

Package ‘MetaComp’

January 20, 2025

Version 1.1.2

Title EDGE Taxonomy Assignments Visualization

Description Implements routines for metagenome sample taxonomy assignments collection, aggregation, and visualization. Accepts the EDGE-formatted output from GOTTCHA/GOTTCHA2, BWA, Kraken, MetaPhlAn, DIAMOND, and Pangia. Produces SVG and PDF heatmap-like plots comparing taxa abundances across projects.

URL <https://github.com/seninp-bioinfo/MetaComp>

BugReports <https://github.com/seninp-bioinfo/MetaComp/issues>

Depends R (>= 3.1.0)

Imports reshape2, plyr, dplyr, data.table, ggplot2, Cairo

Suggests testthat

LazyData true

License GPL-2

RoxygenNote 6.0.1

NeedsCompilation no

Author Pavel Senin [aut, cre]

Maintainer Pavel Senin <senin@hawaii.edu>

Repository CRAN

Date/Publication 2018-06-18 20:06:45 UTC

Contents

load_edge_assignment	2
load_edge_assignments	3
merge_edge_assignments	4
merge_edge_counts	5
plot_edge_assignment	5
plot_merged_assignment	6
Index	8

`load_edge_assignment` *Efficiently loads an EDGE-produced taxonomic assignment from a file. An assumption has been made – since EDGE tables are generated in an automated fashion, they should be properly formatted – thus the code doesn't check for any inconsistencies except for the very file existence. Note however, the unassigned to taxa entries are removed. This implementation fully relies on the fread function from data.table package gaining performance over traditional R techniques.*

Description

Efficiently loads an EDGE-produced taxonomic assignment from a file. An assumption has been made – since EDGE tables are generated in an automated fashion, they should be properly formatted – thus the code doesn't check for any inconsistencies except for the very file existence. Note however, the unassigned to taxa entries are removed. This implementation fully relies on the fread function from data.table package gaining performance over traditional R techniques.

Usage

```
load_edge_assignment(filepath, type)
```

Arguments

`filepath` the path to EDGE-generated tab-delimited taxonomy assignment file.

`type` the assignment type. Following types are recognized: 'bwa', 'diamond', 'gottcha', 'gottcha2', 'kraken', 'metaphlan', and 'pangia'.

Value

a data frame containing four columns: TAXA, LEVEL, COUNT, and ABUNDANCE, representing taxonomically anchored sequences from the sample.

Examples

```
pa_fpath <- system.file("extdata", "HMP_even//allReads-pangia.list.txt", package="MetaComp")
pangia_assignment = load_edge_assignment(pa_fpath, type = "pangia")

table(pangia_assignment$LEVEL)

pangia_assignment[pangia_assignment$LEVEL == "phylum",]
```

load_edge_assignments *Efficiently loads a BWA (or other EDGE-like taxonomic assignment) tables from a list of files. Outputs a named list of assignments.*

Description

Efficiently loads a BWA (or other EDGE-like taxonomic assignment) tables from a list of files. Outputs a named list of assignments.

Usage

```
load_edge_assignments(filepath, type)
```

Arguments

filepath the path to tab delimited, two-column file whose first column is a project_id (which will be used to name this assignment) and the second column is the assignment filename.

type the type of assignments to be loaded. Following types are recognized: 'bwa', 'diamond', 'gottcha', 'gottcha2', 'kraken', 'metaphlan', and 'pangia'.

Value

a list of all read assignments.

Examples

```
hmp_even_fp <- system.file("extdata", "HMP_even", package="MetaComp")
hmp_stagger_fp <- system.file("extdata", "HMP_stagger", package="MetaComp")
data_files <- data.frame(V1 = c("HMP_even", "HMP_stagger"),
                        V2 = c(file.path(hmp_even_fp, "allReads-gottcha2-speDB-b.list.txt"),
                              file.path(hmp_stagger_fp, "allReads-gottcha2-speDB-b.list.txt")))
write.table(data_files, file.path(tempdir(), "assignments.txt"),
            row.names = FALSE, col.names = FALSE)
gottcha2_assignments = load_edge_assignments(file.path(tempdir(), "assignments.txt"),
                                           type = "gottcha2")

names(gottcha2_assignments)
table(gottcha2_assignments[[1]]$LEVEL)
```

merge_edge_assignments

Merges two or more EDGE-like taxonomical assignments. The input data frames are assumed to have the following columns: LEVEL, TAXA, and ABUNDANCE – these will be used in the merge procedure, all other columns will be ignored.

Description

Merges two or more EDGE-like taxonomical assignments. The input data frames are assumed to have the following columns: LEVEL, TAXA, and ABUNDANCE – these will be used in the merge procedure, all other columns will be ignored.

Usage

```
merge_edge_assignments(assignments)
```

Arguments

`assignments` A named list of assignments (the list element's name will be used as a resulting data frame column name).

Value

A merged table, which is a data frame whose rows are taxonomical ids and columns are the input assignments ids.

Examples

```
## Not run:
hmp_even_fp <- system.file("extdata", "HMP_even", package="MetaComp")
hmp_stagger_fp <- system.file("extdata", "HMP_stagger", package="MetaComp")
data_files <- data.frame(V1 = c("HMP_even", "HMP_stagger"),
                        V2 = c(file.path(hmp_even_fp, "allReads-gottcha2-speDB-b.list.txt"),
                              file.path(hmp_stagger_fp, "allReads-gottcha2-speDB-b.list.txt")))
write.table(data_files, file.path(tempdir(), "assignments.txt"),
            row.names = FALSE, col.names = FALSE)
gottcha2_assignments = merge_edge_assignments(
  load_edge_assignments(
    file.path(tempdir(), "assignments.txt"), type = "gottcha2"))

## End(Not run)
```

merge_edge_counts	<i>Merges two or more EDGE-like taxonomical assignments. The input data frames are assumed to have the following columns: LEVEL, TAXA, and COUNT – these will be used in the merge procedure, all other columns will be ignored.</i>
-------------------	--

Description

Merges two or more EDGE-like taxonomical assignments. The input data frames are assumed to have the following columns: LEVEL, TAXA, and COUNT – these will be used in the merge procedure, all other columns will be ignored.

Usage

```
merge_edge_counts(assignments)
```

Arguments

assignments	A named list of assignments (the list element's name will be used as a resulting data frame column name).
-------------	---

Value

A merged table, which is a data frame whose rows are taxonomical ids and columns are the input assignments ids.

plot_edge_assignment	<i>Generates a single column ggplot for a taxonomic assignment table and also outputs a PDF.</i>
----------------------	--

Description

This implementation is built upon ggplot geom_tile.

Usage

```
plot_edge_assignment(assignment, level, plot_title, column_title, filename)
```

Arguments

assignment	The EDGE-like assignment table.
level	The taxonomic level to plot (i.e., family, strain, etc...).
plot_title	The plot title, e.g., "Project XX, Run YY".
column_title	The column title.
filename	The PDF file name mask.

Value

the ggplot2 plot.

Examples

```
pa_fpath <- system.file("extdata", "HMP_even//allReads-pangia.list.txt", package="MetaComp")
pangia_assignment = load_edge_assignment(pa_fpath, type = "pangia")

plot_edge_assignment(pangia_assignment, "phylum", "Pangia", "HMP Even",
                    file.path(tempdir(), "assignment.pdf"))
```

plot_merged_assignment

Generates a single column ggplot for a taxonomic assignment table.

Description

This implementation...

Usage

```
plot_merged_assignment(assignment, taxonomy_level,
                      sorting_order = "abundance", row_limit = 60, min_row_abundance = 0,
                      plot_title, filename)
```

Arguments

assignment	The gottcha-like merged assignment table.
taxonomy_level	The level which need to be plotted.
sorting_order	the order in which rows shall be sorted, "abundance" is default, "alphabetical" is an alternative.
row_limit	the max amount of rows to plot (default is 60).
min_row_abundance	the minimal sum of abundances in a row required to plot. Rows whose sum is less than this value are dropped even if row_limit is specified. Ignored for "alphabetical" order. (default 0.0).
plot_title	The plot title.
filename	The output file mask, PDF and SVG files will be produced with Cairo device.

Examples

```
## Not run:
hmp_even_fp <- system.file("extdata", "HMP_even", package="MetaComp")
hmp_stagger_fp <- system.file("extdata", "HMP_stagger", package="MetaComp")
data_files <- data.frame(V1 = c("HMP_even", "HMP_stagger"),
                        V2 = c(file.path(hmp_even_fp, "allReads-gottcha2-speDB-b.list.txt"),
                              file.path(hmp_stagger_fp, "allReads-gottcha2-speDB-b.list.txt")))
write.table(data_files, file.path(tempdir(), "assignments.txt"),
            row.names = FALSE, col.names = FALSE)
gottcha2_assignments = merge_edge_assignments(
  load_edge_assignments(
    file.path(tempdir(), "assignments.txt"), type = "gottcha2"))
plot_merged_assignment(gottcha2_assignments, "family", 'alphabetical', 100, 0,
                      "HMP side-to-side", file.path(tempdir(), "assignment.pdf"))

## End(Not run)
```

Index

load_edge_assignment, [2](#)
load_edge_assignments, [3](#)

merge_edge_assignments, [4](#)
merge_edge_counts, [5](#)

plot_edge_assignment, [5](#)
plot_merged_assignment, [6](#)