Package 'LeyLabRMisc'

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Author Nick Youngblut

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Maintainer Nick Youngblut <nyoungb2@gmail.com>

Description Ley Lab misc R functions, rmd templates, etc.			
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.get_brewer_palette

getting RColorBrewer entire palette

Description

getting RColorBrewer entire palette

Usage

```
.get_brewer_palette()
```

.HFE

supporting function for HFE

Description

supporting function for HFE

Usage

```
.HFE(brk, class_level, corr_cutoff = 0.5, quiet = TRUE)
```

.well384_index

making 384-well plate index

Description

making 384-well plate index

Usage

```
.well384_index()
```

Value

named vector (Well -> location); column-wise location

4 as.Num

 $.well96_index$

making a 96-well plate index

Description

making a 96-well plate index

Usage

```
.well96_index()
```

Value

named vector (Well -> location); column-wise location

as.Num

convert to numeric while avoiding factor conversion issues

Description

convert to numeric while avoiding factor conversion issues

Usage

```
as.Num(x)
```

Arguments

Х

an interable

Value

a numeric object

bash_job 5

basii_job basii job usiiig conaa env	bash_job	bash job using conda env	
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Description

The conda setup is assumed to be in your ~/.bashrc If print_output == TRUE: the stdout/stderr will be printed instead of returned Else: the stdout/stderr with be returned by the function stderr/stdout is printed unless print_output==FALSE

Usage

```
bash_job(cmd, conda_env, stdout = TRUE, stderr = TRUE, print_output = TRUE)
```

Arguments

cmd The bash command in a string format

conda_env The conda env to use

stdout Print the stdout from the command? stderr Print the stderr from the command?

quiet No printing

calculate_rarefaction_curves

Function for rarefaction analysis

Description

Running estimate_richness_phy() at multiple subsampling depths

Usage

```
calculate_rarefaction_curves(psdata, measures, depths, parallel = FALSE)
```

Arguments

psdata phyloseq object

measures Which diversity measures (see vegan package)

depths Which sequencing depths? Example: c(10, 100, 1000)

Value

A dataframe

6 calc_beta_div

age installed to your R project and loaded for this code to run

Description

Faith's Phylogenetic Diversity ("PD") can be calculated only if a tree is provided. The tree can have extra tips, but there must be tip labels for all taxa in the provided table.

Usage

```
calc_alpha_div(df, tree = NULL, index = c("nobs", "shannon", "PD"))
```

Arguments

df sample x taxon abundance table (usual format for vegan)

tree tree with tips matching taxa in the abundance table (only needed for PD)

index which of the indices to calculate? (nobs = no. of observations, shannon = Shan-

non Index, PD = Faith's PD)

Value

a data.frame of alpha diversity values (and sample names)

calc_beta_div

beta-diversity calculation

Description

A wrapper around vegan::vegdist and rbiom (rbiom used for UniFrac calculations). For unifrac: "wunifrac" = weighted unifrac, "unifrac" = unweighted unifrac. The function returns a tidy dataframe of PCoA axes (PC1 & PC2), percent variance explained for each PC.

```
calc_beta_div(
   df,
   tree = NULL,
   method = c("wunifrac", "unifrac", "manhattan", "euclidean", "canberra", "clark",
   "bray", "kulczynski", "jaccard", "gower", "altGower", "morisita", "horn", "mountford",
        "raup", "binomial", "chao", "cao", "mahalanobis"),
   threads = 1
)
```

calc_PCoA 7

Arguments

df sample x taxon dataframe. Colnames (taxa) must match the tree tip labels if the

tree is provided

tree phylogeny with tips matching the df colnames (only needed for wunifrac &

unifrac methods)

method distance method (vegdist distances; wunifrac=Weighted Unifrac; unifrac=Unweighted

Unifrac)

threads used for UniFrac calculations with rbiom

Details

Unifrac is calculated with the https://github.com/cmmr/rbiom package (requires bioconductor packages).

If the goal is PCoA, then see the "tidy_PCoA" function.

Value

data.frame

	Wrapper for cmdscale	calc_PCoA
--	----------------------	-----------

Description

Simple wrapper for cmdscale to provide data.frame formatted table. If the distance matrices contain NAs, the samples containing NAs will be removed (with a warning).

Usage

```
calc_PCoA(dist_mtx, k = 2)
```

Arguments

dist_mtx distance matrix object

Value

data.frame

8 clustermq_get_logs

cat_file

pretty printing of a text file via cat

Description

This is most useful for working with IRkernl in Jupyter notebooks

Usage

```
cat_file(file_name)
```

Arguments

file_name

the name of the file to print

clustermq_get_logs

Get/read clustermq cluster job log files

Description

If you use "log_file = clustermq_logfile()" in your template, then you can use this function to get the log file paths or directly read the contents of the log files.

Usage

```
clustermq_get_logs(lines = 0, logfile_dir = NULL)
```

Arguments

lines

The number of lines of each log file to read. If 0, then the log file paths will be returned; if >0 then the first N lines will be printed; if <0 then the last N lines

will be printed.

logfile_dir

The base directory containing all of the logfiles. If not provided, then this is obtained by getOption('clustermq.logfile')

Value

logfile paths or NULL

Examples

clustermq_logfile 9

clustermq_logfile

Set a path for clustermq cluster job log files

Description

Log files are optional for clustermq. The must be set in the template. This function will create a unique directory within the "base_dir". It will also return a path that you MUST use for the "log_file" parameter in the Q template. Moreover, the function will set the "clustermq.logfile" option to that directory (used by clustermq_get_logs).

Usage

```
clustermq_logfile(base_dir = "/ebio/abt3_scratch/")
```

Arguments

base_dir

The base directory will the logfiles will be located.

Details

The function requires the uuid package.

Value

logfile path

Examples

```
clustermq_setup()
tmpl = list(job_mem = '8G', log_file = clustermq_logfile())
fx = function(x, y) x * 2 + y
Q(fx, x=1:3, const=list(y=10), n_jobs=10, job_size=1, template=tmpl)
```

clustermq_setup

Set clustermq options

Description

These options must be set before running clustermq

```
clustermq_setup(
  scheduler = c("sge", "multicore"),
  template = file.path(Sys.getenv("HOME"), ".clustermq.tmpl")
)
```

10 df.dims

Arguments

scheduler The clustermq.scheduler option. Use "multicore" for local jobs. template The clustermq.template option. It defaults to ~.clustermq.tmpl

Examples

condaInfo

"conda list" in R

Description

This is most useful for working with IRkernl in Jupyter notebooks

Usage

```
condaInfo(conda_env)
```

Arguments

conda_env

The name of the conda env to list

df.dims

Changing number of rows/columns shown when printing a data frame

Description

This is most useful for working with IRkernl in Jupyter notebooks

Usage

```
df.dims(nrows = 4, ncols = 20)
```

Arguments

nrows number of rows to print ncols number of columns to print dfhead 11

dfhead

A simple dataframe summary

Description

A simple dataframe summary

Usage

```
dfhead(df, n = 3)
```

Arguments

df dataframe object

n Number of lines to print

Value

a dataframe object

 $dist_format$

creating a string with distance & percent explained

Description

creating a string with distance & percent explained

Usage

```
dist_format(dist, PC1_perc_exp, PC2_perc_exp, label1 = 1, label2 = 2)
```

Arguments

dist str, distance metric

PC1_perc_exp float, percent variance explained for PC1
PC2_perc_exp float, percent variance explained for PC2

label1 First PC labellabel2 Seconda PC label

Value

```
str, formatted as "metric, <PC1_perc_exp>
```

estimate_rarified_richness

Helper Function for rarefaction analysis

Description

Helper Function for rarefaction analysis

Usage

```
estimate_rarified_richness(psdata, measures, depth)
```

Arguments

psdata phyloseq object

measures Which diversity measures depth The sampling depth

Value

molten alpha diversity object

 $\verb|estimate_richness_phy|| phyloseq::estimate_richness, but includes Faith's PD|$

Description

See physeq::estimate richness for full details

Usage

```
estimate_richness_phy(physeq, split = TRUE, measures = NULL)
```

Arguments

physeq Phyloseq object

split Splitting the OTU table

measures Which diversity measures (Faith's PD = "FaithPD)

Value

Dataframe can calculate faith's PD (using Picante, "FaithPD")

extract_pltdt 13

extract_pltdt

Extract data from ggplot object

Description

The data is written to files

Usage

```
extract_pltdt(plot_object, output_path)
```

Arguments

plot_object

A ggplot object

output_path

Where to write the output

fig_uuid

create UUID for figure file name

Description

create UUID for figure file name

Usage

```
fig_uuid(full = FALSE)
```

Arguments

full

Full length uuid or trimmed to just 24 char?

Value

character object

14 Fread

	list

convert a vector of file paths into a named list

Description

convert a vector of file paths into a named list

Usage

```
files_to_list(files, label_index = -1)
```

Arguments

files Vector of file paths (eg., by using "list_files()")

label_index Which item in the path to return? 1-indexing. If <1, samples selected from the

end.

Examples

```
files = c('/path/to/project/Sample1/table.txt', '/path/to/project/Sample2/table.txt') \\ files_to_list(files, -1) \\ files = c('/path/to/project/Sample1.txt', '/path/to/project/Sample2.txt') \\ files_to_list(files, 0) \\
```

Fread

Simple wrapper around data.table::fread

Description

Simple wrapper around data.table::fread

Usage

```
Fread(infile = NULL, cmd = NULL, sep = "\t", check.names = TRUE, ...)
```

Arguments

infile input file name

cmd command instead of input file (eg., "gunzip -c INFILE")

sep value delimiter

check.names format check column names
... passed to data.table::fread

Value

data.table

hello 15

hello

Hello, World!

Description

```
Prints 'Hello, world!'.
```

Usage

hello()

Examples

hello()

itol_boxplot

create itol boxplot file

Description

https://itol.embl.de/help.cgi#boxplot

Usage

```
itol_boxplot(
   df,
   dataset_label,
   out_file,
   out_dir = NULL,
   key_color = "#ff0000",
   WIDTH = 200
)
```

Arguments

df	Dataframe, in which the rownames should correspond with the tree labels; the columns must specify: minimum,q1,median,q3,maximum,extreme_value1,extreme_value2
dataset_label	What to label the itol dataset
out_file	Name of the output file
out_dir	Where to write the output
key_color	The color for the legend key
WIDTH	Maximum width

16 itol_colorstrip

itol_colorstrip	create itol colorstrip file
-----------------	-----------------------------

Description

https://itol.embl.de/help.cgi#strip

Usage

```
itol_colorstrip(df, dataset_label, out_file, out_dir = NULL, legend = NULL)
```

Arguments

df	Dataframe, in which the rownames should correspond with the tree labels; the plotting parameter should be column 1
dataset_label	What to label the itol dataset
out_file	Name of the output file
out_dir	Where to write the output
legend	Custom legend (see the function description)

Details

Custom Legend: requires a data.frame with the number of rows equaling the number of unique values in the legend.

```
• "shapes" => numeric (see the itol docs)
```

- "colors" => hexidecimal (see this website for examples)
- "labels" => legend labels

Examples

```
# creating a custom legend
legend = data.frame(unique(iris$Species),
colors = c('#00FF00', '#FFCC33', '#FF0000'),
shapes = rep(1, length(unique(iris$Species))))
legend
```

itol_externalshape 17

itol_externalshape

create itol external shape file

Description

https://itol.embl.de/help.cgi#shapes

Usage

```
itol_externalshape(
   df,
   dataset_label,
   out_file,
   out_dir = NULL,
   legend = NULL,
   WIDTH = 200
)
```

Arguments

df	Dataframe, in which the rownames should correspond with the tree labels; other columns should be values corresponding to symbol size
dataset_label	What to label the itol dataset
out_file	Name of the output file
out_dir	Where to write the output
legend	Specify particular legend (see itol_colorstrip)

itol_heatmap

create itol heatmap file

Description

https://itol.embl.de/help.cgi#heatmap

```
itol_heatmap(
   df,
   dataset_label,
   out_file,
   out_dir = NULL,
   tree = NULL,
   dist_method = "bray",
   color_scheme = c("color", "bw")
)
```

18 itol_multibar

Arguments

df Dataframe, in which the rownames should correspond with the tree labels; all

columns should be numeric values for the heatmap

dataset_label What to label the itol dataset
out_file Name of the output file
out_dir Where to write the output

tree Tree object used for ordering the heatmap columns; if NULL, the dist_method

will be used to create the tree

dist_method vegan::vegdist method for creating the correlation dendrogram

color_scheme Heatmap color scheme. color = blue-orange-yellow; bw=white-grey-black

itol_multibar

create itol multi-bar file

Description

https://itol.embl.de/help.cgi#multibar

Usage

```
itol_multibar(
   df,
   dataset_label,
   out_file,
   out_dir = NULL,
   legend = NULL,
   WIDTH = 200,
   COLOR = "#ff0000")
```

Arguments

df Dataframe, in which the rownames should correspond with the tree labels

dataset_label What to label the itol dataset
out_file Name of the output file
out_dir Where to write the output

legend A list that includes shapes, colors, and labels (see itol_colorstrip)

WIDTH Bar width
COLOR Legend color

itol_simplebar 19

itol_simplebar

create itol simple-bar file

Description

https://itol.embl.de/help.cgi#bar

Usage

```
itol_simplebar(
   df,
   dataset_label,
   out_file,
   out_dir = NULL,
   legend = NULL,
   WIDTH = 200
)
```

Arguments

df Dataframe, the rownames should correspond with the tree labels
dataset_label What to label the itol dataset
out_file Name of the output file
out_dir Where to write the output
legend Specify particular legend (see itol_colorstrip)
WIDTH Bar width

itol_symbol

create itol symbol file

Description

https://itol.embl.de/help.cgi#symbols

```
itol_symbol(
   df,
   dataset_label,
   out_file,
   out_dir = NULL,
   MAXIMUM_SIZE = 50,
   COLOR = "#ff0000"
)
```

20 list_files

Arguments

df Dataframe, in which the rownames should correspond with the tree internal node

labels, and other columns should be: symbol,size,color,fill,position,(label)

dataset_label What to label the itol dataset

out_file Name of the output file
out_dir Where to write the output
MAXIMUM_SIZE The max size of the symbols

COLOR Legend color

Description

list.files with full.names=TRUE & recursive=TRUE

Usage

```
list_files(path, pattern = NULL, full.names = TRUE, recursive = TRUE, ...)
```

Arguments

path a character vector of full path names; the default corresponds to the working

directory,

pattern an optional regular expression. Only file names which match the regular expres-

sion will be returned.

full.names a logical value. If TRUE, the directory path is prepended to the file names to

give a relative file path. If FALSE, the file names (rather than paths) are returned

recursive logical. Should the listing recurse into directories?

Value

A character vector containing the names of the files in the specified directories

make_dir 21

make_dir

A helper function for creating a directory (recursively)

Description

A helper function for creating a directory (recursively)

Usage

```
make_dir(dir, quiet = FALSE)
```

Arguments

dir path for the new directory (will create recursively)
quite quite output

mlr_boruta_filter

Custom mlr filter for Boruta

Description

A custom mlr filter that uses Boruta to select important features This function registers the "boruta.filter" filter to be used with makeFilterWrapper and other mlr filter functions.

Usage

```
mlr_boruta_filter()
```

Details

- target str; what is the target variable in the task object (default: 'Class')
- pValue float; see Boruta docs (default: 0.01)
- maxRuns int; see Boruta docs (default: 200)
- hostHistory bool; see Boruta docs (default: FALSE)
- with Tentative bool; keep tentative features (default: TRUE)
- verbose bool; list features selected? (default: FALSE)
- mustKeep vector; features that cannot be filtered (default: NULL)
- threads int; number of threads to use for Boruta (default: 1)

Value

Nothing, but "boruta.filter" filter will be registered

22 ml_tax_HFE

 $mlr_getNestedTuneResultsOptPathDf$

Version of getNestedTuneResultsOptPathDf that actually works

Description

For main docs, see ?getNestedTuneResultsOptPathDf

Usage

```
mlr_getNestedTuneResultsOptPathDf(r, trafo = FALSE)
```

Arguments

r The result of resampling of a tuning wrapper

trafo Should the units of the hyperparameter path be converted to the transformed

scale?

Value

data.frame

ml_tax_HFE

Hierachical Feature Selection

Description

For each clade (defined by tax_level), aggregate species abundances at each taxonomic level up to the user-defined "tax_level", then filter out taxa that correlate strongly (just one taxon is selected of those that correlate).

Usage

```
ml_tax_HFE(brk, tax_level, corr_cutoff = 0.7, threads = 2, quiet = TRUE)
```

Arguments

brk data.table generated by read_bracken()

tax_level which taxonmoic level to use?

corr_cutoff features with >cutoff will be filtered to just one

Value

data.table of filtered features

overlap 23

overlap	Determine counts of setdiff, intersect, & union of 2 vectors (or data.tables)

Description

The output is printed text of intersect, each-way setdiff, and union. Data.table compatible! Just make sure to provide sel_col_x and/or sel_col_y

Usage

```
overlap(
    x,
    y,
    sel_col_x = NULL,
    sel_col_y = NULL,
    to_return = c("counts", "diff_x", "diff_y", "diff_fuzzy"),
    diff = c(NA, "x", "y", "int", "union", "fuzzy")
)
```

Arguments

p.dims

Global change of plot size options

Description

This is most useful for working with IRkernl in Jupyter notebooks

```
p.dims(w = 5, h = 5, res = 200)
```

phyloseq2df

Arguments

w figure widthh figure height

res figure resolution (DPI)

path_get_label

splitting path and returning just one item in the vector

Description

This is useful for merging tables in which the individual table ID is within the file path.

Usage

```
path_get_label(file_path, index)
```

Arguments

file_path File path(s). If vector or list of paths provided, then a list will be returned index Which item in the path to return? 1-indexing. If <1, samples selected from the

end. "O" will select the file name.

Value

string if 1 path, else list

phyloseq2df

Convert a sub-object of a phyloseq object to a dataframe

Description

A helper function for converting OTU, taxonomy, and metadata to dataframes

Usage

```
phyloseq2df(physeq_obj, physeq_func, long = FALSE, flip = FALSE)
```

Arguments

physeq_obj The phyloseq object

physeq_func Which object do you want ('otu_table', 'tax_table', or 'sample_data')

long Do you want the table in "long" format ("gathered")

flip Flip (transpose) the table?

Value

A tibble

phyloseq_rel_abund 25

phyloseq_rel_abund

Transform abundances to relative

Description

A simple wrapper for transform_sample_counts()

Usage

```
phyloseq_rel_abund(physeq_obj, percent_abund = TRUE)
```

Arguments

physeq_obj

The phyloseq object

percent_abund

Fractional or percent abundance?

Value

A phyloseq object

pipelineInfo

pipeline sessionInfo

Description

sessionInfo for LeyLab snakemake pipelines

Usage

```
pipelineInfo(pipeline_path, head_n = 10)
```

Arguments

pipeline_path The path to the pipeline directory

head_n The number of lines to print from the readme

26 qsave_obj

Plot

plot figure and save the figure grob object to a file at the same time

Description

This is most useful for working with IRkernl in Jupyter notebooks

Usage

```
Plot(
  p,
  file = NULL,
  path = NULL,
  suffix = "",
  saveObj = TRUE,
  saveImg = FALSE,
  width = NA,
  height = NA,
  ...
)
```

Arguments

p	Plot object (ggplot2, base, etc)
file	File name to write
path	Path to write to
suffix	File name suffix (eg., '.png')
saveObj	Write the Robj to a file?
saveImg	Write the image to a file?
width	Figure width. If NA, uses global options
height	Figure height. If NA, uses global options

qsave_obj Simple function for serializing a distance matrix or list of distance matrices

Description

Serializing done with the "qs" R package.

```
qsave_obj(x, file, msg = "Writing file to: ", threads = 1)
```

readLinesTail 27

Arguments

x a distance matrix or list of distance matrices

file file name to save to

threads number of threads used for serializing

Value

the input distance matrix or list of distance matrices

readLinesTail

Read the last N lines of a file

Description

Read the last N lines of a file

Usage

```
readLinesTail(x, n, ...)
```

Arguments

x The file name

n The last N lines to read

... Passed to scan()

read_bracken

Function for reading in a bracken taxonomy table

Description

The table will be converted to long form (sample ~ abundance). Only "_frac" or "_num" columns will be kept (see "keep_frac"). Taxonomy will be split into separate levels (see "tax_levs"). tidytable (w/ data.table) used to speed the process up.

```
read_bracken(
  infile,
  nrows = Inf,
  keep_frac = TRUE,
  tax_levs = c("Domain", "Phylum", "Class", "Order", "Family", "Genus", "Species"),
  ...
)
```

Arguments

infile Path to bracken table file

nrows Number of table rows to read. If Inf, all lines will be read.

keep_frac If TRUE, keep all columns ending in "_frac"; otherwise, keep "_num" columns.

tax_levs Taxonomic levels to separate the taxonomy column into.

... Params passed to fread()

Value

data.table

read_eggnog_mapper Function for reading in eggnog-mapper annotations and returning tidy

subsets of the info

Description

Many of the data in the eggnog-mapper annotation table (eg., generated by the LLG pipeline) is encoded as comma-delimited lists within a single column (eg., KEGG pathways). This makes it challenging to "tidy" the table.

Usage

```
read_eggnog_mapper(
  infile = NULL,
  cmd = NULL,
  sep = "\t",
  nrows = Inf,
  to_keep = c("COG", "KEGG pathway", "CAZy"),
  column_names = c("query_name", "seed_eggNOG_ortholog", "seed_ortholog_evalue",
    "seed_ortholog_score", "Predicted_taxonomic_group", "Predicted_protein_name",
    "Gene_Ontology_terms", "EC_number", "KEGG_ko", "KEGG_Pathway", "KEGG_Module",
    "KEGG_Reaction", "KEGG_rclass", "BRITE", "KEGG_TC", "CAZy", "BiGG_Reaction",
    "tax_scope__eggNOG_taxonomic_level_used_for_annotation", "eggNOG_OGs", "bestOG",
    "COG_Functional_Category", "eggNOG_free_text_description")
)
```

Arguments

infile Path to eggnog-annotation table file

cmd command instead of input file (eg., "gunzip -c INFILE")

sep table value delimiter

nrows Number of table rows to read. If Inf, all lines will be read. to_keep Which functional grouping to keep (eg., KEGG pathways)?

column_names The column names to use for the table (use NULL if the input table has column

names)

Robj_md5sum 29

Details

This function will read in the table and output a tidy table of one part of the table (eg., COG functional categories or KEGG pathways).

The function will also provide info on how to obtain metadata for function groupings.

Value

data.table

Robj_md5sum

Dump an R object as text to a temp file and get the md5sum of the file

Description

Dump an R object as text to a temp file and get the md5sum of the file

Usage

```
Robj_md5sum(Robj)
```

Arguments

Robj

Any R object

Value

md5sum

row_means

rowMeans that works inside a dplyr::mutate() call

Description

rowMeans that works inside a dplyr::mutate() call

```
row_means(..., na.rm = TRUE)
```

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row_sums

rowSums that works inside a dplyr::mutate() call

Description

rowSums that works inside a dplyr::mutate() call

Usage

```
row_sums(..., na.rm = TRUE)
```

scale_color_all

Great a better coloring scheme for taxon abundance barcharts

Description

The default coloring scheme for ggplot2 makes it hard to distinguish among data points in complex bar charts (eg., taxa plots). This function is a wrapper around scale_color_continuous() which changes the color scheme used.

Usage

```
scale_color_all(return_hex = FALSE, ...)
```

Arguments

return_hex

Return a vector of color hexidecimals instead of a plotting object.

Value

ScaleContinuous/ggproto object or vector

Examples

```
ggplot(mpg, aes(cty, hwy, color=class)) +
  geom_point() +
  scale_color_all()
```

scale_fill_all 31

scale_fill_all

Great a better coloring scheme for taxon abundance barcharts

Description

The default coloring scheme for ggplot2 makes it hard to distinguish among data points in complex bar charts (eg., taxa plots). This function is a wrapper around scale_color_continuous() which changes the color scheme used.

Usage

```
scale_fill_all(return_hex = FALSE, ...)
```

Arguments

return_hex

Return a vector of color hexidecimals instead of a plotting object.

Value

ScaleContinuous/ggproto object or vector

Examples

```
ggplot(mpg, aes(fl, hwy, fill=model)) +
  geom_bar(stat='identity') +
  scale_fill_all()
```

send_email

A helper function to send an email via the mail bash cmd

Description

A helper function to send an email via the mail bash cmd

```
send_email(
  body,
  subject = "R job complete",
  email = NULL,
  email_ext = "tuebingen.mpg.de"
)
```

32 snakemakeInfo

Arguments

body The email body

subject The email subject line

email The email address. If NULL, then username used

email_ext The part after the "at" symbol

Value

The output of the system() call

size_objects

Returns the sizes of R objects

Description

Returns the sizes of R objects

Usage

```
size_objects(Robj)
```

Arguments

Robj

Vector with the names of R objects as characters

Value

A list with the name of R objects as names and the formatted size of the objects

snakemakeInfo

snakemake conda info

Description

snakemake conda info

Usage

```
snakemakeInfo(config_file, pipeline_dir, conda_env)
```

Arguments

config_file The path to the config file

pipeline_dir The path to the pipeline_directory

conda_env The conda env that has snakemake installed

split_path 33

Value

The environment info

split_path

python's os.path.split() for R

Description

```
python's os.path.split() for R
```

Usage

```
split_path(x)
```

Arguments

Χ

The full file path

Value

A vector of all path parts

summary_x

Summary for numeric vectors that includes sd and stderr

Description

```
sd = standard deviation stderr = standard error of the mean (<math>sd(x) / sqrt(length(x)))
```

Usage

```
summary_x(x, label = NULL, sel_col = NULL, rnd = 3)
```

Arguments

x a numeric vector

label row name label for the output. If NULL, then the label will be the input object

label.

sel_col If "x" is data.table or data.frame, which column to assess?

rnd number of digits to round sd and stderr to

Value

a matrix

34 tidy_pcoa

taxonomy_levels

A simple function that returns a vector of taxonomy levels

Description

This just saves some typing, since I find myself constantly typing out: c('Domain', 'Phylum', 'Class', 'Order', 'Family', 'Genus', 'Species')

Usage

```
taxonomy_levels()
```

tidy_pcoa

PCoA on a 'long' (tidy) tibble, and a long tibble is returned

Description

Perform PCoA in a "tidy" way. If multiple diversity metrics are provided (eg., "bray" and "jaccard"), all PCoA results will be combined into one data.frame.

Usage

```
tidy_pcoa(
    df,
    taxon_col,
    sample_col,
    abundance_col,
    dists = c("bray", "jaccard", "wunifrac", "unifrac"),
    tree = NULL,
    threads = 1,
    threads_unifrac = 1,
    k = 2,
    dist_mtx_file = NULL,
    pcoa_file = NULL
)
```

Arguments

df	data.frame or tibble
taxon_col	the column specifying taxa or OTUs (no quotes needed)
sample_col	the column specifying sample names (no quotes needed)
abundance_col	the column specifying the taxon abundances in each sample (no quotes needed)
dists	vector of beta-diversity distances ('wunifrac' = weighted UniFrac, 'unifrac' = unweighted Unifrac; see vegan::vegsist for others)

unique_n 35

tree phylogeny for UniFrac calculations. It can have more tips that what is in the

data.frame

threads number of parallel calculations of each distance metric (1 thread per distance)

threads_unifrac

number of threads to use for wunifrac & unifrac calculations

k passed to cmdscale

dist_mtx_file file name for saving the distance matrices (qs serialization; use ".qs" for the file

extension)

pcoa_file file name for saving the raw pcoa results

Details

Weighted/Unweighted UniFrac is calculated via the rbiom R package. All other beta-diversity metrics are calculated via the vegan R package.

Value

a tibble of PCoA info for all selected "dists"

unique_n

Pretty print number of unique elements in a vector

Description

The result will be cat'ed to the screen. tidytable compatable. Maje

Usage

```
unique_n(x, label = "items", sel_col = NULL, ret = FALSE)
```

Arguments

x a vector or data.table. If data.table, sel_col must not be NULL

label what to call the items in the vector (eg., "samples")

sel_col If x is data.table or data.frame, which column to assess?

ret Return the unique values?

36 write_table

well2index Convert between wellID and column-num
--

Description

Useful for converting between WellIDs (eg., "A2") and well position in a plate (eg., 9)

Usage

```
well2index(x, plate_type = "96-well")
```

Arguments

x A vector of well IDs plate_type Either 96-well or 384-well

Value

A vector of plate positions

|--|

Description

This is most useful for working with IRkernl in Jupyter notebooks. If a data.table is provided, then fwrite is used; otherwise, write.table is used.

Usage

```
write_table(df, file, sep = "\t", quote = FALSE, row.names = FALSE, ...)
```

Arguments

df	data.frame or data.table to write out
file	Output file path
sep	the field separator string. Values within each row of x are separated by this string
quote	a logical value (TRUE or FALSE) or a numeric vector. If TRUE, any character or factor columns will be surrounded by double quotes.
row.names	either a logical value indicating whether the row names of x are to be written along with x, or a character vector of row names to be written.
	Passed to write.table (if data.frame) or fwrite (if data.table)

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