1.
axis.text.x.angle
  numeric | Degree to rotate the X labels. One of: 0, 45, 90.
grid.color    character | Color of the grid in the plot. In heatmaps, color of the border of the
cells.
border.color   character | Color for the border of the heatmap body.
plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
  character | Controls the style of the font for the corresponding theme element.
  One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.

```

Value

A ggplot2 object.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_CorrelationPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Default values.
  p <- SCpubr:::do_CorrelationPlot(sample = sample)
  p

} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_DimPlot *Wrapper for DimPlot.*

Description

Wrapper for [DimPlot](#).

Usage

```
do_DimPlot(  
  sample,  
  reduction = NULL,  
  group.by = NULL,  
  split.by = NULL,  
  split.by.combined = TRUE,  
  colors.use = NULL,  
  shuffle = TRUE,  
  order = NULL,  
  raster = FALSE,  
  pt.size = 1,  
  label = FALSE,  
  label.color = "black",  
  label.fill = "white",  
  label.size = 4,  
  label.box = TRUE,  
  repel = FALSE,  
  cells.highlight = NULL,  
  idents.highlight = NULL,  
  idents.keep = NULL,  
  sizes.highlight = 1,  
  ncol = NULL,  
  plot.title = NULL,  
  plot.subtitle = NULL,  
  plot.caption = NULL,  
  legend.title = NULL,  
  legend.position = "bottom",  
  legend.title.position = "top",  
  legend.ncol = NULL,  
  legend.nrow = NULL,  
  legend.icon.size = 4,  
  legend.byrow = FALSE,  
  legend.dot.border = TRUE,  
  raster.dpi = 2048,  
  dims = c(1, 2),  
  font.size = 14,  
  font.type = "sans",  
  na.value = "grey75",
```

```

plot_cell_borders = TRUE,
border.size = 2,
border.color = "black",
border.density = 1,
plot_marginal_distributions = FALSE,
marginal.type = "density",
marginal.size = 5,
marginal.group = TRUE,
plot.axes = FALSE,
plot_density_contour = FALSE,
contour.position = "bottom",
contour.color = "grey90",
contour.lineend = "butt",
contour.linejoin = "round",
contour_expand_axes = 0.25,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
reduction	character Reduction to use. Can be the canonical ones such as "umap", "pca", or any custom ones, such as "diffusion". If you are unsure about which reductions you have, use <code>Seurat::Reductions(sample)</code> . Defaults to "umap" if present or to the last computed reduction if the argument is not provided.
group.by	character Metadata variable to group the output by. Has to be a character or factor column.
split.by	character Secondary metadata variable to further group (split) the output by. Has to be a character of factor column.
split.by.combined	logical Adds a combined view of all the values before splitting them by split.by. Think of this as a regular DimPlot added in front. This is set to TRUE if split.by is used in combination with group.by.
colors.use	named_vector Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of group.by. If group.by is not provided, defaults to the unique values of Idents . If not provided, a color scale will be set by default.
shuffle	logical Whether to shuffle the cells or not, so that they are not plotted cluster-wise. Recommended.
order	character Vector of identities to be plotted. Either one with all identities or just some, which will be plotted last.

raster	<code>logical</code> Whether to raster the resulting plot. This is recommendable if plotting a lot of cells.
pt.size	<code>numeric</code> Size of the dots.
label	<code>logical</code> Whether to plot the cluster labels in the UMAP. The cluster labels will have the same color as the cluster colors.
label.color	<code>character</code> Color of the labels in the plot.
label.fill	<code>character</code> Color to fill the labels. Has to be a single color, that will be used for all labels. If NULL, the colors of the clusters will be used instead.
label.size	<code>numeric</code> Size of the labels in the plot.
label.box	<code>logical</code> Whether to plot the plot labels as <code>geom_text</code> (FALSE) or <code>geom_label</code> (TRUE).
repel	<code>logical</code> Whether to repel the text labels.
cells.highlight, idents.highlight	<code>character</code> Vector of cells/identities to focus into. The identities have to much those in <code>Seurat::Idents(sample)</code> The rest of the cells will be grayed out. Both parameters can be used at the same time.
idents.keep	<code>character</code> Vector of identities to keep. This will effectively set the rest of the cells that do not match the identities provided to NA, therefore coloring them according to na.value parameter.
sizes.highlight	<code>numeric</code> Point size of highlighted cells using cells.highlight parameter.
ncol	<code>numeric</code> Number of columns used in the arrangement of the output plot using "split.by" parameter.
plot.title, plot.subtitle, plot.caption	<code>character</code> Title, subtitle or caption to use in the plot.
legend.title	<code>character</code> Title for the legend.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • top: Top of the figure. • bottom: Bottom of the figure. • left: Left of the figure. • right: Right of the figure. • none: No legend is displayed.
legend.title.position	<code>character</code> Position for the title of the legend. One of: <ul style="list-style-type: none"> • top: Top of the legend. • bottom: Bottom of the legend. • left: Left of the legend. • right: Right of the legend.
legend.ncol	<code>numeric</code> Number of columns in the legend.
legend.nrow	<code>numeric</code> Number of rows in the legend.

```

legend.icon.size      numeric | Size of the icons in legend.
legend.byrow         logical | Whether the legend is filled by row or not.
legend.dot.border    logical | Adds a black border around the dots in the legend.
raster.dpi          numeric | Pixel resolution for rasterized plots. Defaults to 1024. Only activates on Seurat versions higher or equal than 4.1.0.
dims                numeric | Vector of 2 numerics indicating the dimensions to plot out of the selected reduction. Defaults to c(1, 2) if not specified.
font.size            numeric | Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type            character | Base font family for the plot. One of:
                           • mono: Mono spaced font.
                           • serif: Serif font family.
                           • sans: Default font family.
na.value             character | Color value for NA.
plot_cell_borders   logical | Whether to plot border around cells.
border.size          numeric | Width of the border of the cells.
border.color          character | Color for the border of the heatmap body.
border.density        numeric | Controls the number of cells used when plot_cell_borders = TRUE. Value between 0 and 1. It computes a 2D kernel density and based on this cells that have a density below the specified quantile will be used to generate the cluster contour. The lower this number, the less cells will be selected, thus reducing the overall size of the plot but also potentially preventing all the contours to be properly drawn.
plot_marginal_distributions
                           logical | Whether to plot marginal distributions on the figure or not.
marginal.type        character | One of:
                           • density: Compute density plots on the margins.
                           • histogram: Compute histograms on the margins.
                           • boxplot: Compute boxplot on the margins.
                           • violin: Compute violin plots on the margins.
                           • densigram: Compute densigram plots on the margins.
marginal.size         numeric | Size ratio between the main and marginal plots. A value of 5 means that the main plot is 5 times bigger than the marginal plots.
marginal.group        logical | Whether to group the marginal distribution by group.by or current identities.
plot.axes             logical | Whether to plot axes or not.
plot_density_contour
                           logical | Whether to plot density contours in the UMAP.

```

```

contour.position
  character | Whether to plot density contours on top or at the bottom of the
  visualization layers, thus overlapping the clusters/cells or not.
contour.color  character | Color of the density lines.
contour.lineend
  character | Line end style (round, butt, square).
contour.linejoin
  character | Line join style (round, mitre, bevel).
contour_expand_axes
  numeric | To make the contours fit the plot, the limits of the X and Y axis are
  expanding a given percentage from the min and max values for each axis. This
  controls such percentage.
plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
  character | Controls the style of the font for the corresponding theme element.
One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.

```

Value

A ggplot2 object containing a DimPlot.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_DimPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Basic DimPlot.
  p <- SCpubr::do_DimPlot(sample = sample)

  # Restrict the amount of identities displayed.
  p <- SCpubr::do_DimPlot(sample = sample,
                         idents.keep = c("1", "3", "5"))

  # Group by another variable rather than `Seurat::Idents(sample)~
  p <- SCpubr::do_DimPlot(sample = sample,
                         group.by = "seurat_clusters")

  # Split the output in as many plots as unique identities.
}

```

```

p <- SCpubr::do_DimPlot(sample = sample,
                         split.by = "seurat_clusters")

# Highlight given identities
p <- SCpubr::do_DimPlot(sample,
                         idents.highlight = c("1", "3"))

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_DotPlot

This function is a wrapper for [DotPlot](#). It provides most of its functionalities while adding extra. You can

Description

This function is a wrapper for [DotPlot](#). It provides most of its functionalities while adding extra.
You can

Usage

```

do_DotPlot(
  sample,
  features,
  assay = NULL,
  slot = "data",
  group.by = NULL,
  split.by = NULL,
  zscore.data = FALSE,
  min.cutoff = NA,
  max.cutoff = NA,
  enforce_symmetry = ifelse(base::isTRUE(zscore.data), TRUE, FALSE),
  legend.title = NULL,
  legend.type = "colorbar",
  legend.position = "bottom",
  legend.framewidth = 0.5,
  legend.tickwidth = 0.5,
  legend.length = 20,
  legend.width = 1,
  legend.framecolor = "grey50",
  legend.tickcolor = "white",
  legend.ncol = NULL,

```

```

legend.nrow = NULL,
legend.byrow = FALSE,
dot.scale = 8,
plot.title = NULL,
plot.subtitle = NULL,
plot.caption = NULL,
xlab = NULL,
ylab = NULL,
font.size = 14,
font.type = "sans",
cluster = FALSE,
flip = FALSE,
axis.text.x.angle = 45,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
diverging.palette = "RdBu",
diverging.direction = -1,
na.value = "grey75",
plot.grid = TRUE,
grid.color = "grey75",
grid.type = "dashed",
number.breaks = 5,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
features	character Features to represent.
assay	character Assay to use. Defaults to the current assay.
slot	character Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
group.by	character Metadata variable to group the output by. Has to be a character of factor column.
split.by	character Secondary metadata variable to further group (split) the output by. Has to be a character of factor column.
zscore.data	logical Whether to compute Z-scores instead of showing average expression values. This allows to see, for each gene, which group has the highest average

expression, but prevents you from comparing values across genes. Can not be used with slot = "scale.data" or with split.by.

<code>min.cutoff, max.cutoff</code>	<code>numeric</code> Set the min/max ends of the color scale. Any cell/group with a value lower than min.cutoff will turn into min.cutoff and any cell with a value higher than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
<code>enforce_symmetry</code>	<code>logical</code> Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.
<code>legend.title</code>	<code>character</code> Title for the legend.
<code>legend.type</code>	<code>character</code> Type of legend to display. One of: <ul style="list-style-type: none"> • <code>normal</code>: Default legend displayed by <code>ggplot2</code>. • <code>colorbar</code>: Redefined colorbar legend, using <code>guide_colorbar</code>.
<code>legend.position</code>	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • <code>top</code>: Top of the figure. • <code>bottom</code>: Bottom of the figure. • <code>left</code>: Left of the figure. • <code>right</code>: Right of the figure. • <code>none</code>: No legend is displayed.
<code>legend.framewidth, legend.tickwidth</code>	<code>numeric</code> Width of the lines of the box in the legend.
<code>legend.length, legend.width</code>	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.
<code>legend.framecolor</code>	<code>character</code> Color of the lines of the box in the legend.
<code>legend.tickcolor</code>	<code>character</code> Color of the ticks of the box in the legend.
<code>legend.ncol</code>	<code>numeric</code> Number of columns in the legend.
<code>legend.nrow</code>	<code>numeric</code> Number of rows in the legend.
<code>legend.byrow</code>	<code>logical</code> Whether the legend is filled by row or not.
<code>dot.scale</code>	<code>numeric</code> Scale the size of the dots.
<code>plot.title, plot.subtitle, plot.caption</code>	<code>character</code> Title, subtitle or caption to use in the plot.
<code>xlab, ylab</code>	<code>character</code> Titles for the X and Y axis.
<code>font.size</code>	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
<code>font.type</code>	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • <code>mono</code>: Mono spaced font. • <code>serif</code>: Serif font family.

- sans: Default font family.

cluster `logical` | Whether to cluster the identities based on the expression of the features.

flip `logical` | Whether to invert the axis of the displayed plot.

axis.text.x.angle `numeric` | Degree to rotate the X labels. One of: 0, 45, 90.

use_viridis `logical` | Whether to use viridis color scales.

viridis.palette `character` | A capital letter from A to H or the scale name as in `scale_fill_viridis`.

viridis.direction `numeric` | Either 1 or -1. Controls how the gradient of viridis scale is formed.

sequential.palette `character` | Type of sequential color palette to use. Out of the sequential palettes defined in `brewer.pal`.

sequential.direction `numeric` | Direction of the sequential color scale. Either 1 or -1.

diverging.palette `character` | Type of symmetrical color palette to use. Out of the diverging palettes defined in `brewer.pal`.

diverging.direction `numeric` | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.

na.value `character` | Color value for NA.

plot.grid `logical` | Whether to plot grid lines.

grid.color `character` | Color of the grid in the plot. In heatmaps, color of the border of the cells.

grid.type `character` | One of the possible linetype options:

- blank.
- solid.
- dashed.
- dotted.
- dotdash.
- longdash.
- twodash.

number.breaks `numeric` | Controls the number of breaks in continuous color scales of ggplot2-based plots.

plot.title.face, plot.subtitle.face, plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face

`character` | Controls the style of the font for the corresponding theme element.
One of:

- plain: For normal text.
- italic: For text in italic.
- bold: For text in bold.
- bold.italic: For text both in italic and bold.

Value

A ggplot2 object containing a Dot Plot.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_DotPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  # sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Basic Dot plot.
  # genes <- rownames(sample)[1:14]
  # p <- SCpubr:::do_DotPlot(sample = sample,
  #                           features = genes)

} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

do_EnrichmentHeatmap *Create enrichment scores heatmaps.*

Description

This function computes the enrichment scores for the cells using [AddModuleScore](#) and then aggregates the scores by the metadata variables provided by the user and displays it as a heatmap, computed by [Heatmap](#).

Usage

```
do_EnrichmentHeatmap(
  sample,
  input_gene_list,
  features.order = NULL,
  groups.order = NULL,
  cluster = TRUE,
  scale_scores = FALSE,
  assay = NULL,
  slot = NULL,
  reduction = NULL,
  group.by = NULL,
  verbose = FALSE,
```

```
na.value = "grey75",
legend.position = "bottom",
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = 1,
legend.framewidth = 0.5,
legend.tickwidth = 0.5,
legend.length = 20,
legend.width = 1,
legend.framecolor = "grey50",
legend.tickcolor = "white",
legend.type = "colorbar",
font.size = 14,
font.type = "sans",
axis.text.x.angle = 45,
enforce_symmetry = FALSE,
nbin = 24,
ctrl = 100,
flavor = "Seurat",
legend.title = NULL,
ncores = 1,
storeRanks = TRUE,
min.cutoff = NA,
max.cutoff = NA,
pt.size = 1,
plot_cell_borders = TRUE,
border.size = 2,
return_object = FALSE,
number.breaks = 5,
sequential.palette = "YlGnBu",
diverging.palette = "RdBu",
diverging.direction = -1,
sequential.direction = 1,
flip = FALSE,
grid.color = "white",
border.color = "black",
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)
```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
--------	---

```

input_gene_list
  named_list | Named list of lists of genes to be used as input.

features.order character | Should the gene sets be ordered in a specific way? Provide it as a vector of characters with the same names as the names of the gene sets.

groups.order named_list | Should the groups in theheatmaps be ordered in a specific way? Provide it as a named list (as many lists as values in group.by) with the order for each of the elements in the groups.

cluster logical | Whether to perform clustering of rows and columns.

scale_scores logical | Whether to transform the scores to a range of 0-1 for plotting.

assay character | Assay to use. Defaults to the current assay.

slot character | Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".

reduction character | Reduction to use. Can be the canonical ones such as "umap", "pca", or any custom ones, such as "diffusion". If you are unsure about which reductions you have, use Seurat::Reductions(sample). Defaults to "umap" if present or to the last computed reduction if the argument is not provided.

group.by character | Metadata variable to group the output by. Has to be a character of factor column.

verbose logical | Whether to show extra comments, warnings,etc.

na.value character | Color value for NA.

legend.position
  character | Position of the legend in the plot. One of:
    • top: Top of the figure.
    • bottom: Bottom of the figure.
    • left: Left of the figure.
    • right: Right of the figure.
    • none: No legend is displayed.

use_viridis logical | Whether to use viridis color scales.

viridis.palette
  character | A capital letter from A to H or the scale name as in scale_fill_viridis.

viridis.direction
  numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.

legend.framewidth, legend.tickwidth
  numeric | Width of the lines of the box in the legend.

legend.length, legend.width
  numeric | Length and width of the legend. Will adjust automatically depending on legend side.

legend.framecolor
  character | Color of the lines of the box in the legend.

legend.tickcolor
  character | Color of the ticks of the box in the legend.

legend.type character | Type of legend to display. One of:

```

	<ul style="list-style-type: none"> • normal: Default legend displayed by ggplot2. • colorbar: Redefined colorbar legend, using guide_colorbar.
font.size	numeric Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	character Base font family for the plot. One of: <ul style="list-style-type: none"> • mono: Mono spaced font. • serif: Serif font family. • sans: Default font family.
axis.text.x.angle	numeric Degree to rotate the X labels. One of: 0, 45, 90.
enforce_symmetry	logical Whether the geyser and feature plot has a symmetrical color scale.
nbin	numeric Number of bins to use in AddModuleScore .
ctrl	numeric Number of genes in the control set to use in AddModuleScore .
flavor	character One of: Seurat, UCell. Compute the enrichment scores using AddModuleScore or AddModuleScore_UCell .
legend.title	character Title for the legend.
ncores	numeric Number of cores used to run UCell scoring.
storeRanks	logical Whether to store the ranks for faster UCell scoring computations. Might require large amounts of RAM.
min.cutoff, max.cutoff	numeric Set the min/max ends of the color scale. Any cell/group with a value lower than min.cutoff will turn into min.cutoff and any cell with a value higher than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
pt.size	numeric Size of the dots.
plot_cell_borders	logical Whether to plot border around cells.
border.size	numeric Width of the border of the cells.
return_object	logical Return the Seurat object with the enrichment scores stored.
number.breaks	numeric Controls the number of breaks in continuous color scales of ggplot2-based plots.
sequential.palette	character Type of sequential color palette to use. Out of the sequential palettes defined in brewer.pal .
diverging.palette	character Type of symmetrical color palette to use. Out of the diverging palettes defined in brewer.pal .
diverging.direction	numeric Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.
sequential.direction	numeric Direction of the sequential color scale. Either 1 or -1.

flip `logical` | Whether to invert the axis of the displayed plot.

grid.color `character` | Color of the grid in the plot. In heatmaps, color of the border of the cells.

border.color `character` | Color for the border of the heatmap body.

plot.title.face, **plot.subtitle.face**, **plot.caption.face**,
axis.title.face, **axis.text.face**, **legend.title.face**, **legend.text.face**
`character` | Controls the style of the font for the corresponding theme element.
One of:

- **plain**: For normal text.
- **italic**: For text in italic.
- **bold**: For text in bold.
- **bold.italic**: For text both in italic and bold.

Value

A ggplot2 object.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_EnrichmentHeatmap", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Genes have to be unique.
  genes <- list("A" = rownames(sample)[1:5],
                "B" = rownames(sample)[6:10],
                "C" = rownames(sample)[11:15])

  # Default parameters.
  p <- SCpubr::do_EnrichmentHeatmap(sample = sample,
                                      input_gene_list = genes,
                                      nbin = 1,
                                      ctrl = 10)
  p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`.")
}
```

do_ExpressionHeatmap *Create heatmaps of averaged expression by groups.*

Description

This function generates a heatmap with averaged expression values by the unique groups of the metadata variables provided by the user.

Usage

```
do_ExpressionHeatmap(  
  sample,  
  features,  
  group.by = NULL,  
  assay = NULL,  
  cluster = TRUE,  
  features.order = NULL,  
  groups.order = NULL,  
  slot = "data",  
  legend.title = "Avg. Expression",  
  na.value = "grey75",  
  legend.position = "bottom",  
  legend.width = 1,  
  legend.length = 20,  
  legend.framewidth = 0.5,  
  legend.tickwidth = 0.5,  
  legend.framecolor = "grey50",  
  legend.tickcolor = "white",  
  legend.type = "colorbar",  
  font.size = 14,  
  font.type = "sans",  
  axis.text.x.angle = 45,  
  enforce_symmetry = FALSE,  
  min.cutoff = NA,  
  max.cutoff = NA,  
  diverging.palette = "RdBu",  
  diverging.direction = -1,  
  sequential.palette = "YlGnBu",  
  sequential.direction = 1,  
  number.breaks = 5,  
  use_viridis = FALSE,  
  viridis.palette = "G",  
  viridis.direction = -1,  
  flip = FALSE,  
  grid.color = "white",  
  border.color = "black",  
  plot.title.face = "bold",
```

```

plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
features	<code>character</code> Features to represent.
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
assay	<code>character</code> Assay to use. Defaults to the current assay.
cluster	<code>logical</code> Whether to perform clustering of rows and columns.
features.order	<code>character</code> Should the gene sets be ordered in a specific way? Provide it as a vector of characters with the same names as the names of the gene sets.
groups.order	<code>named_list</code> Should the groups in theheatmaps be ordered in a specific way? Provide it as a named list (as many lists as values in group.by) with the order for each of the elements in the groups.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
legend.title	<code>character</code> Title for the legend.
na.value	<code>character</code> Color value for NA.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • top: Top of the figure. • bottom: Bottom of the figure. • left: Left of the figure. • right: Right of the figure. • none: No legend is displayed.
legend.length, legend.width	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.
legend.framewidth, legend.tickwidth	<code>numeric</code> Width of the lines of the box in the legend.
legend.framecolor	<code>character</code> Color of the lines of the box in the legend.
legend.tickcolor	<code>character</code> Color of the ticks of the box in the legend.
legend.type	<code>character</code> Type of legend to display. One of: <ul style="list-style-type: none"> • normal: Default legend displayed by <code>ggplot2</code>.

- **colorbar**: Redefined colorbar legend, using `guide_colorbar`.
- font.size** `numeric` | Overall font size of the plot. All plot elements will have a size relationship with this font size.
- font.type** `character` | Base font family for the plot. One of:
 - `mono`: Mono spaced font.
 - `serif`: Serif font family.
 - `sans`: Default font family.
- axis.text.x.angle** `numeric` | Degree to rotate the X labels. One of: 0, 45, 90.
- enforce_symmetry** `logical` | Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.
- min.cutoff, max.cutoff** `numeric` | Set the min/max ends of the color scale. Any cell/group with a value lower than min.cutoff will turn into min.cutoff and any cell with a value higher than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
- diverging.palette** `character` | Type of symmetrical color palette to use. Out of the diverging palettes defined in `brewer.pal`.
- diverging.direction** `numeric` | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.
- sequential.palette** `character` | Type of sequential color palette to use. Out of the sequential palettes defined in `brewer.pal`.
- sequential.direction** `numeric` | Direction of the sequential color scale. Either 1 or -1.
- number.breaks** `numeric` | Controls the number of breaks in continuous color scales of ggplot2-based plots.
- use_viridis** `logical` | Whether to use viridis color scales.
- viridis.palette** `character` | A capital letter from A to H or the scale name as in `scale_fill_viridis`.
- viridis.direction** `numeric` | Either 1 or -1. Controls how the gradient of viridis scale is formed.
- flip** `logical` | Whether to invert the axis of the displayed plot.
- grid.color** `character` | Color of the grid in the plot. In heatmaps, color of the border of the cells.
- border.color** `character` | Color for the border of the heatmap body.
- plot.title.face, plot.subtitle.face, plot.caption.face, axis.title.face, axis.text.face, legend.title.face, legend.text.face** `character` | Controls the style of the font for the corresponding theme element. One of:
 - `plain`: For normal text.

- **italic**: For text in italic.
- **bold**: For text in bold.
- **bold.italic**: For text both in italic and bold.

Value

A ggplot2 object.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_ExpressionHeatmap", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Define list of genes.
  genes <- rownames(sample)[1:10]

  # Default parameters.
  p <- SCpubr:::do_ExpressionHeatmap(sample = sample,
                                       features = genes,
                                       viridis.direction = -1)

  p
} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

do_FeaturePlot

Wrapper for FeaturePlot.

Description

Wrapper for [FeaturePlot](#).

Usage

```
do_FeaturePlot(
  sample,
  features,
  assay = NULL,
  reduction = NULL,
  slot = NULL,
```

```
order = FALSE,
group.by = NULL,
group.by.colors.use = NULL,
group.by.legend = NULL,
group.by.show.dots = TRUE,
group.by.dot.size = 8,
group.by.cell_borders = FALSE,
group.by.cell_borders.alpha = 0.1,
split.by = NULL,
idents.keep = NULL,
cells.highlight = NULL,
idents.highlight = NULL,
dims = c(1, 2),
enforce_symmetry = FALSE,
symmetry.type = "absolute",
symmetry.center = NA,
pt.size = 1,
font.size = 14,
font.type = "sans",
legend.title = NULL,
legend.type = "colorbar",
legend.position = "bottom",
legend.framewidth = 0.5,
legend.tickwidth = 0.5,
legend.length = 20,
legend.width = 1,
legend.framecolor = "grey50",
legend.tickcolor = "white",
legend.ncol = NULL,
legend.nrow = NULL,
legend.byrow = FALSE,
plot.title = NULL,
plot.subtitle = NULL,
plot.caption = NULL,
individual.titles = NULL,
individual.subtitles = NULL,
individual.captions = NULL,
ncol = NULL,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = 1,
raster = FALSE,
raster.dpi = 1024,
plot_cell_borders = TRUE,
border.size = 2,
border.color = "black",
border.density = 1,
na.value = "grey75",
```

```

verbose = TRUE,
plot.axes = FALSE,
min.cutoff = rep(NA, length(features)),
max.cutoff = rep(NA, length(features)),
plot_density_contour = FALSE,
contour.position = "bottom",
contour.color = "grey90",
contour.lineend = "butt",
contour.linejoin = "round",
contour_expand_axes = 0.25,
label = FALSE,
label.color = "black",
label.size = 4,
number.breaks = 5,
diverging.palette = "RdBu",
diverging.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
features	<code>character</code> Features to represent.
assay	<code>character</code> Assay to use. Defaults to the current assay.
reduction	<code>character</code> Reduction to use. Can be the canonical ones such as "umap", "pca", or any custom ones, such as "diffusion". If you are unsure about which reductions you have, use <code>Seurat::Reductions(sample)</code> . Defaults to "umap" if present or to the last computed reduction if the argument is not provided.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
order	<code>logical</code> Whether to order the cells based on expression.
group.by	<code>character</code> Metadata variable based on which cells are grouped. This will effectively introduce a big dot in the center of each cluster, colored using a categorical color scale or with the values provided by the user in <code>group.by.colors.use</code> . It will also displays a legend.
group.by.colors.use	<code>character</code> Colors to use for the group dots.
group.by.legend	<code>character</code> Title for the legend when <code>group.by</code> is used. Use NA to disable it and NULL to use the default column title provided in <code>group.by</code> .

```

group.by.show.dots
  logical | Controls whether to place in the middle of the groups.

group.by.dot.size
  numeric | Size of the dots placed in the middle of the groups.

group.by.cell_borders
  logical | Plots another border around the cells displaying the same color code
  of the dots displayed with group.by. Legend is shown always with alpha = 1
  regardless of the alpha settings.

group.by.cell_borders.alpha
  numeric | Controls the transparency of the new borders drawn by group.by.cell_borders.

split.by
  character | Secondary metadata variable to further group (split) the output by.
  Has to be a character of factor column.

idents.keep
  character | Vector of identities to plot. The gradient scale will also be subset
  to only the values of such identities.

cells.highlight, idents.highlight
  character | Vector of cells/identities to focus into. The identities have to much
  those in Seurat::Idents(sample) The rest of the cells will be grayed out.
  Both parameters can be used at the same time.

dims
  numeric | Vector of 2 numerics indicating the dimensions to plot out of the
  selected reduction. Defaults to c(1, 2) if not specified.

enforce_symmetry
  logical | Return a symmetrical plot axes-wise or continuous color scale-wise,
  when applicable.

symmetry.type
  character | Type of symmetry to be enforced. One of:
    • absolute: The highest absolute value will be taken into account to generate
      the color scale. Works after min.cutoff and max.cutoff.
    • centered: Centers the scale around the provided value in symmetry.center.
      Works after min.cutoff and max.cutoff.

symmetry.center
  numeric | Value upon which the scale will be centered.

pt.size
  numeric | Size of the dots.

font.size
  numeric | Overall font size of the plot. All plot elements will have a size relationship
  with this font size.

font.type
  character | Base font family for the plot. One of:
    • mono: Mono spaced font.
    • serif: Serif font family.
    • sans: Default font family.

legend.title
  character | Title for the legend.

legend.type
  character | Type of legend to display. One of:
    • normal: Default legend displayed by ggplot2.
    • colorbar: Redefined colorbar legend, using guide_colorbar.

legend.position
  character | Position of the legend in the plot. One of:

```

- **top**: Top of the figure.
 - **bottom**: Bottom of the figure.
 - **left**: Left of the figure.
 - **right**: Right of the figure.
 - **none**: No legend is displayed.
- legend.framework**, **legend.tickwidth**
 numeric | Width of the lines of the box in the legend.
- legend.length**, **legend.width**
 numeric | Length and width of the legend. Will adjust automatically depending on legend side.
- legend.framecolor**
 character | Color of the lines of the box in the legend.
- legend.tickcolor**
 character | Color of the ticks of the box in the legend.
- legend.ncol** **numeric** | Number of columns in the legend.
- legend.nrow** **numeric** | Number of rows in the legend.
- legend.byrow** **logical** | Whether the legend is filled by row or not.
- plot.title**, **plot.subtitle**, **plot.caption**
 character | Title, subtitle or caption to use in the plot.
- individual.titles**, **individual.subtitles**, **individual.captions**
 character | Titles or subtitles. for each feature if needed. Either NULL or a vector of equal length of features.
- ncol** **numeric** | Number of columns used in the arrangement of the output plot using "split.by" parameter.
- use_viridis** **logical** | Whether to use viridis color scales.
- viridis.palette**
 character | A capital letter from A to H or the scale name as in `scale_fill_viridis`.
- viridis.direction**
 numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.
- raster** **logical** | Whether to raster the resulting plot. This is recommendable if plotting a lot of cells.
- raster.dpi** **numeric** | Pixel resolution for rasterized plots. Defaults to 1024. Only activates on Seurat versions higher or equal than 4.1.0.
- plot_cell_borders**
 logical | Whether to plot border around cells.
- border.size** **numeric** | Width of the border of the cells.
- border.color** **character** | Color for the border of the heatmap body.
- border.density** **numeric** | Controls the number of cells used when `plot_cell_borders = TRUE`. Value between 0 and 1. It computes a 2D kernel density and based on this cells that have a density below the specified quantile will be used to generate the cluster contour. The lower this number, the less cells will be selected, thus reducing the overall size of the plot but also potentially preventing all the contours to be properly drawn.

na.value **character** | Color value for NA.
 verbose **logical** | Whether to show extra comments, warnings,etc.
 plot.axes **logical** | Whether to plot axes or not.
 min.cutoff, max.cutoff
 numeric | Set the min/max ends of the color scale. Any cell/group with a value lower than min.cutoff will turn into min.cutoff and any cell with a value higher than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
 plot_density_contour
 logical | Whether to plot density contours in the UMAP.
 contour.position
 character | Whether to plot density contours on top or at the bottom of the visualization layers, thus overlapping the clusters/cells or not.
 contour.color **character** | Color of the density lines.
 contour.lineend
 character | Line end style (round, butt, square).
 contour.linejoin
 character | Line join style (round, mitre, bevel).
 contour_expand_axes
 numeric | To make the contours fit the plot, the limits of the X and Y axis are expanding a given percentage from the min and max values for each axis. This controls such percentage.
 label **logical** | Whether to plot the cluster labels in the UMAP. The cluster labels will have the same color as the cluster colors.
 label.color **character** | Color of the labels in the plot.
 label.size **numeric** | Size of the labels in the plot.
 number.breaks **numeric** | Controls the number of breaks in continuous color scales of ggplot2-based plots.
 diverging.palette
 character | Type of symmetrical color palette to use. Out of the diverging palettes defined in [brewer.pal](#).
 diverging.direction
 numeric | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.
 sequential.palette
 character | Type of sequential color palette to use. Out of the sequential palettes defined in [brewer.pal](#).
 sequential.direction
 numeric | Direction of the sequential color scale. Either 1 or -1.
 plot.title.face, plot.subtitle.face, plot.caption.face,
 axis.title.face, axis.text.face, legend.title.face, legend.text.face
 character | Controls the style of the font for the corresponding theme element.
 One of:
 • plain: For normal text.

- **italic**: For text in italic.
- **bold**: For text in bold.
- **bold.italic**: For text both in italic and bold.

Value

A ggplot2 object containing a Feature Plot.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_FeaturePlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Regular FeaturePlot.
  p <- SCpubr:::do_FeaturePlot(sample = sample,
                                 features = "nCount_RNA")

  # FeaturePlot with a subset of identities
  # (in Seurat::Idents(sample)) maintaining the original UMAP shape.
  idents.use <- levels(sample)[!(levels(sample) %in% c("2", "5", "8"))]
  p <- SCpubr:::do_FeaturePlot(sample = sample,
                                 idents.highlight = idents.use,
                                 features = c("EPC1"))

  # Splitting the FeaturePlot by a variable and
  # maintaining the color scale and the UMAP shape.
  p <- SCpubr:::do_FeaturePlot(sample = sample,
                                 features = "EPC1",
                                 split.by = "seurat_clusters")

} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

Description

A Geyser plot is a custom plot in which we plot continuous values on the Y axis grouped by a categorical value in the X. This is plotted as a dot plot, jittered so that the dots span all the way

to the other groups. On top of this, the mean and .66 and .95 of the data is plotted, depicting the overall distribution of the dots. The cells can, then, be colored by a continuous variable (same as Y axis or different) or a categorical one (same as X axis or different).

Usage

```
do_GeyserPlot(  
  sample,  
  features,  
  assay = NULL,  
  slot = "data",  
  group.by = NULL,  
  split.by = NULL,  
  enforce_symmetry = FALSE,  
  scale_type = "continuous",  
  order = TRUE,  
  plot_cell_borders = TRUE,  
  jitter = 0.45,  
  pt.size = 1,  
  border.size = 2,  
  border.color = "black",  
  legend.position = "bottom",  
  legend.width = 1,  
  legend.length = 20,  
  legend.framewidth = 0.5,  
  legend.tickwidth = 0.5,  
  legend.framecolor = "grey50",  
  legend.tickcolor = "white",  
  legend.type = "colorbar",  
  font.size = 14,  
  font.type = "sans",  
  axis.text.x.angle = 45,  
  viridis.palette = "G",  
  viridis.direction = 1,  
  colors.use = NULL,  
  na.value = "grey75",  
  legend.ncol = NULL,  
  legend.nrow = NULL,  
  legend.icon.size = 4,  
  legend.byrow = FALSE,  
  legend.title = NULL,  
  plot.title = NULL,  
  plot.subtitle = NULL,  
  plot.caption = NULL,  
  xlab = "Groups",  
  ylab = feature,  
  flip = FALSE,  
  min.cutoff = rep(NA, length(features)),  
  max.cutoff = rep(NA, length(features)),
```

```

number.breaks = 5,
diverging.palette = "RdBu",
diverging.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = -1,
use_viridis = TRUE,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
features	<code>character</code> Features to represent.
assay	<code>character</code> Assay to use. Defaults to the current assay.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character of factor column.
split.by	<code>character</code> Secondary metadata variable to further group (split) the output by. Has to be a character of factor column.
enforce_symmetry	<code>logical</code> Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.
scale_type	<code>character</code> Type of color scale to use. One of: <ul style="list-style-type: none"> • <code>categorical</code>: Use a categorical color scale based on the values of "group.by". • <code>continuous</code>: Use a continuous color scale based on the values of "feature".
order	<code>logical</code> Whether to order the groups by the median of the data (highest to lowest).
plot_cell_borders	<code>logical</code> Whether to plot border around cells.
jitter	<code>numeric</code> Amount of jitter in the plot along the X axis. The lower the value, the more compacted the dots are.
pt.size	<code>numeric</code> Size of the dots.
border.size	<code>numeric</code> Width of the border of the cells.
border.color	<code>character</code> Color for the border of the heatmap body.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • <code>top</code>: Top of the figure.

- **bottom**: Bottom of the figure.
 - **left**: Left of the figure.
 - **right**: Right of the figure.
 - **none**: No legend is displayed.
- legend.length, legend.width**
`numeric` | Length and width of the legend. Will adjust automatically depending on legend side.
- legend.framewidth, legend.tickwidth**
`numeric` | Width of the lines of the box in the legend.
- legend.framecolor**
`character` | Color of the lines of the box in the legend.
- legend.tickcolor**
`character` | Color of the ticks of the box in the legend.
- legend.type**
`character` | Type of legend to display. One of:
 - **normal**: Default legend displayed by `ggplot2`.
 - **colorbar**: Redefined colorbar legend, using `guide_colorbar`.
- font.size**
`numeric` | Overall font size of the plot. All plot elements will have a size relationship with this font size.
- font.type**
`character` | Base font family for the plot. One of:
 - **mono**: Mono spaced font.
 - **serif**: Serif font family.
 - **sans**: Default font family.
- axis.text.x.angle**
`numeric` | Degree to rotate the X labels. One of: 0, 45, 90.
- viridis.palette**
`character` | A capital letter from A to H or the scale name as in `scale_fill_viridis`.
- viridis.direction**
`numeric` | Either 1 or -1. Controls how the gradient of viridis scale is formed.
- colors.use**
`character` | Named vector of colors to use. Has to match the unique values of group.by when scale_type is set to categorical.
- na.value**
`character` | Color value for NA.
- legend.ncol**
`numeric` | Number of columns in the legend.
- legend.nrow**
`numeric` | Number of rows in the legend.
- legend.icon.size**
`numeric` | Size of the icons in legend.
- legend.byrow**
`logical` | Whether the legend is filled by row or not.
- legend.title**
`character` | Title for the legend.
- plot.title, plot.subtitle, plot.caption**
`character` | Title, subtitle or caption to use in the plot.
- xlab, ylab**
`character` | Titles for the X and Y axis.
- flip**
`logical` | Whether to invert the axis of the displayed plot.

```

min.cutoff, max.cutoff
  numeric | Set the min/max ends of the color scale. Any cell/group with a value
  lower than min.cutoff will turn into min.cutoff and any cell with a value higher
  than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many
  values as features. Use NAs to skip a feature.

number.breaks  numeric | Controls the number of breaks in continuous color scales of ggplot2-
  based plots.

diverging.palette
  character | Type of symmetrical color palette to use. Out of the diverging
  palettes defined in brewer.pal.

diverging.direction
  numeric | Either 1 or -1. Direction of the diverging palette. This basically flips
  the two ends.

sequential.palette
  character | Type of sequential color palette to use. Out of the sequential
  palettes defined in brewer.pal.

sequential.direction
  numeric | Direction of the sequential color scale. Either 1 or -1.

use_viridis    logical | Whether to use viridis color scales.

plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
  character | Controls the style of the font for the corresponding theme element.
  One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.

```

Details

Special thanks to Christina Blume for coming up with the name of the plot.

Value

Either a plot or a list of plots, depending on the number of features provided.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_GeyserPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Geyser plot with categorical color scale.

```

```

p <- SCpubr::do_GeyserPlot(sample = sample,
                            features = "nCount_RNA",
                            scale_type = "categorical")
p

# Geyser plot with continuous color scale.
p <- SCpubr::do_GeyserPlot(sample = sample,
                            features = "nCount_RNA",
                            scale_type = "continuous")

p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`.")
}

```

do_GroupwiseDEPlot *Compute a dotplot with the results of a group-wise DE analysis.*

Description

Compute a dotplot with the results of a group-wise DE analysis.

Usage

```

do_GroupwiseDEPlot(
  sample,
  de_genes,
  group.by = NULL,
  assay = NULL,
  slot = "data",
  number.breaks = 5,
  dot.scale = 8,
  top_genes = 5,
  p.cutoff = 0.05,
  flip = FALSE,
  plot.title = NULL,
  plot.subtitle = NULL,
  plot.caption = NULL,
  xlab = NULL,
  ylab = NULL,
  use_viridis = FALSE,
  colors.use = NULL,
  viridis.direction = -1,
  viridis.palette = "G",

```

```

sequential.direction = 1,
sequential.palette = "YlGnBu",
diverging.palette = "RdBu",
diverging.direction = -1,
legend.position = "bottom",
legend.title = NULL,
legend.width = 1,
legend.length = 7.5,
legend.framewidth = 0.5,
legend.tickwidth = 0.5,
legend.framecolor = "grey50",
legend.tickcolor = "white",
legend.ncol = NULL,
legend.nrow = NULL,
legend.byrow = FALSE,
legend.type = "colorbar",
font.size = 14,
font.type = "sans",
axis.text.x.angle = 45,
min.cutoff = NA,
max.cutoff = NA,
enforce_symmetry = FALSE,
na.value = "grey75",
border.color = "black",
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
de_genes	<code>tibble</code> DE genes matrix resulting of running <code>Seurat::FindAllMarkers()</code> .
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
assay	<code>character</code> Assay to use. Defaults to the current assay.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
number.breaks	<code>numeric</code> Controls the number of breaks in continuous color scales of ggplot2-based plots.
dot.scale	<code>numeric</code> Scale the size of the dots.
top_genes	<code>numeric</code> Top N differentially expressed (DE) genes by group to retrieve.

p.cutoff **numeric** | Cutoff to use for adjusted p.value to filter significant genes.

flip **logical** | Whether to invert the axis of the displayed plot.

plot.title, plot.subtitle, plot.caption
 character | Title, subtitle or caption to use in the plot.

xlab, ylab **character** | Titles for the X and Y axis.

use_viridis **logical** | Whether to use viridis color scales.

colors.use **named_vector** | Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of group.by. If group.by is not provided, defaults to the unique values of **Idents**. If not provided, a color scale will be set by default.

viridis.direction
 numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.

viridis.palette
 character | A capital letter from A to H or the scale name as in **scale_fill_viridis**.

sequential.direction
 numeric | Direction of the sequential color scale. Either 1 or -1.

sequential.palette
 character | Type of sequential color palette to use. Out of the sequential palettes defined in **brewer.pal**.

diverging.palette
 character | Type of symmetrical color palette to use. Out of the diverging palettes defined in **brewer.pal**.

diverging.direction
 numeric | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.

legend.position
 character | Position of the legend in the plot. One of:

- top: Top of the figure.
- bottom: Bottom of the figure.
- left: Left of the figure.
- right: Right of the figure.
- none: No legend is displayed.

legend.title **character** | Title for the legend.

legend.length, legend.width
 numeric | Length and width of the legend. Will adjust automatically depending on legend side.

legend.framewidth, legend.tickwidth
 numeric | Width of the lines of the box in the legend.

legend.framecolor
 character | Color of the lines of the box in the legend.

legend.tickcolor
 character | Color of the ticks of the box in the legend.

legend.ncol **numeric** | Number of columns in the legend.

legend.nrow **numeric** | Number of rows in the legend.
 legend.byrow **logical** | Whether the legend is filled by row or not.
 legend.type **character** | Type of legend to display. One of:
 • normal: Default legend displayed by **ggplot2**.
 • colorbar: Redefined colorbar legend, using **guide_colorbar**.
 font.size **numeric** | Overall font size of the plot. All plot elements will have a size relationship with this font size.
 font.type **character** | Base font family for the plot. One of:
 • mono: Mono spaced font.
 • serif: Serif font family.
 • sans: Default font family.
 axis.text.x.angle
 numeric | Degree to rotate the X labels. One of: 0, 45, 90.
 min.cutoff, max.cutoff
 numeric | Set the min/max ends of the color scale. Any cell/group with a value lower than min.cutoff will turn into min.cutoff and any cell with a value higher than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
 enforce_symmetry
 logical | Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.
 na.value **character** | Color value for NA.
 border.color **character** | Color for the border of the heatmap body.
 plot.title.face, plot.subtitle.face, plot.caption.face,
 axis.title.face, axis.text.face, legend.title.face, legend.text.face
 character | Controls the style of the font for the corresponding theme element. One of:
 • plain: For normal text.
 • italic: For text in italic.
 • bold: For text in bold.
 • bold.italic: For text both in italic and bold.

Value

A dotplot composed of 3 main panels: -log10(adjusted p-value), log2(FC) and mean expression by cluster.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_GroupwiseDEPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
}
  
```

```

# Define your Seurat object.
sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

# Compute DE genes and transform to a tibble.
de_genes <- readRDS(system.file("extdata/de_genes_example.rds", package = "SCpubr"))

# Default output.
p <- SCpubr::do_GroupwiseDEPlot(sample = sample,
                                   de_genes = de_genes)

p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_LigandReceptorPlot *Visualize Ligand-Receptor analysis output.*

Description

This function makes use of **liana** package to run Ligand-Receptor analysis. Takes the output of liana and generates a dot-plot visualization according to the user's specifications.

Usage

```

do_LigandReceptorPlot(
  liana_output,
  split.by = NULL,
  keep_source = NULL,
  keep_target = NULL,
  top_interactions = 25,
  top_interactions_by_group = FALSE,
  dot_border = TRUE,
  magnitude = "sca.LRscore",
  specificity = "aggregate_rank",
  sort.by = "E",
  sorting.type.specificity = "descending",
  sorting.type.magnitude = "descending",
  border.color = "black",
  axis.text.x.angle = 45,
  legend.position = "bottom",
  legend.type = "colorbar",
  legend.length = 20,
  legend.width = 1,
  legend.framecolor = "grey50",

```

```

legend.tickcolor = "white",
legend.framewidth = 0.5,
legend.tickwidth = 0.5,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = 1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
font.size = 14,
dot.size = 1,
font.type = "sans",
plot.grid = TRUE,
grid.color = "grey90",
grid.type = "dotted",
compute_ChordDiagrams = FALSE,
sort_interactions_alphabetically = FALSE,
number.breaks = 5,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain",
return_interactions = FALSE,
invert_specificity = TRUE,
invert_magnitude = FALSE,
verbose = TRUE
)

```

Arguments

<code>liana_output</code>	<code>tibble</code> Object resulting from running <code>liana_wrap</code> and <code>liana_aggregate</code> .
<code>split.by</code>	<code>character</code> Whether to further facet the plot on the y axis by common ligand.complex or receptor.complex. Values to provide: NULL, ligand.complex, receptor.complex.
<code>keep_source, keep_target</code>	<code>character</code> Identities to keep for the source/target of the interactions. NULL otherwise.
<code>top_interactions</code>	<code>numeric</code> Number of unique interactions to retrieve ordered by magnitude and specificity. It does not necessarily mean that the output will contain as many, but rather an approximate value.
<code>top_interactions_by_group</code>	<code>logical</code> Enforce the value on <code>top_interactions</code> to be applied to each group in source column.
<code>dot_border</code>	<code>logical</code> Whether to draw a black border in the dots.

specificity, magnitude
`character` | Which columns to use for specificity and magnitude.

sort.by
`character` | How to arrange the top interactions. Interactions are sorted and then the top N are retrieved and displayed. This takes place after subsetting for `keep_source` and `keep_target`. One of:

- A: Sorts by specificity.
- B: Sorts by magnitude.
- C: Sorts by specificity, then magnitude (gives extra weight to specificity).
- D: Sorts by magnitude, then specificity (gives extra weight to magnitude). Might lead to the display of non-significant results.
- E: Sorts by specificity and magnitude equally.

sorting.type.specificity, sorting.type.magnitude
`character` | Whether the sorting of e magnitude or specificity columns is done in ascending or descending order. This synergises with the value of e `invert_specificity` and e `invert_magnitude` parameters.

border.color `character` | Color for the border of the heatmap body.

axis.text.x.angle
`numeric` | Degree to rotate the X labels. One of: 0, 45, 90.

legend.position
`character` | Position of the legend in the plot. One of:

- top: Top of the figure.
- bottom: Bottom of the figure.
- left: Left of the figure.
- right: Right of the figure.
- none: No legend is displayed.

legend.type `character` | Type of legend to display. One of:

- normal: Default legend displayed by `ggplot2`.
- colorbar: Redefined colorbar legend, using `guide_colorbar`.

legend.length, legend.width
`numeric` | Length and width of the legend. Will adjust automatically depending on legend side.

legend.framecolor
`character` | Color of the lines of the box in the legend.

legend.tickcolor
`character` | Color of the ticks of the box in the legend.

legend.framewidth, legend.tickwidth
`numeric` | Width of the lines of the box in the legend.

use_viridis `logical` | Whether to use viridis color scales.

viridis.palette
`character` | A capital letter from A to H or the scale name as in `scale_fill_viridis`.

viridis.direction
`numeric` | Either 1 or -1. Controls how the gradient of viridis scale is formed.

```

sequential.palette
  character | Type of sequential color palette to use. Out of the sequential
  palettes defined in brewer.pal.
sequential.direction
  numeric | Direction of the sequential color scale. Either 1 or -1.
font.size
  numeric | Overall font size of the plot. All plot elements will have a size relationship with this font size.
dot.size
  numeric | Size aesthetic for the dots.
font.type
  character | Base font family for the plot. One of:
    • mono: Mono spaced font.
    • serif: Serif font family.
    • sans: Default font family.
plot.grid
  logical | Whether to plot grid lines.
grid.color
  character | Color of the grid in the plot. In heatmaps, color of the border of the
  cells.
grid.type
  character | One of the possible linetype options:
    • blank.
    • solid.
    • dashed.
    • dotted.
    • dotdash.
    • longdash.
    • twodash.
compute_ChordDiagrams
  logical | Whether to also compute Chord Diagrams for both the number of
  interactions between source and target but also between ligand.complex and re-
  ceptor.complex.
sort_interactions_alphabetically
  logical | Sort the interactions to be plotted alphabetically (TRUE) or keep them
  in their original order in the matrix (FALSE).
number.breaks
  numeric | Controls the number of breaks in continuous color scales of ggplot2-
  based plots.
plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
  character | Controls the style of the font for the corresponding theme element.
  One of:
    • plain: For normal text.
    • italic: For text in italic.
    • bold: For text in bold.
    • bold.italic: For text both in italic and bold.
return_interactions
  logical | Whether to return the data.frames with the interactions so that they
  can be plotted as chord plots using other package functions.
invert_specificity, invert_magnitude
  logical | Whether to  $-\log_{10}$  transform specificity and magnitude columns.
verbose
  logical | Whether to show extra comments, warnings,etc.

```

Value

A ggplot2 plot with the results of the Ligand-Receptor analysis.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_LigandReceptorPlot", passive = TRUE)

if (isTRUE(value)){
  liana_output <- readRDS(system.file("extdata/liana_output_example.rds", package = "SCpubr"))
  # Ligand Receptor analysis plot.
  p <- SCpubr::do_LigandReceptorPlot(liana_output = liana_output)
  p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`.")
}
```

do_LoadingsPlot

Compute a heatmap summary of the top and bottom genes in the PCA loadings for the desired PCs in a Seurat object.

Description

Compute a heatmap summary of the top and bottom genes in the PCA loadings for the desired PCs in a Seurat object.

Usage

```
do_LoadingsPlot(
  sample,
  group.by = NULL,
  subsample = NA,
  dims = 1:10,
  top_loadings = 5,
  assay = "SCT",
  slot = "data",
  grid.color = "white",
  border.color = "black",
  number.breaks = 5,
  na.value = "grey75",
  legend.position = "bottom",
  legend.title = "Expression",
  legend.type = "colorbar",
  legend.framewidth = 0.5,
```

```

legend.tickwidth = 0.5,
legend.length = 20,
legend.width = 1,
legend.framecolor = "grey50",
legend.tickcolor = "white",
font.size = 14,
font.type = "sans",
axis.text.x.angle = 45,
use_viridis = FALSE,
sequential.direction = 1,
sequential.palette = "YlGnBu",
viridis.palette = "G",
viridis.direction = -1,
diverging.palette = "RdBu",
diverging.direction = -1,
flip = FALSE,
min.cutoff.loadings = NA,
max.cutoff.loadings = NA,
min.cutoff.expression = NA,
max.cutoff.expression = NA,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

<code>sample</code>	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
<code>group.by</code>	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
<code>subsample</code>	<code>numeric</code> Number of cells to subsample the Seurat object to increase computational speed. Use NA to include the Seurat object as is.
<code>dims</code>	<code>numeric</code> PCs to include in the analysis.
<code>top_loadings</code>	<code>numeric</code> Number of top and bottom scored genes in the PCA Loadings for each PC.
<code>assay</code>	<code>character</code> Assay to use. Defaults to the current assay.
<code>slot</code>	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
<code>grid.color</code>	<code>character</code> Color of the grid in the plot. In heatmaps, color of the border of the cells.
<code>border.color</code>	<code>character</code> Color for the border of the heatmap body.
<code>number.breaks</code>	<code>numeric</code> Controls the number of breaks in continuous color scales of ggplot2-based plots.

na.value **character** | Color value for NA.
 legend.position
 character | Position of the legend in the plot. One of:
 • top: Top of the figure.
 • bottom: Bottom of the figure.
 • left: Left of the figure.
 • right: Right of the figure.
 • none: No legend is displayed.
 legend.title **character** | Title for the legend.
 legend.type **character** | Type of legend to display. One of:
 • normal: Default legend displayed by **ggplot2**.
 • colorbar: Redefined colorbar legend, using **guide_colorbar**.
 legend.framewidth, legend.tickwidth
 numeric | Width of the lines of the box in the legend.
 legend.length, legend.width
 numeric | Length and width of the legend. Will adjust automatically depending on legend side.
 legend.framecolor
 character | Color of the lines of the box in the legend.
 legend.tickcolor
 character | Color of the ticks of the box in the legend.
 font.size **numeric** | Overall font size of the plot. All plot elements will have a size relationship with this font size.
 font.type **character** | Base font family for the plot. One of:
 • mono: Mono spaced font.
 • serif: Serif font family.
 • sans: Default font family.
 axis.text.x.angle
 numeric | Degree to rotate the X labels. One of: 0, 45, 90.
 use_viridis **logical** | Whether to use viridis color scales.
 sequential.direction
 numeric | Direction of the sequential color scale. Either 1 or -1.
 sequential.palette
 character | Type of sequential color palette to use. Out of the sequential palettes defined in **brewer.pal**.
 viridis.palette
 character | A capital letter from A to H or the scale name as in **scale_fill_viridis**.
 viridis.direction
 numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.
 diverging.palette
 character | Type of symmetrical color palette to use. Out of the diverging palettes defined in **brewer.pal**.

```

diverging.direction
  numeric | Either 1 or -1. Direction of the diverging palette. This basically flips
  the two ends.

flip      logical | Whether to invert the axis of the displayed plot.

min.cutoff.loadings, max.cutoff.loadings
  numeric | Cutoff to subset the scale of the Loading score heatmap. NA will use
  quantiles 0.05 and 0.95.

min.cutoff.expression, max.cutoff.expression
  numeric | Cutoff to subset the scale of the expression heatmap. NA will use 0
  (no quantile) and quantile 0.95.

plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
  character | Controls the style of the font for the corresponding theme element.
  One of:
    • plain: For normal text.
    • italic: For text in italic.
    • bold: For text in bold.
    • bold.italic: For text both in italic and bold.

```

Value

A ggplot2 object.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_LoadingsPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  p <- SCpubr::do_LoadingsPlot(sample = sample,
                                dims = 1:2)
  p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_MetadataPlot	<i>Compute a heatmap of categorical variables.</i>
-----------------	--

Description

The main use of this function is to generate a metadata heatmap of your categorical data, normally targeted to the different patient samples one has in the Seurat object. It requires that the metadata columns chosen have one and only one possible value for each of the values in group.by.

Usage

```
do_MetadataPlot(  
  sample = NULL,  
  group.by = NULL,  
  metadata = NULL,  
  from_df = FALSE,  
  df = NULL,  
  colors.use = NULL,  
  cluster = TRUE,  
  flip = TRUE,  
  heatmap.gap = 1,  
  axis.text.x.angle = 45,  
  legend.position = "bottom",  
  font.size = 14,  
  legend.font.size = NULL,  
  legend.symbol.size = NULL,  
  legend.ncol = NULL,  
  legend.nrow = NULL,  
  legend.byrow = FALSE,  
  na.value = "grey75",  
  font.type = "sans",  
  grid.color = "white",  
  border.color = "black",  
  plot.title.face = "bold",  
  plot.subtitle.face = "plain",  
  plot.caption.face = "italic",  
  axis.title.face = "bold",  
  axis.text.face = "plain",  
  legend.title.face = "bold",  
  legend.text.face = "plain",  
  xlab = "",  
  ylab = "")
```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
--------	---

group.by	<code>character</code> Metadata column to use as basis for the plot.
metadata	<code>character</code> Metadata columns that will be used to plot the heatmap on the basis of the variable provided to group.by.
from_df	<code>logical</code> Whether to provide a data frame with the metadata instead.
df	<code>data.frame</code> Data frame containing the metadata to plot. Rows contain the unique values common to all columns (metadata variables). The columns must be named.
colors.use	<code>named_list</code> A named list of named vectors. The names of the list correspond to the names of the values provided to metadata and the names of the items in the named vectors correspond to the unique values of that specific metadata variable. The values are the desired colors in HEX code for the values to plot. The used are pre-defined by the package but, in order to get the most out of the plot, please provide your custom set of colors for each metadata column!
cluster	<code>logical</code> Whether to perform clustering of rows and columns.
flip	<code>logical</code> Whether to invert the axis of the displayed plot.
heatmap.gap	<code>numeric</code> Size of the gap between heatmaps in mm.
axis.text.x.angle	<code>numeric</code> Degree to rotate the X labels. One of: 0, 45, 90.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none">• <code>top</code>: Top of the figure.• <code>bottom</code>: Bottom of the figure.• <code>left</code>: Left of the figure.• <code>right</code>: Right of the figure.• <code>none</code>: No legend is displayed.
font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
legend.font.size	<code>numeric</code> Size of the font size of the legend. NULL uses default theme font size for legend according to the <code>strongfont.size</code> parameter.
legend.symbol.size	<code>numeric</code> Size of symbols in the legend in mm. NULL uses the default size.
legend.ncol	<code>numeric</code> Number of columns in the legend.
legend.nrow	<code>numeric</code> Number of rows in the legend.
legend.byrow	<code>logical</code> Whether the legend is filled by row or not.
na.value	<code>character</code> Color value for NA.
font.type	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none">• <code>mono</code>: Mono spaced font.• <code>serif</code>: Serif font family.• <code>sans</code>: Default font family.
grid.color	<code>character</code> Color of the grid in the plot. In heatmaps, color of the border of the cells.

```

border.color      character | Color for the border of the heatmap body.
plot.title.face,       plot.subtitle.face,       plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
                           character | Controls the style of the font for the corresponding theme element.
                           One of:
                           • plain: For normal text.
                           • italic: For text in italic.
                           • bold: For text in bold.
                           • bold.italic: For text both in italic and bold.

xlab, ylab      character | Titles for the X and Y axis.

```

Value

A ggplot2 object.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_MetadataPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Can also use a Seurat object.
  df <- data.frame(row.names = letters[1:5],
                    "A" = as.character(seq(1, 5)),
                    "B" = rev(as.character(seq(1, 5))))}

p <- SCpubr:::do_MetadataPlot(from_df = TRUE,
                                df = df)

p

} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_NebulosaPlot	<i>Wrapper for Nebulosa::plot_density in Seurat.</i>
-----------------	--

Description

Wrapper for Nebulosa::plot_density in Seurat.

Usage

```
do_NebulosaPlot(  
  sample,  
  features,  
  slot = NULL,  
  dims = c(1, 2),  
  pt.size = 1,  
  reduction = NULL,  
  combine = TRUE,  
  method = c("ks", "wkde"),  
  joint = FALSE,  
  return_only_joint = FALSE,  
  plot.title = NULL,  
  plot.subtitle = NULL,  
  plot.caption = NULL,  
  legend.type = "colorbar",  
  legend.framewidth = 0.5,  
  legend.tickwidth = 0.5,  
  legend.length = 20,  
  legend.width = 1,  
  legend.framecolor = "grey50",  
  legend.tickcolor = "white",  
  font.size = 14,  
  font.type = "sans",  
  legend.position = "bottom",  
  plot_cell_borders = TRUE,  
  border.size = 2,  
  border.color = "black",  
  viridis.palette = "G",  
  viridis.direction = 1,  
  verbose = TRUE,  
  na.value = "grey75",  
  plot.axes = FALSE,  
  number.breaks = 5,  
  use_viridis = FALSE,  
  sequential.palette = "YlGnBu",  
  sequential.direction = 1,  
  plot.title.face = "bold",  
  plot.subtitle.face = "plain",  
  plot.caption.face = "italic",  
  axis.title.face = "bold",  
  axis.text.face = "plain",  
  legend.title.face = "bold",  
  legend.text.face = "plain"  
)
```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
features	<code>character</code> Features to represent.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
dims	<code>numeric</code> Vector of 2 numerics indicating the dimensions to plot out of the selected reduction. Defaults to c(1, 2) if not specified.
pt.size	<code>numeric</code> Size of the dots.
reduction	<code>character</code> Reduction to use. Can be the canonical ones such as "umap", "pca", or any custom ones, such as "diffusion". If you are unsure about which reductions you have, use <code>Seurat::Reductions(sample)</code> . Defaults to "umap" if present or to the last computed reduction if the argument is not provided.
combine	<code>logical</code> Whether to create a single plot out of multiple features.
method	Kernel density estimation method: <ul style="list-style-type: none"> • <code>ks</code>: Computes density using the <code>kde</code> function from the <code>ks</code> package. • <code>wkde</code>: Computes density using a modified version of the <code>kde2d</code> function from the <code>MASS</code> package to allow weights. Bandwidth selection from the <code>ks</code> package is used instead.
joint	<code>logical</code> Whether to plot different features as joint density.
return_only_joint	<code>logical</code> Whether to only return the joint density panel.
plot.title, plot.subtitle, plot.caption	<code>character</code> Title, subtitle or caption to use in the plot.
legend.type	<code>character</code> Type of legend to display. One of: <ul style="list-style-type: none"> • <code>normal</code>: Default legend displayed by <code>ggplot2</code>. • <code>colorbar</code>: Redefined colorbar legend, using guide_colorbar.
legend.framewidth, legend.tickwidth	<code>numeric</code> Width of the lines of the box in the legend.
legend.length, legend.width	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.
legend.framecolor	<code>character</code> Color of the lines of the box in the legend.
legend.tickcolor	<code>character</code> Color of the ticks of the box in the legend.
font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • <code>mono</code>: Mono spaced font. • <code>serif</code>: Serif font family. • <code>sans</code>: Default font family.

```

legend.position
  character | Position of the legend in the plot. One of:
  • top: Top of the figure.
  • bottom: Bottom of the figure.
  • left: Left of the figure.
  • right: Right of the figure.
  • none: No legend is displayed.

plot_cell_borders
  logical | Whether to plot border around cells.

border.size
  numeric | Width of the border of the cells.

border.color
  character | Color for the border of the heatmap body.

viridis.palette
  character | A capital letter from A to H or the scale name as in scale\_fill\_viridis.

viridis.direction
  numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.

verbose
  logical | Whether to show extra comments, warnings,etc.

na.value
  character | Color value for NA.

plot.axes
  logical | Whether to plot axes or not.

number.breaks
  numeric | Controls the number of breaks in continuous color scales of ggplot2-based plots.

use_viridis
  logical | Whether to use viridis color scales.

sequential.palette
  character | Type of sequential color palette to use. Out of the sequential palettes defined in brewer.pal.

sequential.direction
  numeric | Direction of the sequential color scale. Either 1 or -1.

plot.title.face, plot.subtitle.face, plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
  character | Controls the style of the font for the corresponding theme element. One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.

```

Value

A ggplot2 object containing a Nebulosa plot.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_NebulosaPlot", passive = TRUE)

```

```

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Basic Nebulosa plot.
  p <- SCpubr::do_NebulosaPlot(sample = sample,
                                 features = "EPC1")

  # Compute joint density.
  p <- SCpubr::do_NebulosaPlot(sample = sample,
                                 features = c("EPC1", "TOX2"),
                                 joint = TRUE)

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_PathwayActivityPlot

Plot Pathway Activities from decoupleR using Progeny prior knowledge.

Description

Plot Pathway Activities from decoupleR using Progeny prior knowledge.

Usage

```

do_PathwayActivityPlot(
  sample,
  activities,
  group.by = NULL,
  split.by = NULL,
  slot = "scale.data",
  statistic = "norm_wmean",
  pt.size = 1,
  border.size = 2,
  na.value = "grey75",
  legend.position = "bottom",
  legend.width = 1,
  legend.length = 20,
  legend.framewidth = 0.5,
  legend.tickwidth = 0.5,
  legend.framecolor = "grey50",

```

```

legend.tickcolor = "white",
legend.type = "colorbar",
font.size = 14,
font.type = "sans",
axis.text.x.angle = 45,
enforce_symmetry = TRUE,
min.cutoff = NA,
max.cutoff = NA,
number.breaks = 5,
diverging.palette = "RdBu",
diverging.direction = -1,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
flip = FALSE,
return_object = FALSE,
grid.color = "white",
border.color = "black",
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

<code>sample</code>	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
<code>activities</code>	<code>tibble</code> Result of running decoupleR method with progeny regulon prior knowledge.
<code>group.by</code>	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
<code>split.by</code>	<code>character</code> Secondary metadata variable to further group (split) the output by. Has to be a character of factor column.
<code>slot</code>	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
<code>statistic</code>	<code>character</code> DecoupleR statistic to use. One of: <ul style="list-style-type: none"> • <code>wmean</code>: For weighted mean. • <code>norm_wmean</code>: For normalized weighted mean. • <code>corr_wmean</code>: For corrected weighted mean.
<code>pt.size</code>	<code>numeric</code> Size of the dots.
<code>border.size</code>	<code>numeric</code> Width of the border of the cells.

na.value **character** | Color value for NA.

legend.position
 character | Position of the legend in the plot. One of:
 • top: Top of the figure.
 • bottom: Bottom of the figure.
 • left: Left of the figure.
 • right: Right of the figure.
 • none: No legend is displayed.

legend.length, legend.width
 numeric | Length and width of the legend. Will adjust automatically depending on legend side.

legend.framewidth, legend.tickwidth
 numeric | Width of the lines of the box in the legend.

legend.framecolor
 character | Color of the lines of the box in the legend.

legend.tickcolor
 character | Color of the ticks of the box in the legend.

legend.type **character** | Type of legend to display. One of:
 • normal: Default legend displayed by **ggplot2**.
 • colorbar: Redefined colorbar legend, using **guide_colorbar**.

font.size **numeric** | Overall font size of the plot. All plot elements will have a size relationship with this font size.

font.type **character** | Base font family for the plot. One of:
 • mono: Mono spaced font.
 • serif: Serif font family.
 • sans: Default font family.

axis.text.x.angle
 numeric | Degree to rotate the X labels. One of: 0, 45, 90.

enforce_symmetry
 logical | Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.

min.cutoff, max.cutoff
 numeric | Set the min/max ends of the color scale. Any cell/group with a value lower than min.cutoff will turn into min.cutoff and any cell with a value higher than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many values as features. Use NAs to skip a feature.

number.breaks **numeric** | Controls the number of breaks in continuous color scales of ggplot2-based plots.

diverging.palette
 character | Type of symmetrical color palette to use. Out of the diverging palettes defined in **brewer.pal**.

diverging.direction
 numeric | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.

```

use_viridis      logical | Whether to use viridis color scales.
viridis.palette
                  character | A capital letter from A to H or the scale name as in scale_fill_viridis.
viridis.direction
                  numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.
sequential.palette
                  character | Type of sequential color palette to use. Out of the sequential
                           palettes defined in brewer.pal.
sequential.direction
                  numeric | Direction of the sequential color scale. Either 1 or -1.
flip              logical | Whether to invert the axis of the displayed plot.
return_object     logical | Returns the Seurat object with the modifications performed in the
                           function. Normally, this contains a new assay with the data that can then be used
                           for any other visualization desired.
grid.color        character | Color of the grid in the plot. In heatmaps, color of the border of the
                           cells.
border.color      character | Color for the border of the heatmap body.
plot.title.face,   plot.subtitle.face,   plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
                  character | Controls the style of the font for the corresponding theme element.
                           One of:
                           • plain: For normal text.
                           • italic: For text in italic.
                           • bold: For text in bold.
                           • bold.italic: For text both in italic and bold.

```

Value

A ggplot2 object.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_PathwayActivityPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/

  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds",
                                package = "SCpubr"))

  # Define your activities object.
  progeny_activities <- readRDS(system.file("extdata/progeny_activities_example.rds",
                                             package = "SCpubr"))

  # General heatmap.

```

```

out <- SCpubr::do_PathwayActivityPlot(sample = sample,
                                         activities = progeny_activities)
p <- out$heatmaps$average_scores
p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_RankedEnrichmentPlot

Compute a heatmap of enrichment of gene sets on the context of a diffusion component.

Description

Compute a heatmap of enrichment of gene sets on the context of a diffusion component.

Usage

```

do_RankedEnrichmentPlot(
  sample,
  input_gene_list,
  assay = NULL,
  slot = NULL,
  scale.enrichment = TRUE,
  dims = 1:2,
  subsample = 2500,
  reduction = NULL,
  group.by = NULL,
  colors.use = NULL,
  raster = FALSE,
  interpolate = FALSE,
  nbin = 24,
  ctrl = 100,
  flavor = "Seurat",
  main.heatmap.size = 0.95,
  enforce_symmetry = ifelse(isTRUE(scale.enrichment), TRUE, FALSE),
  use_viridis = FALSE,
  viridis.palette = "G",
  viridis.direction = -1,
  sequential.palette = "YlGnBu",
  sequential.direction = 1,
  font.size = 14,
  font.type = "sans",
  na.value = "grey75",

```

```

legend.width = 1,
legend.length = 20,
legend.framewidth = 0.5,
legend.tickwidth = 0.5,
legend.framecolor = "grey50",
legend.tickcolor = "white",
legend.type = "colorbar",
legend.position = "bottom",
legend.nrow = NULL,
legend.ncol = NULL,
legend.byrow = FALSE,
number.breaks = 5,
diverging.palette = "RdBu",
diverging.direction = -1,
axis.text.x.angle = 45,
border.color = "black",
return_object = FALSE,
verbose = FALSE,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

<code>sample</code>	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
<code>input_gene_list</code>	<code>named_list</code> Named list of lists of genes to be used as input.
<code>assay</code>	<code>character</code> Assay to use. Defaults to the current assay.
<code>slot</code>	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
<code>scale.enrichment</code>	<code>logical</code> Should the enrichment scores be scaled (z-scored) for better comparison in between gene sets? Setting this to TRUE should make intra- gene set comparisons easier at the cost of not being able to compare inter- gene sets in absolute values.
<code>dims</code>	<code>numeric</code> Vector of 2 numerics indicating the dimensions to plot out of the selected reduction. Defaults to c(1, 2) if not specified.
<code>subsample</code>	<code>numeric</code> Number of cells to subset for the analysis. NA will use all. Cells are selected at random.
<code>reduction</code>	<code>character</code> Reduction to use. Can be the canonical ones such as "umap", "pca", or any custom ones, such as "diffusion". If you are unsure about which reductions you have, use <code>Seurat::Reductions(sample)</code> . Defaults to "umap" if present or to the last computed reduction if the argument is not provided.

group.by	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
colors.use	<code>list</code> A named list of named vectors. The names of the list correspond to the names of the values provided to metadata and the names of the items in the named vectors correspond to the unique values of that specific metadata variable. The values are the desired colors in HEX code for the values to plot. The used are pre-defined by the package but, in order to get the most out of the plot, please provide your custom set of colors for each metadata column!
raster	<code>logical</code> Whether to raster the resulting plot. This is recommendable if plotting a lot of cells.
interpolate	<code>logical</code> Smoothes the output heatmap, saving space on disk when saving the image. However, the image is not as crisp.
nbin	<code>numeric</code> Number of bins to use in <code>AddModuleScore</code> .
ctrl	<code>numeric</code> Number of genes in the control set to use in <code>AddModuleScore</code> .
flavor	<code>character</code> One of: Seurat, UCell. Compute the enrichment scores using <code>AddModuleScore</code> or <code>AddModuleScore_UCell</code> .
main.heatmap.size	<code>numeric</code> A number from 0 to 1 corresponding to how big the main heatmap plot should be with regards to the rest (corresponds to the proportion in size).
enforce_symmetry	<code>logical</code> Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.
use_viridis	<code>logical</code> Whether to use viridis color scales.
viridis.palette	<code>character</code> A capital letter from A to H or the scale name as in <code>scale_fill_viridis</code> .
viridis.direction	<code>numeric</code> Either 1 or -1. Controls how the gradient of viridis scale is formed.
sequential.palette	<code>character</code> Type of sequential color palette to use. Out of the sequential palettes defined in <code>brewer.pal</code> .
sequential.direction	<code>numeric</code> Direction of the sequential color scale. Either 1 or -1.
font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • mono: Mono spaced font. • serif: Serif font family. • sans: Default font family.
na.value	<code>character</code> Color value for NA.
legend.length, legend.width	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.
legend.framewidth, legend.tickwidth	<code>numeric</code> Width of the lines of the box in the legend.

```

legend.framecolor
  character | Color of the lines of the box in the legend.
legend.tickcolor
  character | Color of the ticks of the box in the legend.
legend.type    character | Type of legend to display. One of:
  • normal: Default legend displayed by ggplot2.
  • colorbar: Redefined colorbar legend, using guide\_colorbar.
legend.position
  character | Position of the legend in the plot. One of:
  • top: Top of the figure.
  • bottom: Bottom of the figure.
  • left: Left of the figure.
  • right: Right of the figure.
  • none: No legend is displayed.
legend.nrow     numeric | Number of rows in the legend.
legend.ncol     numeric | Number of columns in the legend.
legend.byrow    logical | Whether the legend is filled by row or not.
number.breaks   numeric | Controls the number of breaks in continuous color scales of ggplot2-based plots.
diverging.palette
  character | Type of symmetrical color palette to use. Out of the diverging palettes defined in brewer.pal.
diverging.direction
  numeric | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.
axis.text.x.angle
  numeric | Degree to rotate the X labels. One of: 0, 45, 90.
border.color    character | Color for the border of the heatmap body.
return_object   logical | Returns the Seurat object with the modifications performed in the function. Normally, this contains a new assay with the data that can then be used for any other visualization desired.
verbose         logical | Whether to show extra comments, warnings,etc.
plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
  character | Controls the style of the font for the corresponding theme element. One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.

```

Value

A list of ggplot2 objects and a Seurat object if desired.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_RankedEnrichmentPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Genes have to be unique.
  genes <- list("A" = rownames(sample)[1:5],
                "B" = rownames(sample)[6:10],
                "C" = rownames(sample)[11:15])

  # This will query, for the provided components, the enrichment of the gene
  # sets for all cells and plot them in the context of the cells reordered by
  # the position alongside each dimensional reduction component.
  p <- SCpubr:::do_RankedEnrichmentPlot(sample = sample,
                                         input_gene_list = genes,
                                         nbin = 1,
                                         ctrl = 5,
                                         flavor = "Seurat",
                                         subsample = NA,
                                         dims = 1:2,
                                         verbose = FALSE)

  p

} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_RankedExpressionPlot

Compute a heatmap of enrichment of gene sets on the context of a diffusion component.

Description

Compute a heatmap of enrichment of gene sets on the context of a diffusion component.

Usage

```
do_RankedExpressionPlot(
  sample,
```

```
features,
assay = NULL,
slot = NULL,
dims = 1:2,
subsample = 2500,
reduction = NULL,
group.by = NULL,
colors.use = NULL,
raster = FALSE,
interpolate = FALSE,
nbin = 24,
ctrl = 100,
main.heatmap.size = 0.95,
enforce_symmetry = TRUE,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
font.size = 14,
font.type = "sans",
na.value = "grey75",
legend.width = 1,
legend.length = 20,
legend.framewidth = 0.5,
legend.tickwidth = 0.5,
legend.framecolor = "grey50",
legend.tickcolor = "white",
legend.type = "colorbar",
legend.position = "bottom",
legend.nrow = NULL,
legend.ncol = NULL,
legend.byrow = FALSE,
number.breaks = 5,
diverging.palette = "RdBu",
diverging.direction = -1,
axis.text.x.angle = 45,
border.color = "black",
return_object = FALSE,
verbose = FALSE,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)
```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
features	<code>character</code> Features to represent.
assay	<code>character</code> Assay to use. Defaults to the current assay.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
dims	<code>numeric</code> Vector of 2 numerics indicating the dimensions to plot out of the selected reduction. Defaults to c(1, 2) if not specified.
subsample	<code>numeric</code> Number of cells to subset for the analysis. NA will use all. Cells are selected at random.
reduction	<code>character</code> Reduction to use. Can be the canonical ones such as "umap", "pca", or any custom ones, such as "diffusion". If you are unsure about which reductions you have, use <code>Seurat::Reductions(sample)</code> . Defaults to "umap" if present or to the last computed reduction if the argument is not provided.
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character of factor column.
colors.use	<code>list</code> A named list of named vectors. The names of the list correspond to the names of the values provided to metadata and the names of the items in the named vectors correspond to the unique values of that specific metadata variable. The values are the desired colors in HEX code for the values to plot. The used are pre-defined by the package but, in order to get the most out of the plot, please provide your custom set of colors for each metadata column!
raster	<code>logical</code> Whether to raster the resulting plot. This is recommendable if plotting a lot of cells.
interpolate	<code>logical</code> Smoothes the output heatmap, saving space on disk when saving the image. However, the image is not as crisp.
nbin	<code>numeric</code> Number of bins to use in AddModuleScore .
ctrl	<code>numeric</code> Number of genes in the control set to use in AddModuleScore .
main.heatmap.size	<code>numeric</code> A number from 0 to 1 corresponding to how big the main heatmap plot should be with regards to the rest (corresponds to the proportion in size).
enforce_symmetry	<code>logical</code> Return a symmetrical plot axes-wise or continuous color scale-wise, when applicable.
use_viridis	<code>logical</code> Whether to use viridis color scales.
viridis.palette	<code>character</code> A capital letter from A to H or the scale name as in <code>scale_fill_viridis</code> .
viridis.direction	<code>numeric</code> Either 1 or -1. Controls how the gradient of viridis scale is formed.
sequential.palette	<code>character</code> Type of sequential color palette to use. Out of the sequential palettes defined in <code>brewer.pal</code> .

```

sequential.direction
  numeric | Direction of the sequential color scale. Either 1 or -1.

font.size
  numeric | Overall font size of the plot. All plot elements will have a size relationship with this font size.

font.type
  character | Base font family for the plot. One of:
    • mono: Mono spaced font.
    • serif: Serif font family.
    • sans: Default font family.

na.value
  character | Color value for NA.

legend.length, legend.width
  numeric | Length and width of the legend. Will adjust automatically depending on legend side.

legend.framewidth, legend.tickwidth
  numeric | Width of the lines of the box in the legend.

legend.framecolor
  character | Color of the lines of the box in the legend.

legend.tickcolor
  character | Color of the ticks of the box in the legend.

legend.type
  character | Type of legend to display. One of:
    • normal: Default legend displayed by ggplot2.
    • colorbar: Redefined colorbar legend, using guide_colorbar.

legend.position
  character | Position of the legend in the plot. One of:
    • top: Top of the figure.
    • bottom: Bottom of the figure.
    • left: Left of the figure.
    • right: Right of the figure.
    • none: No legend is displayed.

legend.nrow
  numeric | Number of rows in the legend.

legend.ncol
  numeric | Number of columns in the legend.

legend.byrow
  logical | Whether the legend is filled by row or not.

number.breaks
  numeric | Controls the number of breaks in continuous color scales of ggplot2-based plots.

diverging.palette
  character | Type of symmetrical color palette to use. Out of the diverging palettes defined in brewer.pal.

diverging.direction
  numeric | Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.

axis.text.x.angle
  numeric | Degree to rotate the X labels. One of: 0, 45, 90.

border.color
  character | Color for the border of the heatmap body.

```

`return_object` **logical** | Returns the Seurat object with the modifications performed in the function. Normally, this contains a new assay with the data that can then be used for any other visualization desired.

`verbose` **logical** | Whether to show extra comments, warnings,etc.

`plot.title.face, plot.subtitle.face, plot.caption.face,`
`axis.title.face, axis.text.face, legend.title.face, legend.text.face`

character | Controls the style of the font for the corresponding theme element.
 One of:

- `plain`: For normal text.
- `italic`: For text in italic.
- `bold`: For text in bold.
- `bold.italic`: For text both in italic and bold.

Value

A list of ggplot2 objects and a Seurat object if desired.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_RankedExpressionPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Genes have to be unique.
  genes <- rownames(sample)[1:15]

  # This will query, for the provided components, the expression of the genes
  # for all cells and plot them in the context of the cells reordered by
  # the position alongside each dimensional reduction component.
  p <- SCpubr::do_RankedExpressionPlot(sample = sample,
                                         features = genes,
                                         nbin = 1,
                                         ctrl = 5,
                                         subsample = NA,
                                         dims = 1:2,
                                         verbose = FALSE)

  p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

do_RidgePlot *Create ridge plots.*

Description

This function computes ridge plots based on the **ggridges** package.

Usage

```
do_RidgePlot(  
  sample,  
  feature,  
  group.by = NULL,  
  split.by = NULL,  
  assay = "SCT",  
  slot = "data",  
  continuous_scale = FALSE,  
  legend.title = NULL,  
  legend.ncol = NULL,  
  legend.nrow = NULL,  
  legend.byrow = FALSE,  
  legend.position = NULL,  
  legend.width = 1,  
  legend.length = 20,  
  legend.framewidth = 0.5,  
  legend.tickwidth = 0.5,  
  legend.framecolor = "grey50",  
  legend.tickcolor = "white",  
  legend.type = "colorbar",  
  colors.use = NULL,  
  font.size = 14,  
  font.type = "sans",  
  axis.text.x.angle = 45,  
  plot.title = NULL,  
  plot.subtitle = NULL,  
  plot.caption = NULL,  
  xlab = NULL,  
  ylab = NULL,  
  compute_quantiles = FALSE,  
  compute_custom_quantiles = FALSE,  
  quantiles = c(0.25, 0.5, 0.75),  
  compute_distribution_tails = FALSE,  
  prob_tails = 0.025,  
  color_by_probabilities = FALSE,  
  use_viridis = TRUE,  
  viridis.palette = "G",  
  viridis.direction = 1,
```

```

sequential.palette = "YlGnBu",
sequential.direction = 1,
plot.grid = TRUE,
grid.color = "grey75",
grid.type = "dashed",
flip = FALSE,
number.breaks = 5,
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
feature	<code>character</code> Feature to represent.
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character of factor column.
split.by	<code>character</code> Secondary metadata variable to further group (split) the output by. Has to be a character of factor column.
assay	<code>character</code> Assay to use. Defaults to the current assay.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
continuous_scale	<code>logical</code> Whether to color the ridges depending on a categorical or continuous scale.
legend.title	<code>character</code> Title for the legend.
legend.ncol	<code>numeric</code> Number of columns in the legend.
legend.nrow	<code>numeric</code> Number of rows in the legend.
legend.byrow	<code>logical</code> Whether the legend is filled by row or not.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • top: Top of the figure. • bottom: Bottom of the figure. • left: Left of the figure. • right: Right of the figure. • none: No legend is displayed.
legend.length, legend.width	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.

```

legend.framewidth, legend.tickwidth
    numeric | Width of the lines of the box in the legend.
legend.framecolor
    character | Color of the lines of the box in the legend.
legend.tickcolor
    character | Color of the ticks of the box in the legend.
legend.type    character | Type of legend to display. One of:
    • normal: Default legend displayed by ggplot2.
    • colorbar: Redefined colorbar legend, using guide_colorbar.
colors.use     character | Named vector of colors to use. Has to match the unique values of
                    group_by or color_by (if used) when scale_type is set to categorical.
font.size      numeric | Overall font size of the plot. All plot elements will have a size rela-
                    tionship with this font size.
font.type       character | Base font family for the plot. One of:
    • mono: Mono spaced font.
    • serif: Serif font family.
    • sans: Default font family.
axis.text.x.angle
    numeric | Degree to rotate the X labels. One of: 0, 45, 90.
plot.title, plot.subtitle, plot.caption
    character | Title, subtitle or caption to use in the plot.
xlab, ylab      character | Titles for the X and Y axis.
compute_quantiles
    logical | Whether to compute quantiles of the distribution and color the ridge
              plots by them.
compute_custom_quantiles
    logical | Whether to compute custom quantiles.
quantiles        numeric | Numeric vector of quantiles.
compute_distribution_tails
    logical | Whether to compute distribution tails and color them.
prob_tails       numeric | The accumulated probability that the tails should contain.
color_by_probabilities
    logical | Whether to color the ridges depending on the probability.
use_viridis      logical | Whether to use viridis color scales.
viridis.palette
    character | A capital letter from A to H or the scale name as in scale_fill_viridis.
viridis.direction
    numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.
sequential.palette
    character | Type of sequential color palette to use. Out of the sequential
              palettes defined in brewer.pal.
sequential.direction
    numeric | Direction of the sequential color scale. Either 1 or -1.

```

plot.grid **logical** | Whether to plot grid lines.
 grid.color **character** | Color of the grid in the plot. In heatmaps, color of the border of the cells.
 grid.type **character** | One of the possible linetype options:
 • blank.
 • solid.
 • dashed.
 • dotted.
 • dotdash.
 • longdash.
 • twodash.
 flip **logical** | Whether to invert the axis of the displayed plot.
 number.breaks **numeric** | Controls the number of breaks in continuous color scales of ggplot2-based plots.
 plot.title.face, plot.subtitle.face, plot.caption.face,
 axis.title.face, axis.text.face, legend.title.face, legend.text.face
 character | Controls the style of the font for the corresponding theme element.
 One of:
 • plain: For normal text.
 • italic: For text in italic.
 • bold: For text in bold.
 • bold.italic: For text both in italic and bold.

Value

A ggplot2 object.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_RidgePlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Compute the most basic ridge plot.
  p <- SCpubr::do_RidgePlot(sample = sample,
                            feature = "nFeature_RNA")
  p

  # Use continuous color scale.
  p <- SCpubr::do_RidgePlot(sample = sample,
                            feature = "nFeature_RNA",
                            continuous_scale = TRUE,

```

```

    viridis.direction = 1)
p

# Draw quantiles of the distribution.
p <- SCpubr::do_RidgePlot(sample = sample,
                           feature = "nFeature_RNA",
                           continuous_scale = TRUE,
                           compute_quantiles = TRUE,
                           compute_custom_quantiles = TRUE)

p

# Draw probability tails.
p <- SCpubr::do_RidgePlot(sample = sample,
                           feature = "nFeature_RNA",
                           continuous_scale = TRUE,
                           compute_quantiles = TRUE,
                           compute_distribution_tails = TRUE)

p

# Draw probability tails.
p <- SCpubr::do_RidgePlot(sample = sample,
                           feature = "nFeature_RNA",
                           continuous_scale = TRUE,
                           compute_quantiles = TRUE,
                           color_by_probabilities = TRUE)

p
} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_SCEnrichmentHeatmap

Perform a single-cell-based heatmap showing the enrichment in a list of gene sets.

Description

This function is heavily inspired by [DoHeatmap](#).

Usage

```
do_SCEnrichmentHeatmap(
  sample,
  input_gene_list,
  assay = NULL,
  slot = NULL,
  group.by = NULL,
```

```
features.order = NULL,
metadata = NULL,
metadata.colors = NULL,
subsample = NA,
cluster = TRUE,
flavor = "Seurat",
return_object = FALSE,
ncores = 1,
storeRanks = TRUE,
interpolate = FALSE,
nbin = 24,
ctrl = 100,
xlab = "Cells",
ylab = "Genes",
font.size = 14,
font.type = "sans",
plot.title = NULL,
plot.subtitle = NULL,
plot.caption = NULL,
legend.position = "bottom",
legend.title = NULL,
legend.type = "colorbar",
legend.framewidth = 0.5,
legend.tickwidth = 0.5,
legend.length = 20,
legend.width = 1,
legend.framecolor = "grey50",
legend.tickcolor = "white",
strip.text.color = "black",
strip.text.angle = 0,
strip.spacing = 10,
legend.ncol = NULL,
legend.nrow = NULL,
legend.byrow = FALSE,
min.cutoff = NA,
max.cutoff = NA,
number.breaks = 5,
main.heatmap.size = 0.95,
enforce_symmetry = FALSE,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = -1,
na.value = "grey75",
diverging.palette = "RdBu",
diverging.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
proportional.size = TRUE,
```

```

verbose = FALSE,
border.color = "black",
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

<code>sample</code>	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
<code>input_gene_list</code>	<code>named_list</code> Named list of lists of genes to be used as input.
<code>assay</code>	<code>character</code> Assay to use. Defaults to the current assay.
<code>slot</code>	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
<code>group.by</code>	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
<code>features.order</code>	<code>character</code> Should the gene sets be ordered in a specific way? Provide it as a vector of characters with the same names as the names of the gene sets.
<code>metadata</code>	<code>character</code> Categorical metadata variables to plot alongside the main heatmap.
<code>metadata.colors</code>	<code>named_list</code> Named list of valid colors for each of the variables defined in metadata.
<code>subsample</code>	<code>numeric</code> Number of cells to subset for the analysis. NA will use all. Cells are selected at random.
<code>cluster</code>	<code>logical</code> Whether to perform clustering of rows and columns.
<code>flavor</code>	<code>character</code> One of: Seurat, UCell. Compute the enrichment scores using AddModuleScore or AddModuleScore_UCell .
<code>return_object</code>	<code>logical</code> Returns the Seurat object with the modifications performed in the function. Normally, this contains a new assay with the data that can then be used for any other visualization desired.
<code>ncores</code>	<code>numeric</code> Number of cores used to run UCell scoring.
<code>storeRanks</code>	<code>logical</code> Whether to store the ranks for faster UCell scoring computations. Might require large amounts of RAM.
<code>interpolate</code>	<code>logical</code> Smoothes the output heatmap, saving space on disk when saving the image. However, the image is not as crisp.
<code>nbin</code>	<code>numeric</code> Number of bins to use in AddModuleScore .
<code>ctrl</code>	<code>numeric</code> Number of genes in the control set to use in AddModuleScore .
<code>xlab, ylab</code>	<code>character</code> Titles for the X and Y axis.

font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • <code>mono</code>: Mono spaced font. • <code>serif</code>: Serif font family. • <code>sans</code>: Default font family.
plot.title, plot.subtitle, plot.caption	<code>character</code> Title, subtitle or caption to use in the plot.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • <code>top</code>: Top of the figure. • <code>bottom</code>: Bottom of the figure. • <code>left</code>: Left of the figure. • <code>right</code>: Right of the figure. • <code>none</code>: No legend is displayed.
legend.title	<code>character</code> Title for the legend.
legend.type	<code>character</code> Type of legend to display. One of: <ul style="list-style-type: none"> • <code>normal</code>: Default legend displayed by <code>ggplot2</code>. • <code>colorbar</code>: Redefined colorbar legend, using <code>guide_colorbar</code>.
legend.framewidth, legend.tickwidth	<code>numeric</code> Width of the lines of the box in the legend.
legend.length, legend.width	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.
legend.framecolor	<code>character</code> Color of the lines of the box in the legend.
legend.tickcolor	<code>character</code> Color of the ticks of the box in the legend.
strip.text.color	<code>character</code> Color of the strip text.
strip.text.angle	<code>numeric</code> Rotation of the strip text (angles).
strip.spacing	<code>numeric</code> Controls the size between the different facets.
legend.ncol	<code>numeric</code> Number of columns in the legend.
legend.nrow	<code>numeric</code> Number of rows in the legend.
legend.byrow	<code>logical</code> Whether the legend is filled by row or not.
min.cutoff, max.cutoff	<code>numeric</code> Set the min/max ends of the color scale. Any cell/group with a value lower than <code>min.cutoff</code> will turn into <code>min.cutoff</code> and any cell with a value higher than <code>max.cutoff</code> will turn into <code>max.cutoff</code> . In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
number.breaks	<code>numeric</code> Controls the number of breaks in continuous color scales of ggplot2-based plots.

```

main.heatmap.size
    numeric | Controls the size of the main heatmap (proportion-wise, defaults to
    0.95).
enforce_symmetry
    logical | Return a symmetrical plot axes-wise or continuous color scale-wise,
    when applicable.
use_viridis      logical | Whether to use viridis color scales.
viridis.palette
    character | A capital letter from A to H or the scale name as in scale_fill_viridis.
viridis.direction
    numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.
na.value         character | Color value for NA.
diverging.palette
    character | Type of symmetrical color palette to use. Out of the diverging
    palettes defined in brewer.pal.
diverging.direction
    numeric | Either 1 or -1. Direction of the diverging palette. This basically flips
    the two ends.
sequential.palette
    character | Type of sequential color palette to use. Out of the sequential
    palettes defined in brewer.pal.
sequential.direction
    numeric | Direction of the sequential color scale. Either 1 or -1.
proportional.size
    logical | Whether the groups should take the same space in the plot or not.
verbose          logical | Whether to show extra comments, warnings,etc.
border.color     character | Color for the border of the heatmap body.
plot.title.face,   plot.subtitle.face,   plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
    character | Controls the style of the font for the corresponding theme element.
    One of:
        • plain: For normal text.
        • italic: For text in italic.
        • bold: For text in bold.
        • bold.italic: For text both in italic and bold.

```

Value

A ggplot2 object.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_SCEnrichmentHeatmap", passive = TRUE)

if (isTRUE(value)){

```

```

# Consult the full documentation in https://enblacar.github.io/SCpubr-book/

# Define your Seurat object.
sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

# Genes have to be unique.
genes <- list("A" = rownames(sample)[1:5],
              "B" = rownames(sample)[6:10],
              "C" = rownames(sample)[11:15])

p <- SCpubr::do_SCEnrichmentHeatmap(sample = sample,
                                      input_gene_list = genes,
                                      nbin = 1,
                                      ctrl = 5,
                                      flavor = "Seurat",
                                      subsample = NA)

p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_SCExpressionHeatmap*Perform a single-cell-based heatmap showing the expression of genes.***Description**

This function is heavily inspired by [DoHeatmap](#).

Usage

```
do_SCExpressionHeatmap(
  sample,
  features,
  assay = NULL,
  slot = NULL,
  group.by = NULL,
  features.order = NULL,
  metadata = NULL,
  metadata.colors = NULL,
  subsample = NA,
  cluster = TRUE,
  interpolate = FALSE,
  xlab = "Cells",
  ylab = "Genes",
  font.size = 14,
```

```

font.type = "sans",
plot.title = NULL,
plot.subtitle = NULL,
plot.caption = NULL,
legend.position = "bottom",
legend.title = "Expression",
legend.type = "colorbar",
legend.framewidth = 0.5,
legend.tickwidth = 0.5,
legend.length = 20,
legend.width = 1,
legend.framecolor = "grey50",
legend.tickcolor = "white",
strip.text.color = "black",
strip.text.angle = 0,
strip.spacing = 10,
legend.ncol = NULL,
legend.nrow = NULL,
legend.byrow = FALSE,
min.cutoff = NA,
max.cutoff = NA,
number.breaks = 5,
main.heatmap.size = 0.95,
enforce_symmetry = FALSE,
use_viridis = FALSE,
viridis.palette = "G",
viridis.direction = -1,
na.value = "grey75",
diverging.palette = "RdBu",
diverging.direction = -1,
sequential.palette = "YlGnBu",
sequential.direction = 1,
proportional.size = TRUE,
verbose = TRUE,
border.color = "black",
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain"
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
features	character Features to represent.

assay	<code>character</code> Assay to use. Defaults to the current assay.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character or factor column.
features.order	<code>character</code> Should the gene sets be ordered in a specific way? Provide it as a vector of characters with the same names as the names of the gene sets.
metadata	<code>character</code> Categorical metadata variables to plot alongside the main heatmap.
metadata.colors	<code>named_list</code> Named list of valid colors for each of the variables defined in metadata.
subsample	<code>numeric</code> Number of cells to subset for the analysis. NA will use all. Cells are selected at random.
cluster	<code>logical</code> Whether to perform clustering of rows and columns.
interpolate	<code>logical</code> Smoothes the output heatmap, saving space on disk when saving the image. However, the image is not as crisp.
xlab, ylab	<code>character</code> Titles for the X and Y axis.
font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • mono: Mono spaced font. • serif: Serif font family. • sans: Default font family.
plot.title, plot.subtitle, plot.caption	<code>character</code> Title, subtitle or caption to use in the plot.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • top: Top of the figure. • bottom: Bottom of the figure. • left: Left of the figure. • right: Right of the figure. • none: No legend is displayed.
legend.title	<code>character</code> Title for the legend.
legend.type	<code>character</code> Type of legend to display. One of: <ul style="list-style-type: none"> • normal: Default legend displayed by <code>ggplot2</code>. • colorbar: Redefined colorbar legend, using <code>guide_colorbar</code>.
legend.framewidth, legend.tickwidth	<code>numeric</code> Width of the lines of the box in the legend.
legend.length, legend.width	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.

```

legend.framecolor
  character | Color of the lines of the box in the legend.
legend.tickcolor
  character | Color of the ticks of the box in the legend.
strip.text.color
  character | Color of the strip text.
strip.text.angle
  numeric | Rotation of the strip text (angles).
strip.spacing  numeric | Controls the size between the different facets.
legend.ncol    numeric | Number of columns in the legend.
legend.nrow    numeric | Number of rows in the legend.
legend.byrow   logical | Whether the legend is filled by row or not.
min.cutoff, max.cutoff
  numeric | Set the min/max ends of the color scale. Any cell/group with a value
  lower than min.cutoff will turn into min.cutoff and any cell with a value higher
  than max.cutoff will turn into max.cutoff. In FeaturePlots, provide as many
  values as features. Use NAs to skip a feature.
number.breaks  numeric | Controls the number of breaks in continuous color scales of ggplot2-
  based plots.
main.heatmap.size
  numeric | Controls the size of the main heatmap (proportion-wise, defaults to
  0.95).
enforce_symmetry
  logical | Return a symmetrical plot axes-wise or continuous color scale-wise,
  when applicable.
use_viridis    logical | Whether to use viridis color scales.
viridis.palette
  character | A capital letter from A to H or the scale name as in scale\_fill\_viridis.
viridis.direction
  numeric | Either 1 or -1. Controls how the gradient of viridis scale is formed.
na.value       character | Color value for NA.
diverging.palette
  character | Type of symmetrical color palette to use. Out of the diverging
  palettes defined in brewer.pal.
diverging.direction
  numeric | Either 1 or -1. Direction of the diverging palette. This basically flips
  the two ends.
sequential.palette
  character | Type of sequential color palette to use. Out of the sequential
  palettes defined in brewer.pal.
sequential.direction
  numeric | Direction of the sequential color scale. Either 1 or -1.
proportional.size
  logical | Whether the groups should take the same space in the plot or not.

```

```

verbose      logical | Whether to show extra comments, warnings,etc.
border.color character | Color for the border of the heatmap body.
plot.title.face,    plot.subtitle.face,    plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
character | Controls the style of the font for the corresponding theme element.
One of:
  • plain: For normal text.
  • italic: For text in italic.
  • bold: For text in bold.
  • bold.italic: For text both in italic and bold.

```

Value

A ggplot2 object.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_SCExpressionHeatmap", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/

  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  p <- SCpubr::do_SCExpressionHeatmap(sample = sample,
                                         features = rownames(sample)[1:2],
                                         subsample = NA)
  p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

`do_TermEnrichmentPlot` *Display the enriched terms for a given list of genes.*

Description

Display the enriched terms for a given list of genes.

Usage

```
do_TermEnrichmentPlot(
  mat,
  n.chars = 40,
  n.terms = 25,
  font.size = 14,
  font.type = "sans",
  plot.title = NULL,
  plot.subtitle = NULL,
  plot.caption = NULL,
  use_viridis = FALSE,
  viridis.palette = "G",
  viridis.direction = -1,
  sequential.palette = "YlGnBu",
  sequential.direction = 1,
  dot.scale = 8,
  legend.type = "colorbar",
  legend.position = "bottom",
  legend.framewidth = 0.5,
  legend.tickwidth = 0.5,
  legend.length = 20,
  legend.width = 1,
  legend.framecolor = "grey50",
  legend.tickcolor = "white",
  number.breaks = 5,
  xlab = NULL,
  ylab = NULL,
  na.value = "grey75",
  grid.color = "grey90",
  grid.type = "dashed",
  plot.title.face = "bold",
  plot.subtitle.face = "plain",
  plot.caption.face = "italic",
  axis.title.face = "bold",
  axis.text.face = "plain",
  axis.text.x.angle = 45,
  legend.title.face = "bold",
  legend.text.face = "plain"
)
```

Arguments

mat	list Result of over-representation test with clusterProfiler. Accepts only one result, be aware of that if you compute the test for all GO ontologies. Accessed through mat@result.
n.chars	numeric Number of characters to use as a limit to wrap the term names. The higher this value, the longer the lines would be for each term in the plots. Defaults to 40.

n.terms **numeric** | Number of terms to display. Defaults to 25.

font.size **numeric** | Overall font size of the plot. All plot elements will have a size relationship with this font size.

font.type **character** | Base font family for the plot. One of:

- mono: Mono spaced font.
- serif: Serif font family.
- sans: Default font family.

plot.title, plot.subtitle, plot.caption **character** | Title, subtitle or caption to use in the plot.

use_viridis **logical** | Whether to use viridis color scales.

viridis.palette **character** | A capital letter from A to H or the scale name as in `scale_fill_viridis`.

viridis.direction **numeric** | Either 1 or -1. Controls how the gradient of viridis scale is formed.

sequential.palette **character** | Type of sequential color palette to use. Out of the sequential palettes defined in `brewer.pal`.

sequential.direction **numeric** | Direction of the sequential color scale. Either 1 or -1.

dot.scale **numeric** | Scale the size of the dots.

legend.type **character** | Type of legend to display. One of:

- normal: Default legend displayed by `ggplot2`.
- colorbar: Redefined colorbar legend, using `guide_colorbar`.

legend.position **character** | Position of the legend in the plot. One of:

- top: Top of the figure.
- bottom: Bottom of the figure.
- left: Left of the figure.
- right: Right of the figure.
- none: No legend is displayed.

legend.framewidth, legend.tickwidth **numeric** | Width of the lines of the box in the legend.

legend.length, legend.width **numeric** | Length and width of the legend. Will adjust automatically depending on legend side.

legend.framecolor **character** | Color of the lines of the box in the legend.

legend.tickcolor **character** | Color of the ticks of the box in the legend.

number.breaks **numeric** | Controls the number of breaks in continuous color scales of ggplot2-based plots.

xlab, ylab **character** | Titles for the X and Y axis.

```

na.value      character | Color value for NA.
grid.color    character | Color of the grid in the plot. In heatmaps, color of the border of the
               cells.
grid.type     character | One of the possible linetype options:
               • blank.
               • solid.
               • dashed.
               • dotted.
               • dotdash.
               • longdash.
               • twodash.
plot.title.face,      plot.subtitle.face,      plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face
               character | Controls the style of the font for the corresponding theme element.
               One of:
               • plain: For normal text.
               • italic: For text in italic.
               • bold: For text in bold.
               • bold.italic: For text both in italic and bold.
axis.text.x.angle
               numeric | Degree to rotate the X labels. One of: 0, 45, 90.

```

Value

A dotplot object with enriched terms.

Examples

```

# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_TermEnrichmentPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  #
  # Define your enriched terms.
  enriched_terms <- readRDS(system.file("extdata/enriched_terms_example.rds", package = "SCpubr"))

  # Default plot.
  p <- SCpubr::do_TermEnrichmentPlot(mat = enriched_terms)

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_TFActivityPlot *Plot TF Activities from decoupleR using Dorothea prior knowledge.*

Description

Plot TF Activities from decoupleR using Dorothea prior knowledge.

Usage

```
do_TFActivityPlot(  
  sample,  
  activities,  
  n_tfs = 25,  
  slot = "scale.data",  
  statistic = "norm_wmean",  
  tfs.use = NULL,  
  group.by = NULL,  
  split.by = NULL,  
  na.value = "grey75",  
  legend.position = "bottom",  
  legend.width = 1,  
  legend.length = 20,  
  legend.framewidth = 0.5,  
  legend.tickwidth = 0.5,  
  legend.framecolor = "grey50",  
  legend.tickcolor = "white",  
  legend.type = "colorbar",  
  font.size = 14,  
  font.type = "sans",  
  axis.text.x.angle = 45,  
  enforce_symmetry = TRUE,  
  diverging.palette = "RdBu",  
  diverging.direction = -1,  
  use_viridis = FALSE,  
  viridis.palette = "G",  
  viridis.direction = -1,  
  sequential.palette = "YlGnBu",  
  sequential.direction = 1,  
  min.cutoff = NA,  
  max.cutoff = NA,  
  number.breaks = 5,  
  flip = FALSE,  
  return_object = FALSE,  
  grid.color = "white",  
  border.color = "black",  
  plot.title.face = "bold",  
  plot.subtitle.face = "plain",
```

```

    plot.caption.face = "italic",
    axis.title.face = "bold",
    axis.text.face = "plain",
    legend.title.face = "bold",
    legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
activities	<code>tibble</code> Result of running decoupleR method with dorothea regulon prior knowledge.
n_tfs	<code>numeric</code> Number of top regulons to consider for downstream analysis.
slot	<code>character</code> Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".
statistic	<code>character</code> DecoupleR statistic to use. One of: <ul style="list-style-type: none"> • wmean: For weighted mean. • norm_wmean: For normalized weighted mean. • corr_wmean: For corrected weighted mean.
tfs.use	<code>character</code> Restrict the analysis to given regulons.
group.by	<code>character</code> Metadata variable to group the output by. Has to be a character of factor column.
split.by	<code>character</code> Secondary metadata variable to further group (split) the output by. Has to be a character of factor column.
na.value	<code>character</code> Color value for NA.
legend.position	<code>character</code> Position of the legend in the plot. One of: <ul style="list-style-type: none"> • top: Top of the figure. • bottom: Bottom of the figure. • left: Left of the figure. • right: Right of the figure. • none: No legend is displayed.
legend.length, legend.width	<code>numeric</code> Length and width of the legend. Will adjust automatically depending on legend side.
legend.framewidth, legend.tickwidth	<code>numeric</code> Width of the lines of the box in the legend.
legend.framecolor	<code>character</code> Color of the lines of the box in the legend.
legend.tickcolor	<code>character</code> Color of the ticks of the box in the legend.
legend.type	<code>character</code> Type of legend to display. One of: <ul style="list-style-type: none"> • normal: Default legend displayed by <code>ggplot2</code>.

	<ul style="list-style-type: none"> • colorbar: Redefined colorbar legend, using <code>guide_colorbar</code>.
<code>font.size</code>	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
<code>font.type</code>	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • mono: Mono spaced font. • serif: Serif font family. • sans: Default font family.
<code>axis.text.x.angle</code>	<code>numeric</code> Degree to rotate the X labels. One of: 0, 45, 90.
<code>enforce_symmetry</code>	<code>logical</code> Whether the geyser and feature plot has a symmetrical color scale.
<code>diverging.palette</code>	<code>character</code> Type of symmetrical color palette to use. Out of the diverging palettes defined in <code>brewer.pal</code> .
<code>diverging.direction</code>	<code>numeric</code> Either 1 or -1. Direction of the diverging palette. This basically flips the two ends.
<code>use_viridis</code>	<code>logical</code> Whether to use viridis color scales.
<code>viridis.palette</code>	<code>character</code> A capital letter from A to H or the scale name as in <code>scale_fill_viridis</code> .
<code>viridis.direction</code>	<code>numeric</code> Either 1 or -1. Controls how the gradient of viridis scale is formed.
<code>sequential.palette</code>	<code>character</code> Type of sequential color palette to use. Out of the sequential palettes defined in <code>brewer.pal</code> .
<code>sequential.direction</code>	<code>numeric</code> Direction of the sequential color scale. Either 1 or -1.
<code>min.cutoff, max.cutoff</code>	<code>numeric</code> Set the min/max ends of the color scale. Any cell/group with a value lower than <code>min.cutoff</code> will turn into <code>min.cutoff</code> and any cell with a value higher than <code>max.cutoff</code> will turn into <code>max.cutoff</code> . In FeaturePlots, provide as many values as features. Use NAs to skip a feature.
<code>number.breaks</code>	<code>numeric</code> Controls the number of breaks in continuous color scales of ggplot2-based plots.
<code>flip</code>	<code>logical</code> Whether to invert the axis of the displayed plot.
<code>return_object</code>	<code>logical</code> Returns the Seurat object with the modifications performed in the function. Normally, this contains a new assay with the data that can then be used for any other visualization desired.
<code>grid.color</code>	<code>character</code> Color of the grid in the plot. In heatmaps, color of the border of the cells.
<code>border.color</code>	<code>character</code> Color for the border of the heatmap body.
<code>plot.title.face, plot.subtitle.face, plot.caption.face,</code> <code>axis.title.face, axis.text.face, legend.title.face, legend.text.face</code>	<code>character</code> Controls the style of the font for the corresponding theme element. One of:

- **plain**: For normal text.
- **italic**: For text in italic.
- **bold**: For text in bold.
- **bold.italic**: For text both in italic and bold.

Value

A ggplot2 object.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_TFActivityPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds",
                                 package = "SCpubr"))

  # Define your activities object.
  dorothea_activities <- readRDS(system.file("extdata/dorothea_activities_example.rds",
                                              package = "SCpubr"))

  # General heatmap.
  out <- SCpubr::do_TFActivityPlot(sample = sample,
                                      activities = dorothea_activities)
  p <- out$heatmaps$average_scores
  p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

do_ViolinPlot *Wrapper for [VlnPlot](#).*

Description

Wrapper for [VlnPlot](#).

Usage

```
do_ViolinPlot(
  sample,
  features,
```

```
assay = NULL,  
slot = NULL,  
group.by = NULL,  
split.by = NULL,  
colors.use = NULL,  
pt.size = 0,  
line_width = 0.5,  
y_cut = rep(NA, length(features)),  
plot_boxplot = TRUE,  
boxplot_width = 0.2,  
legend.position = "bottom",  
plot.title = NULL,  
plot.subtitle = NULL,  
plot.caption = NULL,  
xlab = rep(NA, length(features)),  
ylab = rep(NA, length(features)),  
font.size = 14,  
font.type = "sans",  
axis.text.x.angle = 45,  
plot.grid = TRUE,  
grid.color = "grey75",  
grid.type = "dashed",  
order = TRUE,  
flip = FALSE,  
ncol = NULL,  
share.y.lims = FALSE,  
legend.title = NULL,  
legend.title.position = "top",  
legend.ncol = NULL,  
legend.nrow = NULL,  
legend.byrow = FALSE,  
plot.title.face = "bold",  
plot.subtitle.face = "plain",  
plot.caption.face = "italic",  
axis.title.face = "bold",  
axis.text.face = "plain",  
legend.title.face = "bold",  
legend.text.face = "plain"  
)
```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
features	character Features to represent.
assay	character Assay to use. Defaults to the current assay.
slot	character Data slot to use. Only one of: counts, data, scale.data. Defaults to "data".

group.by	character Metadata variable to group the output by. Has to be a character or factor column.
split.by	character Secondary metadata variable to further group (split) the output by. Has to be a character of factor column.
colors.use	named_vector Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of group.by. If group.by is not provided, defaults to the unique values of Idents . If not provided, a color scale will be set by default.
pt.size	numeric Size of points in the Violin plot.
line_width	numeric Width of the lines drawn in the plot. Defaults to 1.
y_cut	numeric Vector with the values in which the Violins should be cut. Only works for one feature.
plot_boxplot	logical Whether to plot a Box plot inside the violin or not.
boxplot_width	numeric Width of the boxplots. Defaults to 0.2.
legend.position	character Position of the legend in the plot. One of: <ul style="list-style-type: none"> • top: Top of the figure. • bottom: Bottom of the figure. • left: Left of the figure. • right: Right of the figure. • none: No legend is displayed.
plot.title, plot.subtitle, plot.caption	character Title, subtitle or caption to use in the plot.
xlab, ylab	character Titles for the X and Y axis.
font.size	numeric Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	character Base font family for the plot. One of: <ul style="list-style-type: none"> • mono: Mono spaced font. • serif: Serif font family. • sans: Default font family.
axis.text.x.angle	numeric Degree to rotate the X labels. One of: 0, 45, 90.
plot.grid	logical Whether to plot grid lines.
grid.color	character Color of the grid in the plot. In heatmaps, color of the border of the cells.
grid.type	character One of the possible linetype options: <ul style="list-style-type: none"> • blank. • solid. • dashed. • dotted. • dotdash.

	<ul style="list-style-type: none"> • longdash. • twodash.
order	<code>logical</code> Whether to order the boxplots by average values. Can not be used alongside <code>split.by</code> .
flip	<code>logical</code> Whether to invert the axis of the displayed plot.
ncol	<code>numeric</code> Number of columns used in the arrangement of the output plot using "split.by" parameter.
share.y.lims	<code>logical</code> When querying multiple features, force the Y axis of all of them to be on the same range of values (this being the max and min of all features combined).
legend.title	<code>character</code> Title for the legend.
legend.title.position	<code>character</code> Position for the title of the legend. One of: <ul style="list-style-type: none"> • top: Top of the legend. • bottom: Bottom of the legend. • left: Left of the legend. • right: Right of the legend.
legend.ncol	<code>numeric</code> Number of columns in the legend.
legend.nrow	<code>numeric</code> Number of rows in the legend.
legend.byrow	<code>logical</code> Whether the legend is filled by row or not.
plot.title.face, plot.subtitle.face, plot.caption.face, axis.title.face, axis.text.face, legend.title.face, legend.text.face	<code>character</code> Controls the style of the font for the corresponding theme element. One of: <ul style="list-style-type: none"> • plain: For normal text. • italic: For text in italic. • bold: For text in bold. • bold.italic: For text both in italic and bold.

Value

A ggplot2 object containing a Violin Plot.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_ViolinPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Basic violin plot.
```

```

p <- SCpubr::do_ViolinPlot(sample = sample,
                            feature = "nCount_RNA")
p

# Remove the box plots.
p <- SCpubr::do_ViolinPlot(sample = sample,
                            feature = "nCount_RNA",
                            plot_boxplot = FALSE)
p

} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

```

do_VolcanoPlot *Compute a Volcano plot out of DE genes.*

Description

Compute a Volcano plot out of DE genes.

Usage

```

do_VolcanoPlot(
  sample,
  de_genes,
  pval_cutoff = 0.05,
  FC_cutoff = 2,
  pt.size = 2,
  border.size = 1.5,
  border.color = "black",
  font.size = 14,
  font.type = "sans",
  plot.title = NULL,
  plot.subtitle = NULL,
  plot.caption = NULL,
  plot_lines = TRUE,
  line_color = "grey75",
  line_size = 0.5,
  add_gene_tags = TRUE,
  add_tag_side = "both",
  order_tags_by = "both",
  tag_size = 6,
  n_genes = 5,
  use_labels = FALSE,
  colors.use = "steelblue",

```

```

    plot.title.face = "bold",
    plot.subtitle.face = "plain",
    plot.caption.face = "italic",
    axis.title.face = "bold",
    axis.text.face = "plain",
    legend.title.face = "bold",
    legend.text.face = "plain"
)

```

Arguments

sample	<code>Seurat</code> A Seurat object, generated by CreateSeuratObject .
de_genes	<code>tibble</code> Output of <code>Seurat::FindMarkers()</code> .
pval_cutoff	<code>numeric</code> Cutoff for the p-value.
FC_cutoff	<code>numeric</code> Cutoff for the avg_log2FC.
pt.size	<code>numeric</code> Size of the dots.
border.size	<code>numeric</code> Width of the border of the cells.
border.color	<code>character</code> Color for the border of the heatmap body.
font.size	<code>numeric</code> Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	<code>character</code> Base font family for the plot. One of: <ul style="list-style-type: none"> • <code>mono</code>: Mono spaced font. • <code>serif</code>: Serif font family. • <code>sans</code>: Default font family.
plot.title, plot.subtitle, plot.caption	<code>character</code> Title, subtitle or caption to use in the plot.
plot_lines	<code>logical</code> Whether to plot the division lines.
line_color	<code>character</code> Color for the lines.
line_size	<code>numeric</code> Size of the lines in the plot.
add_gene_tags	<code>logical</code> Whether to plot the top genes.
add_tag_side	<code>logical</code> Either "both", "positive" or "negative" to indicate which side of genes to tag
order_tags_by	<code>character</code> Either "both", "pvalue" or "logfc".
tag_size	<code>numeric</code> Size of the text/label for the tags.
n_genes	<code>numeric</code> Number of top genes to plot.
use_labels	<code>logical</code> Whether to use labels instead of text for the tags.
colors.use	<code>character</code> Color to generate a tetradic color scale with.
plot.title.face, plot.subtitle.face, plot.caption.face,	<code>character</code> Controls the style of the font for the corresponding theme element.
axis.title.face, axis.text.face, legend.title.face, legend.text.face	One of: <ul style="list-style-type: none"> • <code>plain</code>: For normal text. • <code>italic</code>: For text in italic. • <code>bold</code>: For text in bold. • <code>bold.italic</code>: For text both in italic and bold.

Value

A volcano plot as a `ggplot2` object.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_VolcanoPlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Retrieve DE genes.
  de_genes <- readRDS(system.file("extdata/de_genes_example.rds", package = "SCpubr"))

  # Generate a volcano plot.
  p <- SCpubr::do_VolcanoPlot(sample = sample,
                                de_genes = de_genes)
  p

} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

do_WafflePlot

Display the enriched terms for a given list of genes.

Description

Display the enriched terms for a given list of genes.

Usage

```
do_WafflePlot(
  sample,
  group.by,
  waffle.size = 2,
  flip = TRUE,
  colors.use = NULL,
  na.value = "grey75",
  font.size = 14,
  font.type = "sans",
  plot.title = NULL,
  plot.subtitle = NULL,
```

```

plot.caption = NULL,
legend.title = NULL,
legend.ncol = NULL,
legend.nrow = NULL,
legend.byrow = FALSE,
legend.position = "bottom",
plot.title.face = "bold",
plot.subtitle.face = "plain",
plot.caption.face = "italic",
axis.title.face = "bold",
axis.text.face = "plain",
legend.title.face = "bold",
legend.text.face = "plain",
strip.text.face = "bold"
)

```

Arguments

sample	Seurat A Seurat object, generated by CreateSeuratObject .
group.by	character Metadata variable to group the output by. Has to be a character or factor column.
waffle.size	numeric Tile border size.
flip	logical Whether to invert the axis of the displayed plot.
colors.use	named_vector Named vector of valid color representations (either name of HEX codes) with as many named colors as unique values of group.by. If group.by is not provided, defaults to the unique values of Idents . If not provided, a color scale will be set by default.
na.value	character Color value for NA.
font.size	numeric Overall font size of the plot. All plot elements will have a size relationship with this font size.
font.type	character Base font family for the plot. One of: <ul style="list-style-type: none"> • <code>mono</code>: Mono spaced font. • <code>serif</code>: Serif font family. • <code>sans</code>: Default font family.
plot.title, plot.subtitle, plot.caption	character Title, subtitle or caption to use in the plot.
legend.title	character Title for the legend.
legend.ncol	numeric Number of columns in the legend.
legend.nrow	numeric Number of rows in the legend.
legend.byrow	logical Whether the legend is filled by row or not.
legend.position	character Position of the legend in the plot. One of: <ul style="list-style-type: none"> • <code>top</code>: Top of the figure. • <code>bottom</code>: Bottom of the figure.

- **left**: Left of the figure.
- **right**: Right of the figure.
- **none**: No legend is displayed.

`plot.title.face, plot.subtitle.face, plot.caption.face,
axis.title.face, axis.text.face, legend.title.face, legend.text.face`
`character` | Controls the style of the font for the corresponding theme element.
One of:

- **plain**: For normal text.
- **italic**: For text in italic.
- **bold**: For text in bold.
- **bold.italic**: For text both in italic and bold.

`strip.text.face`
`character` | Controls the style of the font for the strip text. One of:

- **plain**: For normal text.
- **italic**: For text in italic.
- **bold**: For text in bold.
- **bold.italic**: For text both in italic and bold.

Value

A ggplot2 object with a Waffle Plot.

Examples

```
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "do_WafflePlot", passive = TRUE)

if (isTRUE(value)){
  # Consult the full documentation in https://enblacar.github.io/SCpubr-book/
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Basic Waffle plot.
  p <- SCpubr:::do_WafflePlot(sample = sample,
                                group.by = "seurat_clusters")

} else if (base:::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}
```

human_chr_locations *Chromosome arm locations for human genome GRCh38.*

Description

A tibble containing the chromosome, arm and start and end coordinates.

Usage

```
data(human_chr_locations)
```

Format

A tibble with 48 rows and 4 columns:

chr Chromosome.
arm Chromosome arm.
start Start coordinates.
end End coordinates.

package_report *Generate a status report of SCpubr and its dependencies.*

Description

This function generates a summary report of the installation status of SCpubr, which packages are still missing and which functions can or can not currently be used.

Usage

```
package_report(startup = FALSE, extended = FALSE)
```

Arguments

startup	<code>logical</code> Whether the message should be displayed at startup, therefore, also containing welcoming messages and tips. If FALSE, only the report itself will be printed.
extended	<code>logical</code> Whether the message should also include installed packages, current and available version, and which SCpubr functions can be used with the currently installed packages.

Value

None

Examples

```
# Print a package report.
SCpubr::package_report(startup = FALSE, extended = FALSE)
```

save_Plot

Save a plot as png, pdf and svg.

Description

Save a plot as png, pdf and svg.

Usage

```
save_Plot(
  plot,
  figure_path = NULL,
  create_path = TRUE,
  file_name = NULL,
  dpi = 300,
  output_format = "publication",
  width = 8,
  height = 8
)
```

Arguments

plot	Plot to save.
figure_path	character Path where the figure will be stored.
create_path	logical Whether to create the path.
file_name	character Name of the file (without extension, it will be added automatically).
dpi	numeric Dpi to use.
output_format	character Output format of the saved figure. One of: <ul style="list-style-type: none"> • pdf: Saves the figure as a PDF file. • png: Saves the figure as a PNG file. • jpeg: Saves the figure as a JPEG file. • tiff: Saves the figure as a TIFF file. • svg: Saves the figure as a SVG file. • publication: Saves the figure as PDF, PNG and SVG files. • all: Saves the figure in all possible formats.
width, height	numeric Width and height of the figure (inches).

Value

Nothing.

Examples

```
## Not run:
# Check Suggests.
value <- SCpubr:::check_suggests(function_name = "save_Plot", passive = TRUE)

if (isTRUE(value)){
  # Define your Seurat object.
  sample <- readRDS(system.file("extdata/seurat_dataset_example.rds", package = "SCpubr"))

  # Generate a plot.
  p <- SCpubr::do_DimPlot(sample = sample)

  # Default parameters.
  SCpubr::save_Plot(plot = p)

  # Specifying the name and folder.
  SCpubr::save_Plot(plot = p,
                    figure_path = "/path/to/my/figures/",
                    file_name = "my_figure")

  # Specify to also create a new folder.
  SCpubr::save_Plot(plot = p,
                    figure_path = "/path/to/my/figures/",
                    file_name = "my_figure",
                    create_path = TRUE)

  # Set dimensions for the figure.
  SCpubr::save_Plot(plot = p,
                    figure_path = "/path/to/my/figures/",
                    file_name = "my_figure",
                    create_path = TRUE,
                    width = 8,
                    height = 8)

  # Set quality (dpi).
  SCpubr::save_Plot(plot = p,
                    figure_path = "/path/to/my/figures/",
                    file_name = "my_figure",
                    create_path = TRUE,
                    width = 8,
                    height = 8,
                    dpi = 300)
} else if (base::isFALSE(value)){
  message("This function can not be used without its suggested packages.")
  message("Check out which ones are needed using `SCpubr::state_dependencies()`")
}

## End(Not run)
```


Index

* datasets

- human_chr_locations, 129
- AddModuleScore, 6, 26, 50, 53, 93, 97, 106
- AddModuleScore_UCell, 6, 53, 93, 106
- brewer.pal, 4, 9, 17, 26, 35, 36, 39, 40, 49, 53, 57, 63, 68, 71, 76, 79, 86, 89, 90, 93, 94, 97, 98, 102, 108, 112, 115, 119
- character, 4–6, 8–13, 15–17, 19–21, 24–26, 28, 29, 31, 34–36, 38–40, 42–45, 47–49, 52–54, 56, 57, 60–63, 66–68, 70–72, 74–76, 78–80, 82, 83, 85, 86, 88–90, 92–94, 97–99, 101–103, 106–108, 110–113, 115, 116, 118, 119, 121–123, 125, 127, 128, 130
- chordDiagramFromDataFrame, 29, 30
- chordDiagramFromMatrix, 29, 30
- CreateSeuratObject, 4, 8, 11, 15, 19, 24, 28, 34, 38, 42, 47, 51, 56, 60, 66, 70, 78, 81, 85, 88, 92, 97, 101, 106, 110, 118, 121, 125, 127
- data.frame, 82
- DimPlot, 41
- do_AffinityAnalysisPlot, 3
- do_AlluvialPlot, 7
- do_BarPlot, 10
- do_BeeSwarmPlot, 14
- do_BoxPlot, 18
- do_CellularStatesPlot, 23
- do_ChordDiagramPlot, 28
- do_ColorPalette, 30
- do_CopyNumberVariantPlot, 33
- do_CorrelationPlot, 37
- do_DimPlot, 41
- do_DotPlot, 46
- do_EnrichmentHeatmap, 50
- do_ExpressionHeatmap, 55
- do_FeaturePlot, 58
- do_GeyserPlot, 64
- do_GroupwiseDEPlot, 69
- do_LigandReceptorPlot, 73
- do_LoadingsPlot, 77
- do_MetadataPlot, 81
- do_NebulosaPlot, 83
- do_PathwayActivityPlot, 87
- do_RankedEnrichmentPlot, 91
- do_RankedExpressionPlot, 95
- do_RidgePlot, 100
- do_SCEnrichmentHeatmap, 104
- do_SCExpressionHeatmap, 109
- do_TermEnrichmentPlot, 113
- do_TFActivityPlot, 117
- do_ViolinPlot, 120
- do_VolcanoPlot, 124
- do_WafflePlot, 126
- DoHeatmap, 104, 109
- DotPlot, 46
- FeaturePlot, 58
- geom_alluvium, 9
- geom_flow, 9
- geom_label, 43
- geom_text, 43
- guide_colorbar, 5, 16, 34, 39, 48, 53, 57, 61, 67, 72, 75, 79, 85, 89, 94, 98, 102, 107, 111, 115, 119
- Heatmap, 50
- human_chr_locations, 129
- Idents, 5, 12, 16, 20, 25, 42, 71, 122, 127
- infercnv, 34
- liana_aggregate, 74
- liana_wrap, 74
- list, 93, 97, 114

logical, 4–6, 8, 9, 11–13, 16, 17, 20, 21, 25, 26, 29, 31, 34–36, 38, 39, 42–44, 47–49, 52–54, 56, 57, 60–63, 66–68, 71, 72, 74–76, 79, 80, 82, 85, 86, 89, 90, 92–94, 97–99, 101–103, 106–108, 111–113, 115, 119, 122, 123, 125, 127, 129, 130
named_list, 4, 24, 38, 52, 56, 82, 92, 106, 111
named_vector, 5, 12, 16, 20, 24, 29, 34, 42, 71, 122, 127
numeric, 4–6, 8, 9, 12, 16, 17, 20, 21, 25, 26, 29–31, 35, 36, 38–40, 43–45, 48, 49, 52, 53, 56, 57, 61–63, 66–68, 70–72, 74–76, 78–80, 82, 85, 86, 88–90, 92–94, 97, 98, 101–103, 106–108, 111, 112, 114–116, 118, 119, 122, 123, 125, 127, 130
package_report, 129
position_fill, 12
position_stack, 12
save_Plot, 130
scale_fill_viridis, 4, 9, 17, 26, 35, 39, 49, 52, 57, 62, 67, 71, 75, 79, 86, 90, 93, 97, 102, 108, 112, 115, 119
Seurat, 4, 8, 11, 15, 19, 24, 28, 34, 38, 42, 47, 51, 56, 60, 66, 70, 78, 81, 85, 88, 92, 97, 101, 106, 110, 118, 121, 125, 127
tibble, 34, 70, 74, 88, 118, 125
VlnPlot, 120