# Package 'ggpicrust2'

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Type Package

Title Make 'PICRUSt2' Output Analysis and Visualization Easier

Version 2.1.2

**Description** Provides a convenient way to analyze and visualize 'PICRUSt2' output with predefined plots and functions. Allows for generating statistical plots about microbiome functional predictions and offers customization options. Features a one-click option for creating publication-level plots, saving time and effort in producing professional-grade figures. Streamlines the 'PICRUSt2' analysis and visualization process. For more details, see Yang et al. (2023) <doi:10.1093/bioinformatics/btad470>.

BugReports https://github.com/cafferychen777/ggpicrust2/issues

URL https://github.com/cafferychen777/ggpicrust2,

https://cafferyang.com/ggpicrust2/

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compare\_daa\_results Compare the Consistency of Statistically Significant Features

## Description

This function compares the consistency and inconsistency of statistically significant features obtained using different methods in 'pathway\_daa' from the 'ggpicrust2' package. It creates a report showing the number of common and different features identified by each method, and the features themselves.

#### Arguments

daa\_results\_list

A list of data frames containing statistically significant features obtained using different methods.

method\_names A character vector of names for each method used.

p\_values\_threshold

A numeric value representing the threshold for the p-values. Features with p-values less than this threshold are considered statistically significant. Default is 0.05.

#### Value

A data frame with the comparison results. The data frame has the following columns:

- method: The name of the method.
- num\_features: The total number of statistically significant features obtained by the method.
- num\_common\_features: The number of features that are common to other methods.
- num\_diff\_features: The number of features that are different from other methods.
- diff\_features: The names of the features that are different from other methods.

```
library(magrittr)
library(ggpicrust2)
library(tibble)
data("metacyc_abundance")
data("metadata")
# Run pathway_daa function for multiple methods
methods <- c("DESeq2", "edgeR","Maaslin2")</pre>
daa_results_list <- lapply(methods, function(method) {</pre>
pathway_daa(abundance = metacyc_abundance %>% column_to_rownames("pathway"),
metadata = metadata, group = "Environment", daa_method = method)
})
names(daa_results_list) <- methods</pre>
# Correct Maaslin2 feature names by replacing dots with hyphens.
# Note: When using Maaslin2 as the differential abundance analysis method,
# it modifies the original feature names by replacing hyphens (-) with dots (.).
# This replacement can cause inconsistencies when trying to compare results from Maaslin2
# with those from other methods that do not modify feature names.
# Therefore, this line of code reverses that replacement, converting the dots back into
# hyphens for accurate and consistent comparisons across different methods.
daa_results_list[["Maaslin2"]]$feature <- gsub("\\.", "-", daa_results_list[["Maaslin2"]]$feature)</pre>
# Compare results across different methods
```

```
comparison_results <- compare_daa_results(daa_results_list = daa_results_list,
method_names = c("DESeq2", "edgeR", "Maaslin2"))
```

compare\_gsea\_daa

#### Description

This function compares the results from Gene Set Enrichment Analysis (GSEA) and Differential Abundance Analysis (DAA) to identify similarities and differences.

#### Usage

```
compare_gsea_daa(
  gsea_results,
  daa_results,
  plot_type = "venn",
  p_threshold = 0.05
)
```

## Arguments

gsea_results	A data frame containing GSEA results from the pathway_gsea function
daa_results	A data frame containing DAA results from the pathway_daa function
plot_type	A character string specifying the visualization type: "venn", "upset", "scatter", or "heatmap"
p_threshold	A numeric value specifying the significance threshold

#### Value

A ggplot2 object or a list containing the plot and comparison results

```
## Not run:
# Load example data
data(ko_abundance)
data(metadata)
# Prepare abundance data
abundance_data <- as.data.frame(ko_abundance)
rownames(abundance_data) <- abundance_data[, "#NAME"]
abundance_data <- abundance_data[, -1]
# Run GSEA analysis
gsea_results <- pathway_gsea(
    abundance = abundance_data,
    metadata = metadata,
    group = "Environment",
    pathway_type = "KEGG",
    method = "fgsea"
```

```
)
# Run DAA analysis
daa_results <- pathway_daa(
    abundance = abundance_data,
    metadata = metadata,
    group = "Environment"
)
# Compare results
comparison <- compare_gsea_daa(
    gsea_results = gsea_results,
    daa_results = daa_results,
    plot_type = "venn"
)
## End(Not run)</pre>
```

```
compare_metagenome_results
```

Compare Metagenome Results

## Description

**Compare Metagenome Results** 

## Arguments

metagenomes	A list of metagenomes matrices with rows as KOs and columns as samples. Each matrix in the list should correspond to a different metagenome.
names	A vector of names for the metagenomes in the same order as in the 'metagenomes' list.
daa_method	A character specifying the method for differential abundance analysis (DAA). Possible choices are: "ALDEx2", "DESeq2", "edgeR", "limma voom", "metagenome-Seq", "LinDA", "Maaslin2", and "Lefse". The default is "ALDEx2".
p.adjust	A character specifying the method for p-value adjustment. Possible choices are: "BH" (Benjamini-Hochberg), "holm", "bonferroni", "hochberg", "fdr", and "none". The default is "BH".
reference	A character specifying the reference group level for DAA. This parameter is used when there are more than two groups. The default is NULL.

#### Value

A list containing two elements:

• "daa": a list of results from the 'pathway\_daa' function. Each result is a data frame containing the differential abundance analysis results with columns for the feature ID, the test statistic, the raw p-value, and the adjusted p-value.

• "correlation": a list with two elements: "cor\_matrix" and "p\_matrix", which are matrices of Spearman correlation coefficients and their corresponding p-values, respectively, between every pair of metagenomes.

#### Examples

```
library(dplyr)
library(ComplexHeatmap)
# Generate example data
set.seed(123)
# First metagenome
metagenome1 <- abs(matrix(rnorm(1000), nrow = 100, ncol = 10))</pre>
rownames(metagenome1) <- paste0("KO", 1:100)</pre>
colnames(metagenome1) <- paste0("sample", 1:10)</pre>
# Second metagenome
metagenome2 <- abs(matrix(rnorm(1000), nrow = 100, ncol = 10))</pre>
rownames(metagenome2) <- paste0("KO", 1:100)</pre>
colnames(metagenome2) <- paste0("sample", 1:10)</pre>
# Put the metagenomes into a list
metagenomes <- list(metagenome1, metagenome2)</pre>
# Define names
names <- c("metagenome1", "metagenome2")</pre>
# Call the function
results <- compare_metagenome_results(metagenomes, names, daa_method = "LinDA")
# Print the correlation matrix
print(results$correlation$cor_matrix)
# Print the p-value matrix
print(results$correlation$p_matrix)
```

daa\_annotated\_results\_df Differentially Abundant Analysis Results with Annotation

#### Description

This is a result dataset after processing 'kegg\_abundance' through the 'pathway\_daa' with the LinDA method and further annotation with 'pathway\_annotation'.

#### Usage

daa\_annotated\_results\_df

#### Format

A data frame with 10 variables:

adj\_method Method used for adjusting p-values.

feature Feature being tested.

group1 One group in the comparison.
group2 The other group in the comparison.
method Statistical test used.
p\_adjust Adjusted p-value.
p\_values P-values from the statistical test.
pathway\_class Class of the pathway.
pathway\_description Description of the pathway.
pathway\_map Map of the pathway.
pathway\_name Name of the pathway.

## Source

From ggpicrust2 package demonstration.

#### References

Douglas GM, Maffei VJ, Zaneveld J, Yurgel SN, Brown JR, Taylor CM, Huttenhower C, Langille MGI. PICRUSt2 for prediction of metagenome functions. Nat Biotechnol. 2020.

daa\_results\_df DAA Results Dataset

#### Description

This dataset is the result of processing 'kegg\_abundance' through the 'LinDA' method in the 'pathway\_daa' function. It includes information about the feature, groups compared, p values, and method used.

#### Usage

daa\_results\_df

## Format

A data frame with columns:

adj\_method Method used for p-value adjustment.

feature The feature (pathway) being compared.

group1 The first group in the comparison.

group2 The second group in the comparison.

method The method used for the comparison.

**p\_adjust** The adjusted p-value from the comparison.

**p\_values** The raw p-value from the comparison.

#### Source

From ggpicrust2 package demonstration.

## References

Douglas GM, Maffei VJ, Zaneveld J, Yurgel SN, Brown JR, Taylor CM, Huttenhower C, Langille MGI. PICRUSt2 for prediction of metagenome functions. Nat Biotechnol. 2020.

ggpicrust2	This function integrates pathway name/description annotations, ten of
	the most advanced differential abundance (DA) methods, and visual- ization of DA results.
	- 0

## Description

This function integrates pathway name/description annotations, ten of the most advanced differential abundance (DA) methods, and visualization of DA results.

## Usage

```
ggpicrust2(
  file = NULL,
  data = NULL,
 metadata,
  group,
  pathway,
  daa_method = "ALDEx2",
  ko_to_kegg = FALSE,
  p.adjust = "BH",
  order = "group",
  p_values_bar = TRUE,
  x_{lab} = NULL,
  select = NULL,
  reference = NULL,
  colors = NULL
)
```

# Arguments

file	A character string representing the file path of the input file containing KO abun-
	dance data in picrust2 export format. The input file should have KO identifiers
	in the first column and sample identifiers in the first row. The remaining cells
	should contain the abundance values for each KO-sample pair.
data	An optional data.frame containing KO abundance data in the same format as the input file. If provided, the function will use this data instead of reading from the
	file. By default, this parameter is set to NULL.

## ggpicrust2

metadata	A tibble, consisting of sample information
group	A character, name of the group
pathway	A character, consisting of "EC", "KO", "MetaCyc"
daa_method	a character specifying the method for differential abundance analysis, default is "ALDEx2", choices are: - "ALDEx2": ANOVA-Like Differential Expression tool for high throughput sequencing data - "DESeq2": Differential expression analysis based on the negative binomial distribution using DESeq2 - "edgeR": Exact test for differences between two groups of negative-binomially distributed counts using edgeR - "limma voom": Limma-voom framework for the analy- sis of RNA-seq data - "metagenomeSeq": Fit logistic regression models to test for differential abundance between groups using metagenomeSeq - "LinDA": Linear models for differential abundance analysis of microbiome compositional data - "Maaslin2": Multivariate Association with Linear Models (MaAsLin2) for differential abundance analysis
ko_to_kegg	A character to control the conversion of KO abundance to KEGG abundance
p.adjust	a character specifying the method for p-value adjustment, default is "BH", choices are: - "BH": Benjamini-Hochberg correction - "holm": Holm's correction - "bonferroni": Bonferroni correction - "hochberg": Hochberg's correction - "fdr": False discovery rate correction - "none": No p-value adjustment.
order	A character to control the order of the main plot rows
p_values_bar	A character to control if the main plot has the p_values bar
x_lab	A character to control the x-axis label name, you can choose from "feature", "pathway_name" and "description"
select	A vector consisting of pathway names to be selected
reference	A character, a reference group level for several DA methods
colors	A vector consisting of colors number

## Value

daa.results.df, a dataframe of DA results A list of sub-lists, each containing a ggplot2 plot ('plot') and a dataframe of differential abundance results ('results') for a specific DA method. Each plot visualizes the differential abundance results of a specific DA method, and the corresponding dataframe contains the results used to create the plot.

```
daa_method = "LinDA",
                                  ko_to_kegg = "TRUE",
                                  order = "pathway_class",
                                  p_values_bar = TRUE,
                                  x_lab = "pathway_name")
# Run ggpicrust2 with imported data.frame
abundance_data <- read_delim(abundance_file, delim="\t", col_names=TRUE, trim_ws=TRUE)
# Run ggpicrust2 with input data
results_data_input <- ggpicrust2(data = abundance_data,</pre>
                                  metadata = metadata,
                                  group = "your_group_column",
                                  pathway = "KO",
                                  daa_method = "LinDA",
                                  ko_to_kegg = "TRUE",
                                  order = "pathway_class",
                                  p_values_bar = TRUE,
                                  x_lab = "pathway_name")
# Access the plot and results dataframe for the first DA method
example_plot <- results_file_input[[1]]$plot</pre>
example_results <- results_file_input[[1]]$results</pre>
# Use the example data in ggpicrust2 package
data(ko_abundance)
data(metadata)
results_file_input <- ggpicrust2(data = ko_abundance,</pre>
                                  metadata = metadata,
                                  group = "Environment",
                                  pathway = "KO",
                                  daa_method = "LinDA",
                                  ko_to_kegg = TRUE,
                                  order = "pathway_class",
                                  p_values_bar = TRUE,
                                  x_lab = "pathway_name")
# Analyze the EC or MetaCyc pathway
data(metacyc_abundance)
results_file_input <- ggpicrust2(data = metacyc_abundance,
                                  metadata = metadata,
                                  group = "Environment",
                                  pathway = "MetaCyc",
                                  daa_method = "LinDA",
                                  ko_to_kegg = FALSE,
                                  order = "group",
                                  p_values_bar = TRUE,
                                  x_lab = "description")
## End(Not run)
```

ggpicrust2\_extended Integrated analysis with ggpicrust2 including GSEA

## Description

This function extends the ggpicrust2 functionality to include Gene Set Enrichment Analysis (GSEA).

#### Usage

```
ggpicrust2_extended(..., run_gsea = FALSE, gsea_params = list())
```

## Arguments

	Parameters passed to ggpicrust2()
run_gsea	Logical value indicating whether to perform GSEA analysis
gsea_params	List of parameters to pass to pathway_gsea()

## Value

A list containing ggpicrust2 and GSEA results

```
## Not run:
# Load example data
data(ko_abundance)
data(metadata)
# Run integrated analysis
integrated_results <- ggpicrust2_extended(</pre>
  data = ko_abundance,
  metadata = metadata,
  group = "Environment",
  pathway = "KO",
  daa_method = "LinDA",
  ko_to_kegg = TRUE,
  run_gsea = TRUE,
  gsea_params = list(
    method = "fgsea",
    rank_method = "signal2noise",
    nperm = 1000
  )
)
# Access DAA results
daa_results <- integrated_results$daa_results</pre>
# Access GSEA results
gsea_results <- integrated_results$gsea_results</pre>
# Access plots
daa_plot <- integrated_results$daa_plot</pre>
gsea_plot <- integrated_results$gsea_plot</pre>
## End(Not run)
```

```
gsea_pathway_annotation
```

Annotate GSEA results with pathway information

## Description

This function adds pathway annotations to GSEA results, including pathway names, descriptions, and classifications.

## Usage

```
gsea_pathway_annotation(gsea_results, pathway_type = "KEGG")
```

## Arguments

gsea_results	A data frame containing GSEA results from the pathway_gsea function
pathway_type	A character string specifying the pathway type: "KEGG", "MetaCyc", or "GO"

#### Value

A data frame with annotated GSEA results

```
## Not run:
# Load example data
data(ko_abundance)
data(metadata)
# Prepare abundance data
abundance_data <- as.data.frame(ko_abundance)</pre>
rownames(abundance_data) <- abundance_data[, "#NAME"]</pre>
abundance_data <- abundance_data[, -1]</pre>
# Run GSEA analysis
gsea_results <- pathway_gsea(</pre>
 abundance = abundance_data,
 metadata = metadata,
 group = "Environment",
 pathway_type = "KEGG",
 method = "fgsea"
)
# Annotate results
annotated_results <- gsea_pathway_annotation(</pre>
 gsea_results = gsea_results,
 pathway_type = "KEGG"
)
## End(Not run)
```

## Description

This function imports DAA results from an external platform such as MicrobiomeAnalyst. It can be used to compare the results obtained from different platforms.

## Arguments

file_path	a character string specifying the path to the CSV file containing the DAA re- sults from MicrobiomeAnalyst. If this parameter is NULL and no data frame is provided, an error will be thrown. Default is NULL.
data	a data frame containing the DAA results from MicrobiomeAnalyst. If this parameter is NULL and no file path is provided, an error will be thrown. Default is NULL.
method	a character string specifying the method used for the DAA. This will be added as a new column in the returned data frame. Default is "MicrobiomeAnalyst".
group_levels	a character vector specifying the group levels for the DAA. This will be added as new columns in the returned data frame. Default is c("control", "treatment").

## Value

a data frame containing the DAA results from MicrobiomeAnalyst with additional columns for the method and group levels.

## Examples

```
## Not run:
# Assuming you have a CSV file named "DAA_results.csv" in your current directory
daa_results <- import_MicrobiomeAnalyst_daa_results(file_path = "DAA_results.csv")</pre>
```

## End(Not run)

kegg\_abundance KEGG Abundance Dataset

#### Description

A dataset derived from 'ko\_abundance' by the function 'ko2kegg\_abundance' in the ggpicrust2 package. Each row corresponds to a KEGG pathway, and each column corresponds to a sample.

## Usage

kegg\_abundance

#### Format

A data frame where rownames are KEGG pathways and column names are individual sample names, including: "SRR11393730", "SRR11393731", "SRR11393732", "SRR11393733", "SRR11393734", "SRR11393735", "SRR11393736", "SRR11393737", "SRR11393738", "SRR11393739", "SRR11393740", "SRR11393741", "SRR11393742", "SRR11393743", "SRR11393744", "SRR11393745", "SRR11393746", "SRR11393747", "SRR11393747", "SRR11393748", "SRR11393749", "SRR11393750", "SRR11393751", "SRR11393752", "SRR11393753", "SRR11393754", "SRR11393755", "SRR11393756", "SRR11393757", "SRR11393757", "SRR11393758", "SRR11393759", "SRR11393760", "SRR11393761", "SRR11393762", "SRR11393763", "SRR11393764", "SRR11393765", "SRR11393766", "SRR11393767", "SRR11393768", "SRR11393769", "SRR11393776", "SRR11393776", "SRR11393777", "SRR11393777", "SRR11393777", "SRR11393777", "SRR11393777", "SRR11393777", "SRR11393777", "SRR11393776", "SRR11393777", "SR1393777", "SR1139377

## Source

From ggpicrust2 package demonstration.

## References

Douglas GM, Maffei VJ, Zaneveld J, Yurgel SN, Brown JR, Taylor CM, Huttenhower C, Langille MGI. PICRUSt2 for prediction of metagenome functions. Nat Biotechnol. 2020.

ko2kegg_abundance	Convert KO abundance in picrust2 export files to KEGG pathway
	abundance

## Description

This function takes a file containing KO (KEGG Orthology) abundance data in picrust2 export format and converts it to KEGG pathway abundance data. The input file should be in .tsv, .txt, or .csv format.

#### Usage

```
ko2kegg_abundance(file = NULL, data = NULL)
```

#### Arguments

file	A character string representing the file path of the input file containing KO abun- dance data in picrust2 export format. The input file should have KO identifiers in the first column and sample identifiers in the first row. The remaining cells should contain the abundance values for each KO-sample pair.
data	An optional data.frame containing KO abundance data in the same format as the input file. If provided, the function will use this data instead of reading from the file. By default, this parameter is set to NULL.

#### ko\_abundance

#### Value

A data frame with KEGG pathway abundance values. Rows represent KEGG pathways, identified by their KEGG pathway IDs. Columns represent samples, identified by their sample IDs from the input file. Each cell contains the abundance of a specific KEGG pathway in a given sample, calculated by summing the abundances of the corresponding KOs in the input file.

#### Examples

```
## Not run:
library(ggpicrust2)
library(readr)
# Example 1: Demonstration with a hypothetical input file
# Prepare an input file path
input_file <- "path/to/your/picrust2/results/pred_metagenome_unstrat.tsv"</pre>
# Run ko2kegg_abundance function
kegg_abundance <- ko2kegg_abundance(file = input_file)</pre>
# Alternatively, read the data from a file and use the data argument
file_path <- "path/to/your/picrust2/results/pred_metagenome_unstrat.tsv"</pre>
ko_abundance <- read_delim(file_path, delim = "\t")</pre>
kegg_abundance <- ko2kegg_abundance(data = ko_abundance)</pre>
# Example 2: Working with real data
# In this case, we're using an existing dataset from the ggpicrust2 package.
# Load the data
data(ko_abundance)
# Apply the ko2kegg_abundance function to our real dataset
kegg_abundance <- ko2kegg_abundance(data = ko_abundance)</pre>
## End(Not run)
```

ko\_abundance KO Abundance Dataset

#### Description

This is a demonstration dataset from the ggpicrust2 package, representing the output of PICRUSt2. Each row represents a KO (KEGG Orthology) group, and each column corresponds to a sample.

#### Usage

ko\_abundance

## Format

A data frame where rownames are KO groups and column names include #NAME and individual sample names, such as: "#NAME", "SRR11393730", "SRR11393731", "SRR11393732", "SRR11393733", "SRR11393734", "SRR11393735", "SRR11393736", "SRR11393737", "SRR11393738", "SRR11393749", "SRR11393740", "SRR11393741", "SRR11393742", "SRR11393743", "SRR11393744", "SRR11393745", "SRR11393746", "SRR11393747", "SRR11393748", "SRR11393749", "SRR11393750", "SRR11393751", "SRR11393752", "SRR11393753", "SRR11393754", "SRR11393755", "SRR11393756", "SRR11393757", "SRR11393756", "SRR11393757", "SRR11393756", "SRR11393756", "SRR11393766", "SRR11393761", "SRR11393762", "SRR11393763", "SRR11393764", "SRR11393765", "SRR11393766", "SRR11393767", "SRR11393768", "SRR11393765", "SRR11393766", "SRR11393767", "SRR11393774", "SRR11393767", "SRR11393776", "SRR11393777", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393777", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393776", "SRR11393777", "SRR11393776", "SRR11393777", "SRR11393777", "SRR11393778", "SRR11393779"

#### Source

From ggpicrust2 package demonstration.

#### References

Douglas GM, Maffei VJ, Zaneveld J, Yurgel SN, Brown JR, Taylor CM, Huttenhower C, Langille MGI. PICRUSt2 for prediction of metagenome functions. Nat Biotechnol. 2020.

metacyc\_abundance MetaCyc Abundance Dataset

#### Description

This is a demonstration dataset from the ggpicrust2 package, representing the output of PICRUSt2. Each row represents a MetaCyc pathway, and each column corresponds to a sample.

#### Usage

metacyc\_abundance

## Format

A data frame where rownames are MetaCyc pathways and column names include "pathway" and individual sample names, such as: "pathway", "SRR11393730", "SRR11393731", "SRR11393732", "SRR11393733", "SRR11393734", "SRR11393735", "SRR11393736", "SRR11393737", "SRR11393738", "SRR11393739", "SRR11393740", "SRR11393741", "SRR11393742", "SRR11393743", "SRR11393744", "SRR11393745", "SRR11393746", "SRR11393747", "SRR11393748", "SRR11393749", "SRR11393750", "SRR11393751", "SRR11393752", "SRR11393753", "SRR11393754", "SRR11393755", "SRR11393756", "SRR11393757", "SRR11393758", "SRR11393759", "SRR11393760", "SRR11393761", "SRR11393762", "SRR11393763", "SRR11393764", "SRR11393765", "SRR11393766", "SRR11393767", "SRR11393768", "SRR11393769", "SRR11393770", "SRR11393771", "SRR11393772", "SRR11393773", "SRR11393774", "SRR11393775", "SRR11393776", "SRR11393777", "SRR11393778", "SRR11393779"

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## metadata

## Source

From ggpicrust2 package demonstration.

## References

Douglas GM, Maffei VJ, Zaneveld J, Yurgel SN, Brown JR, Taylor CM, Huttenhower C, Langille MGI. PICRUSt2 for prediction of metagenome functions. Nat Biotechnol. 2020.

metadata

Metadata for ggpicrust2 Demonstration

## Description

This is a demonstration dataset from the ggpicrust2 package. It provides the metadata required for the demonstration functions in the package. The dataset includes environmental information for each sample.

#### Usage

metadata

#### Format

A tibble with each row representing metadata for a sample.

Sample1 Metadata for Sample1, including Environment

Sample2 Metadata for Sample2, including Environment

••• ···

## Source

ggpicrust2 package demonstration.

#### References

Douglas GM, Maffei VJ, Zaneveld J, Yurgel SN, Brown JR, Taylor CM, Huttenhower C, Langille MGI. PICRUSt2 for prediction of metagenome functions. Nat Biotechnol. 2020.

#### Description

This function has two primary use cases: 1. Annotating pathway information using the output file from PICRUSt2. 2. Annotating pathway information from the output of 'pathway\_daa' function, and converting KO abundance to KEGG pathway abundance when 'ko\_to\_kegg' is set to TRUE.

## Usage

```
pathway_annotation(
  file = NULL,
  pathway = NULL,
  daa_results_df = NULL,
  ko_to_kegg = FALSE
)
```

#### Arguments

file	A character, address to store PICRUSt2 export files. Provide this parameter when using the function for the first use case.
pathway	A character, consisting of "EC", "KO", "MetaCyc"
daa_results_df	A data frame, output of pathway_daa. Provide this parameter when using the function for the second use case.
ko_to_kegg	A logical, decide if convert KO abundance to KEGG pathway abundance. Default is FALSE. Set to TRUE when using the function for the second use case.

#### Value

A data frame containing pathway annotation information. The data frame has the following columns:

- feature: The feature ID of the pathway (e.g., KO, EC, or MetaCyc ID).
- description: The description or name of the pathway.
- Other columns depending on the input parameters and type of pathway.

If ko\_to\_kegg is set to TRUE, the output data frame will also include the following columns:

- pathway\_name: The name of the KEGG pathway.
- pathway\_description: The description of the KEGG pathway.
- pathway\_class: The class of the KEGG pathway.
- pathway\_map: The KEGG pathway map ID.

#### pathway\_daa

#### Examples

```
## Not run:
# Prepare the required input files and data frames
# Then, you can use the pathway_annotation function as follows:
# Use case 1: Annotating pathway information using the output file from PICRUSt2
result1 <- pathway_annotation(file = "path/to/picrust2/export/file.txt",</pre>
                               pathway = "KO",
                               daa_results_df = NULL,
                               ko_to_kegg = FALSE)
# Use case 2: Annotating pathway information from the output of pathway_daa function
# and converting KO abundance to KEGG pathway abundance
# This use case will be demonstrated using both a hypothetical example, and a real dataset.
## Hypothetical example
hypothetical_daa_results_df <- data.frame() # Replace this with your actual data frame
result2 <- pathway_annotation(file = NULL,</pre>
                              pathway = "KO",
                              daa_results_df = hypothetical_daa_results_df,
                              ko_to_kegg = TRUE)
## Real dataset example
# Load the real dataset
data(daa_results_df)
result3 <- pathway_annotation(file = NULL,</pre>
                              pathway = "KO",
                               daa_results_df = daa_results_df,
                               ko_to_kegg = TRUE)
## End(Not run)
```

```
pathway_daa
```

Differential Abundance Analysis for Predicted Functional Pathways

## Description

Performs differential abundance analysis on predicted functional pathway data using various statistical methods. This function supports multiple methods for analyzing differences in pathway abundance between groups, including popular approaches like ALDEx2, DESeq2, edgeR, and others.

#### Usage

```
pathway_daa(
   abundance,
   metadata,
   group,
   daa_method = "ALDEx2",
```

```
select = NULL,
p.adjust = "BH",
reference = NULL,
...
)
```

# Arguments

abundance	A data frame or matrix containing predicted functional pathway abundance, with pathways/features as rows and samples as columns. The column names should match the sample names in metadata. Values should be counts or abundance measurements.
metadata	A data frame or tibble containing sample information. Must include a 'sample' column with sample identifiers matching the column names in abundance data.
group	Character string specifying the column name in metadata that contains group information for differential abundance analysis.
daa_method	Character string specifying the method for differential abundance analysis. Available choices are:
	• "ALDEx2": ANOVA-Like Differential Expression tool
	• "DESeq2": Differential expression analysis based on negative binomial dis- tribution
	• "edgeR": Exact test for differences between groups using negative bino- mial model
	• "limma voom": Limma-voom framework for RNA-seq analysis
	<ul> <li>"metagenomeSeq": Zero-inflated Gaussian mixture model</li> </ul>
	• "LinDA": Linear models for differential abundance analysis
	"Maaslin2": Multivariate Association with Linear Models
	• "Lefser": Linear discriminant analysis effect size
	Default is "ALDEx2".
select	Vector of sample names to include in the analysis. If NULL (default), all samples are included.
p.adjust	Character string specifying the method for p-value adjustment. Choices are:
	• "BH": Benjamini-Hochberg procedure (default)
	• "holm": Holm's step-down method
	"bonferroni": Bonferroni correction
	<ul> <li>"hochberg": Hochberg's step-up method</li> </ul>
	• "fdr": False Discovery Rate
	<ul> <li>"none": No adjustment</li> </ul>
reference	Character string specifying the reference level for the group comparison. If NULL (default), the first level is used as reference.
	Additional arguments passed to the specific DAA method

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#### pathway\_daa

#### Value

A data frame containing the differential abundance analysis results. The structure of the results depends on the chosen DAA method. For methods that support multi-group comparisons (like LinDA), when there are more than two groups, the results will contain separate rows for each feature in each pairwise comparison between the reference group and each non-reference group. The data frame includes the following columns:

- feature: Feature/pathway identifier
- method: The DAA method used
- group1: Reference group
- group2: Comparison group
- p\_values: P-values for the comparison
- p\_adjust: Adjusted p-values
- · adj\_method: Method used for p-value adjustment

Some methods may provide additional columns, such as log2FoldChange for effect size information.

## References

- ALDEx2: Fernandes et al. (2014) Unifying the analysis of high-throughput sequencing datasets: characterizing RNA-seq, 16S rRNA gene sequencing and selective growth experiments by compositional data analysis. Microbiome.
- DESeq2: Love et al. (2014) Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biology.
- edgeR: Robinson et al. (2010) edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics.
- limma-voom: Law et al. (2014) voom: precision weights unlock linear model analysis tools for RNA-seq read counts. Genome Biology.
- metagenomeSeq: Paulson et al. (2013) Differential abundance analysis for microbial markergene surveys. Nature Methods.
- Maaslin2: Mallick et al. (2021) Multivariable Association Discovery in Population-scale Meta-omics Studies.

## Examples

results <- pathway\_daa(</pre>

```
# Load example data
data(ko_abundance)
data(metadata)
# Prepare abundance data
abundance_data <- as.data.frame(ko_abundance)
rownames(abundance_data) <- abundance_data[, "#NAME"]
abundance_data <- abundance_data[, -1]
# Run differential abundance analysis using ALDEx2
```

```
abundance = abundance_data,
 metadata = metadata,
 group = "Environment"
)
# Using a different method (DESeq2)
deseq_results <- pathway_daa(</pre>
 abundance = abundance_data,
 metadata = metadata,
 group = "Environment"
 daa_method = "DESeq2"
)
# Create example data with more samples
abundance <- data.frame(</pre>
 sample1 = c(10, 20, 30),
 sample2 = c(20, 30, 40),
 sample3 = c(30, 40, 50),
 sample4 = c(40, 50, 60),
 sample5 = c(50, 60, 70),
 row.names = c("pathway1", "pathway2", "pathway3")
)
metadata <- data.frame(</pre>
 sample = c("sample1", "sample2", "sample3", "sample4", "sample5"),
 group = c("control", "control", "treatment", "treatment", "treatment")
)
# Run differential abundance analysis using ALDEx2
results <- pathway_daa(abundance, metadata, "group")</pre>
# Using a different method (limma voom instead of DESeq2 for this small example)
limma_results <- pathway_daa(abundance, metadata, "group",</pre>
                             daa_method = "limma voom")
# Analyze specific samples only
subset_results <- pathway_daa(abundance, metadata, "group",</pre>
                              select = c("sample1", "sample2", "sample3", "sample4"))
```

pathway_errorbar
------------------

The function pathway\_errorbar() is used to visualize the results of functional pathway differential abundance analysis as error bar plots.

#### Description

The function pathway\_errorbar() is used to visualize the results of functional pathway differential abundance analysis as error bar plots.

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#### Arguments

abundance	A data frame with row names representing pathways and column names repre- senting samples. Each element represents the relative abundance of the corre- sponding pathway in the corresponding sample.
daa_results_df	A data frame containing the results of the differential abundance analysis of the pathways, generated by the pathway_daa function. x_lab should be a column name of daa_results_df.
Group	A data frame or a vector that assigns each sample to a group. The groups are used to color the samples in the figure.
ko_to_kegg	A logical parameter indicating whether there was a convertion that convert ko abundance to kegg abundance.
p_values_thresh	nold
	A numeric parameter specifying the threshold for statistical significance of dif- ferential abundance. Pathways with p-values below this threshold will be con- sidered significant.
order	A parameter controlling the ordering of the rows in the figure. The options are: "p_values" (order by p-values), "name" (order by pathway name), "group" (or- der by the group with the highest mean relative abundance), or "pathway_class" (order by the pathway category).
select	A vector of pathway names to be included in the figure. This can be used to limit the number of pathways displayed. If NULL, all pathways will be displayed.
p_value_bar	A logical parameter indicating whether to display a bar showing the p-value threshold for significance. If TRUE, the bar will be displayed.
colors	A vector of colors to be used to represent the groups in the figure. Each color corresponds to a group.
x_lab	A character string to be used as the x-axis label in the figure. The default value is "description" for KOs'descriptions and "pathway_name" for KEGG pathway names.

## Value

A ggplot2 plot showing the error bar plot of the differential abundance analysis results for the functional pathways. The plot visualizes the differential abundance results of a specific differential abundance analysis method. The corresponding dataframe contains the results used to create the plot.

```
## Not run:
# Example 1: Analyzing KEGG pathway abundance
metadata <- read_delim(
    "path/to/your/metadata.txt",
    delim = "\t",
    escape_double = FALSE,
    trim_ws = TRUE
)
```

```
# data(metadata)
kegg_abundance <- ko2kegg_abundance(</pre>
  "path/to/your/pred_metagenome_unstrat.tsv"
)
# data(kegg_abundance)
# Please change group to "your_group_column" if you are not using example dataset
group <- "Environment"</pre>
daa_results_df <- pathway_daa(</pre>
 abundance = kegg_abundance,
 metadata = metadata,
 group = group,
 daa_method = "ALDEx2",
 select = NULL,
 reference = NULL
)
# Please check the unique(daa_results_df$method) and choose one
daa_sub_method_results_df <- daa_results_df[daa_results_df$method</pre>
== "ALDEx2_Welch's t test", ]
daa_annotated_sub_method_results_df <- pathway_annotation(</pre>
 pathway = "KO",
 daa_results_df = daa_sub_method_results_df,
 ko_to_kegg = TRUE
)
# Please change Group to metadata$your_group_column if you are not using example dataset
Group <- metadata$Environment</pre>
p <- pathway_errorbar(</pre>
 abundance = kegg_abundance,
 daa_results_df = daa_annotated_sub_method_results_df,
 Group = Group,
 p_values_threshold = 0.05,
 order = "pathway_class",
 select = daa_annotated_sub_method_results_df %>%
 arrange(p_adjust) %>%
 slice(1:20) %>%
 select("feature") %>% pull(),
 ko_to_kegg = TRUE,
 p_value_bar = TRUE,
 colors = NULL,
 x_lab = "pathway_name"
)
# Example 2: Analyzing EC, MetaCyc, KO without conversions
metadata <- read_delim(</pre>
  "path/to/your/metadata.txt",
```

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#### pathway\_gsea

```
delim = " \ t'',
  escape_double = FALSE,
  trim_ws = TRUE
)
# data(metadata)
metacyc_abundance <- read.delim("path/to/your/metacyc_abundance.tsv")</pre>
# data(metacyc_abundance)
group <- "Environment"</pre>
daa_results_df <- pathway_daa(</pre>
  abundance = metacyc_abundance %>% column_to_rownames("pathway"),
  metadata = metadata,
  group = group,
  daa_method = "LinDA",
  select = NULL,
  reference = NULL
)
daa_annotated_results_df <- pathway_annotation(</pre>
  pathway = "MetaCyc",
  daa_results_df = daa_results_df,
  ko_to_kegg = FALSE
)
Group <- metadata$Environment</pre>
p <- pathway_errorbar(</pre>
  abundance = metacyc_abundance %>% column_to_rownames("pathway"),
  daa_results_df = daa_annotated_results_df,
  Group = Group,
  p_values_threshold = 0.05,
  order = "group",
  select = NULL,
  ko_to_kegg = FALSE,
  p_value_bar = TRUE,
  colors = NULL,
  x_lab = "description"
)
## End(Not run)
```

pathway\_gsea

# Description

This function performs Gene Set Enrichment Analysis (GSEA) on PICRUSt2 predicted functional data to identify enriched pathways between different conditions.

## Usage

```
pathway_gsea(
   abundance,
   metadata,
   group,
   pathway_type = "KEGG",
   method = "fgsea",
   rank_method = "signal2noise",
   nperm = 1000,
   min_size = 10,
   max_size = 500,
   p.adjust = "BH",
   seed = 42
)
```

## Arguments

abundance	A data frame containing KO/EC/MetaCyc abundance data, with features as rows and samples as columns
metadata	A data frame containing sample metadata
group	A character string specifying the column name in metadata that contains the grouping variable
pathway_type	A character string specifying the pathway type: "KEGG", "MetaCyc", or "GO"
method	A character string specifying the GSEA method: "fgsea", "GSEA", or "cluster-Profiler"
rank_method	A character string specifying the ranking statistic: "signal2noise", "t_test", "log2_ratio", or "diff_abundance"
nperm	An integer specifying the number of permutations
min_size	An integer specifying the minimum gene set size
max_size	An integer specifying the maximum gene set size
p.adjust	A character string specifying the p-value adjustment method
seed	An integer specifying the random seed for reproducibility

## Value

A data frame containing GSEA results

#### pathway\_heatmap

## Examples

```
## Not run:
# Load example data
data(ko_abundance)
data(metadata)
# Prepare abundance data
abundance_data <- as.data.frame(ko_abundance)</pre>
rownames(abundance_data) <- abundance_data[, "#NAME"]</pre>
abundance_data <- abundance_data[, -1]</pre>
# Run GSEA analysis
gsea_results <- pathway_gsea(</pre>
  abundance = abundance_data,
  metadata = metadata,
  group = "Environment",
  pathway_type = "KEGG",
  method = "fgsea"
)
# Visualize results
visualize_gsea(gsea_results, plot_type = "enrichment_plot", n_pathways = 10)
## End(Not run)
```

pathway\_heatmap Create pathway heatmap

# Description

This function creates a heatmap of the predicted functional pathway abundance data. The function first makes the abundance data relative, then converts the abundance data to a long format and orders the samples based on the environment information. The heatmap is then created using the 'ggplot2' library.

#### Arguments

abundance	A matrix or data frame of pathway abundance data, where columns correspond to samples and rows correspond to pathways. Must contain at least two samples.
metadata	A data frame of metadata, where each row corresponds to a sample and each column corresponds to a metadata variable.
group	A character string specifying the column name in the metadata data frame that contains the group variable. Must contain at least two groups.
colors	A vector of colors used for the background of the facet labels in the heatmap. If NULL or not provided, a default color set is used for the facet strips.
font_size	A numeric value specifying the font size for the heatmap.
show_row_names	A logical value indicating whether to show row names in the heatmap.

show\_legendA logical value indicating whether to show the legend in the heatmap.custom\_themeA custom theme for the heatmap.

#### Value

A ggplot heatmap object representing the heatmap of the predicted functional pathway abundance data.

```
## Not run:
library(ggpicrust2)
library(ggh4x)
library(dplyr)
library(tidyr)
library(tibble)
library(magrittr)
# Create example functional pathway abundance data
kegg_abundance_example <- matrix(rnorm(30), nrow = 3, ncol = 10)</pre>
colnames(kegg_abundance_example) <- paste0("Sample", 1:10)</pre>
rownames(kegg_abundance_example) <- c("PathwayA", "PathwayB", "PathwayC")</pre>
# Create example metadata
metadata_example <- data.frame(</pre>
  sample_name = colnames(kegg_abundance_example),
  group = factor(rep(c("Control", "Treatment"), each = 5))
)
# Custom colors for facet strips
custom_colors <- c("skyblue", "salmon")</pre>
# Create a heatmap using custom colors for facet strips
pathway_heatmap(kegg_abundance_example, metadata_example, "group", colors = custom_colors)
# Use real dataset
data("metacyc_abundance")
data("metadata")
metacyc_daa_results_df <- pathway_daa(</pre>
  abundance = metacyc_abundance %>% column_to_rownames("pathway"),
  metadata = metadata,
  group = "Environment",
  daa_method = "LinDA"
)
annotated_metacyc_daa_results_df <- pathway_annotation(</pre>
  pathway = "MetaCyc",
  daa_results_df = metacyc_daa_results_df,
  ko_to_kegg = FALSE
)
feature_with_p_0.05 <- metacyc_daa_results_df %>% filter(p_adjust < 0.05)</pre>
pathway_heatmap(
  abundance = metacyc_abundance %>%
```

## pathway\_pca

```
right_join(
    annotated_metacyc_daa_results_df %>%
    select(all_of(c("feature","description"))),
    by = c("pathway" = "feature")
) %>%
filter(pathway %in% feature_with_p_0.05$feature) %>%
select(-"pathway") %>%
column_to_rownames("description"),
metadata = metadata,
group = "Environment",
colors = custom_colors
```

```
## End(Not run)
```

pathway\_pca

Perform Principal Component Analysis (PCA) on functional pathway abundance data

## Description

)

This function performs PCA analysis on pathway abundance data and creates an informative visualization that includes a scatter plot of the first two principal components (PC1 vs PC2) with density plots for both PCs. The plot helps to visualize the clustering patterns and distribution of samples across different groups.

## Usage

```
pathway_pca(abundance, metadata, group, colors = NULL)
```

#### Arguments

abundance	A numeric matrix or data frame containing pathway abundance data. Rows represent pathways, columns represent samples. Column names must match the sample names in metadata. Values must be numeric and cannot contain missing values (NA).
metadata	A data frame containing sample information. Must include:
	• A column named "sample_name" matching the column names in abun- dance
	• A column for grouping samples (specified by the 'group' parameter)
group	A character string specifying the column name in metadata that contains group information for samples (e.g., "treatment", "condition", "group").
colors	Optional. A character vector of colors for different groups. Length must match the number of unique groups. If NULL, default colors will be used.

## Details

The function performs several validations on input data:

- Abundance matrix must have at least 2 pathways and 3 samples
- All values in abundance matrix must be numeric
- Sample names must match between abundance and metadata
- Group column must exist in metadata
- If custom colors are provided, they must be valid color names or codes

# Value

A ggplot object showing:

- Center: PCA scatter plot with confidence ellipses (95
- Top: Density plot for PC1
- Right: Density plot for PC2

```
# Create example abundance data
abundance_data <- matrix(rnorm(30), nrow = 3, ncol = 10)</pre>
colnames(abundance_data) <- paste0("Sample", 1:10)</pre>
rownames(abundance_data) <- c("PathwayA", "PathwayB", "PathwayC")</pre>
# Create example metadata
metadata <- data.frame(</pre>
  sample_name = paste0("Sample", 1:10),
  group = factor(rep(c("Control", "Treatment"), each = 5))
)
# Basic PCA plot with default colors
pca_plot <- pathway_pca(abundance_data, metadata, "group")</pre>
# PCA plot with custom colors
pca_plot <- pathway_pca(</pre>
  abundance_data,
  metadata,
  "group",
  colors = c("blue", "red") # One color per group
)
# Example with real data
data("metacyc_abundance") # Load example pathway abundance data
data("metadata")
                          # Load example metadata
# Generate PCA plot
# Prepare abundance data
abundance_data <- as.data.frame(metacyc_abundance)</pre>
rownames(abundance_data) <- abundance_data$pathway</pre>
```

## safe\_extract

```
abundance_data <- abundance_data[, -which(names(abundance_data) == "pathway")]
# Create PCA plot
pathway_pca(
    abundance_data,
    metadata,
    "Environment",
    colors = c("green", "purple")
)</pre>
```

safe\_extract Safely Extract Elements from a List

## Description

Safely extracts elements from a list, returning NA if the extraction fails

#### Usage

```
safe_extract(list, field, index = 1)
```

## Arguments

list	A list object from which to extract elements
field	The name of the field to extract from the list
index	The index position to extract from the field. Default is 1

## Value

The extracted element if successful, NA if extraction fails

```
# Create a sample list
my_list <- list(
    a = list(x = 1:3),
    b = list(y = 4:6)
)
# Extract existing element
safe_extract(my_list, "a", 1)
# Extract non-existing element (returns NA)</pre>
```

```
safe_extract(my_list, "c", 1)
```

visualize\_gsea

# Description

This function creates various visualizations for Gene Set Enrichment Analysis (GSEA) results.

# Usage

```
visualize_gsea(
 gsea_results,
 plot_type = "enrichment_plot",
 n_pathways = 20,
 sort_by = "p.adjust",
 colors = NULL,
 abundance = NULL,
 metadata = NULL,
 group = NULL,
 network_params = list(),
 heatmap_params = list()
```

## Arguments

gsea_results	A data frame containing GSEA results from the pathway_gsea function
plot_type	A character string specifying the visualization type: "enrichment_plot", "dot-plot", "barplot", "network", or "heatmap"
n_pathways	An integer specifying the number of pathways to display
sort_by	A character string specifying the sorting criterion: "NES", "pvalue", or "p.adjust"
colors	A vector of colors for the visualization
abundance	A data frame containing the original abundance data (required for heatmap visualization)
metadata	A data frame containing sample metadata (required for heatmap visualization)
group	A character string specifying the column name in metadata that contains the grouping variable (required for heatmap visualization)
network_params	A list of parameters for network visualization
heatmap_params	A list of parameters for heatmap visualization

## Value

A ggplot2 object or ComplexHeatmap object

#### visualize\_gsea

```
## Not run:
# Load example data
data(ko_abundance)
data(metadata)
# Prepare abundance data
abundance_data <- as.data.frame(ko_abundance)</pre>
rownames(abundance_data) <- abundance_data[, "#NAME"]</pre>
abundance_data <- abundance_data[, -1]</pre>
# Run GSEA analysis
gsea_results <- pathway_gsea(</pre>
  abundance = abundance_data,
  metadata = metadata,
  group = "Environment",
  pathway_type = "KEGG",
  method = "fgsea"
)
# Create enrichment plot
visualize_gsea(gsea_results, plot_type = "enrichment_plot", n_pathways = 10)
# Create dotplot
visualize_gsea(gsea_results, plot_type = "dotplot", n_pathways = 20)
# Create barplot
visualize_gsea(gsea_results, plot_type = "barplot", n_pathways = 15)
# Create network plot
visualize_gsea(gsea_results, plot_type = "network", n_pathways = 15)
# Create heatmap
visualize_gsea(
  gsea_results,
  plot_type = "heatmap",
  n_pathways = 15,
  abundance = abundance_data,
  metadata = metadata,
  group = "Environment"
)
## End(Not run)
```

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