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add_alignments

Add an alignment to a file

Description

Add an alignment to a file

Usage

```
add_alignments(filename)
```

Arguments

filename

Parameter filename

Value

Nothing, modifies the parameter file

Author(s)

Richel Bilderbeek

```
add_outgroup_to_phylogeny
```

Adds an outgroup to a phylogeny

Description

Adds an outgroup to a phylogeny

Usage

```
add_outgroup_to_phylogeny(phylogeny, stem_length, outgroup_name = "Outgroup")
```

Arguments

phylogeny a phylogeny

stem_length How long will the length of the outgroup be before the crown?

outgroup_name How is the outgroup named?

Value

A phylogeny, of type phylo

Author(s)

4 add_posteriors

add_pbd_output

Adds a pbd_sim result to a file

Description

Adds a pbd_sim result to a file

Usage

```
add_pbd_output(filename)
```

Arguments

filename

Parameter filename

Value

Nothing, modifies the parameter file

Author(s)

Richel Bilderbeek

add_posteriors

Add BEAST2 posteriors to a file

Description

Add BEAST2 posteriors to a file

Usage

```
add_posteriors(filename, skip_if_output_present = FALSE)
```

Arguments

```
filename Parameter filename skip_if_output_present
```

skip if output files are present, else remove these and start a new BEAST2 run

Value

Nothing, modifies the parameter file

Author(s)

add_species_trees_with_outgroup

Add a species tree with outgroup to a file

Description

Add a species tree with outgroup to a file

Usage

```
add_species_trees_with_outgroup(filename)
```

Arguments

filename

Parameter filename

Value

Nothing, modifies the parameter file

Author(s)

Richel Bilderbeek

branch_sorter

sorts branches for the NRBS analysis

Description

sorts branches for the NRBS analysis

Usage

```
branch_sorter(filename)
```

Arguments

tree1,

tree2, the trees that are to be sorted.

Value

a sorted tree1 and tree2

Author(s)

Femke Thon

6 collect_file_gammas

check_progress

Function that shows how much parameter files are present, and how far the pipeline has gotten

Description

Function that shows how much parameter files are present, and how far the pipeline has gotten

Usage

```
check_progress(folder = ".")
```

Arguments

folder

The folder containing the parameter files

Value

A data table showing the progress

Author(s)

Richel Bilderbeek

collect_file_gammas

Collects the gamm statistics of all phylogenies belonging to a parameter file in the melted/uncast/long form

Description

Collects the gamm statistics of all phylogenies belonging to a parameter file in the melted/uncast/long form

Usage

```
collect_file_gammas(filename, verbose = FALSE)
```

Arguments

filename

name of the parameter file

Value

A dataframe of gamma statistics of each phylogeny in time

collect_gamma_statistics

collect_gamma_statistics

Collects the gamma statistics of all phylogenies in the melted/uncast/long form

Description

Collects the gamma statistics of all phylogenies in the melted/uncast/long form

Usage

```
collect_gamma_statistics(phylogenies)
```

Arguments

phylogenies

the phylogenies, supplied as either a list or a multiPhylo object, where the phylogenies are of type 'phylo'

Value

A dataframe of gamma statistics of each phylogeny in time

 $collect_parameters$

Creates tidy data of all parameter files

Description

Creates tidy data of all parameter files

Usage

```
collect_parameters(filenames, verbose = FALSE)
```

Arguments

filenames name

names of the parameter files

verbose

give verbose output, should be TRUE or FALSE

Author(s)

Examples

```
filenames <- c(
   find_path("article_0_0_0_0_0.RDa"),
   find_path("article_0_1_4_0_2.RDa"),
   find_path("toy_example_1.RDa")
)
df <- collect_parameters(filenames)
testit::assert(nrow(df) == 3)</pre>
```

```
collect_posterior_filenames
```

Collect posterior filenames

Description

Collect posterior filenames

Usage

```
collect_posterior_filenames(parameter_filename)
```

Arguments

```
parameter_filename
```

Name of a parameter file

Value

generates species tree files from the posterior

Author(s)

Richel Bilderbeek and Jolien Gay

Examples

```
parameter_filename <- find_path("toy_example_1.RDa")
posterior_filenames <- collect_posterior_filenames(parameter_filename)
trees_filename <- find_path("toy_example_1_1_1_1.trees")
testit::assert(length(posterior_filenames) == 1)
testit::assert(posterior_filenames[1] == trees_filename)</pre>
```

 $collect_posterior_gammas$

Collect the gamma statistics of the BEAST2 posteriors

Description

Collect the gamma statistics of the BEAST2 posteriors

Usage

```
collect_posterior_gammas(filename)
```

Arguments

filename

name of the file containing the parameters and results

Value

a data frame

Author(s)

Richel Bilderbeek

```
collect_posterior_nltt_values
```

Collect the nLTT values of the BEAST2 posteriors

Description

Collect the nLTT values of the BEAST2 posteriors

Usage

```
collect_posterior_nltt_values(filename, dt = 0.001)
```

Arguments

filename name of the file containing the parameters and results

dt the resolution of the nLTT plot, must be in range <0,1>, default is 0.001

Value

a data frame

Author(s)

```
collect_species_tree_gammas
```

Collect the gamma statistics of the species trees with outgroup

Description

Collect the gamma statistics of the species trees with outgroup

Usage

```
collect_species_tree_gammas(filename)
```

Arguments

filename

name of the file containing the parameters and results

Value

a data frame

Author(s)

Richel Bilderbeek

```
collect_species_tree_nltts
```

Collect the nLTT values of the species trees with outgroup

Description

Collect the nLTT values of the species trees with outgroup

Usage

```
collect_species_tree_nltts(filename, dt = 0.001)
```

Arguments

filename

name of the file containing the parameters and results

dt

the resolution of the nLTT plot, must be in range <0,1>, default is 0.001

Value

a data frame

Author(s)

```
convert_alignment_to_beast_input_file
```

Convert an alignment and parameters to a BEAST XML input file

Description

Convert an alignment and parameters to a BEAST XML input file

Usage

```
convert_alignment_to_beast_input_file(alignment_dnabin, mcmc_chainlength,
  rng_seed = 42, beast_filename, temp_fasta_filename)
```

Arguments

```
alignment_dnabin
```

the DNA alignment of type DNAbin

mcmc_chainlength

the BEAST2 MCMC chain length

rng_seed random number generator seed

beast_filename the filename of the XML BEAST2 input file created

temp_fasta_filename

the name of a temporary file created

Value

Nothing, creates a file called 'beast_filename'

Author(s)

Richel Bilderbeek

```
convert_alignment_to_beast_posterior
```

Convert an alignment to a BEAST2 XML input file

Description

Convert an alignment to a BEAST2 XML input file

Usage

```
convert_alignment_to_beast_posterior(alignment_dnabin, mcmc_chainlength,
  base_filename, rng_seed = 42, beast_bin_path = "",
  beast_jar_path = find_beast_jar_path(), skip_if_output_present = FALSE)
```

Arguments

```
alignment_dnabin
```

The alignmnet

mcmc_chainlength

The length of the MCMC chain BEAST2 will generate

base_filename The base of the filename (the part without the extension)
rng_seed The random number generator seed used by BEAST2

beast_bin_path Where the binary 'beast' can be found beast_jar_path Where the jar 'beast.jar' can be found

skip_if_output_present

skip if output files are present, else remove these and start a new BEAST2 run

Value

A BEAST2 posterior

Author(s)

Richel Bilderbeek

```
convert_alignment_to_fasta
```

Convert an alignment (of type DNAbin) to a FASTA file

Description

Convert an alignment (of type DNAbin) to a FASTA file

Usage

```
convert_alignment_to_fasta(alignment_dnabin, filename)
```

Arguments

alignment_dnabin

DNA alignment of type DNAbin

filename FASTA filename the alignment will be saved to

Value

Nothing, it will create a FASTA file

Author(s)

```
convert_phylogeny_to_alignment
```

Converts a phylogeny to a random DNA alignment

Description

Converts a phylogeny to a random DNA alignment

Usage

```
convert_phylogeny_to_alignment(phylogeny, sequence_length, mutation_rate = 1)
```

Arguments

```
phylogeny a phylogeny sequence_length
```

the number of nucleotides to alignment will have per taxon

mutation_rate the rate per nucleotide to change, per million years

Value

an alignment

Author(s)

Richel Bilderbeek

extract_erg

Extract the ERG ('Extinction Rate of a Good species') parameter value from a file

Description

Extract the ERG ('Extinction Rate of a Good species') parameter value from a file

Usage

```
extract_erg(file)
```

Arguments

file

A loaded parameter file

Value

the value of the ERG parameter

14 extract_eri

Author(s)

Richel Bilderbeek

Examples

```
file <- read_file(find_path("toy_example_1.RDa"))
erg <- extract_erg(file)
testit::assert(erg >= 0.0)
```

extract_eri

Extract the ERI ('Extinction Rate of an Incipient species') parameter value from a file

Description

Extract the ERI ('Extinction Rate of an Incipient species') parameter value from a file

Usage

```
extract_eri(file)
```

Arguments

file

A loaded parameter file

Value

the value of the ERI parameter

Author(s)

Richel Bilderbeek

Examples

```
file <- read_file(find_path("toy_example_1.RDa"))
eri <- extract_eri(file)
testit::assert(eri >= 0.0)
```

extract_scr 15

extract_scr $Extract\ the\ SCR\ (`Speciation\ Completion\ Rate')\ parameter\ value\ from a\ file$

Description

Extract the SCR ('Speciation Completion Rate') parameter value from a file

Usage

```
extract_scr(file)
```

Arguments

file

A loaded parameter file

Value

the value of the SCR parameter

Author(s)

Richel Bilderbeek

Examples

```
file <- read_file(find_path("toy_example_1.RDa"))
scr <- extract_scr(file)
testit::assert(scr >= 0.0)
```

extract_sirg

Extract the SIRG ('Speciation Initial Rate of a Good species') parameter value from a file

Description

Extract the SIRG ('Speciation Initial Rate of a Good species') parameter value from a file

Usage

```
extract_sirg(file)
```

Arguments

file

A loaded parameter file

16 extract_siri

Value

the value of the SIRG parameter

Author(s)

Richel Bilderbeek

Examples

```
file <- read_file(find_path("toy_example_1.RDa"))
sirg <- extract_sirg(file)
testit::assert(sirg >= 0.0)
```

extract_siri

Extract the SIRI ('Speciation Initial Rate of an Incipient species') parameter value from a file

Description

Extract the SIRI ('Speciation Initial Rate of an Incipient species') parameter value from a file

Usage

```
extract_siri(file)
```

Arguments

file

A loaded parameter file

Value

the value of the SIRI parameter

Author(s)

Richel Bilderbeek

Examples

```
file <- read_file(find_path("toy_example_1.RDa"))
siri <- extract_siri(file)
testit::assert(siri >= 0.0)
```

find_beast_bin_path 17

find_beast_bin_path

Find the path of the BEAST2 binary file

Description

Find the path of the BEAST2 binary file

Usage

```
find_beast_bin_path()
```

Value

the path of the BEAST2 binary file

find_beast_jar_path

Find the path of the BEAST2 jar file

Description

Find the path of the BEAST2 jar file

Usage

```
find_beast_jar_path()
```

Value

the path of the BEAST2 jar file

find_beast_posterior_test_filename

Find the path of known-to-be-valid BEAST2 posterior file

Description

Find the path of known-to-be-valid BEAST2 posterior file

Usage

```
find_beast_posterior_test_filename()
```

Value

the path of a known-to-be-valid BEAST2 posterior file

 $find_path$

Finds the full pathof a file

Description

Finds the full pathof a file

Usage

```
find_path(filename)
```

Arguments

filename

the name of a file

Value

the full path of the filename if an existing file could be found, stops otherwise

Author(s)

Richel Bilderbeek

```
get_phylogeny_crown_age
```

Obtain the crown age of a phylony

Description

Obtain the crown age of a phylony

Usage

```
get_phylogeny_crown_age(phylogeny)
```

Arguments

phylogeny

The phylogeny to obtain the crown age of

Value

the age of the phylogeny

Author(s)

is_alignment 19

is_alignment

Determines if the input is an alignment of type DNAbin

Description

Determines if the input is an alignment of type DNAbin

Usage

```
is_alignment(input)
```

Arguments

input

The input to be testes

Value

TRUE or FALSE

Author(s)

Richel Bilderbeek

is_beast_posterior

Determines if the input is a BEAST2 posterior

Description

Determines if the input is a BEAST2 posterior

Usage

```
is_beast_posterior(x)
```

Arguments

Χ

the input

Value

TRUE or FALSE

Author(s)

20 is_phylogeny

is_pbd_sim_output

Does not use pbd_sim()\$stree, but generates these like PBD does

Description

Does not use pbd_sim()\$stree, but generates these like PBD does

Usage

```
is_pbd_sim_output(pbd_sim_output, verbose = FALSE)
```

Arguments

verbose

 $\verb|pbd_sim_output| the argument tested to output of PBD::pbd_sim|$

will the function show why the input is determined not to be output of PBD::pbd_sim

Value

TRUE or FALSE

Author(s)

Richel Bilderbeek

is_phylogeny

Checks if the input is a phylogeny

Description

Checks if the input is a phylogeny

Usage

```
is_phylogeny(x)
```

Arguments

Х

input to be checked

Value

TRUE or FALSE

Author(s)

is_valid_file 21

is_valid_file

Checks if a file is a valid parameter file

Description

Checks if a file is a valid parameter file

Usage

```
is_valid_file(filename, verbose = FALSE)
```

Arguments

filename the name of the file to be checked

verbose give verbose output, should be TRUE or FALSE

Value

TRUE or FALSE

Author(s)

Richel Bilderbeek

Examples

```
testit::assert(is_valid_file(find_path("toy_example_1.RDa")))
testit::assert(!is_valid_file(find_path("toy_example_1_1_1_1.trees")))
```

is_whole_number

Checks if x is a single, whole-number variable

Description

Checks if x is a single, whole-number variable

Usage

```
is_whole_number(x, tolerance = .Machine$double.eps^0.5)
```

Arguments

x the number to check

tolerance the maximum error a number may deviate from a whole number, before it is

labeled as a floating point value

Value

TRUE or FALSE

Author(s)

Richel Bilderbeek

```
load_parameters_from_file
```

Load parameters from file

Description

Load parameters from file

Usage

```
load_parameters_from_file(filename)
```

Arguments

filename

name of the parameter file

Value

a data frame with all parameters

Author(s)

Richel Bilderbeek

```
normalized_rooted_branch_score
```

calculates the normalized rooted branch score

Description

calculates the normalized rooted branch score

Usage

```
normalized_rooted_branch_score(filenames)
```

Arguments

filenames th

the filenames

plot_alignments 23

Value

the normalized rooted branch score

Author(s)

Femke Thon

 ${\tt plot_alignments}$

Plot alignments

Description

Plot alignments

Usage

```
plot_alignments(filename)
```

Arguments

filename

name of the parameter file

Author(s)

Richel Bilderbeek

```
plot_extant_incipient_tree
```

Plot the extant inicipient species tree

Description

Plot the extant inicipient species tree Plot the extant inicipient species tree

Usage

```
plot_extant_incipient_tree(filename)
plot_extant_incipient_tree(filename)
```

Arguments

filename

name of the parameter file

Value

Nothing. It does produce a plot

Author(s)

Richel Bilderbeek

Description

Plot multiple BEAST2 posteriors

Usage

```
plot_posterior_nltts(filename, dt = 0.001)
```

Arguments

filename name of the file containing the parameters and results

dt the resolution of the nLTT plot, must be in range <0,1>, default is 0.001

Value

Nothing, produces a plot

Author(s)

Richel Bilderbeek

```
plot_posterior_samples
```

Plot a random posterior phylogeny

Description

Plot a random posterior phylogeny

Usage

```
plot_posterior_samples(filename)
```

Arguments

filename a filename

Value

Nothing, but it does generate some plots

Author(s)

Richel Bilderbeek

```
plot_posterior_sample_nltts
```

Plot the nLTTs of the posterior

Description

Plot the nLTTs of the posterior

Usage

```
plot_posterior_sample_nltts(filename, dt = 0.001)
```

Arguments

filename a filename

dt the resolution of the nLTT plot, must be in range <0,1>, default is 0.001

Value

Nothing, but it does generate some plots

Author(s)

Richel Bilderbeek

```
{\tt plot\_species\_tree\_with\_outgroup} \\ {\tt \it Plot} \\
```

Description

Plot

Usage

```
plot_species_tree_with_outgroup(filename)
```

26 read_file

Arguments

filename a file name

Value

Nothing, but it does generate plots

Author(s)

Richel Bilderbeek

```
plot\_species\_tree\_with\_outgroup\_nltt\\ Plot\ species\_tree\_with\_outgroup\_nltt\\
```

Description

Plot species_tree_with_outgroup_nltt

Usage

```
plot_species_tree_with_outgroup_nltt(filename, dt = 0.001)
```

Arguments

filename parameter filename

dt the resolution of the nLTT plot, must be in range <0,1>, default is 0.001

Author(s)

Richel Bilderbeek

read_file

Read a parameter (.RDa) file

Description

Read a parameter (.RDa) file

Usage

```
read_file(filename)
```

Arguments

filename name of the parameter (.RDa) file

Value

the file as R data

Author(s)

Richel Bilderbeek

Examples

```
file <- read_file(find_path("toy_example_1.RDa"))
print(names(file))</pre>
```

```
sample_species_trees_from_pbd_sim_output
```

Samples species trees from PBD::pbd_sim its output It does not use pbd_sim()\$stree, but generates these like PBD does.

Description

Samples species trees from PBD::pbd_sim its output It does not use pbd_sim()\$stree, but generates these like PBD does.

Usage

```
sample_species_trees_from_pbd_sim_output(n, pbd_sim_output)
```

Arguments

```
n The number of species trees to sample pbd_sim_output The output generated by PBD::pbd_sim
```

Value

A list of length n, containing the n sampled species trees

Author(s)

Richel Bilderbeek, inspired by Rampal Etienne's PBD::pbd_sim function

```
save_parameters_to_file
```

Creates a valid parameter file

Description

Creates a valid parameter file

Usage

```
save_parameters_to_file(rng_seed, sirg, siri, scr, erg, eri, age,
   n_species_trees_samples, mutation_rate, n_alignments, sequence_length,
   mcmc_chainlength, n_beast_runs, filename)
```

Arguments

rng_seed	the random number generator seed	
sirg	the species initiation rate of the good species	
siri	the speciation-initiation rate of the incipient species	
scr	the rate at which incipient species become good	
erg	the rate at which good species get extinct	
eri	the rate at which incipient species get extinct	
age	crown age of the phylogeny	
n_species_trees_samples		
	the number of species trees that will be sampled from an incipient species tree	
mutation_rate	the probability per nucleotide to mutate at a DNA replication	
n_alignments	the number of alignments simulated per species tree	
sequence_length		
	the simulated DNA sequence length in nucleotides	
mcmc_chainlength		
	the length of the MCMC chain that BEAST2 will run	
n_beast_runs	the number of BEAST2 runs per DNA alignments	
filename	the name of the parameter file that will be created by this function	

Value

Nothing, it will create a file with filename 'filename'

Author(s)

show_parameter_files 29

Description

Creates a nice knitr::table to show one or more parameter files

Usage

```
show_parameter_files(filenames, verbose = FALSE)
```

Arguments

filenames names of the parameter files

verbose give verbose output, should be TRUE or FALSE

Author(s)

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