Package: microViz (via r-universe)

September 14, 2024		
Title Microbiome Data Analysis and Visualization		
Version 0.12.5		
<pre>URL https://david-barnett.github.io/microViz,</pre>		
https://github.com/david-barnett/microViz		
Description Microbiome data visualization and statistics tools built upon phyloseq.		
License GPL (>= 3)		
Encoding UTF-8		
LazyData true		
Roxygen list(markdown = TRUE)		
RoxygenNote 7.3.2		
Depends R (>= $3.5.0$)		
Imports phyloseq, vegan, microbiome, ggiraph, ggplot2, dplyr (>= 1.0.0), rlang (>= 1.0.0), shiny (>= 1.5.0), ComplexHeatmap (>= 2.0.0), circlize, seriation, colorspace, scales, purrr, tibble, tidyr, broom, cowplot		
Suggests DT, ggraph (>= 2.0.0), tidygraph, corncob (>= 0.2.1), ggrepel, GUniFrac, patchwork (>= 1.0.0), ggtext, viridisLite, stringr, forcats, future, future.apply, ggside, skimr, devtools, testthat, vdiffr, knitr, methods, grDevices, rmarkdown, shinytest2		
Repository https://david-barnett.r-universe.dev		
RemoteUrl https://github.com/david-barnett/microViz		
RemoteRef HEAD		
RemoteSha f78c67193640c5e7b81e75db873e2c18a38f2e43		
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add_p	aths Add paths connecting points on a ggplot scatterplot

Description

Useful for tracing a few select individuals over time on an ordination plot. Samples in phyloseq must be arranged in order of timepoint for the path connections to be drawn in the correct order! You can arrange the samples in timepoint order with ps_arrange.

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Usage

```
add_paths(
  ggplot,
  id_var,
  id_values,
  mapping = NULL,
  arrow = grid::arrow(length = grid::unit(2, units = "mm")),
  ...
)
```

Arguments

```
ggplot ggplot scatterplot such as the output of ord_plot

id_var name of variable used to identify grouping of points

id_values values of id_var variable used to identify groups of points to draw

ggplot aesthetics created by aes(), e.g. aes(colour = ?) - group is already set to id_var internally!

arrow arrowhead to add to path, NULL for none

additional arguments passed to geom_path
```

Value

ggplot with added layer with geom_path

```
library(ggplot2)
data("dietswap", package = "microbiome")
# arrange by timepoint first (or whatever your own time variable is)
dietswap %>%
 ps_arrange(timepoint) %>%
 tax_fix() %>%
 tax_transform("clr", rank = "Genus") %>%
 ord_calc(method = "PCA") %>%
 ord_plot(colour = "timepoint", alpha = 0.5, size = 2) %>%
 add_paths(
   id_var = "subject", id_values = c("azl", "byn"),
   mapping = aes(colour = timepoint), linewidth = 1.5
   # size = 1.5 # size instead of linewidth in older ggplot2 versions
 )
# paths do NOT connect points in the correct order without arranging first
dietswap %>%
 tax_fix() %>%
 tax_transform("clr", rank = "Genus") %>%
 ord_calc(method = "PCA") %>%
 ord_plot(colour = "timepoint", alpha = 0.5) %>%
 add_paths(
```

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```
id_var = "subject", id_values = c("azl", "byn"),
mapping = aes(colour = timepoint), linewidth = 1.5
# size = 1.5 # size instead of linewidth in older ggplot2 versions
) +
ggtitle("WRONG PATH ORDER", "use ps_arrange first!")
```

adjacent_side

Simple heatmap helper to get a default adjacent side for another annotation

Description

Simple heatmap helper to get a default adjacent side for another annotation

Usage

```
adjacent_side(side = c("top", "right", "bottom", "left"))
```

Arguments

```
side one of "top", "right", "bottom", or "left"
```

Value

character

Examples

```
adjacent_side("top")
```

anno_cat

Create colored rectangle annotations for categorical data

Description

Similar to anno_simple but with individual boxes!

```
anno_cat(
    x,
    which,
    renamer = identity,
    col = distinct_palette(),
    width = NULL,
    height = NULL,
    box_col = "white",
```

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```
box_1wd = 0.5,
  border_col = NA,
  border_lwd = 1,
  legend = TRUE,
  legend_title = ""
)
```

Arguments

data vector, treated as categorical Х which

Whether it is a column annotation or a row annotation?

renamer function renaming variable values for legend

col colors vector, at least as long as unique(x), optionally named by x levels

width grid unit object or NULL grid unit object or NULL height

box_col colour of boxes around individual cells box_lwd line width of boxes around individual cells

border_col colour of border around all cells border_lwd line width of border around all cells

legend generate legend for this annotation (attached as attribute of heatmap, and not

automatically included in plot)

legend_title title for legend, if drawn

Value

AnnotationFunction

```
library(ComplexHeatmap)
# draw the annotation without a heatmap, you will never normally do this!
vp <- viewport(width = 0.75, height = 0.75)</pre>
grid::grid.newpage()
pushViewport(vp)
cats <- letters[1:4]</pre>
draw(anno_cat(cats, which = "row"))
grid::grid.newpage()
pushViewport(vp)
draw(
 anno_cat(
   x = cats, col = structure(names = cats, 1:4), which = "column",
    box_col = "black", box_lwd = 5
 )
)
```

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```
# developer note #
# list of annotations can be split and ordered (adding NULL makes a list)
# https://jokergoo.github.io/ComplexHeatmap-reference/book/a-list-of-heatmaps.html
# (section #4.8 concatenate-only-the-annotations)
grid::grid.newpage()
pushViewport(vp)
annoList <- rowAnnotation(
    hi = anno_cat(cats, which = "row", border_col = "black")
) +
    NULL
draw(object = annoList, row_split = c(1, 1:3), row_order = 4:1)
pushViewport(viewport(x = 0.6))
draw(anno_cat(cats, "row", legend_title = "abcd") %>% attr("Legend"))
```

anno_cat_legend

Convenience function for generating a legend for anno_cat annotations.

Description

Convenience function for generating a legend for anno_cat annotations.

Usage

```
anno_cat_legend(col, x = NULL, renamer = identity, title = "", ...)
```

Arguments

vector of colors, named by all levels of data (e.g. x) or not named

optional: vector of data to pair with unnamed col or check against named col

renamer function applied to generate labels: from names(col) or levels of x

title title of legend

Arguments passed on to ComplexHeatmap::Legend

labels Labels corresponding to at. If it is not specified, the values of at are taken as labels.

nrow For legend which is represented as grids, nrow controls number of rows of the grids if the grids are arranged into multiple rows.

ncol Similar as nrow, ncol controls number of columns of the grids if the grids are arranged into multiple columns. Note at a same time only one of nrow and ncol can be specified.

by_row Are the legend grids arranged by rows or by columns?

grid_height The height of legend grid. It can also control the height of the continuous legend if it is horizontal.

grid_width The width of legend grid. It can also control the width of the continuous legend if it is vertical.

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gap If legend grids are put into multiple rows or columns, this controls the gap between neighbouring rows or columns, measured as a unit object.

labels_gp Graphic parameters for labels.

labels_rot Text rotation for labels. It should only be used for horizontal continuous legend.

border Color of legend grid borders. It also works for the ticks in the continuous legend.

type Type of legends. The value can be one of grid, points, lines and boxplot.

direction Direction of the legend, vertical or horizontal?

title_position Position of title relative to the legend. topleft, topcenter, leftcenter-rot and lefttop-rot are only for vertical legend and leftcenter, lefttop are only for horizontal legend.

title_gap Gap between title and the legend body.

Value

a ComplexHeatmap Legend class object

Examples

```
grid::grid.newpage()
ComplexHeatmap::draw(
   anno_cat_legend(
   col = c("ibd" = "blue", "nonibd" = "grey90"),
    renamer = toupper, title = "Hi there, I'm a title"
   )
)
```

anno_sample

Helper to specify simple comp_heatmap annotation for other sample data

Description

Use this as an argument to sampleAnnotation(), which itself is used by comp_heatmap() as sample_anno argument.

This creates a vector, which sampleAnnotation() interprets as a simple annotation, so then you set colours and legend parameters for each simple annotation as further arguments in sampleAnnotation.

```
anno_sample(var, fun = identity, data = NULL, samples = NULL)
```

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Arguments

var name of variable to use for annotation data

fun function to transform variable var

data OPTIONAL phyloseq or psExtra, only set this to override use of same data as

in heatmap

samples OPTIONAL selection vector of sample names, only set this if providing data

argument to override default

Value

vector of values

See Also

```
sampleAnnotation()
```

Examples

```
# see `?sampleAnnotation()`
```

anno_sample_cat

Helper to specify comp_heatmap annotation for categorical sample data

Description

Use this as an argument to sampleAnnotation(), which itself is used by comp_heatmap() as sample_anno argument.

```
anno_sample_cat(
  var,
  col = distinct_palette(),
  renamer = identity,
  size = grid::unit(5, "mm"),
  legend = TRUE,
  legend_title = "",
  box_col = "white",
  box_lwd = 0.5,
  border_col = NA,
  border_lwd = 1,
  data = NULL,
  samples = NULL,
  which = NULL,
  ...
)
```

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Arguments

var name of variable to use for annotation data

col colors vector, at least as long as unique(x), optionally named by x levels

renamer function to rename levels of variable var size width or height as a grid unit object

legend generate legend for this annotation (attached as attribute of heatmap, and not

automatically included in plot)

legend_title title for legend, if drawn

box_col colour of boxes around individual cells
box_lwd line width of boxes around individual cells

border_col colour of border around all cells
border_lwd line width of border around all cells

data OPTIONAL phyloseq or psExtra, only set this to override use of same data as

in heatmap

samples OPTIONAL selection vector of sample names, only set this if providing data

argument to override default

which OPTIONAL indicating if it is a 'column' or a 'row' annotation, only set this if

providing data argument to override default

... Arguments passed on to anno_cat

x data vector, treated as categorical width grid unit object or NULL height grid unit object or NULL

Value

vector of values

```
library("ComplexHeatmap")
data("ibd", package = "microViz")
psq <- ibd
samples <- phyloseq::sample_names(psq)

# makes a function that takes data, taxa and which (at minimum)
fun <- anno_sample_cat(var = "ibd")

# manually specify the prevalence barplot function by giving it data etc.
heatmapAnnoFunction <- fun(data = psq, which = "row", samples = samples)

# draw the barplot without a heatmap, you will never normally do this!
vp <- viewport(width = 0.75, height = 0.75)

grid::grid.newpage()
pushViewport(vp)</pre>
```

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```
draw(heatmapAnnoFunction)
# A legend is attached by default to anno_cat() output, let's plot that.
pushViewport(viewport(x = 0.75))
draw(attr(heatmapAnnoFunction, "Legend"))

# change some options and specify the data up front
grid::grid.newpage()
pushViewport(vp)
anno_sample_cat(
   data = psq, var = "DiseaseState", samples = samples, which = "column",
   size = grid::unit(5, "cm"), col = distinct_palette(pal = "kelly")
) %>%
   draw()
```

anno_tax_box

Helper to specify heatmap annotation for showing taxa abundance on boxplot

Description

Use this as an argument to taxAnnotation(), which itself is used by cor_heatmap and comp_heatmap as tax_anno argument.

```
anno_tax_box(
  undetected = 0,
  only_detected = TRUE,
  trans = "compositional",
  zero_replace = 0,
 use_counts = TRUE,
  size = grid::unit(30, "mm"),
 border = TRUE,
  gp = grid::gpar(fill = "#CCCCCC"),
 ylim = NULL,
  extend = 0.05,
  outline = TRUE,
  box_width = 0.6,
  pch = 1,
  pointsize = grid::unit(0.5, "mm"),
  axis = TRUE,
 data = NULL,
  taxa = NULL,
  which = NULL
)
```

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Arguments

undetected the value above which taxa are classed as detected/present in a sample only_detected only plot values for samples where the taxon abundance is > undetected name of transformation suitable for tax transform, or a function calling tax transform, trans and/or tax_scale, (a function must take a phyloseq or psExtra, and return one) zero_replace value for for tax_transform, ignored if trans is a function zero_replace use_counts try to retrieve counts from data object? width or height as a grid unit object size border Wether draw borders of the annotation region? Graphic parameters for the boxes. The length of the graphic parameters should gp be one or the number of observations. ylim Data ranges. The extension to both side of ylim. The value is a percent value corresponding extend to ylim[2] - ylim[1]. outline Whether draw outline of boxplots? box_width Relative width of boxes. The value should be smaller than one. Point style. pch pointsize size of outlier points, as grid::unit() object Whether to add axis? axis Arguments passed on to ComplexHeatmap::anno_boxplot axis_param parameters for controlling axis. See default_axis_param for all possible settings and default parameters.

data

OPTIONAL phyloseq or psExtra, only set this to override use of same data as

in heatmap

taxa OPTIONAL selection vector of taxa (names, numbers or logical), only set this

if providing data argument to override default

which OPTIONAL indicating if it is a 'column' or a 'row' annotation, only set this if

providing data argument to override default

Value

function or ComplexHeatmap AnnotationFunction object

```
library("ComplexHeatmap")
data("ibd", package = "microViz")
psq <- tax_filter(ibd, min_prevalence = 5)
psq <- tax_mutate(psq, Species = NULL)
psq <- tax_fix(psq)
psq <- tax_agg(psq, rank = "Family")
taxa <- tax_top(psq, n = 15, rank = "Family")
# makes a function that takes data, taxa and which (at minimum)</pre>
```

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```
fun <- anno_tax_box()</pre>
# manually specify the prevalence barplot function by giving it data etc.
heatmapAnnoFunction <- fun(data = psq, which = "column", taxa = taxa)
# draw the barplot without a heatmap, you will never normally do this!
vp <- viewport(width = 0.75, height = 0.75)</pre>
grid.newpage()
pushViewport(vp)
draw(heatmapAnnoFunction)
# let's change some style options and specify the data up front
grid::grid.newpage()
pushViewport(vp)
draw(anno_tax_box(
 data = psq, taxa = taxa, which = "row", pointsize = grid::unit(1, "mm"),
 gp = grid::gpar(fill = "red"), border = FALSE, box_width = 0.2
))
# clear drawings
grid::grid.newpage()
```

anno_tax_density

Helper to specify heatmap annotation for showing taxa abundance density plot

Description

Use this as an argument to taxAnnotation(), which itself is used by cor_heatmap and comp_heatmap as tax_anno argument.

```
anno_tax_density(
  undetected = 0,
  only_detected = TRUE,
  trans = "log10p",
  zero_replace = 0,
  use_counts = TRUE,
  size = grid::unit(30, "mm"),
  type = c("lines", "violin", "heatmap"),
  xlim = NULL,
  heatmap_colors = c("white", "forestgreen"),
  joyplot_scale = 1.5,
  border = TRUE,
  gp = grid::gpar(fill = "lightgrey"),
  axis = TRUE,
  data = NULL,
  taxa = NULL,
  which = NULL
)
```

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Arguments

undetected the value above which taxa are classed as detected/present in a sample only_detected only plot values for samples where the taxon abundance is > undetected name of transformation suitable for tax transform, or a function calling tax transform, trans and/or tax_scale, (a function must take a phyloseq or psExtra, and return one) zero_replace zero_replace value for for tax_transform, ignored if trans is a function try to retrieve counts from data object? use_counts width or height as a grid unit object size Type of graphics to represent density distribution. "lines" for normal density type plot; "violine" for violin plot and "heatmap" for heatmap visualization of density distribution. xlim Range on x-axis. heatmap_colors A vector of colors for interpolating density values. Relative height of density distribution. A value higher than 1 increases the height joyplot_scale of the density distribution and the plot will represented as so-called "joyplot". border Wether draw borders of the annotation region? Graphic parameters for the boxes. The length of the graphic parameters should gp be one or the number of observations. Whether to add axis? axis Arguments passed on to ComplexHeatmap::anno_density axis_param parameters for controlling axis. See default_axis_param for all possible settings and default parameters. data OPTIONAL phyloseq or psExtra, only set this to override use of same data as in heatmap taxa OPTIONAL selection vector of taxa (names, numbers or logical), only set this if providing data argument to override default which OPTIONAL indicating if it is a 'column' or a 'row' annotation, only set this if providing data argument to override default

Value

function or ComplexHeatmap AnnotationFunction object

```
library("ComplexHeatmap")
data("ibd", package = "microViz")
psq <- tax_filter(ibd, min_prevalence = 5)
psq <- tax_mutate(psq, Species = NULL)
psq <- tax_fix(psq)
psq <- tax_agg(psq, rank = "Family")
taxa <- tax_top(psq, n = 15, rank = "Family")
# makes a function that takes data, taxa and which (at minimum)
fun <- anno_tax_density()</pre>
```

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```
# manually specify the density plot function by giving it data etc.
heatmapAnnoFunction <- fun(data = psq, which = "column", taxa = taxa)
# draw the density plot without a heatmap, you will never normally do this!
vp <- viewport(width = 0.75, height = 0.75)</pre>
grid.newpage()
pushViewport(vp)
draw(heatmapAnnoFunction)
# let's change some style options and specify the data up front
grid.newpage()
pushViewport(vp)
draw(anno_tax_density(
 data = psq, taxa = taxa, which = "row",
 gp = grid::gpar(fill = "red"), border = FALSE
))
# heatmap type, with alternative transformation and axis_param
grid.newpage()
pushViewport(vp)
draw(anno_tax_density(
 data = psq, taxa = taxa, which = "row", type = "heatmap",
 trans = "log2", zero_replace = "halfmin", axis_param = list(labels_rot = 0)
))
grid.newpage()
```

anno_tax_prev

Helper to specify heatmap annotation for showing taxa prevalence as barplot

Description

Use this as an argument to taxAnnotation(), which itself is used by cor_heatmap and comp_heatmap as tax_anno argument.

```
anno_tax_prev(
  undetected = 0,
  use_counts = TRUE,
  size = grid::unit(20, "mm"),
  baseline = 0,
  border = TRUE,
  bar_width = 0.6,
  gp = grid::gpar(fill = "#CCCCCC"),
  ylim = NULL,
  extend = 0.05,
  axis = TRUE,
```

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```
data = NULL,
taxa = NULL,
which = NULL
```

Arguments

undetected	the value above which taxa are classed as detected/present in a sample
use_counts	try to retrieve counts from data object?
size	width or height as a grid unit object
baseline	baseline of bars. The value should be "min" or "max", or a numeric value. It is enforced to be zero for stacked barplots.
border	Wether draw borders of the annotation region?
bar_width	Relative width of the bars. The value should be smaller than one.
gp	Graphic parameters for bars. The length of each graphic parameter can be 1, length of x if x is a vector, or number of columns of x is x is a matrix.
ylim	Data ranges. By default it is $range(x)$ if x is a vector, or $range(rowSums(x))$ if x is a matrix.
extend	The extension to both side of ylim. The value is a percent value corresponding to ylim[2] - ylim[1].
axis	Whether to add axis?
	Arguments passed on to ComplexHeatmap::anno_barplot
	axis_param parameters for controlling axis. See default_axis_param for all possible settings and default parameters.
data	OPTIONAL phyloseq or psExtra, only set this to override use of same data as in heatmap
taxa	OPTIONAL selection vector of taxa (names, numbers or logical), only set this if providing data argument to override default
which	OPTIONAL indicating if it is a 'column' or a 'row' annotation, only set this if providing data argument to override default

Value

function or ComplexHeatmap AnnotationFunction object

```
library("ComplexHeatmap")
data("ibd", package = "microViz")
psq <- tax_filter(ibd, min_prevalence = 5)
psq <- tax_mutate(psq, Species = NULL)
psq <- tax_fix(psq)
psq <- tax_agg(psq, rank = "Family")
taxa <- tax_top(psq, n = 15, rank = "Family")</pre>
```

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```
# makes a function that takes data, taxa and which (at minimum)
fun <- anno_tax_prev()</pre>
# manually specify the prevalence barplot function by giving it data etc.
heatmapAnnoFunction <- fun(data = psq, which = "row", taxa = taxa)</pre>
# draw the barplot without a heatmap, you will never normally do this!
vp <- viewport(width = 0.75, height = 0.75)</pre>
grid::grid.newpage()
pushViewport(vp)
draw(heatmapAnnoFunction)
# let's change some style options and specify the data up front
grid::grid.newpage()
pushViewport(vp)
anno_tax_prev(
 data = psq, taxa = taxa, which = "column",
 gp = grid::gpar(fill = "red", lwd = 3, alpha = 0.5),
 border = FALSE, bar_width = 1
) %>%
 draw()
# clear drawings
grid::grid.newpage()
```

anno_var_box

Helper to specify heatmap annotation for variable distribution boxplots

Description

Use this as an argument to varAnnotation(), which itself is used by cor_heatmap as var_anno() argument.

```
anno_var_box(
  fun = identity,
  size = grid::unit(30, "mm"),
  border = TRUE,
  gp = grid::gpar(fill = "#CCCCCC"),
  ylim = NULL,
  extend = 0.05,
  outline = TRUE,
  box_width = 0.6,
  pch = 1,
  pointsize = grid::unit(0.5, "mm"),
```

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```
axis = TRUE,
...,
data = NULL,
vars = NULL,
which = NULL
```

Arguments

fun function applied to all variables, with apply() size width or height as a grid unit object border Wether draw borders of the annotation region? Graphic parameters for the boxes. The length of the graphic parameters should gp be one or the number of observations. ylim Data ranges. The extension to both side of ylim. The value is a percent value corresponding extend to ylim[2] - ylim[1]. outline Whether draw outline of boxplots? box_width Relative width of boxes. The value should be smaller than one. pch Point style. pointsize size of outlier points, as grid::unit() object axis Whether to add axis? Arguments passed on to ComplexHeatmap::anno_boxplot axis_param parameters for controlling axis. See default_axis_param for all possible settings and default parameters. data OPTIONAL phyloseq or psExtra, only set this to override use of same data as in heatmap OPTIONAL selection vector of variable names, only set this if providing data vars argument to override default

OPTIONAL indicating if it is a 'column' or a 'row' annotation, only set this if

Value

function or ComplexHeatmap AnnotationFunction object

Examples

which

```
library(ComplexHeatmap)
set.seed(123)
fakeData <- as.data.frame.matrix(matrix(rnorm(500, 10, 3), ncol = 10))
names(fakeData) <- paste0("var_", 1:10)

# draw the boxplot without a heatmap, you will never normally do this!
vp <- viewport(width = 0.75, height = 0.75)</pre>
```

providing data argument to override default

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```
grid.newpage()
pushViewport(vp)
draw(
   anno_var_box(data = fakeData, vars = names(fakeData), which = "column")
)
grid.newpage()
pushViewport(vp)
draw(
   anno_var_box(
    data = fakeData, fun = function(x) log(x + 1),
    vars = rev(names(fakeData)),
    which = "row"
   )
)
```

anno_var_density

Helper to specify heatmap annotation for variable distribution density plot

Description

Use this as an argument to varAnnotation(), which itself is used by cor_heatmap var_anno argument.

Usage

```
anno_var_density(
  fun = identity,
  size = grid::unit(30, "mm"),
  type = c("lines", "violin", "heatmap"),
  xlim = NULL,
  heatmap_colors = c("white", "forestgreen"),
  joyplot_scale = 1.5,
  border = TRUE,
  gp = grid::gpar(fill = "lightgrey"),
  axis = TRUE,
  ...,
  data = NULL,
  vars = NULL,
  which = NULL
)
```

Arguments

fun function applied to all variables, with apply() size width or height as a grid unit object

20 anno_var_density

type	Type of graphics to represent density distribution. "lines" for normal density plot; "violine" for violin plot and "heatmap" for heatmap visualization of density distribution.
xlim	Range on x-axis.
heatmap_colors	A vector of colors for interpolating density values.
joyplot_scale	Relative height of density distribution. A value higher than 1 increases the height of the density distribution and the plot will represented as so-called "joyplot".
border	Wether draw borders of the annotation region?
gp	Graphic parameters for the boxes. The length of the graphic parameters should be one or the number of observations.
axis	Whether to add axis?
	Arguments passed on to ComplexHeatmap::anno_density
	axis_param parameters for controlling axis. See default_axis_param for all possible settings and default parameters.
data	OPTIONAL phyloseq or psExtra, only set this to override use of same data as in heatmap
vars	OPTIONAL selection vector of variable names, only set this if providing data argument to override default
which	OPTIONAL indicating if it is a 'column' or a 'row' annotation, only set this if providing data argument to override default

Value

function or ComplexHeatmap AnnotationFunction object

```
library(ComplexHeatmap)
set.seed(123)
fakeData <- as.data.frame.matrix(matrix(rnorm(500, 10, 3), ncol = 10))</pre>
names(fakeData) <- paste0("var_", 1:10)</pre>
# draw the plots without a heatmap, you will never normally do this!
vp <- viewport(width = 0.75, height = 0.75)</pre>
grid.newpage()
pushViewport(vp)
  anno_var_density(data = fakeData, vars = names(fakeData), which = "row")
grid.newpage()
pushViewport(vp)
draw(
  anno_var_density(
    data = fakeData, fun = function(x) log(x + 1),
    vars = rev(names(fakeData)), type = "heatmap",
    which = "column"
 )
)
```

anno_var_hist 21

anno_var_hist	Helper to specify heatmap annotation for variable distribution histograms

Description

Use this as an argument to varAnnotation(), which itself is used by cor_heatmap var_anno argument.

Usage

```
anno_var_hist(
  fun = identity,
  size = grid::unit(30, "mm"),
  n_breaks = 11,
  border = FALSE,
  gp = grid::gpar(fill = "#CCCCCC"),
  axis = TRUE,
  ...,
  data = NULL,
  vars = NULL,
  which = NULL
)
```

Arguments

fun	function applied to all variables, with apply()
size	width or height as a grid unit object
n_breaks	number of breaks
border	Wether draw borders of the annotation region?
gp	Graphic parameters for the boxes. The length of the graphic parameters should be one or the number of observations.
axis	Whether to add axis?
	Arguments passed on to ComplexHeatmap::anno_density
	axis_param parameters for controlling axis. See default_axis_param for all possible settings and default parameters.
data	OPTIONAL phyloseq or psExtra, only set this to override use of same data as in heatmap
vars	OPTIONAL selection vector of variable names, only set this if providing data argument to override default
which	OPTIONAL indicating if it is a 'column' or a 'row' annotation, only set this if providing data argument to override default

Value

function or ComplexHeatmap AnnotationFunction object

Examples

```
library(ComplexHeatmap)
set.seed(123)
fakeData <- as.data.frame.matrix(matrix(rnorm(500, 10, 3), ncol = 10))</pre>
names(fakeData) <- paste0("var_", 1:10)</pre>
# draw the histograms without a heatmap, you will never normally do this!
vp <- viewport(width = 0.75, height = 0.75)</pre>
grid.newpage()
pushViewport(vp)
draw(
  anno_var_hist(data = fakeData, vars = names(fakeData), which = "row")
)
grid.newpage()
pushViewport(vp)
draw(
  anno_var_hist(
    data = fakeData, fun = sqrt,
    vars = rev(names(fakeData)), n_breaks = 5,
    which = "column", gp = grid::gpar(fill = 2:6, lwd = c(0.9, 2.5))
)
```

comp_barplot

Plot (grouped and ordered) compositional barplots

Description

Stacked barplots showing composition of phyloseq samples for a specified number of coloured taxa. Normally your phyloseq object should contain counts data, as by default comp_barplot() performs the "compositional" taxa transformation for you, and requires count input for some sample_order methods!

```
comp_barplot(
   ps,
   tax_level,
   n_taxa = 8,
   tax_order = sum,
   merge_other = TRUE,
   taxon_renamer = function(x) identity(x),
   sample_order = "bray",
   order_with_all_taxa = FALSE,
   label = "SAMPLE",
   group_by = NA,
   facet_by = NA,
```

```
bar_width = 1,
bar_outline_colour = "grey5",
bar_outline_width = 0.1,
palette = distinct_palette(n_taxa),
tax_transform_for_ordering = "identity",
tax_transform_for_plot = "compositional",
seriate_method = "OLO_ward",
keep_all_vars = TRUE,
interactive = FALSE,
max_taxa = 10000,
other_name = "Other",
x = "SAMPLE",
counts_warn = TRUE,
...
)
```

Arguments

palette

ps phyloseq object taxonomic aggregation level (from rank_names(ps)) tax_level how many taxa to show distinct colours for (all others grouped into "Other") n taxa order of taxa within the bars, either a function for tax_sort (e.g. sum), or a vector tax_order of (all) taxa names at tax_level to set order manually merge_other if FALSE, taxa coloured/filled as "other" remain distinct, and so can have bar outlines drawn around them taxon_renamer function that takes taxon names and returns modified names for legend sample_order vector of sample names; or any distance measure in dist_calc that doesn't require phylogenetic tree; or "asis" for the current order as is returned by phyloseq::sample_names(ps) order_with_all_taxa if TRUE, this will always use all taxa (not just the top n_taxa) to calculate any distances needed for sample ordering name of variable to use for labelling samples, or "SAMPLE" for sample names label splits dataset by this variable (must be categorical) group_by • resulting in a list of plots, one for each level of the group_by variable. facet_by facets plots by this variable (must be categorical). If group_by is also set the faceting will occur separately in the plot for each group. bar_width default 1 avoids random gapping otherwise seen with many samples (set to less than 1 to introduce gaps between samples) bar_outline_colour line colour separating taxa and samples (use NA for no outlines) bar_outline_width width of line separating taxa and samples (for no outlines set bar_outline_colour = NA)

palette for taxa fill colours

tax_transform_for_ordering

transformation of taxa values used before ordering samples by similarity

tax_transform_for_plot

default "compositional" draws proportions of total counts per sample, but you could reasonably use another transformation, e.g. "identity", if you have truly

quantitative microbiome profiling data

seriate_method name of any ordering method suitable for distance matrices (see ?seriation::seriate)

keep_all_vars FALSE may speed up internal melting with ps_melt for large phyloseq objects

but TRUE is required for some post-hoc plot customisation

interactive creates plot suitable for use with ggiraph

max_taxa maximum distinct taxa groups to show (only really useful for limiting complex-

ity of interactive plots e.g. within ord_explore)

other_name name for other taxa after N

name of variable to use as x aesthetic: it probably only makes sense to change

this when also using facets (check only one sample is represented per bar!)

counts_warn should a warning be issued if counts are unavailable? TRUE, FALSE, or "error"

(passed to ps_get)

... extra arguments passed to facet_wrap() (if facet_by is not NA)

Details

• sample_order: Either specify a list of sample names to order manually, or the bars/samples can/will be sorted by similarity, according to a specified distance measure (default 'bray'-curtis).

- seriate_method specifies a seriation/ordering algorithm (default Ward hierarchical clustering with optimal leaf ordering, see seriation::list_seriation_methods())
- group_by: You can group the samples on distinct plots by levels of a variable in the phyloseq object. The list of ggplots produced can be arranged flexibly with the patchwork package functions. If you want to group by several variables you can create an interaction variable with interaction(var1, var2) in the phyloseq sample_data BEFORE using comp_barplot.
- facet_by can allow faceting of your plot(s) by a grouping variable. Using this approach is less flexible than using group_by but means you don't have to arrange a list of plots yourself like with the group_by argument. Using facet_by is equivalent to adding a call to facet_wrap(facets = facet_by, scales = "free") to your plot(s). Calling facet_wrap() yourself is itself a more flexible option as you can add other arguments like the number of rows etc. However you must use keep_all_vars = TRUE if you will add faceting manually.
- bar_width: No gaps between bars, unless you want them (decrease width argument to add gaps between bars).
- bar_outline_colour: Bar outlines default to "grey5" for almost black outlines. Use NA if you
 don't want outlines.
- merge_other: controls whether bar outlines can be drawn around individual (lower abundance) taxa that are grouped in "other" category. If you want to see the diversity of taxa in "other" use merge_taxa = FALSE, or use TRUE if you prefer the cleaner merged look
- palette: Default colouring is consistent across multiple plots if created with the group_by argument, and the defaults scheme retains the colouring of the most abundant taxa irrespective of n_taxa

Value

ggplot or list of harmonised ggplots

```
library(ggplot2)
data(dietswap, package = "microbiome")
# illustrative simple customised example
dietswap %>%
 ps_filter(timepoint == 1) %>%
 comp_barplot(
    tax_level = "Family", n_taxa = 8,
   bar_outline_colour = NA,
   sample_order = "bray",
   bar_width = 0.7,
   taxon_renamer = toupper
 ) + coord_flip()
# change colour palette with the distinct_palette() function
# remember to set the number of colours to the same as n_taxa argument!
dietswap %>%
 ps_filter(timepoint == 1) %>%
 comp_barplot(
   tax_level = "Family", n_taxa = 8,
   bar_outline_colour = NA,
   sample_order = "bray",
   bar_width = 0.7
   palette = distinct_palette(8, pal = "kelly"),
    taxon_renamer = toupper
 ) + coord_flip()
# Order samples by the value of one of more sample_data variables.
# Use ps_arrange and set sample_order = "default" in comp_barplot.
# ps_mutate is also used here to create an informative variable for axis labelling
dietswap %>%
 ps_mutate(subject_timepoint = interaction(subject, timepoint)) %>%
 ps_filter(nationality == "AAM", group == "DI", sex == "female") %>%
 ps_arrange(desc(subject), desc(timepoint)) %>%
 comp_barplot(
    tax_level = "Genus", n_taxa = 12,
    sample_order = "default",
   bar_width = 0.7,
   bar_outline_colour = "black",
   order_with_all_taxa = TRUE,
    label = "subject_timepoint"
 ) + coord_flip()
# Order taxa differently:
# By default, taxa are ordered by total sum across all samples
# You can set a different function for calculating the order, e.g. median
dietswap %>%
```

```
ps_filter(timepoint == 1) %>%
 comp_barplot(tax_level = "Genus", tax_order = median) +
 coord_flip()
# Or you can set the taxa order up front, with tax_sort() and use it as is
dietswap %>%
 ps_filter(timepoint == 1) %>%
 tax_sort(at = "Genus", by = sum) %>%
 comp_barplot(tax_level = "Genus", tax_order = "asis") +
 coord_flip()
# how many taxa are in those light grey "other" bars?
# set merge_other to find out (& remember to set a bar_outline_colour)
dietswap %>%
 ps_filter(timepoint == 1) %>%
 comp_barplot(
   tax_level = "Genus", n_taxa = 12, merge_other = FALSE, bar_outline_colour = "grey50",
 ) +
 coord_flip()
# Often to compare groups, average compositions are presented
p1 <- phyloseq::merge_samples(dietswap, group = "group") %>%
 comp_barplot(
    tax_level = "Genus", n_taxa = 12,
    sample_order = c("ED", "HE", "DI"),
   bar_width = 0.8
 coord_flip() + labs(x = NULL, y = NULL)
р1
# However that "group-averaging" approach hides a lot of within-group variation
p2 <- comp_barplot(dietswap,</pre>
 tax_level = "Genus", n_taxa = 12, group_by = "group",
 sample_order = "euclidean", bar_outline_colour = NA
) %>%
 patchwork::wrap_plots(nrow = 3, guides = "collect") &
 coord_flip() & labs(x = NULL, y = NULL) &
 theme(axis.text.y = element_blank(), axis.ticks.y = element_blank())
p2
# Only from p2 you can see that the apparently higher average relative abundance
# of Oscillospira in group DI is probably driven largely by a subgroup
# of DI samples with relatively high Oscillospira.
# make a list of 2 harmonised composition plots (grouped by sex)
p <- comp_barplot(dietswap,</pre>
 n_taxa = 15, tax_level = "Genus",
 bar_outline_colour = "black", merge_other = TRUE,
 sample_order = "aitchison", group_by = "sex"
)
# plot them side by side with patchwork package
```

```
patch <- patchwork::wrap_plots(p, ncol = 2, guides = "collect")
patch & coord_flip() # make bars in all plots horizontal (note: use & instead of +)

# beautifying tweak #
# modify one plot in place (flip the order of the samples in the 2nd plot)
# notice that the scaling is for the x-axis
# (that's because coord_flip is used afterwards when displaying the plots
patch[[2]] <- patch[[2]] + scale_x_discrete(limits = rev)
# Explainer: rev() function takes current limits and reverses them.
# You could also pass a completely arbitrary order, naming all samples

# you can theme all plots with the & operator
patch & coord_flip() &
    theme(axis.text.y = element_text(size = 5), legend.text = element_text(size = 6))
# See https://patchwork.data-imaginist.com/index.html</pre>
```

comp_heatmap

Draw heatmap of microbiome composition across samples

Description

Heatmap made with ComplexHeatmap::Heatmap(), with optional annotation of taxa prevalence/abundance, and/or other sample data.

Transform your data with tax_transform() prior to plotting (and/or scale with tax_scale()).

See the heatmaps vignette for more examples of use.

Plotting "compositional" data can give an idea of the dominant taxa in each sample. Plotting some form of log or clr transformed (or scaled) microbial features can highlight other patterns.

The data will be ordered via your selected seriation methods and distances on either the transformed data (default) or the original count data (or with any other transformation).

Any cell numbers printed can be transformed independently of the colour scheme, and do not affect ordering.

```
comp_heatmap(
  data,
  taxa = NA,
  taxa_side = "right",
  tax_anno = NULL,
  taxon_renamer = identity,
  samples = NA,
  sample_side = adjacent_side(taxa_side),
  sample_anno = NULL,
  sample_names_show = FALSE,
  colors = heat_palette(palette = "Rocket", rev = TRUE),
  numbers = NULL,
```

```
sample_seriation = "OLO_ward",
  sample_ser_dist = "euclidean",
 sample_ser_counts = !sample_ser_dist %in% c("euclidean", "maximum", "manhattan",
    "canberra", "binary"),
  sample_ser_trans = NULL,
  tax_seriation = "OLO_ward",
  tax_ser_dist = "euclidean",
  tax_ser_counts = FALSE,
  tax_ser_trans = NULL,
  numbers_trans = NULL,
  numbers_zero_replace = 0,
  numbers_use_counts = TRUE,
  grid_col = "white",
  grid_lwd = 0.5,
  name = "Abd.",
  anno_tax = NULL,
)
```

Arguments

data phyloseq or phyloseq extra taxa list of taxa to include, or NA for all "top"/"right"/"bottom"/"left": controls heatmap orientation and where any antaxa_side notations specified in tax_anno are placed tax_anno NULL or annotation function for taxa: taxAnnotation() output. function to rename taxa before plotting taxon_renamer samples list of samples to include on plot sample_side which side to show any sample annotation on, must be adjacent to taxa_side sample_anno NULL or annotation function for samples: sampleAnnotation() output. sample_names_show show sample names? (you can control side and rotation of names with other ComplexHeatmap::Heatmap arguments) colors output of heat_palette() to set heatmap fill color scheme output of heat_numbers() to draw numbers on heatmap cells numbers sample_seriation name of method used to order the samples (from seriation::seriate) sample_ser_dist name of distance to use with sample seriation method (if needed) sample_ser_counts insist on using count data for sample seriation? sample_ser_trans function for transformation of data used for sample seriation (such as a call to tax_transform()) name of method used to order the taxa (from seriation::seriate) tax_seriation

```
tax_ser_dist
                  name of distance to use with tax_seriation method (if needed)
tax_ser_counts insist on using count data for taxa seriation?
                 function for transformation of data used for taxa seriation (such as a call to
tax_ser_trans
                  tax_transform())
numbers_trans
                 name of tax_transform transformation, or a function for transformation of data
                 used for drawing numbers on cells
numbers_zero_replace
                 zero replacement method used if named transformation given to number_trans
numbers_use_counts
                 insist on using count data for number drawing? (if TRUE, any numbers_trans
                  transformation would be applied to count data)
grid_col
                 colour of gridlines, or NA for none
grid_lwd
                  width of gridlines
name
                 used as legend title (colourbar)
anno_tax
                 DEPRECATED: optional annotation of taxa distributions: tax_anno() list out-
                 put, or a pre-made ComplexHeatmap HeatmapAnnotation
                  Arguments passed on to ComplexHeatmap::Heatmap
                  row_dend_side Should the row dendrogram be put on the left or right of the
                      heatmap?
                  row_dend_width Width of the row dendrogram, should be a unit object.
                  show_row_dend Whether show row dendrogram?
                  row_dend_gp Graphic parameters for the dendrogram segments. If users al-
                      ready provide a dendrogram object with edges rendered, this argument will
                      be ignored.
                  show_row_names Whether show row names.
                  row_names_gp Graphic parameters for row names.
                  row names rot Rotation of row names.
                  row_names_centered Should row names put centered?
```

See Also

cor_heatmap()

```
library(dplyr)
data("dietswap", package = "microbiome")
# create a couple of numerical variables to use
psq <- dietswap %>%
  ps_mutate(
    weight = recode(bmi_group, obese = 3, overweight = 2, lean = 1),
    female = if_else(sex == "female", true = 1, false = 0),
    african = if_else(nationality == "AFR", true = 1, false = 0)
)
psq <- tax_filter(psq, min_prevalence = 1 / 10, min_sample_abundance = 1 / 10)
psq <- tax_agg(psq, "Genus")</pre>
```

```
# randomly select 20 taxa from the 40 top taxa, and 40 random samples
set.seed(123)
taxa <- sample(tax_top(psq, n = 40), size = 20)
samples \leftarrow sample(1:122, size = 40)
comp_heatmap(data = psq, taxa = taxa, samples = samples)
# transforming taxon abundances #
# NOTE: if you plan on transforming taxa (e.g. to compositional data or clr)
# but only want to plot a subset of the taxa (e.g. most abundant)
# you should NOT subset the original phyloseq before transformation!
# Instead, choose the subset of taxa plotted with:
# Note 2, choose a symmetrical palette for clr-transformed data
psq %>%
 tax_transform("clr", zero_replace = "halfmin") %>%
 comp_heatmap(
   taxa = taxa, samples = samples, colors = heat_palette(sym = TRUE)
# Almost all the taxa have high values (>> 0) because they are a highly
# abundant subset taken after clr transformation was calculated on all taxa
# See how just taking the first 30 taxa from the dataset gives more balance
psq %>%
 tax_transform("clr", zero_replace = "halfmin") %>%
 comp_heatmap(
   taxa = 1:30, samples = samples, colors = heat_palette(sym = TRUE)
 )
# annotating taxa #
# Notes:
# - Unlike cor_heatmap, taxa are not annotated by default
# - Detection threshold set to 50 as HITchip example data seems to have background noise
comp_heatmap(
 data = psq, taxa = taxa, samples = samples,
 tax_anno = taxAnnotation(Prev = anno_tax_prev(undetected = 50))
# annotating samples #
htmp <- psq %>%
 tax_transform("clr", zero_replace = "halfmin") %>%
 comp_heatmap(
   taxa = taxa, samples = samples, colors = heat_palette(sym = TRUE),
   sample_anno = sampleAnnotation(
     Nation. = anno_sample_cat("nationality", legend_title = "Nation.")
    )
```

```
)
htmp

# legends from `anno_sample_cat()` are stored as an attribute of the Heatmap
ComplexHeatmap::draw(
  object = htmp,
    annotation_legend_list = attr(htmp, "AnnoLegends"), merge_legends = TRUE
)
```

cor_heatmap

Microbe-to-sample-data correlation heatmap

Description

Plot correlations between (transformed) microbial abundances and (selected) numeric-like sample_data variables from a phyloseq object.

Lots of customisation options available through the listed arguments, and you can pass any other argument from ComplexHeatmap::Heatmap() too.

```
cor_heatmap(
  data,
  taxa = NA,
  tax_anno = taxAnnotation(Prev. = anno_tax_prev(), Abun. = anno_tax_box()),
  taxon_renamer = identity,
  vars = NA,
  var_anno = NULL,
  cor = c("pearson", "kendall", "spearman"),
  cor_use = "everything",
  colors = heat_palette(palette = "Blue-Red 2", sym = TRUE),
  numbers = heat_numbers(decimals = 1, col = "black", fontface = "plain"),
  taxa_side = "right",
  vars_side = adjacent_side(taxa_side),
  seriation_method = "OLO_ward",
  seriation_dist = "euclidean",
  seriation_method_col = seriation_method,
  seriation_dist_col = seriation_dist,
  var_fun = "identity",
  grid_col = "white",
  grid_lwd = 0.5,
  anno_tax = NULL,
 anno_vars = NULL,
)
```

Arguments

data phyloseq or phyloseq extra

taxa list of taxa to include, or NA for all

tax_anno NULL or annotation function for taxa: taxAnnotation() output.

taxon_renamer function to rename taxa before plotting vars selection of variable names from sample_data

var_anno NULL or annotation function for variables: varAnnotation() output.

cor correlation coefficient. pearson/kendall/spearman, can be abbreviated (used as

legend title)

cor_use passed to cor(use = cor_use)

colors output of heat_palette() to set heatmap fill color scheme output of heat_numbers() to draw numbers on heatmap cells

taxa_side "top"/"right"/"bottom"/"left": controls heatmap orientation and where any an-

notations specified in tax_anno are placed

vars_side which side to place any variable annotations specified in var_anno, must be an

adjacent side to taxa_side

seriation_method

method to order the rows (in seriation::seriate)

seriation_dist distance to use in seriation_method (if needed)

seriation_method_col

method to order the columns (in seriation::seriate)

seriation_dist_col

distance to use in seriation_method_col (if needed)

var_fun a function (or name of) to be applied to columns of a matrix of vars before

correlating (but not used in any variable annotations)

grid_col colour of gridlines, or NA for none

grid_lwd width of gridlines

anno_tax DEPRECATED: optional annotation of taxa distributions: tax anno() list out-

put, or a pre-made ComplexHeatmap HeatmapAnnotation

anno_vars DEPRECATED: use var_anno argument instead. Optional annotation of vari-

able distributions: var_anno() list output, or a pre-made ComplexHeatmap Heatma-

pAnnotation

... Arguments passed on to ComplexHeatmap::Heatmap

row_dend_width Width of the row dendrogram, should be a unit object.

show_row_dend Whether show row dendrogram?

row_dend_gp Graphic parameters for the dendrogram segments. If users already provide a dendrogram object with edges rendered, this argument will be ignored

be ignored.

show_row_names Whether show row names.

row_names_gp Graphic parameters for row names.

row_names_rot Rotation of row names.

row_names_centered Should row names put centered? show_heatmap_legend Whether show heatmap legend?

Details

Using a data.frame for the data argument is also possible, in which case the (selected) numeric-like variables will be correlated with each other, and all arguments relating to taxa will be ignored.

See Also

```
taxAnnotation() varAnnotation()
comp_heatmap()
ComplexHeatmap::Heatmap()
```

```
library(dplyr)
data("dietswap", package = "microbiome")
# create a couple of numerical variables to use
psq <- dietswap %>%
  ps_mutate(
    weight = recode(bmi_group, obese = 3, overweight = 2, lean = 1),
    female = if_else(sex == "female", true = 1, false = 0),
    african = if_else(nationality == "AFR", true = 1, false = 0)
psq <- tax_filter(psq, min_prevalence = 1 / 10, min_sample_abundance = 1 / 10)</pre>
psq <- tax_agg(psq, "Genus")</pre>
# randomly select 20 taxa from the 50 most abundant taxa
set.seed(123)
taxa \leftarrow sample(tax_top(psq, n = 50), size = 20)
# NOTE: detection threshold set to 50 as HITchip example data seems to have background noise
ud <- 50
# make simple correlation heatmap with all numeric-like variables
cor_heatmap(
  data = psq, taxa = taxa,
  tax_anno = taxAnnotation(
    Prv. = anno_tax_prev(undetected = ud),
    Abd. = anno_tax_box(undetected = ud)
  )
)
# You can create an annotation object separately in advance
taxAnno <- taxAnnotation(</pre>
  Prv. = anno_tax_prev(undetected = ud), Abd. = anno_tax_box(undetected = ud)
class(taxAnno) # "function"
# You can select which numeric-like variables to correlate taxa with
cor_heatmap(
  psq, taxa,
  vars = c("african", "female", "weight"), tax_anno = taxAnno
```

```
)
# Also you can choose alternative correlation measures
cor_heatmap(psq, taxa, cor = "spearman", tax_anno = taxAnno)
# Annotating variables is possible, and easy with varAnnotation()
cor_heatmap(
 data = psq, taxa = taxa, tax_anno = taxAnno,
 var_anno = varAnnotation(Val. = anno_var_box(size = grid::unit(2, "cm")))
# you can transform the variables before correlating by var_fun
# notice this does not affect the data used for annotations
cor_heatmap(
 data = psq, taxa = taxa, tax_anno = taxAnno, var_fun = "exp",
 var_anno = varAnnotation(Val. = anno_var_box(size = grid::unit(2, "cm")))
)
# other and multiple annotations
cor_heatmap(
 data = psq, taxa = taxa[1:10], vars = c("african", "weight", "female"),
 tax_anno = taxAnno,
 var_anno = varAnnotation(
   value = anno_var_hist(size = grid::unit(15, "mm")),
   log10p = anno_var_box(function(x) log10(x + 1))
 )
)
# make the same heatmap, but rotated
cor_heatmap(
 data = psq, taxa = taxa[1:10], vars = c("african", "weight", "female"),
 tax_anno = taxAnno, taxa_side = "top",
 var_anno = varAnnotation(
   value = anno_var_hist(size = grid::unit(15, "mm")),
   log10p = anno_var_box(function(x) log10(x + 1))
 )
)
# You can change the colour scheme used, using heat_palette()
cor_heatmap(
 data = psq, taxa = taxa, tax_anno = taxAnno,
 colors = heat_palette("Green-Orange", rev = TRUE, sym = TRUE)
)
# You can hide or change the style of the numbers with heat_numbers()
cor_heatmap(data = psq, taxa = taxa, tax_anno = taxAnno, numbers = NULL)
cor_heatmap(
 data = psq, taxa = taxa, tax_anno = taxAnno,
 colors = heat_palette("Berlin", rev = TRUE, sym = TRUE),
 numbers = heat_numbers(decimals = 2, col = "white", fontface = "bold")
)
# You can hide or change the style of the grid lines with grid_col & grid_lwd
```

cor_test 35

```
cor_heatmap(psq, taxa = taxa, tax_anno = taxAnno, grid_col = NA) # hidden
cor_heatmap(psq, taxa = taxa, tax_anno = taxAnno, grid_lwd = 3) # bigger

# You can pass any other argument from `ComplexHeatmap::Heatmap()` to `...`

# e.g. You can set the absolute width and height of the heatmap body
cor_heatmap(
    data = psq, taxa = taxa, tax_anno = taxAnno,
    width = grid::unit(40, "mm"), height = grid::unit(10, "cm")
)

# e.g. You can suppress the legend
cor_heatmap(
    data = psq, taxa = taxa, tax_anno = taxAnno, show_heatmap_legend = FALSE,
    width = grid::unit(40, "mm"), height = grid::unit(10, "cm")
)
```

cor_test

Simple wrapper around cor.test for $y \sim x$ style formula input

Description

Intended for use within the function tax_model

Usage

```
cor_test(formula, data, ...)
```

Arguments

```
formula a formula in form y \sim x data dataframe passed to cor.test
```

```
data("shao19")
ps <- shao19 %>%
    ps_filter(family_role == "mother") %>%
    tax_filter(min_prevalence = 20) %>%
    tax_agg("family")

cors <- ps %>% tax_model(
    rank = "family", variables = list("age", "number_reads"), type = cor_test
)

tax_models_get(cors)
```

deprecated-heatmap-annotations

DEPRECATED Heatmap annotations helpers

Description

Functions to easily define ComplexHeatmap annotations for taxa and/or variables

- tax_anno creates list describing taxa annotation (for cor_heatmap or comp_heatmap)
- var_anno creates list describing variable annotation (for cor_heatmap)

```
tax_anno(
  undetected = 0,
 which = NA,
 prev = 1,
 abund = 2,
  size = 30,
  gap = 2,
  rel_sizes = NA,
 args = NULL,
)
anno_prev(
 data,
  taxa,
 undetected = 0,
 which = "row",
 size = 15,
 bar_width = 0.6,
 gp = grid::gpar(fill = "grey85"),
)
anno_abund(
  data,
  taxa,
  undetected = 0,
 which = "row",
  size = 15,
 point_size = 0.75,
 box_width = 0.6,
 gp = grid::gpar(fill = "grey85"),
)
```

```
var_anno(
  annos = "var_box",
  funs = "identity",
  names = NA,
  which = "column",
  size = 15 * length(annos),
  gap = 2,
  rel_sizes = NA,
  args = NULL,
  ...
)

old_anno_var_hist(data, vars = NA, which = "column", size = 15, ...)

old_anno_var_box(data, vars = NA, which = "column", size = 15, ...)
```

Arguments

undetected

	value de l'e vinien dans de constactes present de cettes in a sample
which	"row" or "column" annnotation
prev	order in which prevalence annotation shown (number, or NA to not show)
abund	order in which abundance annotation shown (number, or NA to not show)
size	total size (mm) of annotations (width/height depending on which)
gap	gap in mm between annotations
rel_sizes	relative sizes of annotations (NA for equal sizes, or same length as annos)
args	extra args passed to each annotation: give as list of lists (one inner list per arg, named, e.g. list(prev = list(whatever = whatever))
	further named args to be passed on (to list)
data	phyloseq or ps-extra (or a data.frame or matrix for anno_var_* functions)
taxa	names of taxa to plot
bar_width	relative width of barchart bars
gp	a grid::gpar() object for graphics parameter settings like fill or lwd
point_size	size of outlier points in mm
box_width	relative width of boxplot boxes
annos	name(s) of annotation(s) to show, in order (e.g. 'var_box', 'var_hist')
funs	$function(s) \ to \ transform \ matrix \ of \ variable \ values \ before \ plotting \ (length \ must be \ 1 \ or \ same \ length \ as \ annos)$
names	names to use for each annotation in annos
vars	names of variables to plot

value above which taxa are considered present/detected in a sample

38 distinct_palette

distinct_palette

Colour palettes suitable for 20+ categories

Description

Available palettes (max colors) are "brewerPlus" (41), "kelly" (20) and "greenArmytage" (25).

- "brewerPlus" is an arbitrary expansion of the "Paired" and "Dark2" colorbrewer palettes. The philosophy behind this expansion was to ensure that similar colours are far apart, and the earlier colours are attractive.
- "kelly" is based on the 22-colour palette developed by Kenneth Kelly but with white and black starting colours removed. This palette is ordered such that the first colours are most distinct.
- "greenArmytage" is based on a 26-colour palette proposed by Paul Green-Armytage, with black removed. This palette is not ordered by maximum contrast.

Usage

```
distinct_palette(n = NA, pal = "brewerPlus", add = "lightgrey")
```

Arguments

n	number of colours to return
pal	palette name, one of "brewerPlus", "kelly", "greenArmytage"
add	colour to append to end of palette, as colour n+1, lightgrey by default for the use
	as "other" taxa in comp_barplot, or NA for no additional colour.

Details

Hex color codes for 'kelly' and 'greenArmytage' palettes are copied and slightly modified from the Polychrome R package: i.e. Polychrome::kelly.colors() and Polychrome::green.armytage.colors()

Please consider also citing Coombes 2019 doi:10.18637/jss.v090.c01 if you use either of these palettes.

See the Polychrome reference manual for more information: https://CRAN.R-project.org/package=Polychrome

Value

vector of colours

```
brewerPlus <- distinct_palette()
scales::show_col(brewerPlus)

kelly <- distinct_palette(pal = "kelly")
scales::show_col(kelly)</pre>
```

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```
greenArmytage <- distinct_palette(pal = "greenArmytage")
scales::show_col(greenArmytage)</pre>
```

dist_bdisp

Wrapper for vegan::betadisper()

Description

Takes the output of dist_calc function. Or use with the result of the permanova function to ensure the results correspond to exactly the same input data. Runs betadisper for all categorical variables in variables argument. See help('betadisper', package = 'vegan').

Usage

```
dist_bdisp(
  data,
  variables,
  method = c("centroid", "median")[[1]],
  complete_cases = TRUE,
  verbose = TRUE
)
```

Arguments

data psExtra output from dist_calc
variables list of variables to use as group
method centroid or median
complete_cases drop samples with NAs in any of the variables listed

verbose sends messages about progress if true

Value

psExtra containing betadisper results

```
library(phyloseq)
library(vegan)
data("dietswap", package = "microbiome")

# add some missings to demonstrate automated removal
sample_data(dietswap)$sex[3:6] <- NA

# create a numeric variable to show it will be skipped with a warning
dietswap <- ps_mutate(dietswap, timepoint = as.numeric(timepoint))

# straight to the betadisp</pre>
```

40 dist_calc

```
bd1 <- dietswap %>%
  tax_agg("Genus") %>%
  dist_calc("aitchison") %>%
  dist_bdisp(variables = c("sex", "bmi_group", "timepoint")) %>%
  bdisp_get()
bd1$sex
# quick vegan plotting methods
plot(bd1$sex$model, label.cex = 0.5)
boxplot(bd1$sex$model)
# compute distance and use for both permanova and dist_bdisp
testDist <- dietswap %>%
  tax_agg("Genus") %>%
  dist_calc("bray")
PERM <- testDist %>%
  dist_permanova(
    variables = c("sex", "bmi_group"),
    n_processes = 1, n_perms = 99
str(PERM, max.level = 1)
bd <- PERM %>% dist_bdisp(variables = c("sex", "bmi_group"))
bd
```

dist_calc

Calculate distances between pairs of samples in phyloseq object

Description

Can compute various sample-sample distances using the microbiota composition of your samples:

- Bray Curtis ('bray') or any other ecological distance from phyloseq::distance() / vegan::vegdist()
- UniFrac distances (using the GUniFrac package)
 - generalised: 'gunifrac' (optionally set weighting alpha in gunifrac alpha)
 - unweighted: 'unifrac'
 - weighted: 'wunifrac'
- Aitchison distance (Euclidean distance after centered log ratio transform clr, see details)
- · Euclidean distance

Use dist_calc with psExtra output of tax_transform (or tax_agg). It returns a psExtra object containing the phyloseq and the name of the distance used in addition to the distance matrix itself. The resulting object is intended to be piped into ord_calc or dist_permanova functions. Alternatively you can directly access the distance matrix with dist_get().

Usage

```
dist_calc(data, dist = "bray", gunifrac_alpha = 0.5, ...)
```

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Arguments

```
data psExtra object, e.g. output from tax_transform()

dist name of distance to calculate between pairs of samples

gunifrac_alpha setting alpha value only relevant if gunifrac distance used

... optional distance-specific named arguments passed to phyloseq::distance()
```

Value

psExtra object including distance matrix and name of distance used

Aitchison distance note

You should EITHER:

- 1. skip the dist_calc function and call ord_calc(method = "PCA") directly on an object with taxa transformed with tax_transform(trans = "clr")
- 2. pass an object with untransformed (or 'identity' transformed) taxa to the data argument of dist_calc() and specify dist = "aitchison".

If ordination plots with taxon loading vectors are desired, users require option 1. If the distance matrix is required for permanova, users require option 2.

Binary Jaccard distance note

Jaccard distance can be computed on abundances, but often in microbiome research it is the Binary Jaccard distance that is desired. So remember to first perform a "binary" transformation with tax_transform("binary"), OR pass an additional argument to dist_calc("jaccard", binary = TRUE)

See Also

```
tax_transform for the function to use before dist_calc
ord_calc
ord_plot
dist_permanova
phyloseq::distance
vegan::vegdist
```

```
# bray curtis distance on genera-level features
data("dietswap", package = "microbiome")
bc <- dietswap %>%
  tax_agg("Genus") %>%
  dist_calc("bray")
bc
class(bc)
```

42 dist_calc_seq

```
# gunifrac distance using phyloseq input
data("esophagus", package = "phyloseq")
gunifrac <- esophagus %>%
   dist_calc("gunifrac") %>%
   dist_get()
class(gunifrac)
```

dist_calc_seq

Calculate distances between sequential samples in ps_extra/phyloseq object

Description

Calculate distances between sequential samples in ps_extra/phyloseq object

Usage

```
dist_calc_seq(
  data,
  dist,
  group,
  seq,
  unequal = "warn",
  start_value = NaN,
  return = "data",
  var_name = paste0(dist, "_DistFromLast")
)
```

Arguments

data	psExtra object, e.g. output from tax_transform()
dist	name of distance to calculate between pairs of sequential samples
group	name of variable in phyloseq sample_data used to define groups of samples
seq	name of variable in phyloseq sample_data used to define order of samples within groups $% \left(1\right) =\left(1\right) \left(1\right) \left$
unequal	"error" or "warn" or "ignore" if groups of samples, defined by group argument, are of unequal size
start_value	value returned for the first sample in each group, which has no preceding sample in the group's sequence, and so has no obvious value
return	format of return object: "data" returns psExtra with sorted samples and additional variable. "vector" returns only named vector of sequential distances.
var_name	name of variable created in psExtra if return arg = "data"

Value

psExtra object sorted and with new sequential distance variable or a named vector of that variable

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See Also

```
dist_calc
```

```
library(ggplot2)
library(dplyr)
data("dietswap", package = "microbiome")
pseq <- dietswap %>%
 tax_transform("identity", rank = "Genus") %>%
 dist_calc_seq(
   dist = "aitchison", group = "subject", seq = "timepoint",
   # group sizes are unequal because some subjects are missing a timepoint
   unequal = "ignore"
 )
pseq %>%
 samdat_tbl() %>%
 dplyr::select(1, subject, timepoint, dplyr::last_col())
# ggplot heatmap - unsorted
pseq %>%
 samdat_tbl() %>%
 filter(timepoint != 1) %>%
 ggplot(aes(x = timepoint, y = subject)) +
 geom_tile(aes(fill = aitchison_DistFromLast)) +
 scale_fill_viridis_c(na.value = NA, name = "dist") +
 theme_minimal(base_line_size = NA) +
 scale_y_discrete(limits = rev(levels(samdat_tbl(pseq)$subject)))
# ComplexHeatmap plotting with clustering #
library(tidyr)
library(ComplexHeatmap)
# make data matrix
heatmat <- pseq %>%
 samdat_tbl() %>%
 filter(timepoint != 1) %>%
 pivot_wider(
    id_cols = subject,
   names_from = timepoint, names_prefix = "t",
   values_from = aitchison_DistFromLast
 ) %>%
 tibble::column_to_rownames("subject")
heatmat <- as.matrix.data.frame(heatmat)</pre>
heatmap <- Heatmap(</pre>
 name = "dist",
 matrix = heatmat, col = viridisLite::viridis(12), na_col = "white",
 cluster_columns = FALSE,
```

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```
cluster_rows = hclust(dist(heatmat), method = "ward.D"),
  width = unit(1.5, "in"), rect_gp = gpar(col = "black"),
  row_names_side = "left", row_names_gp = gpar(fontsize = 8)
)
heatmap
# comparison with subject tracking on PCA
pca <- pseq %>%
  # already sorted data
  dist_calc("aitchison") %>%
  ord_calc("PCoA") %>%
  ord_plot(alpha = 0.1, shape = "nationality", size = 2) %>%
  add_paths(
    mapping = aes(colour = subject, alpha = timepoint, size = timepoint),
    id_var = "subject", id_values = c(
      "eve", "hsf", # low variation
      "vem", # medium
      "ufm", # high variation
      "pku" # starts high
   )
  ) +
  scale_alpha_continuous(range = c(0.3, 1), breaks = c(2, 4, 6)) +
  scale\_size\_continuous(range = c(1, 2), breaks = c(2, 4, 6))
heatmap
рса
```

dist_permanova

Calculate PERMANOVA after dist_calc()

Description

dist_permanova runs a Permutational Multivariate ANOVA (aka Non-parametric MANOVA). This is a way to test for the statistical significance of (independent) associations between variables in your phyloseq::sample_data(), and a microbiota distance matrix you have already calculated with dist_calc().

This function is a wrapper around vegan's adonis2() function. See ?vegan::adonis2() for more insight.

You can also read this excellent book chapter on PERMANOVA by Marti Anderson: doi:10.1002/9781118445112.stat07841

Or this NPMANOVA page on GUSTA ME: https://sites.google.com/site/mb3gustame/hypothesis-tests/manova/npmanova

Usage

```
dist_permanova(
  data,
  variables = NULL,
```

dist_permanova 45

```
interactions = NULL,
complete_cases = TRUE,
n_processes = 1,
n_perms = 999,
seed = NULL,
by = "margin",
verbose = TRUE,
...
)
```

Arguments

data psExtra output from dist_calc()

variables character vector of variables to include in model or character representation of

the right-hand side of a formula, e.g "varA + varB + varA:varB"

interactions optional argument to define any interactions between variables, written in the

style of e.g. "var_a * var_b"

complete_cases if TRUE, drops observations if they contain missing values (otherwise stops if

missings are detected)

n_processes how many parallel processes to use? (on windows this uses parallel::makePSOCKcluster())

n_perms how many permutations? e.g. 9999. Less is faster but more is better!

seed set a random number generator seed to ensure you get the same results each run by passed to vegan::adonis2() by argument: what type of sums of squares to calcu-

late? "margin" or "terms"

verbose sends messages about progress if TRUE

... additional arguments are passed directly to vegan::adonis2() (e.g. strata, add,

sqrt.dist etc.)

Details

The variables argument will be collapsed into one string (if length > 1) by pasting together, separated by "+". Any interaction terms described in the interactions argument will be pasted onto the end of the pasted variables argument. Alternatively, you can supply the complete right hand side of the formula yourself e.g variables = "varA + varB + varC*varD"

Watch out, if any of your variable names contain characters that would normally separate variables in a formula then you should rename the offending variable (e.g. avoid any of "+" "*" "|" or ":") otherwise permanova will split that variable into pieces.

Value

psExtra list containing permanova results and (filtered) input objects

See Also

```
dist_calc for calculating the required distance matrix input
ord_plot with constraints as a way to visualise the microbial associations of significant predictors
vegan::adonis2
```

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```
data("dietswap", package = "microbiome")
# add some missings to demonstrate automated removal
phyloseq::sample_data(dietswap)$sex[3:6] <- NA</pre>
# compute distance
testDist <- dietswap %>%
  tax_agg("Genus") %>%
  tax_transform("identity") %>%
  dist_calc("bray")
PERM <- testDist %>%
  dist_permanova(
    seed = 1,
   variables = c("sex", "bmi_group"),
   n_{processes} = 1,
   n_perms = 99 # only 99 perms used in examples for speed (use 9999+!)
PERM
str(PERM, max.level = 1)
# try permanova with interaction terms
PERM2 <- testDist %>%
  dist_permanova(
   seed = 1,
   variables = "nationality + sex * bmi_group",
   n_processes = 1, n_perms = 99
  )
perm_get(PERM2)
# specify the same model in alternative way
PERM3 <- testDist %>%
  dist_permanova(
    seed = 1,
   variables = c("nationality", "sex", "bmi_group"),
   interactions = "sex * bmi_group",
   n_processes = 1, n_perms = 99
  )
perm_get(PERM3)
identical(PERM3, PERM2) # TRUE
# take same distance matrix used for the permanova and plot an ordination
PERM2 %>%
  ord_calc(method = "PCoA") %>%
  ord_plot(color = "bmi_group")
# this trick ensures any samples dropped from the permanova
# for having missing values in the covariates are NOT included
# in the corresponding ordination plot
```

heat_grid 47

heat_grid

set options for drawing gridlines on heatmaps

Description

set options for drawing gridlines on heatmaps

Usage

```
heat_grid(
   col = "white",
   alpha = 1,
   lty = 1,
   lwd = 0.5,
   lex = 1,
   lineend = "round",
   linejoin = "round")
```

Arguments

col	Colour for lines and borders.
alpha	Alpha channel for transparency
lty	Line type
lwd	Line width
lex	Multiplier applied to line width
lineend	Line end style (round, butt, square)
linejoin	Line join style (round, mitre, bevel)

 $heat_numbers$

Aesthetic settings for drawing numbers on heatmap tiles

Description

Works with comp_heatmap() and cor_heatmap(). See the help for those functions.

Usage

```
heat_numbers(
  decimals = 0,
  fontsize = 7,
  col = "darkgrey",
  fontface = "bold",
  fmt = NULL,
  ...
)
```

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Arguments

decimals number of decimal places to print

fontsize fontsize specification,

col colour of font fontface plain, bold, italic

fmt NULL or number print format, see ?sprintf, overrides decimals arg if set

... passed to grid::gpar() for grid.text

Value

list

heat_palette

Easy palettes for ComplexHeatmap

Description

Pass a named colorspace hel palette to circlize::colorRamp2.

- If you do not specify a range this function returns a function and the heatmap color palette will use the range of the data automatically
- If you do specify a range, this returns a colour palette with that range

Usage

```
heat_palette(
  palette = ifelse(sym, "Blue-Red 3", "Rocket"),
  breaks = "auto",
  range = NA,
  sym = FALSE,
  rev = FALSE
)
```

Arguments

palette named palette from colorspace::hcl_palettes() diverging/sequential or a vector

of colour names/hexcodes

breaks number of breaks, "auto" is 11 for a named palette, or uses palette length

range NA to return palette generating function that takes range or numeric vector in-

dicating the range, to return a palette

sym makes palette range symmetrical around 0 if TRUE

rev reverse the palette?

Value

circlize::colorRamp2 palette if range = NA, or function returning a palette when given a range

ibd 49

ibd

IBD study data in phyloseq object.

Description

A phyloseq object with an OTU table and sample data from an IBD microbiome study. Originally released as ibd example data in the corncob package.

Usage

ibd

Format

A phyloseq-class experiment-level object with an OTU table and sample data.

References

Papa, E., Docktor, M., Smillie, C., Weber, S., Preheim, S. P., Gevers, D., Giannoukos, G., Ciulla, D., Tabbaa, D., Ingram, J., Schauer, D. B., Ward, D. V., Korzenik, J. R., Xavier, R. J., Bousvaros, A., Alm, E. J. & Schauer, D. B. (2012). *Non-invasive mapping of the gastrointestinal microbiota identifies children with inflammatory bowel disease*. PloS One, 7(6), e39242. <doi.org/10.1371/journal.pone.0039242>.

Duvallet, C., Gibbons, S., Gurry, T., Irizarry, R., & Alm, E. (2017). *MicrobiomeHD: the human gut microbiome in health and disease [Data set]*. Zenodo. <doi.org/10.5281/zenodo.1146764>.

microViz

microViz: microbiome data analysis and visualization

Description

microViz provides functions for statistics and visualization of microbiome sequencing data. microViz wraps, extends and complements popular microbial ecology packages like phyloseq, vegan, and microbiome.

Check out the website for tutorials and illustrated help pages.

https://david-barnett.github.io/microViz/

Author(s)

David Barnett (ORCID) (GitHub)

See Also

Useful links:

- https://david-barnett.github.io/microViz
- https://github.com/david-barnett/microViz

50 Ordination-arrows

Ordination-arrows

Create ordination plot vector styling lists

Description

Used by ord_plot, see examples there.

Usage

```
vec_constraint(
  linewidth = 1,
  alpha = 0.8,
  colour = "brown",
 arrow = grid::arrow(length = grid::unit(0.005, units = "npc"), type = "closed", angle =
 lineend = "round",
 linejoin = "mitre",
)
vec_tax_sel(
  linewidth = 0.5,
 alpha = 1,
 colour = "black",
 arrow = grid::arrow(length = grid::unit(0.005, units = "npc"), type = "closed", angle =
    30),
 lineend = "round",
 linejoin = "mitre",
)
vec_tax_all(linewidth = 0.5, alpha = 0.25, arrow = NULL, ...)
```

Arguments

linewidth	width of vector
alpha	opacity of vector
colour	colour of vector
arrow	arrow style specified with grid::arrow() or NULL for no arrow
lineend	Line end style (round, butt, square).
linejoin	Line join style (round, mitre, bevel).
	further arguments passed to geom_segment

Value

list

Ordination-labels 51

Description

Customise taxa and constraint labels on your ordination plots. Choose 'text' or 'label' type, rotate and/or justify the text/labels and set aesthetic appearances using tax_lab_style() or constraint_lab_style().

Usage

```
tax_lab_style(
  type = "label",
 max\_angle = 0,
 perpendicular = FALSE,
 aspect_ratio = 1,
  justify = "auto",
  size = 2,
  alpha = 1,
  colour = "black",
)
constraint_lab_style(
  type = "label",
 max\_angle = 0,
 perpendicular = FALSE,
 aspect_ratio = 1,
  justify = "auto",
  size = 2.5,
 alpha = 1,
  colour = "brown",
)
```

Arguments

type	'label', 'text' or 'richtext' ('richtext' also used if 'label' type are rotated, when max_angle > 0)
max_angle	maximum angle of rotation to allow to match vector angle (requires ggtext package to rotate "label" type)
perpendicular	if TRUE, sets rotated labels perpendicular to desired angle, not parallel
aspect_ratio	aspect ratio of plot (y/x) must also be used in coord_fixed() ratio argument (must be set when rotated labels are used, to ensure match to arrow angles)
justify	"center", "side", or "auto"? Should the text/label align with the arrows at the text center or text sides (uses hjust, if 'auto', picks based on whether max_angle is greater than 0)

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```
size fixed size of text or label

alpha fixed alpha of text or label

colour fixed colour of text or label

... further named arguments passed to geom_text, geom_label or geom_richtext
```

Value

named list

```
# These examples show styling of taxa labels with tax_lab_style().
# The same options are available for constraint labels in constrained
# ordinations. constraint_lab_style() just has different default settings.
library(ggplot2)
# get example inflammatory bowel disease stool dataset from corncob package
data("ibd", package = "microViz")
# filter out rare taxa and clean up names etc
ibd <- ibd %>%
 tax_filter(min_prevalence = 3) %>%
 tax_fix() %>%
 phyloseq_validate()
# calculate a centered-log-ratio transformed PCA ordination
ibd_ord <- ibd %>%
 tax_transform("clr", rank = "Genus") %>%
 ord_calc("PCA")
# basic plot with default label style
ibd_ord %>% ord_plot(color = "ibd", plot_taxa = 1:10)
# Rotating labels: requires the ggtext package #
# A fixed coordinate ratio must be set to ensure label rotation
# matches the vectors. It is also helpful to set the vector and label length
# multipliers manually for a good look. Rotated labels are justified to the
# 'sides' automatically by tax_lab_style() with justify = 'auto'
ibd_ord %>%
 ord_plot(
   color = "ibd", plot_taxa = 1:7,
    tax_vec_length = 1.3, tax_lab_length = 1.3,
   tax_lab_style = tax_lab_style(max_angle = 90)
 coord_fixed(ratio = 1, clip = "off", xlim = c(-3.5, 3.5))
# You can use text instead of labels
# - a bold fontface helps text to stand out
# - see ?ggplot2::geom_text for all settings available
ibd_ord %>%
```

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```
ord_plot(
   color = "ibd", plot_taxa = 1:7,
    tax_vec_length = 1.3, tax_lab_length = 1.4,
   tax_lab_style = tax_lab_style(
     type = "text", max_angle = 90, size = 2.5, fontface = "bold.italic"
   )
 ) +
 coord_fixed(ratio = 1, clip = "off", xlim = c(-3.5, 3.5))
# With text you can prevent overlaps with check_overlap = TRUE
ibd_ord %>%
 ord_plot(
    color = "ibd", plot_taxa = 1:12,
    tax_vec_length = 1.3, tax_lab_length = 1.4,
   tax_lab_style = tax_lab_style(
     type = "text", max_angle = 90, size = 3, fontface = "bold.italic",
     check_overlap = TRUE
   )
 ) +
 coord_fixed(ratio = 1, clip = "off", xlim = c(-3.5, 3.5))
# With labels, you can reduce the padding and line weight to free space
# but check_overlap is not available
# see ?ggtext::geom_richtext for more possibilities
ibd_ord %>%
 ord_plot(
   color = "ibd", plot_taxa = 1:7,
    tax_vec_length = 1.3, tax_lab_length = 1.35,
   tax_lab_style = tax_lab_style(
     max_angle = 90, fontface = "italic", size = 2.5, fill = "grey95",
     label.size = 0.1, # width outline
     label.padding = unit(0.1, "lines"),
     label.r = unit(0, "lines") # reduces rounding of corners to radius 0
   )
 ) +
 coord_fixed(ratio = 1, clip = "off", xlim = c(-3.5, 3.5))
# Perpendicular angled labels/text are possible
ibd_ord %>%
 ord_plot(
   color = "ibd", plot_taxa = 1:12,
    tax_lab_style = tax_lab_style(
     type = "text", max_angle = 90, perpendicular = TRUE, size = 3,
     check\_overlap = TRUE
   )
 ) +
 coord_fixed(ratio = 1, clip = "off", xlim = c(-3.5, 3.5))
# You can limit and/or attenuate the angle of rotation by:
# - setting a lower max_angle
# - decreasing the aspect_ratio in the tax_lab_style call
ibd_ord %>%
```

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```
ord_plot(
   shape = "circle", color = "ibd", plot_taxa = 1:7,
   tax_vec_length = 1.3, tax_lab_length = 1.3,
   tax_lab_style = tax_lab_style(
     max_angle = 10, size = 2, label.size = 0.1,
     label.padding = unit(0.1, "lines"), label.r = unit(0, "lines")
   )
 ) +
 coord_fixed(ratio = 1, clip = "off", xlim = c(-3.5, 3.5))
ibd_ord %>%
 ord_plot(
   shape = "circle", color = "ibd", plot_taxa = 1:7,
   tax_vec_length = 1.3, tax_lab_length = 1.3,
   tax_lab_style = tax_lab_style(
     max_angle = 90, size = 2, label.size = 0.1, aspect_ratio = 0.5,
     label.padding = unit(0.1, "lines"), label.r = unit(0, "lines")
   )
 ) +
 coord_fixed(ratio = 1, clip = "off", xlim = c(-3.5, 3.5))
# another example with some extras #
ibd_ord %>%
 ord_plot(
   shape = "circle filled", fill = "ibd",
   plot_taxa = 1:10,
   taxon_renamer = function(x) stringr::str_replace_all(x, "_", " "),
   tax_vec_length = 2, tax_lab_length = 2.1,
   tax_lab_style = tax_lab_style(
     type = "text", max_angle = 90, size = 2.5,
     fontface = "bold.italic", check_overlap = TRUE
   )
 ) +
 coord_fixed(1, clip = "off", xlim = c(-5, 5)) +
 theme(legend.position = c(0.8, 0.2), legend.background = element_rect()) +
 stat_chull(mapping = aes(colour = ibd, fill = ibd), alpha = 0.1)
```

ord_calc

Ordinate samples (arrange by similarity in multiple dimensions)

Description

Used before plotting with ord_plot() or explorating interactively with ord_explore(). Use method = "auto" to automatically pick an appropriate method from:

- "PCA" (Principal Components Analysis) combines taxa abundances into new dimensions. The first axes display the greatest variation in your microbial data.
- "RDA" (Redundancy Analysis) is constrained PCA, roughly speaking. It finds variation in your data that can be explained both by the constraints variables, and the microbial data.

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• "PCoA" (Principal Coordinates Analysis) finds a coordinate system that best preserves the original distances between samples.

• "CAP" (Constrained Analysis of Principal Coordinates) is also known as distance-based Redundancy Analysis.

Alternatively to leaving method = "auto", you can explicitly specify any of the above methods, or choose one of the following:

- "CCA" (Canonical Correspondence Analysis) NOT canonical correlation analysis!
- "NMDS" (Non-metric Multidimensional Scaling)

You are strongly recommended to check out this useful website for introductory explanations of these methods the "GUide to STatistical Analysis in Microbial Ecology": https://sites.google.com/site/mb3gustame/

Usage

```
ord_calc(
  data,
  method = "auto",
  constraints = NULL,
  conditions = NULL,
  scale_cc = TRUE,
  verbose = TRUE,
  ...
)
```

Arguments

data	psExtra object: output from dist_calc(), or tax_transform() if no distance calculation required for method e.g. for RDA $$
method	which ordination method to use? "auto" means automatically determined from psExtra and other args. If you really know what you want: manually set one of 'PCoA', 'PCA', 'CCA', 'CAP' or 'RDA'
constraints	(a vector of) valid sample_data name(s) to constrain analyses, or leave as NULL for unconstrained ordination. Non-NULL values are compatible with method = "auto"/"RDA"/"CAP"
conditions	(a vector of) valid sample_data name(s) to partial these out of analyses with Condition(), or leave as $NULL$
scale_cc	If TRUE (default) ensures any constraints and conditions variables are scaled before use, to ensure their effects are comparable. If set to FALSE you must ensure you have already set the variables on a similar scale yourself! If there are no constraints or conditions, scale_cc does nothing.
verbose	If TRUE or "max", show any warnings and messages about constraint and conditions scaling and missings etc. FALSE suppresses warnings!
	optional arguments passed on to phyloseq::ordinate()

ord_calc

Details

Extends functionality of phyloseq::ordinate(). Results can be used directly in ord_plot(). You can extract the ordination object for other analyses with ord_get()

Value

```
psExtra object
```

See Also

```
dist_calc for distance matrix calculation
ord_plot and ord_explore
phyloseq ordinate
```

```
library(phyloseq)
library(vegan)
data("dietswap", package = "microbiome")
# create a couple of numerical variables to use as constraints
dietswap <- ps_mutate(</pre>
  dietswap,
  female = dplyr::if_else(sex == "female", true = 1, false = 0),
  weight = dplyr::recode(bmi_group, obese = 3, overweight = 2, lean = 1)
)
# add a couple of missing values to demo automated dropping of observations with missings
sample_data(dietswap)$female[c(3, 4)] <- NA</pre>
# compute ordination
test <- dietswap %>%
  tax_agg("Genus") %>%
  dist_calc("bray") %>%
  ord_calc(constraints = c("weight", "female"))
# familiarise yourself with the structure of the returned psExtra object
test
str(test, max.level = 1)
# compute RDA with centre-log-ratio transformed taxa
test2 <- dietswap %>%
  tax_agg("Genus") %>%
  tax_transform("clr") %>%
  ord_calc(constraints = c("weight", "female"))
# plot with vegan package graphics to show it returns a standard ordination object
ord_get(test2) %>% vegan::ordiplot()
# This is equivalent to CAP with "aitchison" distance
ord_plot(test2, plot_taxa = 8:1)
```

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```
# but the latter (below) doesn't allow plotting taxa loadings with ord_plot
dietswap %>%
  tax_agg("Genus") %>%
  dist_calc("aitchison") %>%
  ord_calc(constraints = c("weight", "female")) %>%
  ord_plot()
```

ord_explore

Interactively explore microbial compositions of ordinated samples

Description

A Shiny app used to create and explore an interactive version of ord_plot(). You can select samples on an ordination plot to view their composition with stacked barplots.

The ord_explore() data argument takes either:

- the output of ord_calc() (i.e. a psExtra with an ordination)
- a plain phyloseq object: ord_explore() will help you build an ordination

Once the app is running (in your browser), you can:

- 1. Create/edit the ordination if required
 - look at the R console error messages if your chosen options don't build
- 2. Style the ordination plot (e.g. choose dimensions; set colour and size; ...)
 - Taxa loading arrows can be added only to PCA, RDA and CCA plots
 - Convex hulls or ellipses can only be drawn if Colour is set to a variable
 - To track individuals over time with the path plotter, your data MUST already be sorted by time (e.g. with ps_arrange)!
- 3. Click on or use the lasso tool to select 1 or more samples to view their compositions
 - By default samples can be selected individually
 - Set the "Select" option to another variable to select by level of that variable
- 4. Style the taxonomic compositions barplot
 - The samples are ordered using the seriate_method argument and the same transformation and distance as used in the ordination plot
 - The app may lag if you select 100s of samples and ungroup the "Other" category
 - To avoid this lag: either reduce the number of taxa or samples, or deselect "Interactive" barplot
- 5. Stop the app by clicking the red stop button in the R console
 - Closing the web browser window doesn't stop the app, (you can find the app again at the local http address shown in the R console)
 - Don't forget to copy the ordination plot code before you close the app

See the Details section for some known limitations of the app. Please report any other app problems on the microViz GitHub issues page.

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Usage

```
ord_explore(
  data,
  sample_id = NULL,
  seriate_method = "OLO_ward",
  app_options = list(launch.browser = TRUE),
  plot_widths = c(7, 9),
  modal_fade = TRUE,
  notification_durations = list(2, 20),
  counts_warn = TRUE,
  ...
)
```

Arguments

data a phyloseq, or the psExtra output of ord_calc sample_id name of sample ID variable to use as default for selecting samples seriate_method seriation method to order phyloseq samples by similarity app_options passed to shinyApp() options argument plot_widths widths of plots in inches, including any legends (first number is ordination, second is composition barplot) modal_fade should the popover menus (modals) have a fade animation? notification_durations length 2 list giving duration in seconds of short and long notifications or NULL for notifications that do not disappear automatically should a warning be issued if accounts are unavailable? counts_warn additional arguments passed to ord_plot

Details

Limitations:

- If a "Select:" grouping variable is NA for some samples, then that grouping variable cannot be used to select those samples
- "Shape:" can only be mapped to variables with maximum 5 distinct levels, not including NAs. NAs in the shape variable are shown as hollow circles.

On some web browsers, e.g. older versions of Firefox, the numeric inputs' buttons are sometimes hard to click. As a workaround, click the box and type a number or use the arrow keys. This problem occurs in all Shiny apps, not just microViz.

Value

nothing, opens default browser

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```
# example code only runs in interactive R session
if (interactive()) {
 library(phyloseq)
 library(dplyr)
 # example of quickstart approach with interactive ordination calculation #
 microViz::ibd %>%
   # filtering makes subsequent calculations faster
   tax_filter(min_prevalence = 2) %>%
   tax_fix() %>%
   ord_explore()
 # simple example with precalculated ordination #
 data("enterotype")
 taxa_names(enterotype)[1] <- "unclassified" # replaces the "-1" taxon name
 ps <- tax_fix(enterotype) # remove NA taxa</pre>
 ord1 <- ps %>%
   tax_transform("identity", rank = "Genus") %>%
   dist_calc("bray") %>%
   ord_calc(method = "PCoA")
 ord_explore(data = ord1, auto_caption = 6)
 # constrained ordination example #
 data("dietswap", package = "microbiome")
 # create a couple of numerical variables to use as constraints
 dietswap <- dietswap %>%
   ps_mutate(
     weight = recode(bmi_group, obese = 3, overweight = 2, lean = 1),
     female = if_else(sex == "female", true = 1, false = 0)
   ) %>%
   tax_agg("Genus")
 constrained_aitchison_rda <- dietswap %>%
   tax_transform("clr") %>%
   ord_calc(constraints = c("weight", "female"))
 # label style arguments can be passed to ord_explore
 constrained_aitchison_rda %>%
   ord_explore(
     tax_lab_style = list(size = 3),
     constraint_lab_style = list(size = 4), auto_caption = 6
 # Try changing the point colour to bmi_group or similar
 # Style points interactively!
 # (setting colour/shape/etc as arguments doesn't work)
 # dietswap is actually a longitudinal dataset, with multiple samples per
 # subject. If we arrange by timepoint first (!!!), we can use the "paths"
 \mbox{\tt\#} additional plot layer from the ord_explore "Add:" menu to track
```

```
# individual subjects over time.
 dietswap %>%
   ps_arrange(timepoint) %>%
    tax_fix() %>%
   ord_explore()
 # Another dataset, where "size" variable drives gradient on PC1
 # Try setting size and/or alpha to correspond to "size"!
 # Then edit the ordination to use "size" as a condition, see what happens
 # hmp2 <- microbiomeutilities::hmp2</pre>
 hmp2 %>%
    tax_fix() %>%
    tax_transform(rank = "Genus", "identity") %>%
    dist_calc("aitchison") %>%
    ord_calc() %>%
    ord_explore()
 # another dataset
 data("soilrep", package = "phyloseq")
 # test auto creation of SAMPLE var
 ps <- soilrep %>% ps_select(-Sample)
 \ensuremath{\text{\#}} The barplot is actually quite useless with the 16000+ anonymous OTUs
 # in this dataset, but the 1000s of unmerged "Other" categories do render
 phyloseq_validate(ps) %>%
    tax_fix() %>%
    dist_calc("aitchison") %>%
    ord_calc() %>%
   ord_explore()
}
```

ord_plot

Customisable ggplot2 ordination plot

Description

Draw ordination plot. Utilises psExtra object produced by of ord_calc.

- For an extensive tutorial see the ordination vignette.
- For interpretation see the relevant pages on PCA, PCoA, RDA, or CCA on the GUide to STatistical Analysis in Microbial Ecology (GUSTA ME) website: https://sites.google.com/site/mb3gustame/

Usage

```
ord_plot(
  data,
  axes = 1:2,
  plot_taxa = FALSE,
```

```
tax_vec_length = NA,
  tax_vec_style_all = vec_tax_all(),
  tax_vec_style_sel = vec_tax_sel(),
  tax_lab_length = tax_vec_length * 1.1,
  tax_lab_style = list(),
  taxon\_renamer = function(x) identity(x),
  constraint_vec_length = NA,
  constraint_vec_style = vec_constraint(),
  constraint_lab_length = constraint_vec_length * 1.1,
  constraint_lab_style = list(),
  var\_renamer = function(x) identity(x),
  plot_samples = TRUE,
  scaling = 2,
  auto\_caption = 8,
  center = FALSE,
  clip = "off",
  expand = !center,
  interactive = FALSE,
)
```

Arguments

data psExtra object with ordination attached, i.e. output from ord_calc axes which axes to plot: numerical vector of length 2, e.g. 1:2 or c(3,5)if ord_calc method was "PCA/RDA" draw the taxa loading vectors (see details) plot_taxa tax_vec_length taxon arrow vector scale multiplier. NA = auto-scaling, or provide a numeric multiplier yourself. tax_vec_style_all list of named aesthetic attributes for all (background) taxon vectors tax_vec_style_sel list of named aesthetic attributes for taxon vectors for the taxa selected by plot_taxa tax_lab_length scale multiplier for label distance/position for any selected taxa list of style options for the taxon labels, see tax_lab_style() function. tax_lab_style taxon_renamer function that takes any plotted taxon names and returns modified names for labels constraint_vec_length constraint arrow vector scale multiplier. NA = auto-scaling, or provide a numeric multiplier yourself. constraint_vec_style list of aesthetics/arguments (colour, alpha etc) for the constraint vectors constraint_lab_length label distance/position for any constraints (relative to default position which is proportional to correlations with each axis) constraint_lab_style

list of aesthetics/arguments (colour, size etc) for the constraint labels

```
function to rename constraining variables for plotting their labels
var_renamer
plot_samples
                   if TRUE, plot sample points with geom_point
                   Type 2, or type 1 scaling. For more info, see <a href="https://sites.google.com/">https://sites.google.com/</a>
scaling
                   site/mb3gustame/constrained-analyses/redundancy-analysis. Either "species"
                   or "site" scores are scaled by (proportional) eigenvalues, and the other set of
                   scores is left unscaled (from ?vegan::scores.cca)
auto_caption
                   size of caption with info about the ordination, NA for none
                   expand plot limits to center around origin point (0,0)
center
                   clipping of labels that extend outside plot limits?
clip
                   expand plot limits a little bit further than data range?
expand
                   creates plot suitable for use with ggiraph (used in ord_explore)
interactive
                   pass aesthetics arguments for sample points, drawn with geom_point using aes_string
```

Details

How to specify the plot_taxa argument (when using PCA, CCA or RDA):

- FALSE -> plot no taxa vectors or labels
- integer vector e.g. 1:3 -> plot labels for top 3 taxa (by longest line length)
- single numeric value e.g. $0.75 \rightarrow \text{plot}$ labels for taxa with line length > 0.75
- character vector e.g. c('g_Bacteroides', 'g_Veillonella') -> plot labels for the exactly named taxa

Value

ggplot

See Also

```
tax_lab_style / tax_lab_style for styling labels
ord_explore for interactive ordination plots
ord_calc for calculating an ordination to plot with ord_plot
```

```
library(ggplot2)
data("dietswap", package = "microbiome")

# create a couple of numerical variables to use as constraints or conditions
dietswap <- dietswap %>%
    ps_mutate(
        weight = dplyr::recode(bmi_group, obese = 3, overweight = 2, lean = 1),
        female = dplyr::if_else(sex == "female", true = 1, false = 0)
    )

# unconstrained PCA ordination
unconstrained_aitchison_pca <- dietswap %>%
```

```
tax_transform("clr", rank = "Genus") %>%
 ord_calc() # method = "auto" --> picks PCA as no constraints or distances
unconstrained_aitchison_pca %>%
 ord_plot(colour = "bmi_group", plot_taxa = 1:5) +
 stat_ellipse(aes(linetype = bmi_group, colour = bmi_group))
# you can generate an interactive version of the plot by specifying
# interactive = TRUE, and passing a variable name to another argument
# called `data_id` which is required for interactive point selection
interactive_plot <- unconstrained_aitchison_pca %>%
 ord_plot(
    colour = "bmi_group", plot_taxa = 1:5,
    interactive = TRUE, data_id = "sample"
# to start the html viewer, and allow selecting points, we must use a
# ggiraph function called girafe and set some options and css
ggiraph::girafe(
 ggobj = interactive_plot,
 options = list(
   ggiraph::opts_selection(
     css = ggiraph::girafe_css(
       css = "fill:orange;stroke:black;",
       point = "stroke-width:1.5px"
      type = "multiple", # this activates lasso selection (click top-right)
     only_shiny = FALSE # allows interactive plot outside of shiny app
 )
)
# remove effect of weight with conditions arg
# scaling weight with scale_cc is not necessary as only 1 condition is used
dietswap %>%
 tax_transform("clr", rank = "Genus") %>%
 ord_calc(conditions = "weight", scale_cc = FALSE) %>%
 ord_plot(colour = "bmi_group") +
 stat_ellipse(aes(linetype = bmi_group, colour = bmi_group))
# alternatively, constrain variation on weight and female
constrained_aitchison_rda <- dietswap %>%
 tax_transform("clr", rank = "Genus") %>%
 ord_calc(constraints = c("weight", "female")) # constraints --> RDA
constrained_aitchison_rda %>%
 ord_plot(colour = "bmi_group", constraint_vec_length = 2) +
 stat_ellipse(aes(linetype = bmi_group, colour = bmi_group))
# ggplot allows additional customisation of the resulting plot
p <- constrained_aitchison_rda %>%
 ord_plot(colour = "bmi_group", plot_taxa = 1:3) +
```

```
lims(x = c(-5, 6), y = c(-5, 5)) +
 scale_colour_brewer(palette = "Set1")
p + stat_ellipse(aes(linetype = bmi_group, colour = bmi_group))
p + stat_density2d(aes(colour = bmi_group))
# you can rename the taxa on the labels with any function that
# takes and modifies a character vector
constrained_aitchison_rda %>%
 ord_plot(
   colour = "bmi_group",
   plot_taxa = 1:3,
   taxon_renamer = function(x) stringr::str_extract(x, "^.")
 lims(x = c(-5, 6), y = c(-5, 5)) +
 scale_colour_brewer(palette = "Set1")
# You can plot PCoA and constrained PCoA plots too.
# You don't typically need/want to use transformed taxa variables for PCoA
# But it is good practice to call tax_transform("identity") so that
# the automatic caption can record that no transformation was applied
dietswap %>%
 tax_agg("Genus") %>%
 tax_transform("identity") %>%
 # so caption can record (lack of) transform
 dist_calc("bray") %>%
 # bray curtis
 ord_calc() %>%
 # guesses you want an unconstrained PCoA
 ord_plot(colour = "bmi_group")
# it is possible to facet these plots
# (although I'm not sure it makes sense to)
# but only unconstrained ordination plots and with plot_taxa = FALSE
unconstrained_aitchison_pca %>%
 ord_plot(color = "sex", auto_caption = NA) +
 facet_wrap("sex") +
 theme(line = element_blank()) +
 stat_density2d(aes(colour = sex)) +
 guides(colour = "none")
unconstrained_aitchison_pca %>%
 ord_plot(color = "bmi_group", plot_samples = FALSE, auto_caption = NA) +
 facet_wrap("sex") +
 theme(line = element_blank(), axis.text = element_blank()) +
 stat_density2d_filled(show.legend = FALSE) +
 geom_point(size = 1, shape = 21, colour = "black", fill = "white")
```

Description

Use with ord_calc output as data argument. Order of samples extracted from ordination axes in data. Best paired with ordination plot made from same ord_calc output.

Usage

```
ord_plot_iris(
  data,
  tax_level,
  axes = 1:2,
  n_{\text{taxa}} = 10,
  ord_plot = "none",
  taxon\_renamer = function(x) identity(x),
  palette = distinct_palette(n_taxa),
  anno_colour = NULL,
  anno_colour_style = list(),
  anno_binary = NULL,
  anno_binary_style = list(),
  keep_all_vars = FALSE,
  scaling = 2,
  count_warn = TRUE,
)
```

Arguments

data psExtra output of ord_calc taxonomic aggregation level (from rank_names(ps)) tax_level which 2 axes of ordination to use for ordering bars axes how many taxa to colour show distinct colours for (all other taxa grouped into n_taxa "other"). add a matching ordination plot to your iris plot ('list' returns separate plots in a ord_plot list, 'above'/'below' uses patchwork to pair plots together into one) function to rename taxa in the legend taxon_renamer colour palette palette anno_colour name of sample_data variable to use for colouring geom_segment annotation ring anno_colour_style list of further arguments passed to geom_segment e.g. size name(s) of binary sample_data variable(s) (levels T/F or 1/0) to use for filtered anno_binary geom_point annotation ring(s) (annotates at TRUE values) anno_binary_style list of further arguments passed to geom_point e.g. colour, size, y, etc. keep_all_vars slows down processing but is required for any post-hoc plot customisation options

scaling Type 2, or type 1 scaling. For more info, see https://sites.google.com/ site/mb3gustame/constrained-analyses/redundancy-analysis. Either "species" or "site" scores are scaled by (proportional) eigenvalues, and the other set of scores is left unscaled (from ?vegan::scores.cca) warn if count data are not available? i.e. phyloseq otu_table is not positive count_warn integers and psExtra counts slot is NULL Arguments passed on to comp_barplot . . . merge_other if FALSE, taxa coloured/filled as "other" remain distinct, and so can have bar outlines drawn around them bar_width default 1 avoids random gapping otherwise seen with many samples (set to less than 1 to introduce gaps between samples) bar_outline_colour line colour separating taxa and samples (use NA for no outlines) bar_outline_width width of line separating taxa and samples (for no outlines set bar_outline_colour = NA) tax_transform_for_plot default "compositional" draws proportions of total counts per sample, but you could reasonably use another transformation, e.g. "identity", if you have truly quantitative microbiome profiling data interactive creates plot suitable for use with ggiraph max_taxa maximum distinct taxa groups to show (only really useful for limiting complexity of interactive plots e.g. within ord_explore) other_name name for other taxa after N

Details

data must also contain counts table if taxa were transformed (e.g. for clr PCA ordination) (i.e. you must have used tax_transform with keep_counts = TRUE, if transformation was not "identity")

You cannot set a variable fill aesthetic (only fixed) for the annotation points, as the fill is used for the taxonomic composition bars

Value

ggplot

```
library(dplyr)
library(ggplot2)
data("dietswap", package = "microbiome")

# although these iris plots are great for 100s of samples
# we'll take a subset of the data (for speed in this example)
ps <- dietswap %>%
    ps_filter(timepoint %in% c(1, 2)) %>%
    # copy an otu to the sample data
    ps_otu2samdat("Prevotella melaninogenica et rel.") %>%
    # create a couple of useful variables
    ps_mutate(
```

```
female = sex == "female",
    african = nationality == "AFR",
    log_P.melaninogenica = log10(`Prevotella melaninogenica et rel.` + 1)
 )
# define a function for taking the end off the long genus names in this dataset
tax_renamer <- function(tax) {</pre>
 stringr::str_remove(tax, " [ae]t rel.")
}
ord <- ps %>%
 tax_agg("Genus") %>%
 dist_calc("aitchison") %>%
 ord_calc(method = "PCoA")
# ordination plot for comparison
ord %>% ord_plot(color = "log_P.melaninogenica", size = 3)
ord_plot_iris(
 data = ord,
 tax_level = "Genus",
 n_{\text{taxa}} = 10,
 anno_colour = "nationality",
 anno_colour_style = list(size = 3),
 anno_binary = "female",
 anno_binary_style = list(shape = "F", size = 2.5),
 taxon_renamer = tax_renamer
) +
 scale_colour_brewer(palette = "Dark2")
# It is also possible to use comp_barplot customisation arguments
# like bar_width and bar_outline_colour, and to make interactive iris plots
# using ggiraph:
if (interactive()) {
 hover_over_me <- ord_plot_iris(</pre>
    data = ord,
    tax_level = "Genus",
    n_{\text{taxa}} = 10,
    anno_colour = "nationality",
    anno_colour_style = list(size = 3),
    anno_binary = "female",
    anno_binary_style = list(shape = "F", size = 2.5),
    taxon_renamer = tax_renamer,
    interactive = TRUE,
   bar_width = 0.8, bar_outline_colour = "black"
    scale_colour_brewer(palette = "Dark2")
 ggiraph::girafe(ggobj = hover_over_me)
}
```

Using PCA for ordination after transformations (e.g. clr) means the untransformed taxonomic

```
# data are only available for plotting as compositions if you transformed with
# tax_transform(keep_counts = TRUE) and your original data were in fact counts.
# Compositional data will also work, and you can set count_warn to FALSE to avoid the warning
clr_pca <- ps %>%
  tax_agg("Genus") %>%
  tax_transform("clr") %>%
  ord_calc(method = "PCA")
# you can generate a simple paired layout of ord_plot and iris plot
# or separately create and pair the plots yourself, for more control
# simple pairing
ord_plot_iris(
  data = clr_pca, n_taxa = 12,
  tax_level = "Genus",
  taxon_renamer = tax_renamer,
  ord_plot = "below",
  bar_width = 0.8, bar_outline_colour = "black",
  anno_binary = "african",
  anno_binary_style = list(
   y = 1.08, colour = "gray50", shape = "circle open", size = 1, stroke = 1.5
)
# manual pairing
plot1 <- clr_pca %>% ord_plot(
  plot_taxa = 6:1, tax_vec_length = 0.6,
  colour = "gray50", shape = "nationality",
  taxon_renamer = tax_renamer,
  auto_caption = NA, center = TRUE,
) +
  scale_shape_manual(values = c(AFR = "circle", AAM = "circle open"))
iris <- ord_plot_iris(</pre>
  data = clr_pca, n_taxa = 15,
  tax_level = "Genus",
  taxon_renamer = tax_renamer,
  anno_binary = "african",
  anno_binary_style = list(y = 1.05, colour = "gray50", shape = "circle", size = 1)
  # shrink legend text size
  theme(legend.text = element_text(size = 7))
cowplot::plot_grid(plot1, iris, nrow = 1, align = "h", axis = "b", rel_widths = 3:4)
# you can add multiple rings of binary annotations
ord_plot_iris(
  data = clr_pca, n_taxa = 15,
  tax_level = "Genus",
  taxon_renamer = tax_renamer,
  anno_binary = c("african", "female"),
  anno_binary_style = list(
```

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```
colour = c("gray50", "coral"),
    shape = c("circle", "F"), size = c(0.5, 2)
)
) +
theme(legend.text = element_text(size = 7))
```

phyloseq_validate

Check for (and fix) common problems with phyloseq objects

Description

- It checks for, and messages about, common uninformative entries in the tax_table, which often cause unwanted results
- If there is no sample_data, it creates a sample_data dataframe with the sample_names (as "SAMPLE" variable)
- If there is no tax_table, it creates a 1-column tax_table matrix with the taxa_names, and calls the rank "unique"
- If remove_undetected = TRUE, it removes taxa where phyloseq::taxa_sums() is equal to zero, with a warning

Usage

```
phyloseq_validate(
   ps,
   remove_undetected = FALSE,
   min_tax_length = 4,
   verbose = TRUE
)
```

Arguments

```
ps phyloseq object

remove_undetected

if TRUE, removes taxa that sum to zero across all samples

min_tax_length minimum number of characters to not consider a tax_table entry suspiciously short

verbose print informative messages if true
```

Value

possibly modified phyloseq object

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Examples

```
data(dietswap, package = "microbiome")
# expect warning about taxa summing to zero
phyloseq_validate(dietswap, remove_undetected = TRUE, verbose = TRUE)
# verbose = FALSE will suppress messages and warnings but still:
# replace NULL sample_data and remove taxa that sum to 0 across all samples
# (if remove_undetected = TRUE)
phyloseq_validate(dietswap, verbose = FALSE)
# Sometimes you might have a phyloseq with no sample_data
# This isn't compatible with some microViz functions, like comp_barplot
# So some functions internally use phyloseq_validate to fix this
dietswap@sam_data <- NULL
phyloseq_validate(dietswap)
# Sometimes you might have a phyloseq with no tax_table
# This isn't compatible with some microViz functions, like tax_top,
# so this is another reason to start your analyses with phyloseq_validate!
data("soilrep", package = "phyloseq")
soilrep # has NULL tax_table
phyloseq_validate(soilrep)
# If no messages or warnings are emitted,
# this means no problems were detected, and nothing was changed
# (but only if verbose = TRUE)
```

prev

Calculate prevalence from numeric vector

Description

Useful as helper for taxon prevalence calculation

Usage

```
prev(x, undetected = 0)
```

Arguments

x numeric vector (of taxon counts or proportions)undetected value above which a taxon is considered present or detected

Value

numeric value

print.psExtraInfo 71

Examples

```
prev(c(0, 0, 1, 2, 4))
prev(c(0, 0, 1, 2, 4), undetected = 1.5)
```

print.psExtraInfo

Print method for psExtraInfo object

Description

Print method for psExtraInfo object

Usage

```
## S3 method for class 'psExtraInfo'
print(
    X,
    ...,
    which = c("tax_agg", "tax_trans", "tax_scale", "dist_method", "ord_info")
)
```

Arguments

```
x psExtraInfo object
... ignored
which which elements of psExtraInfo list to print
```

psExtra-accessors

Extract elements from psExtra class

Description

- ps_get returns phyloseq
- info_get returns psExtraInfo object
- dist_get returns distance matrix (or NULL)
- ord_get returns ordination object (or NULL)
- perm_get returns adonis2() permanova model (or NULL)
- bdisp_get returns results of betadisper() (or NULL)
- otu_get returns phyloseq otu_table matrix with taxa as columns
- tt_get returns phyloseq tax_table
- tax_models_get returns list generated by tax_model or NULL
- tax_stats_get returns dataframe generated by tax_models2stats or NULL
- taxatree_models_get returns list generated by taxatree_models or NULL
- taxatree_stats_get returns dataframe generated by taxatree_models2stats or NULL
- samdat_tb1 returns phyloseq sample_data as a tibble with sample_names as new first column called .sample_name

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Usage

```
ps_get(psExtra, ps_extra, counts = FALSE, warn = TRUE)
dist_get(psExtra, ps_extra)
ord_get(psExtra, ps_extra)
info_get(psExtra, ps_extra)
perm_get(psExtra, ps_extra)
bdisp_get(psExtra, ps_extra)
tax_models_get(psExtra)
tax_stats_get(psExtra)
taxatree_models_get(psExtra)
taxatree_stats_get(psExtra)
otu_get(data, taxa = NA, samples = NA, counts = FALSE, warn = TRUE)
tt_get(data)
samdat_tbl(data, sample_names_col = ".sample_name")
```

Arguments

psExtra

psExtra S4 class object ps_extra deprecated! don't use this should ps_get or otu_get attempt to return counts? if present in object counts if counts = TRUE, should a warning be emitted if counts are not available? set warn warn = "error" to stop if counts are not available data phyloseq or psExtra subset of taxa to return, NA for all (default) taxa subset of samples to return, NA for all (default) samples sample_names_col

> name of column where sample_names are put. if NA, return data.frame with rownames (sample_names)

Value

```
element(s) from psExtra object (or NULL)
```

psExtra-class 73

Examples

```
data("dietswap", package = "microbiome")
psx <- tax_transform(dietswap, "compositional", rank = "Genus")</pre>
psx
ps_get(psx)
ps_get(psx, counts = TRUE)
info_get(psx)
dist_get(psx) # this psExtra has no dist_calc result
ord_get(psx) # this psExtra has no ord_calc result
perm_get(psx) # this psExtra has no dist_permanova result
bdisp_get(psx) # this psExtra has no dist_bdisp result
# these can be returned from phyloseq objects too
otu_get(psx, taxa = 6:9, samples = c("Sample-9", "Sample-1", "Sample-6"))
otu_get(psx, taxa = 6:9, samples = c(9, 1, 6), counts = TRUE)
tt_get(psx) %>% head()
samdat_tbl(psx)
samdat_tbl(psx, sample_names_col = "SAMPLENAME")
```

psExtra-class

Define psExtra class S4 object

Description

Define psExtra class S4 object

Slots

```
info list.
counts otu_table.
dist dist.
ord ANY.
permanova ANY.
bdisp ANY.
```

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```
taxatree_models list.

taxatree_stats data.frame.

tax_models list.

tax_stats data.frame.
```

Examples

```
library(phyloseq)
data("shao19")

ps <- shao19 %>% ps_filter(infant_age == 12)
ps %>% tax_agg("genus")
```

ps_arrange

Arrange samples in phyloseq by sample_data variables or taxon abundance

Description

Uses information in the sample_data or tax_table of phyloseq object to set the order of the samples (sample_data or tax_table specified by .target arg)

Give this function arguments in the same way you would use dplyr::arrange()

Usage

```
ps_arrange(ps, ..., .target = "sample_data")
```

Arguments

ps phyloseq object

... dots passed directly to dplyr::arrange()

. target arrange samples by "sample_data" variables or "otu_table" taxa abundances

Value

phyloseq

See Also

arrange

ps_calc_diversity 75

Examples

```
data("dietswap", package = "microbiome")

dietswap %>%
    ps_arrange(subject, timepoint) %>%
    phyloseq::sample_data() %>%
    head(8)

ps <- dietswap %>% ps_arrange(subject, desc(timepoint))
phyloseq::sample_data(ps) %>% head(8)
phyloseq::otu_table(ps)[1:8, 1:8]

# you can also arrange samples by the abundances of taxa in the otu tables
pst <- dietswap %>% ps_arrange(desc(Akkermansia), .target = "otu_table")
phyloseq::otu_table(pst)[1:8, 1:8]
phyloseq::sample_data(pst) %>% head(8)
```

ps_calc_diversity

Calculate diversity index and add to phyloseq sample data

Description

Wrapper around microbiome::diversity() function. Takes and returns a phyloseq object. Calculates an alpha diversity index at a given taxonomic rank. Returns phyloseq unaggregated, with an additional variable. Variable name is by default created by pasting the index and rank.

Usage

```
ps_calc_diversity(
   ps,
   rank,
   index = "shannon",
   exp = FALSE,
   varname = paste0(ifelse(exp, "exp_", ""), paste0(index, "_", rank))
)
```

Arguments

ps	phyloseq
rank	taxonomic rank name, or "unique"
index	name of diversity index from microbiome::diversity(). One of: "inverse_simpson", "gini_simpson", "shannon", "fisher", "coverage"
exp	exponentiate the result? (i.e. return e^index) - see details
varname	name of the variable to be added to phyloseq sample data

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Details

Don't filter taxa before calculating alpha diversity.

See the following resources for a discussion of exponentiated diversity indices http://www.loujost.com/Statistics%20and%20Inttp://www.loujost.com/Statistics%20and%20Physics/Diversity%20and%20Similarity/EffectiveNumberOfSpecies.htm

Value

phyloseq

Examples

```
data(ibd, package = "microViz")
ibd %>%
    ps_filter(abx == "abx") %>%
    tax_fix() %>%
    ps_calc_diversity("Genus", index = "shannon", exp = TRUE) %>%
    ps_calc_diversity("Family", index = "inverse_simpson") %>%
    tax_transform(rank = "Genus", transform = "clr") %>%
    ord_calc("PCA") %>%
    ord_plot(
    colour = "exp_shannon_Genus", size = "inverse_simpson_Family"
) +
    ggplot2::scale_colour_viridis_c()
```

ps_calc_dominant

Calculate dominant taxon in each phyloseq sample

Description

Which taxon is most abundant in each sample of your phyloseq object? This function adds this information as a new variable in your phyloseq sample_data.

- If the most abundant taxon is below the proportional abundance threshold, the dominant taxon will be "none" for that sample
- If there are more than n_max dominant taxa across all samples (not including "none") the dominant taxon will be "other" for those samples

Usage

```
ps_calc_dominant(
   ps,
   rank,
   threshold = 0.3,
   n_max = 6,
   var = paste("dominant", rank, sep = "_"),
   none = "none",
   other = "other"
)
```

ps_calc_dominant 77

Arguments

ps	phyloseq object
rank	taxonomic rank to calculate dominance at
threshold	minimum proportion at which to consider a sample dominated by a taxon
n_max	maximum number of taxa that can be listed as dominant taxa
var	name of variable to add to phyloseq object sample data
none	character value to use when no taxon reaches threshold
other	character value to use when another taxon (>n_max) dominates

Details

Thanks to Vitor Heidrich for the idea and a draft implementation

Value

phyloseq object

Examples

```
library(ggplot2)
ps <- microViz::ibd %>%
  tax_filter(min_prevalence = 3) %>%
  tax_fix() %>%
  phyloseq_validate()
ps %>%
  ps_filter(DiseaseState == "CD") %>%
  ps_calc_dominant(rank = "Genus") %>%
  comp_barplot(tax_level = "Genus", label = "dominant_Genus", n_taxa = 12) +
  coord_flip()
ps %>%
  ps_calc_dominant(rank = "Genus") %>%
  tax_transform(rank = "Genus", trans = "clr") %>%
  ord_calc("PCA") %>%
  ord_plot(colour = "dominant_Genus", size = 3, alpha = 0.6) +
  scale_colour_brewer(palette = "Dark2")
# customise function options
ps %>%
  ps_calc_dominant(
    rank = "Family", other = "Other", none = "Not dominated",
    threshold = 0.4, n_max = 3
  tax_transform(rank = "Genus", trans = "clr") %>%
  ord_calc("PCA") %>%
  ord_plot(colour = "dominant_Family", size = 3, alpha = 0.6) +
  scale_colour_manual(values = c(
   Bacteroidaceae = "forestgreen", Lachnospiraceae = "darkblue",
```

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```
Ruminococcaceae = "darkorange", Other = "red", "Not dominated" = "grey"
))
```

ps_calc_richness

Calculate richness estimate and add to phyloseq sample data

Description

Wrapper around microbiome::richness() function. Takes and returns a phyloseq object. Calculates a richness estimate at a given taxonomic rank. Returns phyloseq unaggregated, with an additional variable. Variable name is by default created by pasting the index and rank.

Usage

```
ps_calc_richness(
  ps,
  rank,
  index = "observed",
  detection = 0,
  varname = paste0(index, "_", rank)
)
```

Arguments

ps phyloseq

rank taxonomic rank name, or "unique"

index "observed" or "chao1" - name of richness estimate from microbiome::richness()

detection Detection threshold. Used for the "observed" index.

varname name of the variable to be added to phyloseq sample data

Details

Don't filter taxa before calculating richness.

These richness indices are estimates. For a discussion of the uncertainty and bias of these estimates see e.g. work by Amy Willis https://doi.org/10.3389/fmicb.2019.02407

Value

phyloseq

See Also

```
ps_calc_diversity
microbiome::richness
```

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Examples

```
data(ibd, package = "microViz")
ibd %>%
    ps_filter(abx == "abx") %>%
    tax_fix() %>%
    ps_calc_richness("Genus", index = "observed") %>%
    ps_calc_richness("Family", index = "chao1") %>%
    tax_transform(rank = "Genus", transform = "clr") %>%
    ord_calc("PCA") %>%
    ord_plot(
    colour = "observed_Genus", size = "chao1_Family"
    ) +
    ggplot2::scale_colour_viridis_c()
```

ps_dedupe

De-duplicate phyloseq samples

Description

Use one or more variables in the sample_data to identify and remove duplicate samples (leaving one sample per group).

methods:

- method = "readcount" keeps the one sample in each duplicate group with the highest total number of reads (phyloseq::sample_sums)
- method = "first" keeps the first sample in each duplicate group encountered in the row order of the sample_data
- method = "last" keeps the last sample in each duplicate group encountered in the row order of the sample_data
- method = "random" keeps a random sample from each duplicate group (set.seed for reproducibility)

More than one "duplicate" sample can be kept per group by setting n samples > 1.

Usage

```
ps_dedupe(
   ps,
   vars,
   method = "readcount",
   verbose = TRUE,
   n = 1,
   .keep_group_var = FALSE,
   .keep_readcount = FALSE,
   .message_IDs = FALSE,
   .label_only = FALSE,
   .keep_all_taxa = FALSE
)
```

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Arguments

ps phyloseq object

vars names of variables, whose (combined) levels identify groups from which only 1 sample is desired

method keep sample with max "readcount" or the "first" or "last" or "random" samples encountered in given sample_data order for each duplicate group

verbose message about number of groups, and number of samples dropped?

n number of 'duplicates' to keep per group, defaults to 1

.keep_group_var

keep grouping variable .GROUP. in phyloseq object?

.keep_readcount

keep readcount variable .READCOUNT. in phyloseq object?

.message_IDs message sample names of dropped variables?

.label_only if TRUE, the samples will NOT be filtered, just labelled with a new logical

variable .KEEP SAMPLE.

.keep_all_taxa keep all taxa after removing duplicates? If FALSE, the default, taxa are removed

if they never occur in any of the retained samples

Details

What happens when duplicated samples have exactly equal readcounts in method = "readcount"? The first encountered maximum is kept (in sample_data row order, like method = "first")

Value

phyloseq object

See Also

ps_filter for filtering samples by sample_data variables

Examples

```
data("dietswap", package = "microbiome")

dietswap
# let's pretend the dietswap data contains technical replicates from each subject
# we want to keep only one of them
ps_dedupe(dietswap, vars = "subject", method = "readcount", verbose = TRUE)

# contrived example to show identifying "duplicates" via the interaction of multiple columns
ps1 <- ps_dedupe(
    ps = dietswap, method = "readcount", verbose = TRUE,
    vars = c("timepoint", "group", "bmi_group")
)
phyloseq::sample_data(ps1)</pre>
```

ps_drop_incomplete 81

```
ps2 <- ps_dedupe(
   ps = dietswap, method = "first", verbose = TRUE,
   vars = c("timepoint", "group", "bmi_group")
)
phyloseq::sample_data(ps2)</pre>
```

ps_drop_incomplete

Deselect phyloseq samples with sample_data missings

Description

Check phyloseq object sample_data for missing values (NAs)

- specify which variables to check with vars argument, or check all
- · drop samples with any missings

Usage

```
ps_drop_incomplete(ps, vars = NA, verbose = FALSE)
```

Arguments

ps phyloseq with sample_data

vars vector of variable names to check for missings (or NA, which uses all variables

in sample data)

verbose message about number of samples dropped if verbose not FALSE, (and only if

> 0 samples dropped) and message about number of missing per variable in vars

if verbose = "max" (and message even if 0 samples dropped)

Details

This is a wrapper for stats::complete.cases function.

Value

phyloseq

See Also

```
ps_filter
```

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Examples

```
library(phyloseq)
data("enterotype", package = "phyloseq")
enterotype
ps_drop_incomplete(enterotype)
ps_drop_incomplete(enterotype, vars = "Enterotype", verbose = TRUE)
ps_drop_incomplete(enterotype, vars = "Sample_ID", verbose = TRUE)
ps_drop_incomplete(enterotype, vars = c("Enterotype", "Sample_ID"))
ps_drop_incomplete(enterotype, verbose = "max")
```

ps_filter

Filter phyloseq samples by sample_data variables

Description

Keep only samples with sample_data matching one or more conditions. By default this function also removes taxa which never appear in any of the remaining samples, by running tax_filter(min_prevalence = 1). You can prevent this taxa filtering with .keep_all_taxa = TRUE.

Usage

```
ps_filter(ps, ..., .target = "sample_data", .keep_all_taxa = FALSE)
```

Arguments

ps phyloseq object
... passed directly to dplyr::filter (see examples and ?dplyr::filter)
.target which slot of phyloseq to use for filtering by, currently only "sample_data" supported
.keep_all_taxa if FALSE (the default), remove taxa which are no longer present in the dataset after filtering

Details

Use ps_filter as you would use use dplyr::filter(), but with a phyloseq object!

Value

```
phyloseq object (with filtered sample_data)
```

See Also

```
filter explains better how to give arguments to this function tax_filter for filtering taxa (not samples)
```

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Examples

```
library(phyloseq)
library(dplyr)
data("enterotype", package = "phyloseq")
enterotype
sample_data(enterotype)[1:10, 1:5]
# keep only samples with seqtech not equal to sanger
ps1 <- ps_filter(enterotype, SeqTech != "Sanger")</pre>
ps1
sample_data(ps1)[1:10, 1:5]
# keep only samples with no NAs in any variables
ps2 <- enterotype %>% ps_filter(!if_any(everything(), is.na))
sample_data(ps2)[1:8, 1:8]
# ps2 is equivalent to dropping samples with incomplete sample_variables and tax_filtering 0s
ps3 <- enterotype %>%
  ps_drop_incomplete() %>%
  tax_filter(undetected = 0, use_counts = FALSE)
# we needed to set a low detection threshold because this example data is proportions
identical(ps2, ps3) # TRUE
# function will give warning if some of the otu_values are negative
# (which may happen when filtering data that has e.g. clr-transformed taxa abundances)
# as it attempts to discard any taxa that become always absent/0 after filtering (by default)
# set .keep_all_taxa = TRUE to avoid this filtering behaviour, which is unwanted in this case
enterotype %>%
  tax_transform("clr") %>%
  ps_get() %>%
  ps_filter(SeqTech == "Sanger", .keep_all_taxa = TRUE)
```

ps_join

Join a dataframe to phyloseq sample data

Description

You can use most types of join from the dplyr::*_join function family, including e.g. "inner", "left", "semi", "anti" (see details below). Defaults to type = "left" which calls left_join(), this supports x as a phyloseq and y as a dataframe. Most of the time you'll want "left" (adds variables with no sample filtering), or "inner" (adds variables and filters samples). This function simply:

- 1. extracts the sample_data from the phyloseq as a dataframe
- 2. performs the chosen type of join (with the given arguments)
- 3. filters the phyloseq if type = inner, semi or anti
- 4. reattaches the modified sample_data to the phyloseq and returns the phyloseq

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Usage

```
ps_join(
    x,
    y,
    by = NULL,
    match_sample_names = NULL,
    keep_sample_name_col = TRUE,
    sample_name_natural_join = FALSE,
    type = "left",
    .keep_all_taxa = FALSE
)
```

Arguments

```
Х
                  phyloseq (or dataframe)
                  dataframe (or phyloseq for e.g. type = "right")
У
by
                  A character vector of variables to join by (col must be present in both x and y or
                  paired via a named vector like c("xname" = "yname", etc.))
match_sample_names
                  match against the phyloseq sample_names by naming a variable in the additional
                  dataframe (this is in addition to any variables named in by)
keep_sample_name_col
                  should the column named in match_sample_names be kept in the returned phy-
                  loseq's sample_data? (only relevant if match_sample_names is not NULL)
sample_name_natural_join
                  if TRUE, use sample name AND all shared colnames to match rows (only rele-
                  vant if match_sample_names is not NULL, this arg takes precedence over any-
                  thing also entered in by arg)
                  name of type of join e.g. "left", "right", "inner", "semi" (see dplyr help pages)
type
.keep_all_taxa if FALSE (the default), remove taxa which are no longer present in the dataset
                  after filtering
```

Details

Mutating joins, which will add columns from a dataframe to phyloseq sample data, matching rows based on the key columns named in the by argument:

- "inner": includes all rows in present in both x and y.
- "left": includes all rows in x. (so x must be the phyloseq)
- "right": includes all rows in y. (so y must be the phyloseq)
- "full": includes all rows present in x or y. (will likely NOT work, as additional rows cannot be added to sample_data!)

If a row in x matches multiple rows in y (based on variables named in the by argument), all the rows in y will be added once for each matching row in x. This will cause this function to fail, as additional rows cannot be added to the phyloseq sample_data!

Filtering joins filter rows from x based on the presence or absence of matches in y:

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- "semi": return all rows from x with a match in y.
- "anti": return all rows from x without a match in y.

Value

phyloseq with modified sample_data (and possibly filtered)

See Also

```
ps_mutate for computing new variables from existing sample data
ps_select for selecting only some sample_data variables
https://www.garrickadenbuie.com/project/tidyexplain/ for an animated introduction to
joining dataframes
```

Examples

```
library(phyloseq)
data("enterotype", package = "phyloseq")
x <- enterotype
v <- data.frame(</pre>
  ID_var = sample_names(enterotype)[c(1:50, 101:150)],
  SeqTech = sample_data(enterotype)[c(1:50, 101:150), "SeqTech"],
  arbitrary_info = rep(c("A", "B"), 50)
)
\# simply match the new data to samples that exist in x, as default is a left_join
# where some sample names of x are expected to match variable ID_var in dataframe y
out1A <- ps_join(x = x, y = y, match_sample_names = "ID_var")
out1A
sample_data(out1A)[1:6, ]
# use sample_name and all shared variables to join
# (a natural join is not a type of join per se,
# but it indicates that all shared variables should be used for matching)
out1B <- ps_join(
  x = x, y = y, match_sample_names = "ID_var",
  sample_name_natural_join = TRUE, keep_sample_name_col = FALSE
)
out1B
sample_data(out1B)[1:6, ]
# if you only want to keep phyloseq samples that exist in the new data, try an inner join
# this will add the new variables AND filter the phyloseq
# this example matches sample names to ID_var and by matching the shared SeqTech variable
out1C <- ps_join(x = x, y = y, type = "inner", by = "SeqTech", match_sample_names = "ID_var")
out1C
sample_data(out1C)[1:6, ]
# the id variable is named Sample_ID in x and ID_var in y
```

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```
# semi_join is only a filtering join (doesn't add new variables but just filters samples in x)
out2A <- ps_join(x = x, y = y, by = c("Sample_ID" = "ID_var"), type = "semi")
out2A
sample_data(out2A)[1:6, ]

# anti_join is another type of filtering join
out2B <- ps_join(x = x, y = y, by = c("Sample_ID" = "ID_var"), type = "anti")
out2B
sample_data(out2B)[1:6, ]

# semi and anti joins keep opposite sets of samples
intersect(sample_names(out2A), sample_names(out2B))

# you can mix and match named and unnamed values in the `by` vector
# inner is like a combination of left join and semi join
out3 <- ps_join(x = x, y = y, by = c("Sample_ID" = "ID_var", "SeqTech"), type = "inner")
out3
sample_data(out3)[1:6, ]</pre>
```

ps_melt

Melt phyloseq data object into large data.frame (tibble)

Description

The ps_melt function is a specialized melt function for melting phyloseq objects (instances of the phyloseq class), usually for producing graphics with ggplot2. The naming conventions used in downstream phyloseq graphics functions have reserved the following variable names that should not be used as the names of sample_variables or taxonomic rank_names. These reserved names are c("Sample", "Abundance", "OTU"). Also, you should not have identical names for sample variables and taxonomic ranks. That is, the intersection of the output of the following two functions sample_variables, rank_names should be an empty vector (e.g. intersect(sample_variables(ps), rank_names(ps))). All of these potential name collisions are checked-for and renamed automatically with a warning. However, if you (re)name your variables accordingly ahead of time, it will reduce confusion and eliminate the warnings.

NOTE: Code and documentation copied (and very slightly modified) from an old version of speedyseq by Michael McLaren. speedyseq reimplements psmelt from phyloseq to use functions from the tidyr and dplyr packages. The name in microViz is changed to ps_melt for consistency with other functions.

Usage

```
ps_melt(ps)
```

Arguments

ps

(Required). An otu_table-class or phyloseq-class. Function most useful for phyloseq-class.

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Details

Note that "melted" phyloseq data is stored much less efficiently, and so RAM storage issues could arise with a smaller dataset (smaller number of samples/OTUs/variables) than one might otherwise expect. For common sizes of graphics-ready datasets, however, this should not be a problem. Because the number of OTU entries has a large effect on the RAM requirement, methods to reduce the number of separate OTU entries – for instance by agglomerating OTUs based on phylogenetic distance using tip_glom – can help alleviate RAM usage problems. This function is made user-accessible for flexibility, but is also used extensively by plot functions in phyloseq.

Value

A tibble class data frame.

See Also

psmelt

Examples

```
library(ggplot2)
library(phyloseq)
data("GlobalPatterns")
gp_ch <- subset_taxa(GlobalPatterns, Phylum == "Chlamydiae")</pre>
mdf <- ps_melt(gp_ch)</pre>
mdf2 <- psmelt(gp_ch) # slower
# same dataframe, except with somewhat different row orders
dplyr::all_equal(tibble::as_tibble(mdf), mdf2, convert = TRUE) # TRUE
nrow(mdf2)
ncol(mdf)
colnames(mdf)
head(rownames(mdf))
p <- ggplot(mdf, aes(x = SampleType, y = Abundance, fill = Genus))</pre>
p <- p + geom_bar(color = "black", stat = "identity", position = "stack")</pre>
# This example plot doesn't make any sense
print(p + coord_flip())
# TODO replace this...
```

ps_mutate

Modify or compute new sample_data in phyloseq object

Description

Add or compute new phyloseq sample_data variables. Uses dplyr::mutate() syntax.

Usage

```
ps_mutate(ps, ...)
```

88 ps_otu2samdat

Arguments

```
ps phyloseq object with sample data
... passed straight to dplyr::mutate (see examples and dplyr::mutate help)
```

Value

phyloseq object with modified sample_data

See Also

```
tax_mutate for manipulation of tax_table variables mutate
```

Examples

```
library(phyloseq)
library(dplyr)
data("enterotype")
sample_data(enterotype)[1:10, ]
months_in_year <- 12</pre>
ps <- enterotype %>%
  ps_mutate(
   age_months = Age * months_in_year,
   IDs_match = as.character(Sample_ID) == as.character(SampleID),
    placeholder = "Word"
sample_data(ps)[1:10, ]
# Using the dplyr::across functionality is also possible
ps <- ps %>%
  ps_mutate(
   dplyr::across(where(is.factor), toupper),
   another_var = TRUE,
   SeqTech = NULL # deletes SeqTech variable
  )
head(sample_data(ps))
```

ps_otu2samdat

Copy phyloseq otu_table data to sample_data

Description

Copy phyloseq otu_table data to sample_data

ps_reorder 89

Usage

```
ps_otu2samdat(ps, taxa = NULL)
```

Arguments

Value

phyloseq with augmented sample_data

Examples

```
library(phyloseq)
data("dietswap", package = "microbiome")

ps <- dietswap %>% ps_otu2samdat("Akkermansia")
sample_variables(ps)

# or if you do not specify any taxa, all are copied
ps_all <- dietswap %>% ps_otu2samdat()
sample_variables(ps_all)[1:15]

# this could be useful for colouring ordination plots, for example
ps %>%
    ps_mutate(log_akkermansia = log(Akkermansia)) %>%
    dist_calc("bray") %>%
    ord_calc(method = "PCoA") %>%
    ord_plot(
    colour = "log_akkermansia",
        size = 3, shape = "nationality"
)
```

ps_reorder

Set order of samples in phyloseq object

Description

Manually set order of samples by specifying samples names in desired order.

Usage

```
ps_reorder(ps, sample_order)
```

90 ps_select

Arguments

```
ps phyloseq sample_order names or current numerical indices of samples in desired order
```

Details

Ordering of samples in a phyloseq is controlled from the otu_table slot!

Value

phyloseq

See Also

```
ps_arrange for arranging samples by sample_data variables (or otu_table)
ps_seriate for arranging samples by microbiome similarity
ps_filter for keeping only some samples, based on sample_data
```

Examples

```
library(phyloseq)
data("dietswap", package = "microbiome")
dietswap %>%
  sample_data() %>%
  head(8)
new_order <- rev(sample_names(dietswap))</pre>
dietswap %>%
  ps_reorder(new_order) %>%
  sample_data() %>%
  head(8)
# random ordering with numbers
set.seed(1000)
random_order <- sample(1:nsamples(dietswap))</pre>
dietswap %>%
  ps_reorder(random_order) %>%
  sample_data() %>%
  head(8)
```

ps_select

Select phyloseq sample_data using dplyr::select syntax

Description

Simple selection of phyloseq sample_data variables, might be useful for printing reduced sample_data, or inside other functions

ps_seriate 91

Usage

```
ps_select(ps, ...)
```

Arguments

```
ps phyloseq with sample_data... passed straight to dplyr::select
```

Value

phyloseq object

Examples

```
library(phyloseq)
library(dplyr)
data("enterotype", package = "phyloseq")
head(sample_data(enterotype))
enterotype %>%
   ps_select(!contains("Sample")) %>%
   sample_data() %>%
   head()
```

ps_seriate

Arrange samples in a phyloseq by microbiome similarity

Description

Uses seriation methods from seriation::seriate and often dist_calc (depending on if seriation method requires a distance matrix)

Usage

```
ps_seriate(
   ps,
   method = "OLO_ward",
   dist = "bray",
   tax_transform = "identity",
   add_variable = FALSE,
   rank = NA
)
```

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Arguments

ps phyloseq object

method seriation method for ordering samples, from seriation::seriate

distance method for dist_calc (only used if required for particular seriation method!)

tax_transform transformation to apply before seriation or any distance calculation

add_variable add a variable to the sample data indicating seriation order

rank taxonomic rank to aggregate at, before seriation, NA for no aggregation

Value

phyloseq

See Also

```
ps_arrange ps_reorder
```

Examples

```
library(phyloseq)
data("dietswap", package = "microbiome")

dietswap %>%
    sample_data() %>%
    head(8)

dietswap %>%
    tax_agg("Genus") %>%
    ps_get() %>%
    ps_seriate(method = "OLO_ward", dist = "bray") %>%
    sample_data() %>%
    head(8)
```

ps_sort_ord

Sort phyloseq samples by ordination axes scores

Description

ps_sort_ord reorders samples in a phyloseq object based on their relative position on 1 or 2 ordination axes.

ord_order_samples gets the sample_names in order from the ordination contained in a psExtra. This is used internally by ps_sort_ord

If 2 axes given, the samples are sorted by anticlockwise rotation around the selected ordination axes, starting on first axis given, upper right quadrant. (This is used by ord_plot_iris.)

If 1 axis is given, samples are sorted by increasing value order along this axis. This could be used to arrange samples on a rectangular barplot in order of appearance along a parallel axis of a paired ordination plot.

ps_sort_ord 93

Usage

```
ps_sort_ord(ps, ord, axes = 1:2, scaling = 2)
ord_order_samples(ord, axes = 1:2, scaling = 2)
```

Arguments

ps phyloseq object to be sorted ord psExtra with ordination object

axes which axes to use for sorting? numerical vector of length 1 or 2

scaling Type 2, or type 1 scaling. For more info, see https://sites.google.com/

site/mb3gustame/constrained-analyses/redundancy-analysis. Either "species"
or "site" scores are scaled by (proportional) eigenvalues, and the other set of

scores is left unscaled (from ?vegan::scores.cca)

Value

```
ps_sort_ord returns a phyloseq
ord_order_samples returns a character vector
```

See Also

- These functions were created to support ordering of samples on ord_plot_iris
- tax_sort_ord for ordering taxa in phyloseq by ordination

Examples

```
# attach other necessary packages
library(ggplot2)
# example data
ibd <- microViz::ibd %>%
  tax_filter(min_prevalence = 2) %>%
  tax_fix() %>%
  phyloseq_validate()
# create numeric variables for constrained ordination
ibd <- ibd %>%
  ps_mutate(
    ibd = as.numeric(ibd == "ibd"),
    steroids = as.numeric(steroids == "steroids"),
   abx = as.numeric(abx == "abx"),
    female = as.numeric(gender == "female"),
    # and make a shorter ID variable
    id = stringr::str_remove_all(sample, "^[0]{1,2}|-[A-Z]+")
# create an ordination
ordi <- ibd %>%
```

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```
tax_transform("clr", rank = "Genus") %>%
  ord_calc()
ord_order_samples(ordi, axes = 1) %>% head(8)
ps_sort_ord(ibd, ordi, axes = 1) %>%
  phyloseq::sample_names() %>%
  head(8)
p1 <- ord_plot(ordi, colour = "grey90", plot_taxa = 1:8, tax_vec_length = 1) +
  geom_text(aes(label = id), size = 2.5, colour = "red")
b1 <- ibd %>%
  ps_sort_ord(ord = ordi, axes = 1) %>%
  comp_barplot(
    tax_level = "Genus", n_taxa = 12, label = "id",
   order_taxa = ord_order_taxa(ordi, axes = 1),
   sample_order = "asis"
  ) +
  theme(axis.text.x = element_text(angle = 90, hjust = 1))
patchwork::wrap_plots(p1, b1, ncol = 1)
# constrained ordination example (and match vertical axis) #
cordi <- ibd %>%
  tax_transform("clr", rank = "Genus") %>%
  ord_calc(
    constraints = c("steroids", "abx", "ibd"), conditions = "female",
    scale_cc = FALSE
cordi %>% ord_plot(plot_taxa = 1:6, axes = 2:1)
```

sampleAnnotation

Helper to specify a HeatmapAnnotation for samples in comp_heatmap

Description

Helper to specify a HeatmapAnnotation for samples in comp_heatmap

Usage

```
sampleAnnotation(
    ...,
    name,
    annotation_legend_param = list(),
    show_legend = TRUE,
    gp = grid::gpar(col = NA),
    border = FALSE,
    gap = grid::unit(2, "mm"),
```

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```
show_annotation_name = TRUE,
annotation_label = NULL,
annotation_name_gp = grid::gpar(),
annotation_name_offset = NULL,
annotation_name_rot = NULL,
annotation_name_align = FALSE,
annotation_name_side = "auto",
.data = NULL,
.samples = NULL,
.side = NULL
```

Arguments

... Name-value pairs where the names correspond to annotation names and values

are the output of sample annotation functions such as anno_sample(), or manu-

ally specified AnnotationFunction objects

name Name of the heatmap annotation, optional.

annotation_legend_param

 $A \ list which contains \ parameters \ for \ annotation \ legends. \ See \ color_mapping_legend, ColorMapping_measurements \ color_mapping_measurements \ color_mapping_legend, ColorMapping_measurements \ color_mapping_legend, ColorMapping_measurements \ color_mapping_legend, ColorMapping_measurements \ color_mapping_measurements \ color_mapping_measurement$

for all possible options.

show_legend Whether show annotation legends. The value can be one single value or a vector.

gp Graphic parameters for simple annotations (with fill parameter ignored).

border border of single annotations.

gap Gap between annotations. It can be a single value or a vector of unit objects.

show_annotation_name

Whether show annotation names? For column annotation, annotation names are drawn either on the left or the right, and for row annotations, names are draw

either on top or at the bottom. The value can be a vector.

annotation_label

Labels for the annotations. By default it is the same as individual annotation

names.

annotation_name_gp

Graphic parameters for annotation names. Graphic parameters can be vectors.

annotation_name_offset

Offset to the annotation names, a unit object. The value can be a vector.

annotation_name_rot

Rotation of the annotation names. The value can be a vector.

annotation_name_align

Whether to align the annotation names.

annotation_name_side

Side of the annotation names.

.data OPTIONAL phyloseq or psExtra, only set this to override use of same data as

in heatmap

. samples OPTIONAL selection vector of sample names, only set this if providing .data

argument to override default

.side

OPTIONAL string, indicating the side for the variable annotations: only set this to override default

Value

HeatmapAnnotation object

See Also

taxAnnotation()

Examples

```
library("ComplexHeatmap")
data("ibd", package = "microViz")
psq <- tax_filter(ibd, min_prevalence = 5)</pre>
psq <- tax_mutate(psq, Species = NULL)</pre>
psq <- tax_fix(psq)</pre>
psq <- tax_agg(psq, rank = "Family")</pre>
taxa <- tax_top(psq, n = 15, rank = "Family")</pre>
samples <- phyloseq::sample_names(psq)</pre>
set.seed(42) # random colours used in first example
# sampleAnnotation returns a function that takes data, samples, and which
fun <- sampleAnnotation(</pre>
  gap = grid::unit(2.5, "mm"),
  Dis1 = anno_sample(var = "DiseaseState"),
  IBD = anno_sample_cat(var = "ibd"),
  Dis2 = anno_sample_cat(var = "DiseaseState", col = 1:4)
)
# manually specify the sample annotation function by giving it data etc.
heatmapAnnoFunction <- fun(.data = psq, .side = "top", .samples = samples)
# draw the annotation without a heatmap, you will never normally do this!
grid.newpage()
vp <- viewport(width = 0.65, height = 0.75)</pre>
pushViewport(vp)
draw(heatmapAnnoFunction)
pushViewport(viewport(x = 0.7, y = 0.6))
draw(attr(heatmapAnnoFunction, "Legends"))
```

scale_shape_girafe_filled

Filled shapes for ggiraph interactive plots

Description

Generates a custom ggplot2 shape scale, as used in ord_explore's ordination. Uses filled shapes, therefore fill aesthetic must be set, in addition to colour, to have filled shapes. Points with NA values for the shape variable are shown as hollow circles.

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Usage

```
scale_shape_girafe_filled()
```

Details

Composite shapes e.g. number 7 "square cross" cause ggiraph interactive plots to fail when a variable shape and tooltip is set.

Shapes used are, in order: "circle filled", "triangle filled", "square filled", "diamond filled", and "triangle down filled"

Value

ggplot2 Scale object

Examples

```
microViz::ibd %>%
  tax_fix() %>%
  phyloseq_validate() %>%
  tax_transform(rank = "Genus", trans = "clr") %>%
  ord_calc(
    method = "PCA"
) %>%
  ord_plot(
    axes = c(1, 2),
    plot_taxa = 1:6,
    colour = "DiseaseState", fill = "DiseaseState",
    shape = "circle", alpha = 0.5,
    size = 3
) +
  scale_shape_girafe_filled()
```

shao19

Gut microbiota relative abundance data from Shao et al. 2019

Description

A phyloseq dataset containing 1644 gut microbiome samples. Fecal samples were collected from 596 infants. 175 mothers also provided fecal samples.

Usage

shao19

Format

large phyloseq object

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Data acquisition

The processed (metaphlan3) relative abundance data were obtained using curatedMetagenomic-Data package version 3.4.2. https://waldronlab.io/curatedMetagenomicData/articles/available-studies.html

A small amount of data cleaning was performed after download:

- 1. to make sample metadata more user friendly
- 2. to fill gaps in the taxonomy table.

Source

Stunted microbiota and opportunistic pathogen colonization in caesarean-section birth. doi:10.1038/s4158601915601

stat_chull

Draw convex hull for a set of points on a ggplot

Description

Draws a (convex) polygon around the outermost points of a set of points. Useful as a visual aid for identifying groups of points on a scatterplot, such as an ordination plot.

Usage

```
stat_chull(
  mapping = NULL,
  data = NULL,
  geom = "polygonHollow",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

Arguments

mapping

Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.

data

The data to be displayed in this layer. There are three options:

If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot().

A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created.

stat_chull 99

A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. ~ head(.x, 10)).

geom

The geometric object to use to display the data for this layer. When using a stat_*() function to construct a layer, the geom argument can be used to override the default coupling between stats and geoms. The geom argument accepts the following:

- A Geom ggproto subclass, for example GeomPoint.
- A string naming the geom. To give the geom as a string, strip the function name of the geom_prefix. For example, to use geom_point(), give the geom as "point".
- For more information and other ways to specify the geom, see the layer geom documentation.

position

A position adjustment to use on the data for this layer. This can be used in various ways, including to prevent overplotting and improving the display. The position argument accepts the following:

- The result of calling a position function, such as position_jitter(). This method allows for passing extra arguments to the position.
- A string naming the position adjustment. To give the position as a string, strip the function name of the position_ prefix. For example, to use position_jitter(), give the position as "jitter".
- For more information and other ways to specify the position, see the layer position documentation.

na.rm

If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed.

show.legend

logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

inherit.aes

If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders().

. . .

Other arguments passed on to layer()'s params argument. These arguments broadly fall into one of 4 categories below. Notably, further arguments to the position argument, or aesthetics that are required can *not* be passed through Unknown arguments that are not part of the 4 categories below are ignored.

- Static aesthetics that are not mapped to a scale, but are at a fixed value and apply to the layer as a whole. For example, colour = "red" or linewidth = 3. The geom's documentation has an **Aesthetics** section that lists the available options. The 'required' aesthetics cannot be passed on to the params. Please note that while passing unmapped aesthetics as vectors is technically possible, the order and required length is not guaranteed to be parallel to the input data.
- When constructing a layer using a stat_*() function, the ... argument can be used to pass on parameters to the geom part of the layer. An example of this is stat_density(geom = "area", outline.type = "both"). The geom's documentation lists which parameters it can accept.

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- Inversely, when constructing a layer using a geom_*() function, the ... argument can be used to pass on parameters to the stat part of the layer.
 An example of this is geom_area(stat = "density", adjust = 0.5). The stat's documentation lists which parameters it can accept.
- The key_glyph argument of layer() may also be passed on through This can be one of the functions described as key glyphs, to change the display of the layer in the legend.

Details

This is a ggplot2 extension - slightly modified from the original code found here:

```
https://CRAN.r-project.org/package=ggplot2/vignettes/extending-ggplot2.html
```

See Also

```
ggplot2::stat_ellipse
ord_plot
```

Examples

```
library(ggplot2)
microViz::ibd %>%
  tax_fix() %>%
  tax_transform(rank = "Genus", trans = "clr") %>%
  ord_calc(method = "PCA") %>%
  ord_plot(colour = "DiseaseState", shape = "DiseaseState", alpha = 0.5) +
  stat_chull(aes(colour = DiseaseState))

microViz::ibd %>%
  tax_fix() %>%
  tax_transform(rank = "Genus", trans = "clr") %>%
  ord_calc(method = "PCA") %>%
  ord_plot(colour = "DiseaseState", shape = "DiseaseState", alpha = 0.5) +
  stat_chull(aes(colour = DiseaseState, fill = DiseaseState), alpha = 0.1)
```

taxAnnotation

Helper to specify a HeatmapAnnotation for taxa

Description

Helper to specify a HeatmapAnnotation for taxa

Usage

```
taxAnnotation(
    ...,
    name,
    annotation_legend_param = list(),
```

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```
show_legend = TRUE,
  gp = grid::gpar(col = NA),
  border = FALSE,
  gap = grid::unit(2, "mm"),
  show_annotation_name = TRUE,
  annotation_label = NULL,
  annotation_name_gp = grid::gpar(),
  annotation_name_offset = NULL,
  annotation_name_rot = NULL,
  annotation_name_align = TRUE,
  annotation_name_side = "auto",
  .data = NULL,
  .taxa = NULL,
  .side = NULL
)
```

Arguments

Name-value pairs where the names correspond to annotation names and values are the output of taxon annotation functions such as anno_tax_prev() or manu-

ally specified AnnotationFunction objects

Name of the heatmap annotation, optional. name

annotation_legend_param

A list which contains parameters for annotation legends. See color_mapping_legend, ColorMapping_me

for all possible options.

show_legend Whether show annotation legends. The value can be one single value or a vector.

Graphic parameters for simple annotations (with fill parameter ignored). gp

border border of single annotations.

Gap between annotations. It can be a single value or a vector of unit objects. gap

show_annotation_name

Whether show annotation names? For column annotation, annotation names are drawn either on the left or the right, and for row annotations, names are draw

either on top or at the bottom. The value can be a vector.

annotation_label

Labels for the annotations. By default it is the same as individual annotation names.

annotation_name_gp

Graphic parameters for annotation names. Graphic parameters can be vectors.

annotation_name_offset

Offset to the annotation names, a unit object. The value can be a vector.

annotation_name_rot

Rotation of the annotation names. The value can be a vector.

annotation_name_align

Whether to align the annotation names.

annotation_name_side

Side of the annotation names.

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.data	OPTIONAL phyloseq or psExtra, only set this to override use of same data as in heatmap
.taxa	OPTIONAL selection vector of taxa (names, numbers or logical), only set this if providing .data argument to override default
.side	OPTIONAL string, indicating the side the taxa annotation should be placed: only set this to override default

Value

HeatmapAnnotation object

Examples

```
library("ComplexHeatmap")
data("ibd", package = "microViz")
psq <- tax_filter(ibd, min_prevalence = 5)</pre>
psq <- tax_mutate(psq, Species = NULL)</pre>
psq <- tax_fix(psq)</pre>
psq <- tax_agg(psq, rank = "Family")</pre>
taxa <- tax_top(psq, n = 15, rank = "Family")</pre>
customAxis <- list(labels_rot = 0, at = c(0, 0.5, 1))
# makes a function that takes data, taxa and which (at minimum)
fun <- taxAnnotation(</pre>
  gap = grid::unit(2.5, "mm"),
  Prev. = anno_tax_prev(axis_param = customAxis, ylim = c(0, 1), extend = 0),
  `Prop. Abd.` = anno_tax_box(size = unit(40, "mm"), axis_param = customAxis),
  `Log10p Abd.` = anno_tax_density(type = "heatmap")
)
# manually specify the prevalence barplot function by giving it data etc.
heatmapAnnoFunction <- fun(.data = psq, .side = "top", .taxa = taxa)
# draw the annotation without a heatmap, you will never normally do this!
grid.newpage()
vp <- viewport(width = 0.65, height = 0.75)</pre>
pushViewport(vp)
draw(heatmapAnnoFunction)
# try again as a row annotation
grid.newpage()
pushViewport(vp)
draw(fun(.data = psq, .side = "right", .taxa = rev(taxa)))
```

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Description

Mostly you will not have to use these functions directly: instead call taxatree_plots with the output of taxatree_stats

- taxatree_nodes creates taxon nodes and calculates a summary statistic about each taxon (given by fun). Takes a psExtra or phyloseq object.
- taxatree_edges uses the output of taxatree_nodes to create a dataframe of edges.

Usage

```
taxatree_nodes(
  ps,
  fun = list(sum = sum),
  ranks = "all",
   .sort = NULL,
   .use_counts = TRUE
)
taxatree_edges(nodes_df)
```

Arguments

ps	phyloseq object or psExtra
fun	function to calculate for each taxon/node
ranks	selection of taxonomic ranks to make nodes for ("all", or names)
.sort	sort nodes by "increasing" or "decreasing" values of fun function result
.use_counts	use count data if available (instead of transformed data)
nodes_df	dataframe output from taxatree_nodes

Details

taxatree_nodes makes nodes for taxa at all ranks or for a list of consecutive ranks (plus a root rank if tree is not rooted).

taxatree_label

Add logical label column to taxatree_stats dataframe

Description

 $taxatree_label\ is\ used\ internally\ by\ taxatree_plotkey,\ but\ can\ also\ be\ used\ prior\ to\ taxatree_plots\ to\ label\ those\ plots\ directly$

```
... arguments are passed to dplyr::filter(), so all filter syntax can be used.
```

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Usage

```
taxatree_label(
  data,
  ...,
  .label_var = "label",
  .node_fun = list(prevalence = prev)
)
```

Arguments

data	psExtra (or phyloseq)
	REQUIRED logical conditions for labelling e.g. rank == "Phylum", p.value < 0.1 taxon %in% listOfTaxa
.label_var	name of label indicator variable to be created. If you change this, beware that taxatree_plotkey will not work, you will need to called taxatree_plot_label with
.node_fun	named list of length 1 providing taxatree_nodes fun arg. (name of list iterm is available for use in)

Details

If taxatree_stats missing (or if data is a phyloseq) it will create a plain taxatree_stats dataframe using only taxatree_nodes

node_fun can also be a precalculated dataframe (output of taxatree_nodes) but you should probably not use this option. This is used internally for efficiency inside taxatree_plotkey()

Value

psExtra with (modified) taxatree_stats dataframe

Examples

```
# simple example with plain phyloseq input
data("dietswap", package = "microbiome")
labelled <- dietswap %>%
    tax_prepend_ranks() %>%
    taxatree_label(rank == "Phylum", prevalence > 0.1)

# Note that "prevalence" column was available in data
# because it is created by `taxatree_nodes()` using the named function
# provided to the `node_fun` argument

# psExtra is returned
labelled
# notice how both conditions must be met for label column to be TRUE
labelled %>% taxatree_stats_get()
```

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taxatree_models

Statistical modelling for individual taxa across multiple ranks

Description

taxatree_models runs tax_model on every taxon at multiple taxonomic ranks (you choose which ranks with the plural ranks argument). It returns the results as a named nested list of models attached to a psExtra object. One list per rank, one model per taxon at each rank.

The result can then be used with taxatree_models2stats to extract a dataframe of statistics for use with taxatree_plots.

Usage

```
taxatree_models(
  ps,
  ranks = NULL,
  type = "lm",
  variables = NULL,
  formula = NULL,
  use_future = FALSE,
  checkVars = TRUE,
  checkNA = "warning",
  verbose = "rank",
  trans = "identity",
  trans_args = list(),
  ...
)
```

Arguments

ps	phyloseq object or psExtra
ranks	vector of rank names at which to aggregate taxa for modelling
type	name of regression modelling function, or the function itself
variables	vector of variable names, to be used as model formula right hand side. If variables is a list, not a vector, a model is fit for each entry in list.
formula	Right hand side of a formula, as a formula object or character string. Or a list of these. (alternative to variables argument, do not provide both)
use_future	if TRUE parallel processing with future is possible, see details of ?tax_model.
checkVars	check variance of variables?
checkNA	check variables for NAs?
verbose	message about progress: "rank" only notifies which rank is being processed; TRUE notifies you about each taxon being processed; FALSE for no messages.
trans	name of tax_transform transformation to apply to aggregated taxa before fitting statistical models

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```
trans_args named list of any additional arguments to tax_transform e.g. list(zero_replace = "halfmin")

... extra arguments are passed directly to modelling function
```

See Also

```
tax_model for more details and examples
taxatree_plots for how to plot the output of taxatree_models
```

Description

Runs a function e.g. broom::tidy on a list of models, i.e. the output of $taxatree_models$ or tax_model

Usage

```
taxatree_models2stats(data, fun = "auto", ..., .keep_models = FALSE)
tax_models2stats(data, rank = NULL, fun = "auto", ..., .keep_models = FALSE)
```

Arguments

data	psExtra with attached tax_models or taxatree_models list, or just the list of models
fun	function to assist extraction of stats dataframe from models, or "auto"
	extra arguments passed to fun
.keep_models	should the models list be kept in the psExtra output?
rank	string naming rank at which tax_model was run (needed if data is a list)

Value

data.frame, attached to psExtra

Functions

- taxatree_models2stats(): Extract stats from list or psExtra output of taxatree_models
- tax_models2stats(): Extract stats from list or psExtra output of tax_model

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Examples

```
# This example is an abbreviated excerpt from article on taxon modelling on
# the microViz documentation website
library(dplyr)
data("ibd", package = "microViz")
# We'll keep only the Ulcerative Colitis and Healthy Control samples, to
# simplify the analyses for this example. We'll also remove the Species
# rank information, as most OTUs in this dataset are not assigned to a
# species. We'll also use `tax_fix` to fill any gaps where the Genus is
# unknown, with the family name or whatever higher rank classification is
# known.
phylo <- ibd %>%
 ps_filter(DiseaseState %in% c("UC", "nonIBD")) %>%
 tax_mutate(Species = NULL) %>%
 tax_fix()
# Let's make some sample data variables that are easier to use and compare
# in the statistical modelling ahead. We will convert dichotomous
# categorical variables into similar binary variables (values: 1 for true,
# or 0 for false). We will also scale and center the numeric variable for
# age.
phylo <- phylo %>%
 ps_mutate(
   UC = ifelse(DiseaseState == "UC", yes = 1, no = 0),
   female = ifelse(gender == "female", yes = 1, no = 0),
   antibiotics = ifelse(abx == "abx", yes = 1, no = 0),
   steroids = ifelse(steroids == "steroids", yes = 1, no = 0),
   age_scaled = scale(age, center = TRUE, scale = TRUE)
lm_models <- phylo %>%
 tax_fix() %>%
 tax_prepend_ranks() %>%
 tax_transform("compositional", rank = "Genus") %>%
 tax_filter(min_prevalence = 0.1, use_counts = TRUE) %>%
 taxatree_models(
   type = lm,
   trans = "log2", trans_args = list(zero_replace = "halfmin"),
   ranks = c("Phylum", "Class", "Genus"),
   variables = c("UC", "female", "antibiotics", "steroids", "age_scaled")
lm_stats <- lm_models %>% taxatree_models2stats()
# inspect the psExtra returned, now with taxatree_stats dataframe added
lm_stats
# inspect the dataframe itself
```

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```
lm_stats %>% taxatree_stats_get()

# keep the models on the psExtra object
lm_models %>% taxatree_models2stats(.keep_models = TRUE)

# you can adjust the p values with taxatree_stats_p_adjust

# you can plot the results with taxatree_plots
```

taxatree_plotkey

Draw labelled key to accompany taxatree_plots

Description

Draw labelled key to accompany taxatree_plots

Usage

```
taxatree_plotkey(
  data,
  . . . ,
 size_stat = list(prevalence = prev),
 node\_size\_range = c(1.5, 5),
  edge_width_range = node_size_range * 0.8,
  size_guide = "none",
  size_trans = "identity",
  colour = "lightgrey",
  edge_alpha = 0.7,
  title = "Key",
  title_size = 14,
  taxon_renamer = identity,
  .combine_label = any,
  .draw_label = TRUE,
  .calc_label = TRUE,
  layout = "tree",
  layout_seed = NA,
  circular = identical(layout, "tree"),
  node_sort = NULL,
 add_circles = isTRUE(circular),
  drop_ranks = TRUE
)
```

Arguments

```
data psExtra (or phyloseq)
... logical conditions for labelling e.g. rank == "Phylum", p.value < 0.1 | taxon %in% listOfTaxa
```

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size_stat named list of length 1, giving function calculated for each taxon, to determine the size of nodes (and edges). Name used as size legend title. node_size_range min and max node sizes, decrease to avoid node overlap edge_width_range min and max edge widths size_guide guide for node sizes, try "none", "legend" or ggplot2::guide_legend() transformation for size scale you can use (the name of) any transformer from the size_trans scales package, such as "identity", "log1p", or "sqrt" fixed colour and fill of nodes and edges colour edge_alpha fixed alpha value for edges title title of plot (NULL for none) font size of title title_size taxon_renamer function that takes taxon names and returns modified names for labels .combine_label all or any: function to combine multiple logical "label" values for a taxon (relevant if taxatree_stats already present in data) .draw_label should labels be drawn, or just the bare tree, set this to FALSE if you want to draw customised labels with taxatree_plot_labels afterwards .calc_label if you already set labels with taxatree_label: set this to FALSE to use / avoid overwriting that data (ignores . . . if FALSE) layout any ggraph layout, default is "tree" layout_seed any numeric, required if a stochastic igraph layout is named circular should the layout be circular? node sort sort nodes by "increasing" or "decreasing" size? NULL for no sorting. Use tax_sort() before taxatree_plots() for finer control.

Value

ggplot

add_circles

drop_ranks

Examples

```
library(ggplot2)
# Normally you make a key to accompany taxatree_plots showing stats results
# So see ?taxatree_plots examples too!
#
# You can also use the taxatree key visualization just to help understand your
# tax_table contents and hierarchical structure
shao19 %>%
    ps_filter(family_id %in% 1:5) %>%
    tax_filter(min_prevalence = 7) %>%
    tax_fix() %>%
```

add faint concentric circles to plot, behind each rank?

drop ranks from tree if not included in stats dataframe

```
tax_agg("genus") %>%
  tax_prepend_ranks() %>%
 taxatree_plotkey(rank %in% c("phylum", "genus"))
# # Let's look at some of the most prevalent Actinobacteria
actinoOnlyPhylo <- shao19 %>%
 tax_select("Actinobacteria", ranks_searched = "phylum") %>%
 tax_filter(min_prevalence = 100)
actinoOnlyPhylo %>%
  tax_mutate(order = NULL) %>%
  tax_sort(by = "prev", at = "species", tree_warn = FALSE) %>%
  taxatree_plotkey(
   circular = FALSE, rank != "genus",
    .draw_label = FALSE, # suppress drawing now so we can make custom labels after
    size_guide = "legend", colour = "skyblue", edge_alpha = 0.2
 ) %>%
  taxatree_plot_labels(
    circular = FALSE, # be sure you match the plotkey layout
    fun = geom_text, fontface = "bold.italic", size = 2.5, hjust = 0,
   nudge_y = 0.1 \# nudge y if you want to nudge x (due to coord_flip!)
 ) +
 coord_flip(clip = "off") +
 scale_y_reverse(expand = expansion(add = c(0.2, 2))) +
 theme(legend.position = c(0.15, 0.25), legend.background = element_rect())
# You can even choose other tree layouts from igraph (e.g. kk)
# and configure multiple styles of custom labels on one plot
set.seed(1) # set seed for reproducibility of ggrepel label positions
actinoOnlyPhylo %>%
 tax_mutate(order = NULL) %>%
 taxatree_label(rank == "family", .label_var = "family_lab") %>%
 taxatree_label(rank == "species", .label_var = "species_lab") %>%
 taxatree_plotkey(
   circular = FALSE, rank != "genus", layout = "kk",
    .draw_label = FALSE, # important not to draw the default labels
    colour = "skyblue", edge_alpha = 0.2
 ) %>%
 taxatree_plot_labels(
   circular = FALSE, label_var = "family_lab", fun = geom_label,
    fontface = "bold.italic", colour = "orange", fill = "black", size = 3
 ) %>%
  taxatree_plot_labels(
    circular = FALSE, label_var = "species_lab", fun = ggrepel::geom_text_repel,
    fontface = "bold.italic", size = 2, force = 20, max.time = 0.1
```

Description

- Uses a psExtra object to make a tree graph structure from the taxonomic table.
- Then adds statistical results stored in "taxatree_stats" of psExtra data
- You must use taxatree_models() first to generate statistical model results.
- You can adjust p-values with taxatree_stats_p_adjust()

Usage

```
taxatree_plots(
  data,
  colour_stat = "estimate",
  palette = "Green-Brown",
  reverse_palette = FALSE,
  colour_lims = NULL,
  colour_oob = scales::oob_squish,
  colour_trans = "abs_sqrt",
  size_stat = list(prevalence = prev),
  node_size_range = c(1, 4),
  edge_width_range = node_size_range * 0.8,
  size_guide = "legend",
  size_trans = "identity",
  sig_stat = "p.value",
  sig_{threshold} = 0.05,
  sig_shape = "circle filled",
  sig_size = 0.75,
  sig_stroke = 0.75,
  sig_colour = "white",
  edge_alpha = 0.7,
  vars = "term",
  var_renamer = identity,
  title_size = 10,
  layout = "tree",
  layout_seed = NA,
  circular = identical(layout, "tree"),
  node_sort = NULL,
  add_circles = isTRUE(circular),
  drop_ranks = TRUE,
 11 = if (palette == "Green-Brown") 10 else NULL,
 12 = if (palette == "Green-Brown") 85 else NULL,
  colour_na = "grey35"
)
```

Arguments

```
data psExtra with taxatree_stats, e.g. output of taxatree_models2stats()
colour_stat name of variable to scale colour/fill of nodes and edges
palette diverging hcl colour palette name from colorspace::hcl_palettes("diverging")
```

reverse_palette

reverse direction of colour palette?

colour_lims limits of colour and fill scale, NULL will infer lims from range of all data

colour_oob scales function to handle colour_stat values outside of colour_lims (default sim-

ply squishes "out of bounds" values into the given range)

colour_trans name of transformation for colour scale: default is "abs_sqrt", the square-root of

absolute values, but you can use the name of any transformer from the scales

package, such as "identity" or "exp"

size_stat named list of length 1, giving function calculated for each taxon, to determine

the size of nodes (and edges). Name used as size legend title.

node_size_range

min and max node sizes, decrease to avoid node overlap

edge_width_range

min and max edge widths

size_guide guide for node sizes, try "none", "legend" or ggplot2::guide_legend()

size_trans transformation for size scale you can use (the name of) any transformer from the

scales package, such as "identity", "log1p", or "sqrt"

sig_stat name of variable indicating statistical significance

sig_threshold value of sig_stat variable indicating statistical significance (below this)

sig_shape fixed shape for significance marker sig_size fixed size for significance marker

sig_stroke fixed stroke width for significance marker

sig_colour fixed colour for significance marker (used as fill for filled shapes)

edge_alpha fixed alpha value for edges

vars name of column indicating terms in models (one plot made per term)

var_renamer function to rename variables for plot titles

title_size font size of title

layout any ggraph layout, default is "tree"

layout_seed any numeric, required if a stochastic igraph layout is named

circular should the layout be circular?

node_sort sort nodes by "increasing" or "decreasing" size? NULL for no sorting. Use

tax_sort() before taxatree_plots() for finer control.

add_circles add faint concentric circles to plot, behind each rank?

drop_ranks drop ranks from tree if not included in stats dataframe

Luminance value at the scale endpoints, NULL for palette's default
Luminance value at the scale midpoint, NULL for palette's default

colour_na colour for NA values in tree. (if unused ranks are not dropped, they will have

NA values for colour_stat)

Details

taxatree_plotkey plots same layout as taxatree_plots, but in a fixed colour

See website article for more examples of use: https://david-barnett.github.io/microViz/articles/web-only/modelling-taxa.html

Uses ggraph, see help for main underlying graphing function with ?ggraph::ggraph

It is possible to provide multiple significance markers for multiple thresholds, by passing vectors to the sig_shape, sig_threshold, etc. arguments. It is critically important that the thresholds are provided in decreasing order of severity, e.g. sig_threshold = c(0.001, 0.01, 0.1) and you must provide a shape value for each of them.

Value

list of ggraph ggplots

See Also

```
taxatree_models() to calculate statistical models for each taxon
taxatree_plotkey() to plot the corresponding labelled key
taxatree_plot_labels() and taxatree_label() to add labels
taxatree_stats_p_adjust() to adjust p-values
```

```
# Limited examples, see website article for more
library(dplyr)
library(ggplot2)
data(dietswap, package = "microbiome")
ps <- dietswap
# create some binary variables for easy visualisation
ps <- ps %>% ps_mutate(
  female = if_else(sex == "female", 1, 0, NaN),
  african = if_else(nationality == "AFR", 1, 0, NaN)
)
# This example dataset has some taxa with the same name for phylum and family...
# We can fix problems like this with the tax_prepend_ranks function
# (This will always happen with Actinobacteria!)
ps <- tax_prepend_ranks(ps)</pre>
# filter out rare taxa
ps <- ps %>% tax_filter(
  min_prevalence = 0.5, prev_detection_threshold = 100
# delete the Family rank as we will not use it for this small example
# this is necessary as taxatree_plots can only plot consecutive ranks
```

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```
ps <- ps %>% tax_mutate(Family = NULL)
# specify variables used for modelling
models <- taxatree_models(</pre>
  ps = ps, type = corncob::bbdml, ranks = c("Phylum", "Genus"),
  formula = ~ female + african, verbose = TRUE
# models list stored as attachment in psExtra
models
# get stats from models
stats <- taxatree_models2stats(models, param = "mu")</pre>
stats
plots <- taxatree_plots(</pre>
  data = stats, colour_trans = "identity",
  size\_stat = list(mean = mean),
  size_guide = "legend", node_size_range = c(1, 6)
)
# if you change the size_stat for the plots, do the same for the key!!
key <- taxatree_plotkey(</pre>
  data = stats,
  rank == "Phylum" | p.value < 0.05, # labelling criteria</pre>
  .combine_label = all, # label only taxa where criteria met for both plots
  size_stat = list(mean = mean),
  node_size_range = c(2, 7), size_guide = "none",
  taxon_renamer = function(x) {
    stringr::str_remove_all(x, "[PG]: | [ae]t rel.")
  }
)
# cowplot is powerful for arranging trees and key and colourbar legend
legend <- cowplot::get_legend(plots[[1]])</pre>
plot_col <- cowplot::plot_grid(</pre>
  plots[[1]] + theme(legend.position = "none"),
  plots[[2]] + theme(legend.position = "none"),
  ncol = 1
cowplot::plot_grid(key, plot_col, legend, nrow = 1, rel_widths = c(4, 2, 1))
```

Description

Finer control over label drawing for taxatree_plotkey (with .draw_label = FALSE), and label drawing for taxatree_plots output too.

taxatree_plot_labels 115

Usage

```
taxatree_plot_labels(
  circular = TRUE,
  taxon_renamer = identity,
  fun = ggrepel::geom_text_repel,
  label_var = "label",
  x_nudge = 0.1,
 y_nudge = 0.025,
  rotate = 0,
  fontface = "bold",
  size = 2.5,
  colour = "grey15",
 max.overlaps = Inf,
 min.segment.length = 0,
  segment.size = 0.15,
  segment.color = "grey15",
  point.padding = 0.05,
  box.padding = 0.1,
  seed = NA,
)
```

Arguments

р	taxatree_plotkey or taxatree_plots output plot
circular	is the plot layout circular? labels are drawn differently for circular trees
taxon_renamer	function that takes taxon names and returns modified names for labels
fun	ggrepel labelling function: geom_text_repel or geom_label_repel
label_var	name of variable in taxatree_stats that indicates which taxa to label
x_nudge	absolute amount by which the initial position of taxon labels is nudged (relevant only for circular layouts, use nudge_x for other layouts)
y_nudge	absolute amount by which the initial position of taxon labels is nudged (relevant only for circular layouts, use nudge_y for other layouts)
rotate	angle to rotate labels' outer edges away from horizontal (relevant only for circular layouts, use angle for other layouts)
fontface	fontface of label text
size	size of labels
colour	colour of label outlines and text
max.overlaps	max number of overlapping labels tolerated
min.segment.length	
	min length of label line segment to bother drawing
segment.size	thickness of line segment
segment.color	colour of line segment

padding around node points (for label positioning) point.padding box.padding padding around labels/text (for label positioning) seed set this for reproducible label positions Arguments passed on to ggrepel::geom_text_repel . . . arrow specification for arrow heads, as created by arrow force Force of repulsion between overlapping text labels. Defaults to 1. force_pull Force of attraction between a text label and its corresponding data point. Defaults to 1. max. time Maximum number of seconds to try to resolve overlaps. Defaults to max.iter Maximum number of iterations to try to resolve overlaps. Defaults to 10000. xlim, ylim Limits for the x and y axes. Text labels will be constrained to these limits. By default, text labels are constrained to the entire plot area. direction "both", "x", or "y" – direction in which to adjust position of labels verbose If TRUE, some diagnostics of the repel algorithm are printed

taxatree_stats_p_adjust

Adjust p values in taxatree stats dataframe

Description

Apply a p value adjustment method from stats::p.adjust.methods, such as false-discovery rate adjustment with "BH", or more conservative family-wise error rate controlling methods such as "holm" or "bonferroni".

Usage

```
taxatree_stats_p_adjust(
  data,
  method,
  grouping = "rank",
  p = "p.value",
  new_var = NULL
)
```

Arguments

data psExtra with taxatree_stats dataframe, or just the dataframe

method any method from stats::p.adjust.methods

grouping defines grouping of p-values into families for adjustment, see details.

p name of variable containing p values for adjustment

new_var name of new variable created for adjusted p values (automatically inferred by

default)

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Details

Define how to group the p values for adjustment with the grouping argument. The default is to adjust the p values in groups at each taxonomic rank, but you could also adjust per "taxon" or per "term". Or even group by a combination of rank and term with c("rank", "term"). You should specify the name of the new variable containing the adjusted p values in the new_var argument. If left as NULL the new variable name will be created by pasting together p.adj, the method, and the grouping variable(s) separated by "."

Value

psExtra with dataframe of statistics, or just the data.frame

See Also

```
taxatree_models2stats
taxatree_models
stats::p.adjust
```

```
# This example is an abbreviated excerpt from article on taxon modelling on
# the microViz documentation website
library(corncob)
library(dplyr)
data("ibd", package = "microViz")
# We'll keep only the Ulcerative Colitis and Healthy Control samples, to
# simplify the analyses for this example. We'll also remove the Species
# rank information, as most OTUs in this dataset are not assigned to a
# species. We'll also use `tax_fix` to fill any gaps where the Genus is
# unknown, with the family name or whatever higher rank classification is
# known.
phylo <- ibd %>%
 ps_filter(DiseaseState %in% c("UC", "nonIBD")) %>%
 tax_mutate(Species = NULL) %>%
 tax_fix()
# Let's make some sample data variables that are easier to use and compare
# in the statistical modelling ahead. We will convert dichotomous
# categorical variables into similar binary variables (values: 1 for true,
# or 0 for false). We will also scale and center the numeric variable for
# age.
phylo <- phylo %>%
 ps_mutate(
   UC = ifelse(DiseaseState == "UC", yes = 1, no = 0),
    female = ifelse(gender == "female", yes = 1, no = 0),
    antibiotics = ifelse(abx == "abx", yes = 1, no = 0),
```

```
steroids = ifelse(steroids == "steroids", yes = 1, no = 0),
    age_scaled = scale(age, center = TRUE, scale = TRUE)
bb_models <- phylo %>%
  tax_fix() %>%
  tax_prepend_ranks() %>%
  tax_filter(min_prevalence = 0.3) %>%
  taxatree_models(
    type = corncob::bbdml,
    ranks = c("Phylum", "Class", "Order"),
    variables = c("UC", "female", "antibiotics", "steroids", "age_scaled")
bb_stats <- bb_models %>%
  taxatree_models2stats(param = "mu") %>%
  taxatree_stats_p_adjust(method = "BH", grouping = "rank")
bb_stats
bb_stats %>% taxatree_stats_get()
# you can also directly modify the dataframe,
# and choose a different variable name
bb_stats %>%
  taxatree_stats_get() %>%
  taxatree_stats_p_adjust(
    method = "holm", grouping = "taxon", new_var = "p_adj_holm"
# see all available adjustment methods
stats::p.adjust.methods
```

tax_agg

Aggregate taxa and track aggregation in psExtra

Description

tax_agg sums the abundances of the phyloseq taxa at the given rank. It records the tax_agg rank argument in the info of the psExtra object output. This psExtra object tracks aggregation, and any further transformations and scaling, to help you keep track of what you have done with your phyloseq object and automatically caption ordination plots.

Instead of tax_agg, consider using tax_transform() with a rank argument instead, to both aggregate and transform the taxa. This is also useful when you want to aggregate but not transform the taxa, and yet still log the "identity" transformation in psExtra for captioning your ordination plots. e.g. tax_transform(rank = "Genus", trans = "identity")

tax_agg allows you to pass NA or "unique" to the rank argument which will NOT aggregate the taxa. If you use rank = "unique" or add_unique = TRUE, it will add a new rank called unique, identical to the taxa_names (after any aggregation)

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Be aware: you should not use the top_N argument yourself without good reason. top_N provides a feature inspired by the deprecated microbiome function aggregate_top_taxa which is primarily useful for decluttering compositional barplots. microViz comp_barplot (and ord_plot_iris) already run tax_agg with a top_N argument for you, so you should not. The tax_table produced when using top N is otherwise INVALID FOR MOST OTHER ANALYSES.

Usage

```
tax_agg(
  ps,
  rank = NA,
  sort_by = NA,
  top_N = NA,
  force = FALSE,
  add_unique = FALSE)
```

Arguments

ps	phyloseq	object

rank NA (for tax_names level) or name of valid taxonomic rank (try phyloseq::rank_names(ps))

or "unique"

sort_by if not NA, how should the taxa be sorted, uses tax sort(), takes same options as

by arg

top_N NA does nothing, but if top_N is a number, it creates an extra tax_table column

called top, which is the same as the unique column for the first top_N number

of taxa, and "other" otherwise.

force If TRUE, this forces aggregation at chosen rank to occur regardless of if the

output will be sensible! This avoids the "Taxa not unique at rank: ..." error, but may allow very inappropriate aggregation to occur. Do not use force = TRUE unless you know why you are doing this, and what the result will be. If you are getting an error with force = FALSE, it is almost certainly better to examine the tax_table and fix the problem. force = TRUE is similar to microbiome::aggregate_taxa, which also does not check that the taxa are uniquely

defined by only the aggregation level.

add_unique if TRUE, adds a rank named unique, identical to the rownames after aggregation

Details

This function is inspired by microbiome::aggregate_taxa. However if microbiome::aggregate_taxa is used, microViz cannot track this aggregation.

Comparing aggregate_taxa and tax_agg:

Except for the ordering of taxa, and the addition of a "unique" rank being optional (in tax_agg), the resulting phyloseq objects are identical for aggregating a phyloseq with no ambiguous taxa. Taxa are ambiguous when the tax_table converges at a lower rank after branching, such as if two different genera share the same species (e.g. "s_"). microbiome::aggregate_taxa handles ambiguous taxa by creating a "unique" rank with all of the taxonomic rank info pasted together into one, often

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very long, name. tax_agg throws an error, and directs the user to tax_fix() to fix the ambiguous taxa before aggregation, which should then result in (much) shorter unique names at the aggregation rank.

Value

psExtra object including phyloseq and tax_agg rank info

See Also

```
tax_fix
tax_fix_interactive
tax_transform
```

```
library(phyloseq)
library(dplyr)
data("dietswap", package = "microbiome")
tax_agg(ps = dietswap, "Phylum") %>%
  ps_get() %>%
  tax_table()
tax_agg(ps = dietswap, "Family") %>%
  ps_get() %>%
  tax_table()
# create some missing values
tax_table(dietswap)[3:7, "Genus"] <- "g_"
# this will produce an error, instructing the user to use tax_fix
# tax_agg(ps = dietswap, "Genus")
# this will then work:
dietswap %>%
  tax_fix() %>%
  tax_agg("Genus")
# you can replace unknown values with `tax_fix()`
# which will fix most problems, like the common "g__" and "s__"
# but default tax_fix settings won't catch this long unknown
tax_table(dietswap)[13:17, "Family"] <- "some_unknown_family"</pre>
dietswap %>%
  tax_fix(unknowns = "some_unknown_family") %>%
  tax_agg("Family")
# try tax_fix_interactive() to help you find and fix all the uninformative
# and converging values in your taxonomy table.
# the code below won't aggregate taxa,
# but just adds a new rank called unique, equal to taxa_names
```

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```
tax_agg(ps = dietswap, rank = NA, add_unique = TRUE)
identical(tax_agg(dietswap, NA, add_unique = TRUE), tax_agg(dietswap, "unique")) # TRUE
```

tax_filter

Filter rare and/or low abundance taxa from a phyloseq object

Description

Removes taxa (from all samples) that do not meet a given criterion or combination of criteria. If a value for min_prevalence, min_total_abundance or min_sample_abundance is 1 or greater, then it is treated as an absolute minimum number of samples/reads. If <1, it is treated as proportion of all samples/reads. This function is designed to work with counts. otu_table must contain counts particularly if you want to set a non-zero value for min_total_abundance.

Usage

```
tax_filter(
  ps,
 min_prevalence = 1,
 prev_detection_threshold = 1,
 min_total_abundance = 0,
 min_sample_abundance = 0,
  tax_level = NA,
  names_only = FALSE,
  use_counts = TRUE,
  undetected = NULL,
  verbose = TRUE
)
```

Arguments

use_counts

phyloseq or psExtra (ideally with count data available) ps min_prevalence number or proportion of samples that a taxon must be present in prev_detection_threshold min required counts (or value) for a taxon to be considered present in that sample (or set undetected arg) min_total_abundance minimum total readcount of a taxon, summed across all samples (can be proportion of all counts) min_sample_abundance taxa must have at least this many reads in one or more samples tax_level if given, aggregates data at named taxonomic rank before filtering, but returns phyloseq at the ORIGINAL level of aggregation! if names_only is true return only names of taxa, not the phyloseq names_only expect count data in phyloseq otu_table? default is TRUE

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undetected e.g. 0, value at (or below) which a taxon is considered not present in that sample.

If set, this overrides prev_detection_threshold.

verbose message about proportional prevalence calculations?

Value

filtered phyloseq object AT ORIGINAL LEVEL OF AGGREGATION (not at the level in tax_level)

Examples

```
data("dietswap", package = "microbiome")
# Dropping rare and low abundance taxa #
# Filter at unique taxa level, keeping only those with a prevalence of 10% or more
# and at least 10 thousand reads when summed across all samples.
# Then aggregate to Family level taxonomy.
dietswap %>%
  tax_filter(min_prevalence = 0.1, min_total_abundance = 10000) %>%
 tax_agg("Family")
# Keeping ubiquitous families #
# keep only families that have at least 1000 counts present in 90% of samples
# then aggregate the remaining taxa at 'Genus' level
dietswap %>%
 tax_filter(
    tax_level = "Family", min_prevalence = 0.90,
   prev_detection_threshold = 1000
 ) %>%
 tax_agg("Genus")
```

tax_fix

Replace unknown, NA, or short tax_table values

Description

Identifies phyloseq tax_table values as unknown or uninformative and replaces them with the first informative value from a higher taxonomic rank.

- Short values in phyloseq tax_table are typically empty strings or " ", or "g__" etc. so it is helpful to replace them. (If this is unwanted: set min_length = 0 to avoid filtering on length.)
- Values in unknowns are also removed, even if longer than min_length. It is up to the user to specify sensible values in unknowns if their dataset has other unwanted values.
- NA values are also replaced.

See this article for an extended discussion of tax_table fixing. https://david-barnett.github.io/microViz/articles/web-only/tax-fixing.html

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Usage

```
tax_fix(
  ps,
  min_length = 4,
  unknowns = NA,
  suffix_rank = "classified",
  sep = " ",
  anon_unique = TRUE,
  verbose = TRUE
)
```

Arguments

ps phyloseq or tax_table (taxonomyTable)

min_length replace strings shorter than this, must be integer > 0

unknowns also replace strings matching any in this vector, NA default vector shown in

details!

suffix_rank "classified" (default) or "current", when replacing an entry, should the suffix be

taken from the lowest classified rank for that taxon, "classified", or the "current"

unclassified rank?

sep character(s) separating new name and taxonomic rank level suffix (see suffix_rank)

anon_unique make anonymous taxa unique by replacing unknowns with taxa_name? other-

wise they are replaced with paste("unknown", first_rank_name), which is therefore the same for every anonymous taxon, meaning they will be merged if tax_agg is used. (anonymous taxa are taxa with all unknown values in their

tax_table row, i.e. cannot be classified even at highest rank available)

verbose emit warnings when cannot replace with informative name?

Details

By default (unknowns = NA), unknowns is set to a vector containing:

```
's__' 'g__' 'f__' 'o__' 'c__' 'p__' 'k__' 'S__' 'G__' 'F__' 'O__' 'C__' 'P__' 'K__' 'NA' 'NaN' 'nan' 'unknown' 'Unknown' 's_unknown' 's_Unknown' 's_NA' 'g_unknown' 'g_Unknown' 'g_NA' 'f_unknown' 'f_NA' 'o_unknown' 'o_Unknown' 'o_NA' 'c_unknown' 'c_Unknown' 'c_NA' 'p_unknown' 'p_Unknown' 'p_NA' 'k_unknown' 'k_Unknown' 'k_NA' 'S_unknown' 'S_Unknown' 'S_NA' 'G_unknown' 'G_NA' 'F_unknown' 'F_Unknown' 'F_NA' 'O_unknown' 'O_NA' 'C_unknown' 'C_Unknown' 'C_NA' 'P_unknown' 'P_NA' 'K_unknown' 'K_Unknown' 'K_NA'
```

Value

object same class as ps

See Also

```
tax_fix_interactive for interactive tax_fix help
```

```
library(dplyr)
library(phyloseq)
data(dietswap, package = "microbiome")
ps <- dietswap
# create unknowns to test filling
tt <- tax_table(ps)</pre>
ntax <- ntaxa(ps)</pre>
set.seed(123)
g <- sample(1:ntax, 30)</pre>
f <- sample(g, 10)
p <- sample(f, 3)
tt[g, 3] <- "g__"
tt[f, 2] <- "f__"
tt[p, 1] <- "p__"
tt[sample(1:ntax, 10), 3] <- "unknown"</pre>
# create a row with only NAs
tt[1, ] <- NA
tax_table(ps) <- tax_table(tt)</pre>
ps
# tax_fix with defaults should solve most problems
tax_table(ps) %>% head(50)
# this will replace "unknown"s as well as short values including "g__" and "f__"
tax_fix(ps) %>%
  tax_table() %>%
  head(50)
# This will only replace short entries, and so won't replace literal "unknown" values
  tax_fix(unknowns = NULL) %>%
  tax_table() %>%
  head(50)
# Change rank suffix and separator settings
tax_fix(ps, suffix_rank = "current", sep = " - ") %>%
  tax_table() %>%
  head(50)
# by default, completely unclassified (anonymous) taxa are named by their
# taxa_names / rownames at all ranks.
# This makes anonymous taxa distinct from each other,
# and so they won't be merged on aggregation with tax_agg.
# If you think your anonymous taxa should merge on tax_agg,
# or you just want them to be named the all same for another reason,
# set anon_unique = FALSE (compare the warning messages)
tax_fix(ps, anon_unique = FALSE)
tax_fix(ps, anon_unique = TRUE)
```

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```
# here's a larger example tax_table shows its still fast with 1000s rows,
# from microbiomeutilities package
# library(microbiomeutilities)
# data("hmp2")
# system.time(tax_fix(hmp2, min_length = 1))
```

tax_fix_interactive

Shiny app to help you use tax_fix

Description

Try this app if you get errors with tax_fix() that are tricky to work past, or suggestions to use tax_fix() that you don't understand.

The app shows you the tax_table of your data (searchable) with unknown values highlighted.

It allows you to interactively modify minimum allowed length and to select further values to be defined as unknown.

It will show you the correct tax_fix() code to copy paste into your script to reproduce the interactive filtering.

Usage

```
tax_fix_interactive(data, app_options = list(launch.browser = TRUE))
```

Arguments

data a phyloseq object app_options options list passed to shinyApp()

nothing

See Also

Value

tax_fix for the non-interactive function to use in your scripts

```
library(dplyr)
library(phyloseq)

# create some problem-filled example tax_table data
data(dietswap, package = "microbiome")
ps <- dietswap
# create unknowns to test filling
tt <- tax_table(ps)
ntax <- ntaxa(ps)</pre>
```

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```
set.seed(123)
g <- sample(1:ntax, 30)
f <- sample(g, 10)
p <- sample(f, 3)
tt[g, 3] <- "g__"
tt[f, 2] <- "f__"
tt[p, 1] <- "p__"
tt[sample(1:ntax, 10), 3] <- "unknown"
# create a row with only NAs
tt[1, ] <- NA
tax_table(ps) <- tax_table(tt)
# function takes a phyloseq and shows code for how to fix the tax_table
# tax_fix_interactive(data = ps)</pre>
```

tax_model

Statistical modelling for individual taxa in a phyloseq

Description

tax_model provides a simple framework to statistically model the abundance of individual taxa in your data. You can choose which type of statistical model you want to fit, and you can choose at which rank and (optionally) which specific taxa to fit statistical models for. tax_model takes a phyloseq and returns a list of statistical models, one model for each taxon. The same independent variables are used for all models, as specified in variables or formula argument (latter takes precedence).

taxatree_models runs tax_model on every taxon at multiple taxonomic ranks (you choose which ranks with the plural ranks argument), and returns the results as a named nested list designed for use with taxatree_plots. One list per rank, one model per taxon at each rank.

type = "bbdml" will run beta binomial regression model(s) using the corncob package. For bbdml the same formula/variables is/are used for modelling both the abundance and dispersion parameters.

Usage

```
tax_model(
   ps,
   rank,
   type = "lm",
   variables = NULL,
   formula = NULL,
   taxa = NULL,
   use_future = FALSE,
   return_psx = TRUE,
   checkVars = TRUE,
   checkNA = "warning",
   verbose = TRUE,
   trans = "identity",
```

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```
trans_args = list(),
...
)
```

Arguments

ps	phyloseq object
rank	name of taxonomic rank to aggregate to and model taxa at
type	name of modelling function to use, or the function itself
variables	vector of variable names, to be used as model formula right hand side. If variables is a list, not a vector, a model is fit for each entry in list.
formula	Right hand side of a formula, as a formula object or character string. Or a list of these. (alternative to variables argument, do not provide both)
taxa	taxa to model (named, numbered, logical selection, or defaulting to all if NULL)
use_future	if TRUE parallel processing with future is possible, see details.
return_psx	if TRUE, list of results will be returned attached to psExtra object
checkVars	should the predictor variables be checked for zero variance?
checkNA	One of "stop", "warning", "message", or "allow", which indicates whether to check predictor variables for NAs, and how to report any NAs if present?
verbose	message about progress and any taxa name modifications
trans	name of tax_transform transformation to apply to aggregated taxa before fitting statistical models
trans_args	named list of any additional arguments to tax_transform e.g. list(zero_replace = "halfmin")
	extra args passed directly to modelling function

Details

tax_model and taxatree_models can use parallel processing with the future package. This can speed up analysis if you have many taxa to model. Set use_future = TRUE and run a line like this before doing your modelling: future::plan(future::multisession, workers = 3) (This requires the future and future.apply packages to be installed.)

Value

psExtra with named list of model objects attached. Or a list of lists of models (if multiple models per taxon). Or just a list (of lists), if return_psx is FALSE.

See Also

```
tax_models_get for the accessor to retrieve model results from psExtra taxatree_models for more details on the underlying approach
```

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```
library(corncob)
library(dplyr)
data(dietswap, package = "microbiome")
ps <- dietswap
# create some binary variables for easy visualization
ps <- ps %>% ps_mutate(
  female = if_else(sex == "female", 1, 0, NaN),
  overweight = if_else(bmi_group == "overweight", 1, 0, NaN),
  obese = if_else(bmi_group == "obese", 1, 0, NaN)
# This example HITChip dataset has some taxa with the same name for phylum and family...
# We can fix problems like this with the tax_prepend_ranks function
ps <- tax_prepend_ranks(ps)</pre>
# filter out rare taxa (it is often difficult to fit multivariable models to rare taxa)
ps <- ps %>% tax_filter(min_prevalence = 0.1, min_total_abundance = 10000)
# specify variables used for modelling
VARS <- c("female", "overweight", "obese")</pre>
# Model first 3 genera using all VARS as predictors (just for a quick test)
models <- tax_model(ps, type = "bbdml", rank = "Genus", taxa = 1:3, variables = VARS)</pre>
# Alternative method using formula arg instead of variables to produce identical results
models2 <- tax_model(</pre>
  ps = ps, rank = "Genus", type = "bbdml",
  taxa = 1:3, formula = ~ female + overweight + obese, return_psx = FALSE
all.equal(models, models2) # should be TRUE
# Model only one genus, NOTE the modified name,
# which was returned by tax_prepend_ranks defaults
models3 <- ps %>%
  tax_model(
    rank = "Genus", type = "bbdml",
    taxa = "G: Bacteroides fragilis et rel.", variables = VARS
# Model all taxa at multiple taxonomic ranks (ranks 1 and 2)
# using only female variable as predictor
models4 <- taxatree_models(</pre>
  ps = ps, type = "bbdml", ranks = 1:2, formula = ~female, verbose = FALSE
# modelling proportions with simple linear regression is also possible via type = lm
# and transforming the taxa to compositional first
models_lm <- ps %>%
  tax_transform("compositional") %>%
```

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```
tax_model(rank = "Genus", taxa = 1:3, variables = VARS, type = "lm")
```

tax_mutate

Modify or compute new taxonomic ranks in phyloseq

Description

Add or overwrite tax_table ranks. Use dplyr::mutate() syntax.

Usage

```
tax_mutate(ps, ...)
```

Arguments

```
ps phyloseq object with a tax_table, or just a tax_table
... passed straight to dplyr::mutate (see examples and dplyr::mutate help)
```

Value

phyloseq object with modified tax_table

See Also

```
mutate
ps_mutate
```

```
library(phyloseq)
library(dplyr)
data("dietswap", package = "microbiome")
# compute new rank
tax_mutate(dietswap, loud_genus = toupper(Genus)) %>%
 tt_get() %>%
 head()
# overwrite a current rank
tax_mutate(dietswap, Genus = toupper(Genus)) %>%
 tt_get() %>%
 head()
# overwrite all ranks
tax_mutate(dietswap, across(everything(), .fns = toupper)) %>%
 tt_get() %>%
 head()
# add a new rank at the beginning
```

tax_name

```
tax_mutate(dietswap, Root = "Bacteria", .before = 1) %>%
   tt_get() %>%
   head()

# this is an error as ranks can't be any other class than character
# tax_mutate(dietswap, Genus = 1:ntaxa(dietswap))
```

tax_name

Simple way to set unique taxa_names for phyloseq object

Description

If your current taxa_names aren't what you want (e.g. they are long DNA sequences), this function will help you set sensible unique names.

It combines:

- a prefix like tax, asy, or otu (pick an appropriate prefix or set your own)
- a unique (sequential) number
- classification information from a chosen taxonomic rank (optional)

Usage

```
tax_name(
  ps,
  prefix = c("tax", "asv", "otu")[1],
  rank = NA,
  pad_number = TRUE,
  sep = "_"
)
```

Arguments

```
prefix e.g. 'tax', 'asv', or 'otu' (or set your own)

rank name of taxonomic rank from which to use classifications in new names

pad_number should unique numbers have zeros added to the front (e.g. 001, 002) to be made the same number of characters?

sep character to separate the unique number and any taxonomic classification info (relevant if rank given)
```

Details

Don't confuse this with the phyloseq function taxa_names() or the newer microViz function tax_rename().

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Value

phyloseq object

See Also

```
tax_rename for a more informative taxon naming tool
phyloseq::taxa_names for accessing and manually setting names
```

Examples

```
library(phyloseq)
# get example data
data("enterotype")
ps <- enterotype
head(taxa_names(ps)) # these are mostly fine (except the -1), but imagine you wanted new names
# consider storing the original names for reference (e.g. if they are DNA sequences)
old_taxa_names <- taxa_names(ps)</pre>
ps <- tax_name(ps)</pre>
taxa_names(ps) %>% head()
# probably better to include the genus info to make these names more informative
ps <- tax_name(ps, rank = "Genus")</pre>
taxa_names(ps) %>% head()
# store new names with old names in dataframe for reference
names_df <- tibble::tibble(old = old_taxa_names, new = taxa_names(ps))</pre>
# alternative settings
tax_name(ps, pad_number = FALSE) %>%
 taxa_names() %>%
 head()
tax_name(ps, prefix = "whateveryoulike") %>%
 taxa_names() %>%
 head()
tax_name(ps, rank = "Genus", sep = "-") %>%
 taxa_names() %>%
 head()
```

tax_names2rank

Add taxa_names as last column in phyloseq tax_table

Description

The taxa names in your phyloseq may specify a further unique classification of your taxa, e.g. ASVs, that is not otherwise represented in the tax_table itself. This function fixes that, and allows you to include this level in taxatree_plots for example.

tax_palette

Usage

```
tax_names2rank(data, colname = "unique")
```

Arguments

data phyloseq object, or psExtra or tax_table (taxonomyTable)
colname name of new rank to add at right side of tax_table

Value

same class object as passed in to data

tax_palette

Make a fixed taxa colour palette e.g. for comp_barplot

Description

Makes a named palette vector from your phyloseq dataset considering overall abundance to assign colours (or some other sorting)

Usage

```
tax_palette(
  data,
  rank,
  n,
  by = sum,
  pal = "brewerPlus",
  add = c(other = "lightgrey"),
  ...
)
```

Arguments

```
data phyloseq or psExtra

rank taxonomic rank name or "unique"

n number of colours / taxa (not including "other")

by tax sorting method for tax_sort e.g. sum

pal palette name from distinct_palette function

add name = value pairs appended to end of output, or NA for none

other args are passed to tax_sort
```

Value

named character vector

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Examples

```
library(ggplot2)
data(dietswap, package = "microbiome")
myPal <- tax_palette(dietswap, rank = "Genus", pal = "brewerPlus", n = 40)</pre>
myPal %>% tax_palette_plot() # just to check the palette
# plot one subset of data
dietswap %>%
  ps_filter(nationality == "AFR", timepoint == 1, sex == "male") %>%
  comp_barplot(
   tax_level = "Genus", n_taxa = 15,
   bar_outline_colour = NA, bar_width = 0.7,
   palette = myPal, label = NULL
  )
# plot a different subset of data (top taxa differ but colours are the same)
dietswap %>%
  ps_filter(nationality != "AFR", timepoint == 1, sex == "male") %>%
  comp_barplot(
    tax_level = "Genus", n_taxa = 15,
   bar_outline_colour = NA, bar_width = 0.7,
   palette = myPal, label = NULL
  )
```

tax_palette_plot

tax_palette plotting helper function

Description

Check the named palette colour vector you created with tax_palette()

Usage

```
tax_palette_plot(named_pal_vec, max_n = NA)
```

Arguments

```
named_pal_vec vector of colours named by taxa (e.g. tax_palette output)

max_n NA to display all colours, or limit this
```

Value

ggplot

tax_prepend_ranks

Examples

```
library(ggplot2)
data(dietswap, package = "microbiome")
myPal <- tax_palette(dietswap, rank = "Genus", pal = "brewerPlus", n = 40)
myPal %>% tax_palette_plot() # just to check the palette
```

tax_prepend_ranks

Add rank prefixes to phyloseq tax_table values

Description

Prepend the start of rank names to each taxon at each rank (useful particularly in case of duplicated taxa names across ranks, e.g. dietswap dataset)

Usage

```
tax_prepend_ranks(ps, sep = ": ", nchar = 1)
```

Arguments

ps phyloseq object

sep characters to paste in between rank initial and taxon name

nchar number of characters to use from start of rank_names

Value

phyloseq

See Also

tax_fix for fixing other tax_table problems

```
data("dietswap", package = "microbiome")
phyloseq::tax_table(dietswap) %>% head()
dietswap %>%
  tax_prepend_ranks() %>%
  phyloseq::tax_table() %>%
  head()
```

tax_rename 135

tax_rename	Make new phyloseq taxa names from classification and taxon abun-
	dance info

Description

Pairs classification at the given rank with a numeric ranking suffix (based on abundance or prevalence data) to automatically create informative taxa names.

Usage

```
tax_rename(
  ps,
  rank,
  sort_by = sum,
  transform_for_sort = "identity",
  pad_digits = "auto",
  sep = " ",
  ...
)
```

Arguments

Details

e.g. "Bacteroides 003" for the third most abundant Bacteroides OTU or ASV. Taxa are returned in original order, and otu_table is returned un-transformed. pad_digits options:

- "auto" -> minimum digits to have equal length numbers within groups
- "max" -> minimum digits to have equal length numbers across all groups
- A number: e.g.

```
- 3 -> 001, 002, ..., 042, ..., 180, ...
- 1 -> 1, 2, ..., 42, ..., 180, ...
```

tax_rename

Value

phyloseq object

See Also

phyloseq::taxa_names for accessing and manually setting names

```
library(phyloseq)
data("ibd", package = "microViz")
ps <- ibd %>%
  tax_filter(min_prevalence = 3) %>%
  tax_fix()
# show a few of the current, uninformative names
taxa_names(ps) %>% head(15)
taxa_names(ps) %>% tail(15)
# change names to genus classification plus number
psNewNames <- ps %>% tax_rename(rank = "Genus")
taxa_names(psNewNames) %>% head(15)
taxa_names(psNewNames) %>% tail(15)
# demonstrate some alternative argument settings
psNewNames2 <- ps %>% tax_rename(
  rank = "Family", sort_by = prev, pad_digits = "max", sep = "-"
taxa_names(psNewNames2) %>% head(15)
taxa_names(psNewNames2) %>% tail(15)
ps %>%
  tax_rename(rank = "Genus", pad_digits = 2) %>%
  taxa_names() %>%
  head(15)
# naming improvement on plots example
library(ggplot2)
library(patchwork)
# Overly aggressive OTU filtering to simplify and speed up example
psExample <- ps %>% tax_filter(min_prevalence = 0.4)
# before OTU renaming
before <- psExample %>%
  ps_filter(activity == "inactive") %>%
  tax_names2rank("Taxon") %>%
  comp_barplot(
    tax_level = "Taxon", n_taxa = 12, other_name = "Other",
```

tax_reorder 137

```
merge_other = FALSE, bar_outline_colour = "grey60"
 coord_flip() +
 ggtitle("Original taxon names :(")
# after OTU renaming
after <- psExample %>%
 ps_filter(activity == "inactive") %>%
 tax_rename(rank = "Genus", pad_digits = "max") %>%
 tax_names2rank("Taxon") %>%
 comp_barplot(
   tax_level = "Taxon", n_taxa = 12, other_name = "Other",
   merge_other = FALSE, bar_outline_colour = "grey60"
 coord_flip() +
 ggtitle("New taxon names :)", "tax_rename(rank = 'Genus', sort_by = sum)")
before + after & theme(legend.text = element_text(size = 8))
# ordination example
psExample %>%
 tax_rename(rank = "Genus", sort_by = sum) %>%
 tax_names2rank("otu") %>%
 tax_transform("clr", rank = "otu") %>%
 ord_calc() %>%
 ord_plot(
   size = 2, colour = "ibd", shape = "circle", alpha = 0.5,
   plot_taxa = 1:10,
   tax_vec_length = 0.5,
   tax_lab_style = tax_lab_style(
     type = "text", max_angle = 90, check_overlap = TRUE,
     size = 2.5, fontface = "bold"
   ),
   tax_vec_style_all = vec_tax_all(alpha = 0.1)
 ) +
 coord_fixed(clip = "off") +
 stat_chull(aes(colour = ibd)) +
 scale_colour_brewer(palette = "Dark2") +
 theme(panel.grid = element_line(size = 0.1))
```

tax_reorder

Reorder taxa in phyloseq object using vector of names

Description

Reorder taxa in phyloseq object using vector of names

Usage

```
tax_reorder(
```

tax_reorder

```
ps,
tax_order,
tree_warn = TRUE,
unmatched_warn = TRUE,
ignore = c("other", "Other")
)
```

Arguments

ps phyloseq object

tax_order Names of taxa in desired order; at least some must match. (Numerical indices

are also possible)

tree_warn If phylogenetic tree is present in phyloseq phy_tree slot, taxa cannot be re-

ordered. Default behaviour of tax_sort is to remove the phylogenetic tree and warn about this. tree_warn = FALSE will suppress the warning message, but

still remove the tree!

unmatched_warn Warn if any names (or indices) given in tax_order are not found within (range

of) taxa_names(ps) - these will be ignored

ignore Values that you do not want to be used for reordering taxa (useful for comp_barplot

when custom palette names are used to set tax_order)

Value

phyloseq object (always without phy_tree)

```
data("dietswap", package = "microbiome")
new_order <- c(</pre>
  "Fusobacteria", "Cyanobacteria", "Verrucomicrobia", "Spirochaetes",
  "Actinobacteria", "Firmicutes", "Proteobacteria", "Bacteroidetes"
)
tax_agg(dietswap, rank = "Phylum") %>%
  ps_get() %>%
  phyloseq::taxa_names()
tax_agg(dietswap, rank = "Phylum") %>%
  ps_get() %>%
  tax_reorder(tax_order = new_order) %>%
  phyloseq::taxa_names()
# partial reordering (of the frontmost positions only) is possible
tax_agg(dietswap, rank = "Phylum") %>%
  ps_get() %>%
  tax_reorder(tax_order = c("Cyanobacteria", "Bacteroidetes")) %>%
  phyloseq::taxa_names()
```

tax_scale 139

tax_scale	Mean-center and SD-scale taxa in phyloseq	

Description

Wrapper for applying base scale function to phyloseq otu_table

Usage

```
tax_scale(data, center = TRUE, scale = TRUE, do = NA, keep_counts = TRUE)
```

Arguments

data	phyloseq or psExtra or otu_table
center	if TRUE: center each taxon by subtracting its mean
scale	if TRUE, divide each centred taxon by its standard deviation (or divide by RMS if not centred!)
do	alternative argument that overrides center and scale options! takes "both", "scale", "center" or "neither"
keep_counts	if TRUE, retain the original count data in psExtra counts slot

```
data("dietswap", package = "microbiome")
ps <- dietswap
ps %>%
  otu_get() %>%
  .[1:6, 1:6]
# standard use (mean center and SD scale)
tax_scale(ps) %>%
  otu_get() %>%
  .[1:6, 1:6] # Aerococcus is NaN as standard deviation = 0 (0 prevalence)
# RMS scale only (directly on otu_table)
otu_get(ps) %>%
  tax_scale(center = FALSE) %>%
  .[1:6, 1:6] # Aerococcus is NaN as standard deviation = 0 (0 prevalence)
# example using alternative `do` argument (to center only, no scaling)
tax_scale(ps, do = "center") %>%
  otu_get() %>%
  .[1:6, 1:6]
# preserves existing info
tax_transform(ps, "compositional", rank = "Genus") %>% tax_scale()
# drops other psExtra objects previously calculated with unscaled data
```

tax_select

```
psxDist <- tax_agg(ps, "Genus") %>% dist_calc()
psxDist
psxDist %>% tax_scale()
tax_scale(psxDist) %>% info_get()
```

tax_select

Subset phyloseq object by (partial) taxa names

Description

Convenient name-based taxa selection/filtering of phyloseq object, including approximate name matching. Takes a phyloseq with tax table and a (partial) taxonomic name, or a list/vector of taxonomic names (full or partial matches).

Usage

```
tax_select(
  ps,
  tax_list,
  ranks_searched = "all",
  strict_matches = FALSE,
  n_typos = 1,
  deselect = FALSE
)
```

Arguments

```
ps phyloseq object

tax_list e.g. c('g__Bifidobacterium', 'g__Akkermansia', 'g__Bacteroides', 'g__Streptococcus')

ranks_searched 'all' or a list of which taxonomic ranks should be searched for the names in tax_list?

strict_matches only perfect full name matches allowed if TRUE

n_typos how many typos to allow in each name? uses agrep approximate matching if > 0

deselect if TRUE, the matching taxa will be REMOVED instead!
```

Details

tax_select will also search the otu names/rownames, BUT only for perfect matches.

Value

phyloseq object with fewer taxa

tax_sort 141

See Also

```
ps_select for selecting variables in phyloseq sample_data
agrep for the function that powers the approximate matching in tax_select
```

Examples

```
# Get example phyloseq object data
data("dietswap", package = "microbiome")
pSeq <- dietswap
# SELECTION EXAMPLES #
a <- pSeq %>% tax_select(tax_list = "Bif", n_typos = 0, ranks_searched = "Genus")
b <- pSeq %>% tax_select(tax_list = "Bifidobacterium", n_typos = 0)
c <- pSeq %>% tax_select(tax_list = "Bif", n_typos = 1)
identical(a, b) # TRUE
identical(a, c) # FALSE
pSeq %>% tax_select(tax_list = "Bifidobactrium") # default 1 typo allowed
one <- pSeq %>% tax_select(tax_list = "Akkarmensia", n_typos = 2)
two <- pSeq %>% tax_select(tax_list = "Akkermansia", n_typos = 0)
identical(one, two) # TRUE
# DESELECTION EXAMPLE # #
pSeq %>% tax_select(tax_list = "Bif", strict_matches = FALSE, deselect = TRUE)
# Incorrect example
# pSeq %>% tax_select(tax_list = "Bif", strict_matches = TRUE) # fails
```

tax_sort

Sort taxa in phyloseq otu_table and tax_table

Description

Multiple ways of sorting taxa are possible and determined by the by argument. The by argument must be one of:

- 'rev' to reverse the current order
- 'name' (sort alphabetically by at)
- 'asis' to keep current order as is
- a sample name (descending abundance sorting within that sample)
- summary stat. function e.g. sum or mean

The at argument must be "names" for sorting unique taxa, or a rank name, for sorting at that rank. at is ignored when by is "rev".

tax_sort

Usage

```
tax_sort(
  data,
  by = "name",
  at = "names",
    ...,
  tree_warn = TRUE,
  verbose = TRUE,
  trans = "identity",
  use_counts = TRUE,
  counts_warn = TRUE
)
```

Arguments

by how to sort, see description at "names" or a taxonomic rank to apply sorting method to, as specified in by. used if summary function given, or pass undetected arg for tax_transform("binary") if by = "prev" or "prevalence" tree_warn If phylogenetic tree is present in phyloseq phy_tree slot, taxa cannot be re- ordered. Default behaviour of tax_sort is to remove the phylogenetic tree and warn about this. tree_warn = FALSE will suppress the warning message, but still remove the tree! verbose passed to phyloseq_validate verbose (if TRUE: message about suspicious values in tax_table, and how to fix)
<pre>used if summary function given, or pass undetected arg for tax_transform("binary") if by = "prev" or "prevalence" tree_warn If phylogenetic tree is present in phyloseq phy_tree slot, taxa cannot be re- ordered. Default behaviour of tax_sort is to remove the phylogenetic tree and warn about this. tree_warn = FALSE will suppress the warning message, but still remove the tree! verbose passed to phyloseq_validate verbose (if TRUE: message about suspicious values</pre>
if by = "prev" or "prevalence" If phylogenetic tree is present in phyloseq phy_tree slot, taxa cannot be reordered. Default behaviour of tax_sort is to remove the phylogenetic tree and warn about this. tree_warn = FALSE will suppress the warning message, but still remove the tree! verbose passed to phyloseq_validate verbose (if TRUE: message about suspicious values
ordered. Default behaviour of tax_sort is to remove the phylogenetic tree and warn about this. tree_warn = FALSE will suppress the warning message, but still remove the tree! verbose passed to phyloseq_validate verbose (if TRUE: message about suspicious values
in un_uoto, und non to in,
trans name of transformation to apply to taxa before sorting (taxa are returned untransformed)
use_counts use count data if available, instead of transformed data
counts_warn warn if count data are not available?

Details

Don't forget to pass na.rm = TRUE to ... if using a summary stat function in by

Value

sorted phyloseq or psExtra

```
library(phyloseq)
data("dietswap", package = "microbiome")
dietswap

# reverse current order
dietswap %>%
```

tax_sort 143

```
tax_sort("rev") %>%
 tax_table() %>%
 head(30)
# sort alphabetically by a taxonomic rank (or "names" for taxa_names)
dietswap %>%
 tax_sort(by = "name", at = "Phylum") %>%
 tax_table() %>%
 head(30)
# sequentially sorting by higher ranks
# sets tax_table in nested alphabetical order
dietswap %>%
 tax_sort(at = "names") %>%
 tax_sort(at = "Genus") %>%
 tax_sort(at = "Family") %>%
 tax_sort(at = "Phylum") %>%
 tax_table() %>%
 head(30)
# sort by function e.g. total sum or median abundance
dietswap %>%
 tax_sort(by = sum) %>%
 taxa_names() %>%
 head(20)
# transform to compositional data (proportions) before sorting
# note that abundances are returned untransformed
dietswap %>%
 tax_sort(by = sum, trans = "compositional") %>%
 taxa_names() %>%
 head(20)
# order by descending abundance in a single named sample
dietswap %>%
 tax_sort(by = "Sample-1") %>%
 otu_table() %>%
 .[1:8, 1:4]
# sum order should always equal mean order if non-negative abundances
# don't forget to add na.rm = TRUE if you expect NAs in otu_table somehow
dietswap %>%
 tax_sort(by = sum, na.rm = TRUE) %>%
 taxa_names() %>%
 head(20)
# if your phyloseq object has a phylogenetic tree,
# tax_sort will remove the tree, and warn you about this
# unless you disable that warning with tree_warn = FALSE
# You can sort by abundance at higher taxonomic ranks,
# without losing lower rank info
```

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```
# e.g. sort (descending) by phyla abundances
dietswap %>%
  tax_sort(by = sum, at = "Phylum") %>%
  tax_table() %>%
  head()

# You can sort by ascending abundance (or prevalence etc) by reversing after
dietswap %>%
  tax_sort(by = "prev", at = "Phylum") %>%
  tax_sort(by = "rev") %>%
  tax_table() %>%
  head()
```

tax_sort_ord

Order taxa in phyloseq by their loading vectors

Description

tax_sort_ord reorders taxa in a phyloseq object based on the relative length of their taxa scores / "loading" vector lengths on 1 or 2 ordination axes.

ord_order_taxa gets the taxa names in order from the ordination contained in a psExtra object. This is used internally by tax_sort_ord.

Usage

```
tax_sort_ord(ps, ord, axes = 1:2, scaling = 2)
ord_order_taxa(ord, axes = 1:2, scaling = 2)
```

phyloseq object to be sorted

Arguments ps

ord psExtra with ordination object

axes which axes to use for sorting? numerical vector of length 1 or 2

scaling Type 2, or type 1 scaling. For more info, see https://sites.google.com/
site/mb3gustame/constrained-analyses/redundancy-analysis. Either "species"
or "site" scores are scaled by (proportional) eigenvalues, and the other set of scores is left unscaled (from ?vegan::scores.cca)

See Also

- These functions were created to support ordering of taxa bars on ord_plot_iris
- ps_sort_ord for ordering samples in phyloseq by ordination

tax_top 145

tax_	+ ~ ~
ıax	1 ()[]

Get names of "top" n taxa

Description

Simple wrapper around tax_sort that:

- 1. optionally aggregates taxa at rank
- 2. sorts the aggregated taxa according to by
- 3. returns the top n number of taxa names

Usage

```
tax_top(data, n = 10, by = sum, rank = "unique", use_counts = FALSE, ...)
```

Arguments

data	phyloseq object or psExtra
n	how many taxa names to return, or NA for all (can return fewer than n values, if there are fewer to return) $\frac{1}{2}$
by	how to sort taxa (see ?tax_sort()), defaults to sum which sorts by total abundance across all samples
rank	taxonomic rank to aggregate at before calculating ("unique" = no aggregation)
use_counts	use count data if available, instead of transformed data
	Arguments passed on to tax_sort
	verbose passed to phyloseq_validate verbose (if TRUE: message about suspicious values in tax_table, and how to fix)
	trans name of transformation to apply to taxa before sorting (taxa are returned un-transformed)

Value

vector of taxa names at chosen rank

See Also

```
tax_agg for more info on taxonomic aggregation
tax_sort for more info on sorting taxa
```

```
data("dietswap", package = "microbiome")
tax_top(dietswap)
tax_top(dietswap, n = 4, by = "prev", rank = "Phylum", undetected = 30)
```

tax_transform

tax_transform

Transform taxa in phyloseq object and record transformation

Description

Transform taxa features, and optionally aggregate at specified taxonomic rank beforehand. You can pipe the results of tax_agg into tax_transform, or equivalently set the rank argument in tax_transform.

Usage

```
tax_transform(
  data,
  trans,
  rank = NA,
  keep_counts = TRUE,
  chain = FALSE,
  zero_replace = 0,
  add = 0,
  transformation = NULL,
  ...
)
```

Arguments

data a phyloseq object or psExtra output from tax_agg

trans any valid taxa transformation (e.g. from microbiome::transform)

rank If data is phyloseq: data are aggregated at this rank before transforming. If NA,

runs tax_agg(data, rank = NA). If rank is NA and data is already psExtra, any

preceding aggregation is left as is.

keep_counts if TRUE, store the pre-transformation count data in psExtra counts slot

chain if TRUE, transforming again is possible when data are already transformed i.e.

multiple transformations can be chained with multiple tax_transform calls

zero_replace Replace any zeros with this value before transforming. Either a numeric, or

"halfmin" which replaces zeros with half of the smallest value across the entire dataset. Beware: the choice of zero replacement is not tracked in the psExtra

output.

add Add this value to the otu_table before transforming. If add != 0, zero_replace

does nothing. Either a numeric, or "halfmin". Beware: this choice is not tracked

in the psExtra output.

transformation deprecated, use trans instead!

... any extra arguments passed to microbiome::transform or pass undetected =

a number when using trans = "binary"

tax_transform 147

Details

This function often uses microbiome::transform internally and can perform the same transformations, including many from vegan::decostand (where the default MARGIN = 2). See below for notes about some of the available transformations.

tax_transform returns a psExtra containing the transformed phyloseq object and extra info (used for annotating ord_plot ordinations):

- tax_transform (a string recording the transformation),
- tax_agg (a string recording the taxonomic aggregation rank if specified here or earlier in tax_agg).

A few commonly used transformations:

- "clr", or "rclr", perform the centered log ratio transformation, or the robust clr, using microbiome::transform
- "compositional" converts the data into proportions, from 0 to 1.
- "identity" does not transform the data, and records this choice for ord_plot
- "binary" can be used to transform tax abundances into presence/abundance data.
- "log2" which performs a log base 2 transformation (don't forget to set zero_replace if there are any zeros in your data)

Value

psExtra object including phyloseq and extra info

(r)clr transformation note

If any values are zero, the clr transform routine first adds a small pseudocount of min(relative abundance)/2 to all values. To avoid this, you can replace any zeros in advance by setting zero_replace to a number > 0.

The rclr transform does not replace zeros. Instead, only non-zero features are transformed, using the geometric mean of non-zero features as denominator.

Binary transformation notes

By default, otu_table values of 0 are kept as 0, and all positive values are converted to 1 (like decostand(method = "pa")). You can set a different threshold, by passing e.g. undetected = 10, for example, in which case all abundances of 10 or below would be converted to 0. All abundances above 10 would be converted to 1s.

Beware that the choice of detection threshold is not tracked in the psExtra.

See Also

microbiome::transform for some more info on available transformations
vegan::decostand for even more transformation options
tax_agg

Examples

```
data("dietswap", package = "microbiome")
# aggregate taxa at Phylum level and center log ratio transform the phyla counts
tax_transform(dietswap, trans = "clr", rank = "Phylum")
# this is equivalent to the two-step method (agg then transform)
tax_agg(dietswap, rank = "Phylum") %>% tax_transform("clr")
# does nothing except record tax_agg as "unique" and tax_transform as "identity" in psExtra info
dietswap %>% tax_transform("identity", rank = NA)
# binary transformation (convert abundances to presence/absence or detected/undetected)
tax_transform(dietswap, trans = "binary", rank = "Genus")
# change detection threshold by setting undetected argument (default is 0)
tax_transform(dietswap, trans = "binary", rank = "Genus", undetected = 50) %>%
 otu_get() %>%
  .[1:6, 1:4]
# log2 transformation after replacing all zeros with a pseudocount of 1
tax_transform(dietswap, trans = "log2", rank = "Family", zero_replace = 1)
# log2 transformation after replacing all zeros with a pseudocount of half
# the minimum non-zero count value in the aggregated dataset
tax_transform(dietswap, trans = "log2", rank = "Family", zero_replace = "halfmin")
```

```
upgrade_ps_extra_to_psExtra
```

Convert old format "ps_extra" objects to new "psExtra" objects

Description

This will only be necessary if you have saved old format "ps_extra" objects generated by an old microViz version (< 0.10.0), and you cannot or do not want to regenerate these old format objects from your original phyloseq object.

Usage

```
upgrade_ps_extra_to_psExtra(ps_extra)
```

Arguments

```
ps_extra an old format "ps_extra" object, as generated by old micro Viz versions (< 0.10.0)
```

Value

```
new format "psExtra" S4 object
```

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Examples

```
# read your old saved 'ps_extra' object that you want to keep using
# oldObject <- readRDS("old-object-path.rds")</pre>
# newObject <- upgrade_ps_extra_to_psExtra(oldObject)</pre>
# continue with your next analysis or plotting steps...
```

varAnnotation

Helper to specify a HeatmapAnnotation for variables in cor_heatmap

Description

Helper to specify a HeatmapAnnotation for variables in cor_heatmap

Usage

```
varAnnotation(
  . . . ,
  name,
  annotation_legend_param = list(),
  show_legend = TRUE,
  gp = grid::gpar(col = NA),
 border = FALSE,
  gap = grid::unit(2, "mm"),
  show_annotation_name = TRUE,
  annotation_label = NULL,
  annotation_name_gp = grid::gpar(),
  annotation_name_offset = NULL,
  annotation_name_rot = NULL,
  annotation_name_align = FALSE,
  annotation_name_side = "auto",
  .data = NULL,
  .vars = NULL,
  .side = NULL
)
```

Arguments

Name-value pairs where the names correspond to annotation names and values are the output of variable annotation functions such as anno_var_box(), or man-

ually specified AnnotationFunction objects

Name of the heatmap annotation, optional. name

annotation_legend_param

A list which contains parameters for annotation legends. See color_mapping_legend, ColorMapping=me for all possible options.

Whether show annotation legends. The value can be one single value or a vector. show_legend

Graphic parameters for simple annotations (with fill parameter ignored). gp

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border border of single annotations.

gap Gap between annotations. It can be a single value or a vector of unit objects.

show_annotation_name

Whether show annotation names? For column annotation, annotation names are drawn either on the left or the right, and for row annotations, names are draw either on top or at the bottom. The value can be a vector.

annotation_label

Labels for the annotations. By default it is the same as individual annotation names.

annotation_name_gp

Graphic parameters for annotation names. Graphic parameters can be vectors.

annotation_name_offset

Offset to the annotation names, a unit object. The value can be a vector.

annotation_name_rot

Rotation of the annotation names. The value can be a vector.

annotation_name_align

Whether to align the annotation names.

annotation_name_side

Side of the annotation names.

.data OPTIONAL phyloseq or psExtra, only set this to override use of same data as

in heatmap

.vars OPTIONAL selection vector of variables (names, numbers or logical), only set

this if providing .data argument to override default

. side OPTIONAL string, indicating the side for the variable annotations: only set this

to override default

Value

HeatmapAnnotation object

See Also

taxAnnotation()

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