Package 'datelife'

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Title Scientific Data on Time of Lineage Divergence for Your Taxa

Version 0.6.8

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Description Methods and workflows to get chronograms (i.e., phylogenetic trees with branch lengths proportional to time), using open, peer-reviewed, state-of-theart scientific data on time of lineage divergence. This package constitutes the main underlying code of the DateLife web service at <https://www.datelife.org>. To obtain a single summary chronogram from a group of relevant chronograms, we implement the Super Distance Matrix (SDM) method described in Criscuolo et al. (2006) <doi:10.1080/10635150600969872>. To find the grove of chronograms with a sufficiently overlapping set of taxa for summarizing, we implement theorem 1.1. from Ané et al. (2009) <doi:10.1007/s00026-009-0017-x>. A given phylogenetic tree can be dated using time of lineage divergence data as secondary calibrations (with caution, see Schenk (2016) <doi:10.1371/journal.pone.0148228>). To obtain and apply secondary calibrations, the package implements the congruification method described in Eastman et al. (2013) <doi:10.1111/2041-210X.12051>. Tree dating can be performed with different methods including BLADJ (Webb et al. (2008) <doi:10.1093/bioinformatics/btn358>), PATHd8 (Britton et al. (2007) <doi:10.1080/10635150701613783>), mrBayes (Huelsenbeck and Ronquist (2001) <doi:10.1093/bioinformatics/17.8.754>), and treePL (Smith and O'Meara (2012) <doi:10.1093/bioinformatics/bts492>). **Depends** R (>= 3.6.0) biocViews Software Imports ape, abind, bold, phangorn, phytools, ips, cluster, compare,

geiger, stats, stringr, rotl, paleotree, knitcitations,

phylobase, taxize, treebase, utils, httr, plyr, phylocomr, BiocManager, data.table, curl

Suggests testthat, knitr, rmarkdown, usethis, devtools, covr, msa, Biostrings

LazyDataCompression xz

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.get_ott_lineage

Get the lineage of a set of taxa. .get_ott_lineage uses rotl::taxonomy_taxon_info() with include_lineage = TRUE.

Description

Get the lineage of a set of taxa. .get_ott_lineage uses rotl::taxonomy_taxon_info() with include_lineage = TRUE.

Usage

.get_ott_lineage(input_ott_match)

Arguments

input_ott_match An Output of check_ott_input function.

Value

A taxonomy_taxon_info object

birds_and_cats	A multiPhylo object with trees resulting from a datelife search of some
	birds and cats species

Description

A multiPhylo object with trees resulting from a datelife search of some birds and cats species

Usage

birds_and_cats

Format

A multiPhylo object

Details

Generated with: taxa <- c("Rhea americana", "Pterocnemia pennata", "Struthio camelus", "Gallus", "Felis") birds_and_cats <- datelife_search(input = taxa, summary_format = "phylo_all", get_spp_from_taxon = TRUE) usethis::use_data(birds_and_cats)

build_grove_list Build grove list

Description

This function implements theorem 1.1 of Ané et al. (2009) doi:10.1007/s000260090017x to find a grove for a given group of chronograms.

Usage

build_grove_list(datelife_result, n = 2)

build_grove_matrix

Arguments

datelife_resul	t
	A datelifeResult object.
n	The degree of taxon name overlap among input chronograms. Defaults to n = 2, i.e., at least two overlapping taxon names.

Value

A list of vectors; each list element is a grove.

build_grove_matrix Find the grove for a group of chronograms and build a matrix.

Description

This function implements theorem 1.1 of Ané et al. (2009) doi:10.1007/s000260090017x to find a grove for a given group of chronograms.

Usage

build_grove_matrix(datelife_result, n = 2)

Arguments

 datelife_result

 A datelifeResult object.

 n
 The degree of taxon name overlap among input chronograms. Defaults to n = 2, i.e., at least two overlapping taxon names.

Value

A matrix. Each cell shows whether n-overlap exists between a pair of inputs.

References

Ané, C., Eulenstein, O., Piaggio-Talice, R., & Sanderson, M. J. (2009). "Groves of phylogenetic trees". Annals of Combinatorics, 13(2), 139-167, doi:10.1007/s000260090017x.

check_conflicting_calibrations

Check for conflicting calibrations.

Description

check_conflicting_calibrations checks if calibrations are younger or older relative to descendants and ancestors, respectively.

Usage

```
check_conflicting_calibrations(phy, calibration_distribution)
```

Arguments

phy A phylo object. calibration_distribution A list of node age distributions, named with phy's node numbers.

Details

It removes conflicting calibrations if needed, but BLADJ works as long as it has an age for the root.

check_ott_input Check input for usage in other datelife functions

Description

check_ott_input is currently used in functions get_ott_clade(), get_ott_children(), and get_otol_synthetic_tree().

Usage

```
check_ott_input(input = NULL, ott_ids = NULL, ...)
```

Arguments

input	Optional. A character vector of names or a datelifeQuery object.
ott_ids	If not NULL, it takes this argument and ignores input. A numeric vector of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().
	Arguments passed on to make_datelife_query
	<pre>use_tnrs Whether to use Open Tree of Life's Taxonomic Name Resolution Service (TNRS) to process input taxon names. Default to TRUE, it corrects misspellings and taxonomic name variations with tnrs_match(), a wrap- per of rot1::tnrs_match_names().</pre>

get_spp_from_taxon Whether to search ages for all species belonging to a given taxon or not. Default to FALSE. If TRUE, it must have same length as input. If input is a newick string with some clades it will be converted to a phylo object, and the order of get_spp_from_taxon will match phy\$tip.label. reference_taxonomy A character vector specifying the reference taxonomy to use for TNRS. Options are "ott", "ncbi", "gbif" or "irmng". The function defaults to "ott".

Details

By default, it uses the ott_id argument if it is not NULL.

Value

A named numeric vector of valid Open Tree Taxonomy (OTT) ids.

choose_cluster	Choose	an	ultrametric	phylo	object	from
	tering meth	nod, or tl	<pre>cmatrix() obtail he next best tree. I m to be ultrametri</pre>	f there are n	1	

Description

Choose an ultrametric phylo object from cluster_patristicmatrix() obtained with a particular clustering method, or the next best tree. If there are no ultrametric trees, it does not force them to be ultrametric.

Usage

```
choose_cluster(phycluster, clustering_method = "nj")
```

Arguments

Value

A phylo object or NA.

classification_paths_from_taxonomy Gets classification paths for a vector of taxa

Description

This uses the taxize package's wrapper of the Global Names Resolver to get taxonomic paths for the vector of taxa you pass in. Sources is a vector of source labels in order (though it works best if everything uses the same taxonomy, so we recommend doing just one source). You can see options by doing taxize::gnr_datasources(). Our default is Catalogue of Life

Usage

```
classification_paths_from_taxonomy(taxa, sources = "Catalogue of Life")
```

Arguments

taxa	Vector of taxon names
sources	Vector of names of preferred sources; see taxize::gnr_datasources(). Currently supports 100 taxonomic resources, see details.

Details

Taxonomies supported by taxize::gnr_datasources()

- 1. Catalogue of Life
- 2. Wikispecies
- 3. ITIS
- 4. NCBI
- 5. Index Fungorum
- 6. GRIN Taxonomy for Plants
- 7. Union 4
- 8. The Interim Register of Marine and Nonmarine Genera
- 9. World Register of Marine Species
- 10. Freebase
- 11. GBIF Backbone Taxonomy
- 12. EOL
- 13. Passiflora vernacular names
- 14. Inventory of Fish Species in the Wami River Basin
- 15. Pheasant Diversity and Conservation in the Mt. Gaoligonshan Region
- 16. Finding Species
- 17. Birds of Lindi Forests Plantation

- 18. Nemertea
- 19. Kihansi Gorge Amphibian Species Checklist
- 20. Mushroom Observer
- 21. TaxonConcept
- 22. Amphibia and Reptilia of Yunnan
- 23. Common names of Chilean Plants
- 24. Invasive Species of Belgium
- 25. ZooKeys
- 26. COA Wildlife Conservation List
- 27. AskNature
- 28. China: Yunnan, Southern Gaoligongshan, Rapid Biological Inventories Report No. 04
- 29. Native Orchids from Gaoligongshan Mountains, China
- 30. Illinois Wildflowers
- 31. Coleorrhyncha Species File
- 32. /home/dimus/files/dwca/zoological names.zip
- 33. Peces de la zona hidrogeográfica de la Amazonia, Colombia (Spreadsheet)
- 34. Eastern Mediterranean Syllidae
- 35. Gaoligong Shan Medicinal Plants Checklist
- 36. birds_of_tanzania
- 37. AmphibiaWeb
- 38. tanzania_plant_sepecimens
- 39. Papahanaumokuakea Marine National Monument
- 40. Taiwanese IUCN species list
- 41. BioPedia
- 42. AnAge
- 43. Embioptera Species File
- 44. Global Invasive Species Database
- Sendoya S., Fernández F. AAT de hormigas (Hymenoptera: Formicidae) del Neotrópico 1.0 2004 (Spreadsheet)
- 46. Flora of Gaoligong Mountains
- 47. ARKive
- 48. True Fruit Flies (Diptera, Tephritidae) of the Afrotropical Region
- 49. 3i Typhlocybinae Database
- 50. CATE Sphingidae
- 51. ZooBank
- 52. Diatoms
- 53. AntWeb

- 54. Endemic species in Taiwan
- 55. Dermaptera Species File
- 56. Mantodea Species File
- 57. Birds of the World: Recommended English Names
- 58. New Zealand Animalia
- 59. Blattodea Species File
- 60. Plecoptera Species File
- 61. /home/dimus/files/dwca/clemens.zip
- 62. Coreoidea Species File
- 63. Freshwater Animal Diversity Assessment Normalized export
- 64. Catalogue of Vascular Plant Species of Central and Northeastern Brazil
- 65. Wikipedia in EOL
- 66. Database of Vascular Plants of Canada (VASCAN)
- 67. Phasmida Species File
- 68. OBIS
- 69. USDA NRCS PLANTS Database
- 70. Catalog of Fishes
- 71. Aphid Species File
- 72. The National Checklist of Taiwan
- 73. Psocodea Species File
- 74. FishBase
- 75. 3i Typhlocybinae Database
- 76. Belgian Species List
- 77. EUNIS
- 78. CU*STAR
- 79. Orthoptera Species File
- 80. Bishop Museum
- 81. IUCN Red List of Threatened Species
- 82. BioLib.cz
- 83. Tropicos Missouri Botanical Garden
- 84. nlbif
- 85. The International Plant Names Index
- 86. Index to Organism Names
- 87. uBio NameBank
- 88. Arctos
- 89. Checklist of Beetles (Coleoptera) of Canada and Alaska. Second Edition.
- 90. The Paleobiology Database

- 91. The Reptile Database
- 92. The Mammal Species of The World
- 93. BirdLife International
- 94. Checklist da Flora de Portugal (Continental, Açores e Madeira)
- 95. FishBase Cache
- 96. Silva
- 97. Open Tree of Life Reference Taxonomy
- 98. iNaturalist
- 99. The Interim Register of Marine and Nonmarine Genera
- 100. Gymno

Value

A list with resolved taxa (a tibble, from taxize::gnr_resolve) and a vector of taxa not resolved

clean_ott_chronogram	Clean up some issues with Open Tree of Life chronograms For now
	<i>it 1) checks unmapped taxa and maps them with tnrs_match.phylo, 2)</i>
	roots the chronogram if unrooted

Description

Clean up some issues with Open Tree of Life chronograms For now it 1) checks unmapped taxa and maps them with tnrs_match.phylo, 2) roots the chronogram if unrooted

Usage

```
clean_ott_chronogram(phy)
```

Arguments

phy A phylo object.

Details

There is no limit to the number of names that can be queried and matched.

The output will preserve all elements from original input phylo object and will add

phy\$mapped A character vector indicating the state of mapping of phy\$tip.labels:

original Tnrs matching was not attempted. Original labeling is preserved.

ott Matching was manually made by a curator in Open Tree of Life.

tnrs Tnrs matching was attempted and successful with no approximate matching. Original label is replaced by the matched name.

- **approximated** Tnrs matching was attempted and successful but with approximate matching. Original labeling is preserved.
- unmatched Tnrs matching was attempted and unsuccessful. Original labeling is preserved.
- phy\$original.tip.label A character vector preserving all original labels.
- **phy\$ott_ids** A numeric vector with ott id numbers of matched tips. Unmatched and original tips will be NaN.

if tips are duplicated, thrs will only be run once (avoiding increases in function running time) but the result will be applied to all duplicated tip labels

Value

An object of class data frame or phylo, with the added class match_names.

NULL

NULL

clean_taxon_info_children

Identify, extract and clean taxonomic children names from a taxonomy_taxon_info() output.

Description

clean_taxon_info_children eliminates all taxa that will give problems when trying to retrieve an induced subtree from Open Tree of Life.

Usage

```
clean_taxon_info_children(
  taxon_info,
  invalid = c("barren", "extinct", "uncultured", "major_rank_conflict", "incertae_sedis",
        "unplaced", "conflict", "environmental", "not_otu", "hidden", "hybrid")
)
```

Arguments

taxon_info	An output of rotl::taxonomy_taxon_info().
invalid	A character vector of "flags", i.e., characteristics that are used by Open Tree of
	Life Taxonomy to detect invalid taxon names.

Value

A list with valid children unique OTT names, OTT ids and taxonomic ranks.

clean_tnrs Eliminates unmatched (NAs) and invalid taxa from a
rotl::tnrs_match_names() or tnrs_match() output Useful to get ott ids to retrieve an induced synthetic Open Tree of
Life. Needed because using include_suppressed = FALSE in
rotl::tnrs_match_names() does not drop all invalid taxa.

Description

Eliminates unmatched (NAs) and invalid taxa from a rotl:::tnrs_match_names() or tnrs_match() output Useful to get ott ids to retrieve an induced synthetic Open Tree of Life. Needed because using include_suppressed = FALSE in rotl::tnrs_match_names() does not drop all invalid taxa.

Usage

```
clean_tnrs(
   tnrs,
   invalid = c("barren", "extinct", "uncultured", "major_rank_conflict", "incertae",
      "unplaced", "conflict", "environmental", "not_otu"),
   remove_nonmatches = FALSE
)
```

Arguments

tnrs	A data frame, usually an output from datelife::tnrs_match or rotl::tnrs_match_names functions, but see details.
invalid	A character string with flags to be removed from final object.
remove_nonmatches	
	Boolean, whether to remove unsuccessfully matched names or not.

Details

Input can be any data frame or named list that relates taxa stored in an element named "unique" to a validity category stored in "flags".

Value

A data frame or named list (depending on the input) with valid taxa only.

```
cluster_patristicmatrix
```

Cluster a patristic matrix into a tree with various methods.

Description

Cluster a patristic matrix into a tree with various methods.

Usage

```
cluster_patristicmatrix(patristic_matrix, variance_matrix = NULL)
```

Arguments

patristic_matrix

A patristic matrix

variance_matrix

A variance matrix from a datelifeResult object, usually an output from datelife_result_variance_u Only used if clustering_method = "mvr".

Details

If clustering method fails, NA is returned.

Value

A list of trees obtained with clustering methods detailed in patristic_matrix_to_phylo().

congruify_and_check Congruify and Check.

Description

Congruify and Check.

Usage

```
congruify_and_check(
  reference,
  target,
  taxonomy = NULL,
  tol = 0.01,
  option = 2,
  scale = "pathd8",
  attempt_fix = TRUE
)
```

Arguments

reference	an ultrametric tree used to time-scale the target
target	a phylogram that is sought to be ultrametricized based on the reference phy- logeny
taxonomy	a linkage table between tips of the phylogeny and clades represented in the tree; rownames of 'taxonomy' should be tips found in the phylogeny
tol	branching time in reference above which secondary constraints will be applied to target
option	an integer (1 or 2; see details).
scale	NA, "PATHd8" or "treePL" (if PATHd8 or "treePL" are available in the R PATH) $% \label{eq:response}$
attempt_fix	Default to TRUE. If congruification results in NA branch lengths, it will attempt to fix them.

congruify_and_mrca_multiPhylo

Congruify nodes of a tree topology to nodes from a source chronogram, and find the mrca nodes

Description

congruify_and_mrca_multiPhylo congruifies a target tree against all source chronograms in a multiPhylo object, and gets nodes of target tree that correspond to the most recent common ancestor (mrca) of taxon pairs in the congruified calibrations. It calls congruify_and_mrca_phylo(), and phytools::findMRCA() to get mrca nodes.

Usage

congruify_and_mrca_multiPhylo(phy, source_chronograms)

Arguments

phy A phylo object.

source_chronograms

A multiPhylo object, output of datelife_search().

Value

a data.frame of node ages from source_chronograms and corresponding mrca nodes in target tree phy. attributes(return)\$phy stores the congruified and mrca matched phylogeny.

```
congruify_and_mrca_phylo
```

Congruify nodes of a tree topology to nodes from a source chronogram, and find the mrca nodes

Description

congruify_and_mrca congruifies a target tree against a single source chronogram, and gets nodes of target tree that correspond to the most recent common ancestor (mrca) of taxon pairs from the congruified calibrations. It uses phytools::findMRCA() to get mrca nodes.

Usage

```
congruify_and_mrca_phylo(phy, source_chronogram, reference)
```

Arguments

phy	A phylo object.	
source_chronogram		
	A phylo object, output of datelife_search().	
reference	A character string indicating the study reference that the source_chronogram comes from.	

Value

a data.frame of node ages from source_chronograms and corresponding mrca nodes in target tree phy.

contributor_cache	Information on contributors, authors, study ids and clades from stud-
	ies with chronograms in Open Tree of Life (Open Tree)

Description

Information on contributors, authors, study ids and clades from studies with chronograms in Open Tree of Life (Open Tree)

Usage

contributor_cache

Format

A list of five data sets.

author.pretty A character vector with the author names from studies with chronograms that are in Open Tree.

author.results A dataframe with three variables: authors, study ids and clades.

curator.pretty A character vector with the names of curators of chronograms that are in Open Tree.

curator.results A data.frame with three variables: curators, study ids and clades.

missed_doi A character vector with study ids whose "doi" could not be retrieved.

Details

Generated with make_contributor_cache().

Source

http://opentreeoflife.org

datelife_authors_tabulate

Return the relevant authors for a set of studies.

Description

Return the relevant authors for a set of studies.

Usage

```
datelife_authors_tabulate(results.index, cache = "opentree_chronograms")
```

Arguments

results.index	A vector from datelife_result_study_index() with the indices of the rele-
	vant studies.
cache	The cached chronogram database.

Value

A vector with counts of each author, with names equal to author names.

```
datelife_result_median
```

Get a median summary chronogram from a datelifeResult object.

Description

Get a median summary chronogram from a datelifeResult object.

Usage

```
datelife_result_median(datelife_result, ...)
```

Arguments

datelife_result	
	A datelifeResult object, usually an output of get_datelife_result().
	Arguments passed on to summary_matrix_to_phylo
	<pre>summ_matrix Any summary patristic distance matrix, such as the ones obtained with datelife_result_sdm_matrix() or datelife_result_median_matrix().</pre>
	total_distance Whether the input summ_matrix stores total age distance (from tip to tip) or distance from node to tip. Default to TRUE, divides the matrix in half, if FALSE it will take it as is.
	use A character vector indicating what type of age to use for summary tree. One of the following:
	"mean" It will use the mean() of the node ages in summ_matrix.
	"median" It uses the stats::median() age of node ages in summ_matrix.
	"min" It will use the min() age from node ages in summ_matrix.
	"max" Choose this if you wanna be conservative; it will use the max() age from node ages in summ_matrix.
	"midpoint" It will use the mean of minimum age and maximum age.
	target_tree A phylo object. Use this in case you want a specific backbone for the output tree.
	datelife_query A datelifeQuery object, usually an output of make_datelife_query().

Value

A phylo object.

datelife_result_median_matrix

Compute a median matrix of a datelifeResult object.

Description

Compute a median matrix of a datelifeResult object.

Usage

```
datelife_result_median_matrix(datelife_result)
```

Arguments

```
datelife_result
A datelifeResult object, usually an output of get_datelife_result().
```

Value

A patristic distance summary matrix from a datelifeResult object.

datelife_result_MRCA	Get a numeric vector of MRCAs from a datelifeResult object. Used
	<pre>in summarize_datelife_result().</pre>

Description

Get a numeric vector of MRCAs from a datelifeResult object. Used in summarize_datelife_result().

Usage

```
datelife_result_MRCA(datelife_result, na_rm = TRUE)
```

Arguments

datelife_resu	ılt
	A datelifeResult object, usually an output of get_datelife_result().
na_rm	If TRUE, it drops rows containing NAs from the datelifeResult patristic matrix;
	if FALSE, it returns NA where there are missing entries.

Value

A named numeric vector of MRCA ages for each element given in datelife_result.

```
datelife_result_sdm_matrix
```

Go from a datelifeResult object to a Super Distance Matrix (SDM) using weighting = "flat"

Description

Go from a datelifeResult object to a Super Distance Matrix (SDM) using weighting = "flat"

Usage

```
datelife_result_sdm_matrix(datelife_result)
```

Arguments

datelife_result

A datelifeResult object, usually an output of get_datelife_result().

Value

A numeric matrix.

```
datelife_result_sdm_phylo
```

Reconstruct a supertree from a datelifeResult *object using the Super Distance Matrix (SDM) method.*

Description

Reconstruct a supertree from a datelifeResult object using the Super Distance Matrix (SDM) method.

Usage

```
datelife_result_sdm_phylo(datelife_result, weighting = "flat", ...)
```

Arguments

datelife_result	t
	A datelifeResult object, usually an output of get_datelife_result().
weighting	A character vector indicating how much weight to give to each tree in input during the SDM analysis. Options are:
	weighting = "flat" All trees have equal weighting.
	weighting = "taxa" Weight is proportional to number of taxa.
	weighting = "inverse" Weight is proportional to 1 / number of taxa.

Defaults to weighting = "flat".

. . .

- Arguments passed on to summary_matrix_to_phylo
- summ_matrix Any summary patristic distance matrix, such as the ones obtained
 with datelife_result_sdm_matrix() or datelife_result_median_matrix().
- total_distance Whether the input summ_matrix stores total age distance (from tip to tip) or distance from node to tip. Default to TRUE, divides the matrix in half, if FALSE it will take it as is.
- use A character vector indicating what type of age to use for summary tree. One of the following:
 - "mean" It will use the mean() of the node ages in summ_matrix.
 - "median" It uses the stats::median() age of node ages in summ_matrix.
 - "min" It will use the min() age from node ages in summ_matrix.
 - "max" Choose this if you wanna be conservative; it will use the max() age from node ages in summ_matrix.
 - "midpoint" It will use the mean of minimum age and maximum age.
- target_tree A phylo object. Use this in case you want a specific backbone for the output tree.
- datelife_query A datelifeQuery object, usually an output of make_datelife_query().

Details

Chronograms given as input in datelife_result are summarized with the Super Distance Matrix (SDM) method described in Criscuolo et al. (2006) doi:10.1080/10635150600969872, implemented with the function ape::SDM(). The resulting summary SDM is clustered with summary_matrix_to_phylo().

Value

A supertree with branch lengths proportional to time, obtained by summarizing individual chronograms given as input in datelife_result. It is returned as an object of class datelifeSDM, which is a phylo object with an additional \$data element storing the input chronograms as a datelifeResult object, and a \$citation element containing citations of studies from input chronograms.

References

Criscuolo A, Berry V, Douzery EJ, Gascuel O. (2006) "SDM: a fast distance-based approach for (super) tree building in phylogenomics" doi:10.1080/10635150600969872.

datelife_result_study_index

Find the index of relevant studies in a cached chronogram database.

Description

datelife_result_study_index is used in summarize_datelife_result().

Usage

```
datelife_result_study_index(datelife_result, cache = "opentree_chronograms")
```

Arguments

datelife_result	t
	A datelifeResult object, usually an output of get_datelife_result().
cache	The cached chronogram database.

Value

A vector of indices of studies that have relevant information.

datelife_result_variance_matrix

Compute a variance matrix of a datelifeResult object.

Description

Compute a variance matrix of a datelifeResult object.

Usage

datelife_result_variance_matrix(datelife_result)

Arguments

```
datelife_result
```

A datelifeResult object, usually an output of get_datelife_result().

Value

A variance matrix from a datelifeResult object.

datelife_search

Get scientific, peer-reviewed information on time of lineage divergence openly available for a given set of taxon names

Description

datelife_search is the core DateLife function to find and get all openly available, peer-reviewed scientific information on time of lineage divergence for a set of input taxon names given as a character vector, a newick character string, a phylo or multiPhylo object or as a an already processed datelifeQuery object obtained with make_datelife_query().

Usage

```
datelife_search(
    input = c("Rhea americana", "Pterocnemia pennata", "Struthio camelus"),
    use_tnrs = FALSE,
    get_spp_from_taxon = FALSE,
    partial = TRUE,
    cache = "opentree_chronograms",
    summary_format = "phylo_all",
    na_rm = FALSE,
    summary_print = c("citations", "taxa"),
    taxon_summary = c("none", "summary", "matrix"),
    criterion = "taxa"
)
```

Arguments

input	One of the following:
	A character vector With taxon names as a single comma separated starting or concatenated with c().
	A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.
	A datelifeQuery object An output from make_datelife_query().
use_tnrs	Whether to use Open Tree of Life's Taxonomic Name Resolution Service (TNRS) to process input taxon names. Default to TRUE, it corrects misspellings and taxonomic name variations with tnrs_match(), a wrapper of rotl::tnrs_match_names().
<pre>get_spp_from_</pre>	taxon
	Whether to search ages for all species belonging to a given taxon or not. De- fault to FALSE. If TRUE, it must have same length as input. If input is a newick string with some clades it will be converted to a phylo object, and the order of get_spp_from_taxon will match phy\$tip.label.
partial	Whether to return or exclude partially matching source chronograms, i.e, those that match some and not all of taxa given in datelife_query. Options are TRUE or FALSE. Defaults to TRUE: return all matching source chronograms.

cache	A character vector of length one, with the name of the data object to cache. Default to "opentree_chronograms", a data object storing Open Tree of Life's database chronograms and other associated information.
summary_format	A character vector of length one, indicating the output format for results of the DateLife search. Available output formats are:
	"citations" A character vector of references where chronograms with some or all of the target taxa are published (source chronograms).
	"mrca" A named numeric vector of most recent common ancestor (mrca) ages of target taxa defined in input, obtained from the source chronograms. Names of mrca vector are equal to citations.
	"newick_all" A named character vector of newick strings corresponding to tar- get chronograms derived from source chronograms. Names of newick_all vector are equal to citations.
	"newick_sdm" Only if multiple source chronograms are available. A character vector with a single newick string corresponding to a target chronogram obtained with SDM supertree method (Criscuolo et al. 2006).
	"newick_median" Only if multiple source chronograms are available. A char- acter vector with a single newick string corresponding to a target chrono- gram from the median of all source chronograms.
	" phylo_sdm " Only if multiple source chronograms are available. A phylo object with a single target chronogram obtained with SDM supertree method (Criscuolo et al. 2006).
	" phylo_median " Only if multiple source chronograms are available. A phylo object with a single target chronogram obtained from source chronograms with median method.
	"phylo_all" A named list of phylo objects corresponding to each target chrono- gram obtained from available source chronograms. Names of phylo_all list correspond to citations.
	" phylo_biggest " The chronogram with the most taxa. In the case of a tie, the chronogram with clade age closest to the median age of the equally large trees is returned.
	"html" A character vector with an html string that can be saved and then opened in any web browser. It contains a 4 column table with data on target taxa: mrca, number of taxa, citations of source chronogram and newick target chronogram.
	"data_frame" A 4 column data.frame with data on target taxa: mrca, number of taxa, citations of source chronograms and newick string.
na_rm	If TRUE, it drops rows containing NAs from the datelifeResult patristic matrix; if FALSE, it returns NA where there are missing entries.
summary_print	A character vector specifying the type of summary information to be printed to screen. Options are:
	"citations" Prints references of chronograms where target taxa are found.
	"taxa" Prints a summary of the number of chronograms where each target taxon is found.
	"none" Nothing is printed to screen.

	Defaults to c("citations", "taxa"), which displays both.
taxon_summary	A character vector specifying if data on target taxa missing in source chrono- grams should be added to the output as a "summary" or as a presence/absence "matrix". Default to "none", no information on taxon_summary added to the output.
criterion	Defaults to criterion = "taxa". Used for chronogram summarizing, i.e., ob- taining a single summary chronogram from a group of input chronograms. For summarizing approaches that return a single summary tree from a group of phy- logenetic trees, it is necessary that the latter form a grove, roughly, a sufficiently overlapping set of taxa between trees, see Ané et al. (2009) doi:10.1007/s00026- 0090017x. In rare cases, a group of trees can have multiple groves. This ar- gument indicates whether to get the grove with the most trees (criterion = "trees") or the most taxa (criterion = "taxa").

Details

If only one taxon name is given as input, get_spp_from_taxon is always set to TRUE.

Value

The output is determined by the argument summary_format:

- If summary_format = "citations" The function returns a character vector of references.
- If summary_format = "mrca" The function returns a named numeric vector of most recent common ancestor (mrca) ages.
- If summary_format = "phylo_[all, sdm, median, or biggest]" The function returns output chronograms as phylo or multiPhylo objects.

Examples

```
## Not run:
```

```
# For this example, we will set a temp working directory, but you can set
# your working directory as needed:
# we will use the tempdir() function to get a temporary directory:
tempwd <- tempdir()
# Obtain median ages from a set of source chronograms in newick format:
ages <- datelife_search(c(
    "Rhea americana", "Pterocnemia pennata", "Struthio camelus",
    "Mus musculus"
), summary_format = "newick_median")
# Save the tree in the temp working directory in newick format:
```

```
write(ages, file = file.path(tempwd, "some.bird.ages.txt"))
```

```
# Obtain median ages from a set of source chronograms in phylo format
# Will produce same tree as above but in "phylo" format:
ages.again <- datelife_search(c(</pre>
  "Rhea americana", "Pterocnemia pennata", "Struthio camelus",
  "Mus musculus"
), summary_format = "phylo_median")
plot(ages.again)
library(ape)
ape::axisPhylo()
mtext("Time (million years ago)", side = 1, line = 2, at = (max(get("last_plot.phylo",
 envir = .PlotPhyloEnv
)$xx) * 0.5))
# Save "phylo" object in newick format
write.tree(ages.again, file = file.path(tempwd, "some.bird.tree.again.txt"))
# Obtain MRCA ages and target chronograms from all source chronograms
# Generate an htm"l output readable in any web browser:
ages.html <- datelife_search(c(</pre>
  "Rhea americana", "Pterocnemia pennata", "Struthio camelus",
  "Mus musculus"
), summary_format = "html")
write(ages.html, file = file.path(tempwd, "some.bird.trees.html"))
system(paste("open", file.path(tempwd, "some.bird.trees.html")))
## End(Not run) # end dontrun
```

datelife_use Generate one or multiple chronograms for a set of given taxon names.

Description

datelife_use gets secondary calibrations available for any pair of given taxon names, mined from the opentree_chronograms object, congruifies them, and uses them to date a given tree topology with the algorithm defined in dating_method. If no tree topology is provided, it will attempt to get one for the given taxon names from Open Tree of Life synthetic tree, using make_bold_otol_tree().

Usage

```
datelife_use(input = NULL, each = FALSE, dating_method = "bladj", ...)
```

Arguments

input	One of the following:
	A character vector With taxon names as a single comma separated starting or
	concatenated with c().
	A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo

A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.

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	A datelifeQuery object An output from make_datelife_query().
each	Boolean, default to FALSE: all calibrations are returned in the same data.frame. If TRUE, calibrations from each chronogram are returned in separate data frames.
dating_method	Tree dating algorithm to use. Options are "bladj" or "pathd8" (Webb et al., 2008, doi:10.1093/bioinformatics/btn358; Britton et al., 2007, doi:10.1080/10635150701613783).
	Arguments passed on to make_datelife_query
	<pre>use_tnrs Whether to use Open Tree of Life's Taxonomic Name Resolution Service (TNRS) to process input taxon names. Default to TRUE, it corrects misspellings and taxonomic name variations with tnrs_match(), a wrap- per of rotl::tnrs_match_names().</pre>
	<pre>get_spp_from_taxon Whether to search ages for all species belonging to a given taxon or not. Default to FALSE. If TRUE, it must have same length as input. If input is a newick string with some clades it will be converted to a phylo object, and the order of get_spp_from_taxon will match phy\$tip.label.</pre>
	reference_taxonomy A character vector specifying the reference taxonomy to use for TNRS. Options are "ott", "ncbi", "gbif" or "irmng". The function defaults to "ott".

Details

If input is a vector of taxon names, the function will attempt to reconstruct a BOLD tree with make_bold_otol_tree() to get a tree with branch lengths. If it fails, it will get an Open Tree of Life synthetic tree topology. The function then calls use_calibrations().

Value

A phylo or multiPhylo object with branch lengths proportional to time.

More

The output object stores the used calibrations and dating_method as attributes(output)\$datelife_calibrations and attributes(output)\$dating_method.

datelife_use_datelifequery

Generate one or multiple chronograms for a set of taxon names given as a datelifeQuery object.

Description

datelife_use gets secondary calibrations available for any pair of given taxon names, mined from the opentree_chronograms object, congruifies them, and uses them to date a given tree topology with the algorithm defined in dating_method. If no tree topology is provided, it will attempt to get one for the given taxon names from Open Tree of Life synthetic tree, using make_bold_otol_tree().

Usage

```
datelife_use_datelifequery(
   datelife_query = NULL,
   dating_method = "bladj",
   each = FALSE
)
```

Arguments

datelife_query	A datelifeQuery object, usually an output of make_datelife_query().
dating_method	Tree dating algorithm to use. Options are "bladj" or "pathd8" (Webb et al., 2008, doi:10.1093/bioinformatics/btn358; Britton et al., 2007, doi:10.1080/10635150701613783).
each	Boolean, default to FALSE: all calibrations are returned in the same data.frame. If TRUE, calibrations from each chronogram are returned in separate data frames.

Details

If phy has no branch lengths, dating_method is ignores, and the function applies secondary calibrations to date the tree with the BLADJ algorithm. See make_bladj_tree() and use_calibrations_bladj(). If phy has branch lengths, the function can use the PATHd8 algorithm. See use_calibrations_pathd8().

Value

A phylo or multiPhylo object with branch lengths proportional to time.

More

The output object stores the used calibrations and dating_method as attributes(output)\$datelife_calibrations and attributes(output)\$dating_method.

date_with_pbdb Date with Paleobiology Database and paleotree.

Description

This will take a topology, look up information about fossils for taxa on the tree, and use paleotree::timePaleoPhy() to compute branch lengths.

Usage

```
date_with_pbdb(phy, recent = FALSE, assume_recent_if_missing = TRUE)
```

Arguments

phy	A phylo object.
recent	If TRUE, forces the minimum age to be zero for any taxon
assume_recent_if_missing	
	If TRUE, any taxon missing from PBDB is assumed to be recent.

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Value

A dated tree.

Examples

```
## Not run: # This is a flag for package development. You are welcome to run the example.
taxa <- c(
    "Archaeopteryx", "Pinus", "Quetzalcoatlus", "Homo sapiens",
    "Tyrannosaurus rex", "Megatheriidae", "Metasequoia", "Aedes", "Panthera"
)
phy <- tree_from_taxonomy(taxa, sources = "The Paleobiology Database")$phy
## End(Not run) # end dontrun
```

extract_calibrations_dateliferesult

Use congruification to extract secondary calibrations from a datelifeResult *object.*

Description

This function extracts node ages for each taxon pair given in input\$tip.labels. It applies the congruification method described in Eastman et al. (2013) doi:10.1111/2041210X.12051, implemented with the function geiger::congruify.phylo(), to create a data.frame of taxon pair node ages that can be used as secondary calibrations.

Usage

```
extract_calibrations_dateliferesult(input = NULL, each = FALSE)
```

Arguments

input	A datelifeResult object.
each	Boolean, default to FALSE: all calibrations are returned in the same data.frame.
	If TRUE, calibrations from each chronogram are returned in separate data frames.

Details

The function takes a datelifeResult object and calls summarize_datelife_result() with summary_format = "phylo_a" eResultobject to aphyloormultiPhylo' object that is passed to extract_calibrations_phylo().

Value

An object of class calibrations, which is a data.frame (if each = FALSE) or a list of data.frames (if each = TRUE) of node ages for each pair of taxon names. You can access the input data from which the calibrations were extracted with attributes(output)\$chronograms.

extract_calibrations_phylo

Use congruification to extract secondary calibrations from a phylo *or* multiPhylo *object with branch lengths proportional to time.*

Description

This function extracts node ages for each taxon pair given in input\$tip.labels. It applies the congruification method described in Eastman et al. (2013) doi:10.1111/2041210X.12051, implemented with the function geiger::congruify.phylo(), to create a data.frame of taxon pair node ages that can be used as secondary calibrations.

Usage

```
extract_calibrations_phylo(input = NULL, each = FALSE)
```

Arguments

input	A phylo or multiPhylo object with branch lengths proportional to time.
each	Boolean, default to FALSE: all calibrations are returned in the same data.frame.
	If TRUE, calibrations from each chronogram are returned in separate data frames.

Value

An object of class calibrations, which is a data.frame (if each = FALSE) or a list of data.frames (if each = TRUE) of node ages for each pair of taxon names. You can access the input data from which the calibrations were extracted with attributes(output)\$chronograms.

References

Eastman et al. (2013) "Congruification: support for time scaling large phylogenetic trees". Methods in Ecology and Evolution, 4(7), 688-691, doi:10.1111/2041210X.12051.

extract_ott_ids *Extract numeric OTT ids from a character vector that combines taxon names and OTT ids.*

Description

Extract numeric OTT ids from a character vector that combines taxon names and OTT ids.

Usage

```
extract_ott_ids(x, na.rm = TRUE)
## Default S3 method:
extract_ott_ids(x, na.rm = TRUE)
```

Arguments

x	A character vector of taxon names, or a phylo object with tree tip labels containing OTT ids.
na.rm	A logical value indicating whether NA values should be stripped from the output.

Value

An object of class numeric containing OTT ids only. NULL

Examples

Not run: # This is a flag for package development. You are welcome to run the example.

```
canis <- rotl::tnrs_match_names("canis")
canis_taxonomy <- rotl::taxonomy_subtree(canis$ott_id)
my_ott_ids <- extract_ott_ids(x = canis_taxonomy$tip_label)
# Get the problematic elements from input
canis_taxonomy$tip_label[attr(my_ott_ids, "na.action")]</pre>
```

End(Not run) # end dontrun

felid_gdr_phylo_all datelifeSummary of a datelifeResult object of all Felidae species.

Description

datelifeSummary of a datelifeResult object of all Felidae species.

Usage

felid_gdr_phylo_all

Format

A list of three elements, containing the summary of a datelifeResult object

phylo_all List of subset chronograms in phylo format

taxon_distribution A data frame with taxon presence across subset chronograms

absent_taxa A dataframe with names of taxon not found in any chronogram

Details

Generated with: felid_spp <- make_datelife_query(input = "felidae", get_spp_from_taxon = TRUE) felid_gdr <- get_datelife_result(input = felid_spp, get_spp_from_taxon = TRUE) felid_gdr_phylo_all <- summarize_datelife_result(datelife_result = felid_gdr, taxon_summary = "summary", summary_format = "phylo_all", datelife_query = felid_spp) usethis::use_data(felid_gdr_phylo_all)

Source

http://opentreeoflife.org

felid_sdm

SDM tree of a datelifeResult object of all Felidae species.

Description

SDM tree of a datelifeResult object of all Felidae species.

Usage

felid_sdm

Format

A list of two elements, containing the summary of a datelifeResult object

phy An ultrametric phylo object with the SDM tree.

data A datelifeResult object with data used to construct phy

Details

Generated with: felid_spp <- make_datelife_query(input = "felidae", get_spp_from_taxon = TRUE) felid_gdr <- get_datelife_result(input = felid_spp, get_spp_from_taxon = TRUE) felid_sdm <datelife_result_sdm_phylo(felid_gdr) usethis::use_data(felid_sdm)

Source

http://opentreeoflife.org

filter_for_grove Filter a datelifeResult object to find the largest grove.

Description

Filter a datelifeResult object to find the largest grove.

Usage

```
filter_for_grove(datelife_result, criterion = "taxa", n = 2)
```

force_ultrametric

Arguments

datelife_result	
	A datelifeResult object. Only needed for criterion = "taxa".
criterion	Defaults to criterion = "taxa". Used for chronogram summarizing, i.e., ob- taining a single summary chronogram from a group of input chronograms. For summarizing approaches that return a single summary tree from a group of phy- logenetic trees, it is necessary that the latter form a grove, roughly, a sufficiently overlapping set of taxa between trees, see Ané et al. (2009) doi:10.1007/s00026- 0090017x. In rare cases, a group of trees can have multiple groves. This ar- gument indicates whether to get the grove with the most trees (criterion = "trees") or the most taxa (criterion = "taxa").
n	The degree of taxon name overlap among input chronograms. Defaults to n = 2, i.e., at least two overlapping taxon names.

Value

A datelifeResult object filtered to only include one grove of trees.

force_ultrametric	Force a non-ultrametric phylo object to be ultrametric wi	th
	<pre>phytools::force.ultrametric().</pre>	

Description

Force a non-ultrametric phylo object to be ultrametric with phytools::force.ultrametric().

Usage

force_ultrametric(phy)

Arguments

phy A phylo object.

Value

A phylo object.

get_all_calibrations Get secondary calibrations from a chronogram database for a set of given taxon names

Description

get_all_calibrations performs a datelife_search() and gets divergence times (i.e., secondary calibrations) from a chronogram database for each taxon name pair given as input.

get_all_calibrations performs a datelife_search() and gets divergence times (i.e., secondary calibrations) from a chronogram database for each taxon name pair given as input.

Usage

```
get_all_calibrations(input = NULL, each = FALSE)
get_all_calibrations(input = NULL, each = FALSE)
```

Arguments

input	One of the following:
	A character vector With taxon names as a single comma separated starting or concatenated with c().
	A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.
	A datelifeQuery object An output from make_datelife_query().
each	Boolean, default to FALSE: all calibrations are returned in the same data.frame. If TRUE, calibrations from each chronogram are returned in separate data frames.

Value

An object of class calibrations, which is a data.frame (if each = FALSE) or a list of data.frames (if each = TRUE) of node ages for each pair of taxon names. You can access the input data from which the calibrations were extracted with attributes(output)\$chronograms.

get_all_descendant_species

Quickly get all species belonging to a taxon from the Open Tree of Life Taxonomy (OTT)

Description

This is less thorough than get_open_tree_species(), but much faster. It uses the fact that something has just two names (genus and species) to assume that something is a single species; if it has more than two names, it is assumed to be a subspecies so it goes up one level in the hierarchy. It will return the subspecies and the species.

get_best_grove

Usage

get_all_descendant_species(taxon_name, ott_id)

Arguments

taxon_name	A character vector providing an inclusive taxonomic name.
ott_id	A numeric vector providig an Open Tree Taxonomic id number for a taxonomic name. If provided, taxon_name is ignored. Used in the context of OTT to detect invalid taxon names.

Value

A list of unique OTT names and OTT ids of species within the provided taxon.

<pre>get_best_grove</pre>	Get grove from a datelifeResult object that can be converted to
	phylo from a median summary matrix

Description

Get grove from a datelifeResult object that can be converted to phylo from a median summary matrix

Usage

```
get_best_grove(datelife_result, criterion = "taxa", n = 2)
```

Arguments

datelife_result

A datelifeResult object. Only needed for criterion = "taxa".

- criterion Defaults to criterion = "taxa". Used for chronogram summarizing, i.e., obtaining a single summary chronogram from a group of input chronograms. For summarizing approaches that return a single summary tree from a group of phylogenetic trees, it is necessary that the latter form a grove, roughly, a sufficiently overlapping set of taxa between trees, see Ané et al. (2009) doi:10.1007/s00026-0090017x. In rare cases, a group of trees can have multiple groves. This argument indicates whether to get the grove with the most trees (criterion = "trees") or the most taxa (criterion = "taxa").
- n The degree of taxon name overlap among input chronograms. Defaults to n = 2, i.e., at least two overlapping taxon names.

Value

A list of two elements:

best_grove A datelifeResult object filtered to only include one grove of trees that can be summarized with median or sdm.

overlap The degree of taxon names overlap among trees in the best grove.

get_biggest_multiphylo

Get the tree with the most tips from a multiPhylo object: the biggest tree.

Description

Get the tree with the most tips from a multiPhylo object: the biggest tree.

Usage

get_biggest_multiphylo(trees)

Arguments

trees A list of trees as multiPhylo or as a generic list object.

Value

The largest tree from those given in trees, as a phylo object with an additional \$citation element containing the reference of the original publication.

get_bold_data	Get genetic data from the Barcode of Life Database (BOLD) for a set
	of taxon names.

Description

get_bold_data uses taxon names from a tree topology, a character vector of names or a datelifeQuery object, to search for genetic markers in the Barcode of Life Database (BOLD).

Usage

```
get_bold_data(
    input = c("Rhea americana", "Struthio camelus", "Gallus gallus"),
    marker = "COI",
    ...
)
```

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input	One of the following:
	A character vector With taxon names as a single comma separated starting or concatenated with c().
	A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.
	A datelifeQuery object An output from make_datelife_query().
marker	A character vector indicating the gene from BOLD system to be used for branch length estimation. It searches "COI" marker by default.
	Arguments passed on to get_otol_synthetic_tree
	<pre>otol_version Version of Open Tree of Life to use resolve Defaults to TRUE. Whether to resolve the tree at random or not. ott_ids If not NULL, it takes this argument and ignores input. A numeric vec- tor of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().</pre>

Details

If input is a phylo object or a newick string, it is used as backbone topology. If input is a character vector of taxon names, an induced synthetic OpenTree subtree is used as backbone.

Value

A phylo object. If there are enough BOLD sequences available for the input taxon names, the function returns a tree with branch lengths proportional to relative substitution rate. If not enough BOLD sequences are available for the input taxon names, the function returns the topology given as input, or a synthetic Open Tree of Life for the taxon names given in input, obtained with get_otol_synthetic_tree().

get_calibrations_datelifequery

Search and extract available secondary calibrations for taxon names in a given datelifeQuery object

Description

The function searches DateLife's local database of phylogenetic trees with branch lengths proportional to time (chronograms) with datelife_search(), and extracts available node ages for each pair of given taxon names with extract_calibrations_phylo().

Usage

```
get_calibrations_datelifequery(datelife_query = NULL, each = FALSE)
```

datelife_query	A datelifeQuery object.
each	Boolean, default to FALSE: all calibrations are returned in the same data.frame.
	If TRUE, calibrations from each chronogram are returned in separate data frames.

Details

The function calls datelife_search() with summary_format = "phylo_all" to get all chronograms in the database containing at least two taxa in input, and generates a phylo or multiPhylo object object that will be passed to extract_calibrations_phylo().

Value

An object of class calibrations, which is a data.frame (if each = FALSE) or a list of data.frames (if each = TRUE) of node ages for each pair of taxon names. You can access the input data from which the calibrations were extracted with attributes(output)\$chronograms.

get_calibrations_vector

Search and extract secondary calibrations for a given character vector of taxon names

Description

The function searches DateLife's local database of phylogenetic trees with branch lengths proportional to time (chronograms) with datelife_search(), and extracts available node ages for each pair of given taxon names with extract_calibrations_phylo().

Usage

get_calibrations_vector(input = NULL, each = FALSE)

Arguments

input	A character vector of taxon names.
each	Boolean, default to FALSE: all calibrations are returned in the same data.frame.
	If TRUE, calibrations from each chronogram are returned in separate data frames.

Details

The function calls datelife_search() with summary_format = "phylo_all" to get all chronograms in the database containing at least two taxa in input, and generates a phylo or multiPhylo object object that will be passed to extract_calibrations_phylo().

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Value

An object of class calibrations, which is a data.frame (if each = FALSE) or a list of data.frames (if each = TRUE) of node ages for each pair of taxon names. You can access the input data from which the calibrations were extracted with attributes(output)\$chronograms.

get_dated_otol_induced_subtree

Get a dated OpenTree induced synthetic subtree from a set of given taxon names, from blackrim's FePhyFoFum service.

Description

Get a dated OpenTree induced synthetic subtree from a set of given taxon names, from blackrim's FePhyFoFum service.

Usage

```
get_dated_otol_induced_subtree(input = NULL, ott_ids = NULL, ...)
```

Arguments

input	Optional. A character vector of names or a datelifeQuery object.
ott_ids	If not NULL, it takes this argument and ignores input. A numeric vector of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().
	Arguments passed on to check_ott_input

Details

OpenTree dated tree from Stephen Smith's OpenTree scaling service at https://github.com/FePhyFoFum/gophy if you want to make an LTT plot of a dated OpenTree tree you'll need to get rid of singleton nodes with ape::collapse.singles() and also probably do phytools::force.ultrametric().

Value

A phylo object with edge length proportional to time in Myrs. It will return NA if any ott_id is invalid.

get_datelife_result

Get a patristic matrix of time of lineage divergence data for a given set of taxon names

Description

get_datelife_result takes as input a vector of taxon names, a newick string, a phylo object, or adatelifeQuery object. It searches the chronogram database specified in cache for chronograms matching two or more given taxon names. For each matching chronogram, it extracts time of lineage divergence data and stores it as a patristic matrix. It then lists all resulting patristic matrices. Each list element is named with the study citation of the source chronogram.

Usage

```
get_datelife_result(
    input = NULL,
    partial = TRUE,
    cache = "opentree_chronograms",
    update_opentree_chronograms = FALSE,
    ...
)
```

Arguments

One of the following:
A character vector With taxon names as a single comma separated starting or concatenated with c().
A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.
A datelifeQuery object An output from make_datelife_query().
Whether to return or exclude partially matching source chronograms, i.e, those that match some and not all of taxa given in datelife_query. Options are TRUE or FALSE. Defaults to TRUE: return all matching source chronograms.
A character vector of length one, with the name of the data object to cache. Default to "opentree_chronograms", a data object storing Open Tree of Life's database chronograms and other associated information.
_chronograms
Whether to update the chronogram database or not. Defaults to FALSE.
Arguments passed on to make_datelife_query
<pre>use_tnrs Whether to use Open Tree of Life's Taxonomic Name Resolution Service (TNRS) to process input taxon names. Default to TRUE, it corrects misspellings and taxonomic name variations with tnrs_match(), a wrap- per of rotl::tnrs_match_names().</pre>

get_spp_from_taxon Whether to search ages for all species belonging to a given taxon or not. Default to FALSE. If TRUE, it must have same length as input. If input is a newick string with some clades it will be converted to a phylo object, and the order of get_spp_from_taxon will match phy\$tip.label. reference_taxonomy A character vector specifying the reference taxonomy to use for TNRS. Options are "ott", "ncbi", "gbif" or "irmng". The function defaults to "ott".

Value

A datelifeResult object – a named list of patristic matrices.

get_datelife_result_datelifequery

Get a list of patristic matrices from a given datelifeQuery object

Description

Get a list of patristic matrices from a given datelifeQuery object

Usage

```
get_datelife_result_datelifequery(
   datelife_query = NULL,
   partial = TRUE,
   cache = "opentree_chronograms",
   update_opentree_chronograms = FALSE,
   ...
)
```

Arguments

datelife_query	A datelifeQuery object, usually an output of make_datelife_query().	
partial	Whether to return or exclude partially matching source chronograms, i.e, those that match some and not all of taxa given in datelife_query. Options are TRUE or FALSE. Defaults to TRUE: return all matching source chronograms.	
cache	A character vector of length one, with the name of the data object to cache. Default to "opentree_chronograms", a data object storing Open Tree of Life's database chronograms and other associated information.	
update_opentree_chronograms		
	Whether to update the chronogram database or not. Defaults to FALSE.	
	Arguments passed on to make_datelife_query	
	input Taxon names as one of the following:	
	A character vector of taxon names With taxon names as a single comma separated starting or concatenated with c().	

- A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.
- use_tnrs Whether to use Open Tree of Life's Taxonomic Name Resolution Service (TNRS) to process input taxon names. Default to TRUE, it corrects misspellings and taxonomic name variations with tnrs_match(), a wrapper of rotl::tnrs_match_names().
- get_spp_from_taxon Whether to search ages for all species belonging to a given taxon or not. Default to FALSE. If TRUE, it must have same length as input. If input is a newick string with some clades it will be converted to a phylo object, and the order of get_spp_from_taxon will match phy\$tip.label.
- reference_taxonomy A character vector specifying the reference taxonomy to use for TNRS. Options are "ott", "ncbi", "gbif" or "irmng". The function defaults to "ott".

Details

If there is just one taxon name in input\$cleaned_names, the function will run make_datelife_query() setting get_spp_from_taxon = TRUE. The datelifeQuery used as input can be accessed with attributes(datelifeResult)\$query.

Value

A datelifeResult object – a named list of patristic matrices.

get_fossil_range Get the ages for a taxon from PBDB

Description

This uses the Paleobiology Database's API to gather information on the ages for all specimens of a taxon. It will also look for all descendants of the taxon. It fixes name misspellings if possible.

Usage

```
get_fossil_range(taxon, recent = FALSE, assume_recent_if_missing = TRUE)
```

Arguments

taxon	The scientific name of the taxon you want the range of occurrences of
recent	If TRUE, forces the minimum age to be zero
assume_recen	t_if_missing
	If TRUE, any taxon missing from pbdb is assumed to be recent

Value

a data.frame of max_ma and min_ma for the specimens

get_goodmatrices Get indices of good matrices to apply Super Distance Matrix (SDM) method with make_sdm().

Description

Get indices of good matrices to apply Super Distance Matrix (SDM) method with make_sdm().

Usage

get_goodmatrices(unpadded.matrices)

Arguments

unpadded.matrices

A list of patristic matrices, a datelifeResult object.

Value

A numeric vector of good matrix indices in unpadded.matrices.

```
get_mrbayes_node_constraints
```

Makes a block of node constraints and node calibrations for a Mr-Bayes run file from a list of taxa and ages, or from a dated tree

Description

Makes a block of node constraints and node calibrations for a MrBayes run file from a list of taxa and ages, or from a dated tree

Usage

```
get_mrbayes_node_constraints(
  constraint = NULL,
  taxa = NULL,
  missing_taxa = NULL,
  ncalibration = NULL,
  age_distribution = "fixed",
  root_calibration = FALSE,
  mrbayes_constraints_file = NULL,
  clockratepr = "prset clockratepr = fixed(1);"
)
```

constraint	The constraint tree: a phylo object or a newick character string, with or without branch lengths.
taxa	A character vector with taxon names to be maintained in tree
missing_taxa	A tree, a data frame or a vector enlisting all missing taxa you want to include.
	A tree Either as a phylo object or as a newick character string. It contains all taxa that you want at the end, both missing and non missing. This tree will be used as a hard constraint.
	A data.frame It contains two columns named "taxon" and "clade". The first one contains a character vector of missing taxon names. The second one contains a character or numeric vector of nodes from a constraint tree to which each taxon will be assigned.
	A character vector It contains the names of the missing taxa. They will be added at random to the constraint tree.
ncalibration	The node calibrations: a phylo object with branch lengths proportional to time; in this case all nodes from ncalibration will be used as calibration points. Alternatively, a list with two elements: the first is a character vector with node names from phy to calibrate; the second is a numeric vector with the corresponding ages to use as calibrations.
age_distributio	n
	A character string specifying the type of calibration. Only "fixed" and "uniform" are implemented for now.
	fixed The age given in ncalibration will be used as fixed age.
	lognormal The age given in ncalibration will be used as mean age. The stan- dard deviation can be provided. # still need to add this option. By default, a 95 CI sd is used.
	uniform The age given in ncalibration will be used as mean age. Where min_age = $0.9 *$ mean age, and max_age = $1.1 *$ mean age.
root_calibratio	n
	Used to set a calibration at the root or not. Default to FALSE. Only relevant if ncalibration is specified.
mrbayes_constraints_file	
	NULL or a character vector indicating the name of mrbayes constraint and/or calibration block file.
clockratepr	A character vector indicating the clockrateprior to be used.

Value

A set of MrBayes constraints and/or calibration commands printed in console as character strings or as a text file specified in mrbayes_constraints_file.

get_opentree_chronograms

Get all chronograms from Open Tree of Life database

Description

Get all chronograms from Open Tree of Life database

Usage

```
get_opentree_chronograms(max_tree_count = "all")
```

```
get_otol_chronograms(max_tree_count = "all")
```

Arguments

max_tree_count Default to "all", it gets all available chronograms. For testing purposes, a numeric value indicating the max number of trees to be cached.

Value

A list of 4 elements:

- **authors** A list of lists of author names of the original studies that published chronograms currently stored in the Open Tree of Life database.
- **curators** A list of lists of curator names that uploaded chronograms to the Open Tree of Life database.
- **studies** A list of study identifiers from original studies that published chronograms currently stored in the Open Tree of Life database.
- trees A multiPhylo object storing the chronograms from Open Tree of Life database.
- update A character vector indicating the time when the database object was last updated.
- version A character vector indicating the datelife package version when the object was last updated.

get_opentree_species Get all species belonging to a taxon from the Open Tree of Life Taxonomy (OTT)

Description

Get all species belonging to a taxon from the Open Tree of Life Taxonomy (OTT)

Usage

```
get_opentree_species(taxon_name, ott_id, synth_tree_only = TRUE)
```

taxon_name	A character vector providing an inclusive taxonomic name.
ott_id	A numeric vector providig an Open Tree Taxonomic id number for a taxonomic name. If provided, taxon_name is ignored. Used in the context of OTT to detect invalid taxon names.
synth_tree_only	Whether to include species that are in the synthetic Open Tree of Life only or not. Default to TRUE.
	not. Dolutin to mot.

Value

A list of unique OTT names and OTT ids of species within the provided taxon.

```
get_otol_synthetic_tree
```

Get an Open Tree of Life synthetic subtree of a set of given taxon names.

Description

Get an Open Tree of Life synthetic subtree of a set of given taxon names.

Usage

```
get_otol_synthetic_tree(
    input = NULL,
    ott_ids = NULL,
    otol_version = "v3",
    resolve = FALSE,
    ...
)
```

Arguments

input	Optional. A character vector of names or a datelifeQuery object.
ott_ids	If not NULL, it takes this argument and ignores input. A numeric vector of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().
otol_version	Version of Open Tree of Life to use
resolve	Defaults to TRUE. Whether to resolve the tree at random or not.
	Arguments passed on to check_ott_input

Value

A phylo object

get_ott_children	Use this instead of rotl::tol_subtree() when taxa are not in synthesis tree and you still need to get all species or an induced OpenTree subtree

Description

Use this instead of rotl::tol_subtree() when taxa are not in synthesis tree and you still need to get all species or an induced OpenTree subtree

Usage

```
get_ott_children(input = NULL, ott_ids = NULL, ott_rank = "species", ...)
```

Arguments

input	Optional. A character vector of names or a datelifeQuery object.
ott_ids	If not NULL, it takes this argument and ignores input. A numeric vector of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().
ott_rank	A character vector with the ranks you wanna get lineage children from.
	Other arguments to pass to get_valid_children().

Value

A data.frame object.

Examples

An example with the dog genus:

```
# It is currently not possible to get an OpenTree subtree of a taxon that is
# missing from the OpenTree synthetic tree.
# The dog genus is not monophyletic in the OpenTree synthetic tree, so in
# practice, it has no node to extract a subtree from.
tnrs <- tnrs_match("Canis")
## Not run: # This is a flag for package development. You are welcome to run the example.
rotl::tol_subtree(tnrs$ott_id[1])
#> Error: HTTP failure: 400
#> [/v3/tree_of_life/subtree] Error: node_id was not found (broken taxon).
## End(Not run) # end dontrun
```

```
ids <- tnrs$ott_id[1]
names(ids) <- tnrs$unique_name
children <- get_ott_children(ott_ids = ids) # or</pre>
```

```
children <- get_ott_children(input = "Canis")
if (!is.na(children)) {
str(children)
ids <- children$Canis$ott_id
names(ids) <- rownames(children$Canis)
tree_children <- datelife::get_otol_synthetic_tree(ott_ids = ids)
plot(tree_children, cex = 0.3)
}
# An example with flowering plants:
## Not run: # This is a flag for package development. You are welcome to run the example.
oo <- get_ott_children(input = "magnoliophyta", ott_rank = "order")
# Get the number of orders of flowering plants that we have
sum(oo$Magnoliophyta$rank == "order")
## End(Not run) # end dontrun</pre>
```

get_	ott_	clade	
------	------	-------	--

Get the Open Tree of Life Taxonomic identifiers (OTT ids) and name of one or several given taxonomic ranks from one or more input taxa.

Description

Get the Open Tree of Life Taxonomic identifiers (OTT ids) and name of one or several given taxonomic ranks from one or more input taxa.

Usage

```
get_ott_clade(input = NULL, ott_ids = NULL, ott_rank = "family")
```

Arguments

input	Optional. A character vector of names or a datelifeQuery object.
ott_ids	If not NULL, it takes this argument and ignores input. A numeric vector of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().
ott_rank	A character vector with the ranks you wanna get lineage children from.

Value

A list of named numeric vectors with OTT ids from input and all requested ranks.

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get_ott_lineage

Get the Open Tree of Life Taxonomic identifier (OTT id) and name of all lineages from one or more input taxa.

Description

Get the Open Tree of Life Taxonomic identifier (OTT id) and name of all lineages from one or more input taxa.

Usage

get_ott_lineage(input = NULL, ott_ids = NULL)

Arguments

input	Optional. A character vector of names or a datelifeQuery object.
ott_ids	If not NULL, it takes this argument and ignores input. A numeric vector of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().

Value

A list of named numeric vectors of ott ids from input and all the clades it belongs to.

Examples

Not run: # This is a flag for package development. You are welcome to run the example.

```
taxa <- c("Homo", "Bacillus anthracis", "Apis", "Salvia")
lin <- get_ott_lineage(taxa)
lin
# Look up an unknown OTT id:
get_ott_lineage(ott_id = 454749)</pre>
```

End(Not run) # end dontrun

 ${\tt get_subset_array_dispatch}$

Figure out which subset function to use.

Description

get_subset_array_dispatch is used inside get_datelife_result()

Usage

```
get_subset_array_dispatch(
  study_element,
  taxa,
  phy = NULL,
  phy4 = NULL,
  dating_method = "PATHd8"
)
```

Arguments

study_element	The thing being passed in: an array or a phylo object to serve as reference for congruification.
taxa	Vector of taxon names to get a subset for.
phy	A user tree to congruify as phylo object (ape).
phy4	A user tree to congruify in phylo4 format (phylobase).
dating_method	The method used for tree dating.

Value

A patristic matrix with ages for the target taxa.

get_taxon_summary Get a taxon summary of a datelifeResult object.

Description

Get a taxon summary of a datelifeResult object.

Usage

```
get_taxon_summary(datelife_result = NULL, datelife_query = NULL)
```

Arguments

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Value

A datelifeTaxonSummary object, which is a list of 4 elements:

\$matrix Data as a presence/absence matrix of taxon names across chronograms.

- **\$summary** A data.frame with taxon names as row.names() and two columns, one with the number of chronograms that contain a taxon name and the other one with the total number of chronograms that have at least 2 taxon names.
- **\$summary2** A data.frame with chronogram citations as row.names() and two columns, one with the number of taxon names found in each chronogram and the other one with the total number of taxon names.
- **\$absent_taxa** A character vector of taxon names that are not found in the chronogram database.

get_tnrs_names Process a character vector of taxon names with TNRS

Description

make_datelife_query2 always uses TNRS (Taxonomic Name Resolution Service to process input taxon names, to correct misspellings and taxonomic name variations with tnrs_match(), a wrapper of rotl::tnrs_match_names()).

Usage

```
get_tnrs_names(
    input = c("Rhea americana", "Pterocnemia pennata", "Struthio camelus"),
    reference_taxonomy = "ott",
    ...
)
```

Arguments

input	Taxon names as a character vector of taxon names. Two or more names can be provided as a single comma separated string or concatenated with $c()$.
reference_taxor	nomy
	A character vector specifying the reference taxonomy to use for TNRS. Options are "ott", "ncbi", "gbif" or "irmng". The function defaults to "ott".
	Arguments passed on to rotl::tnrs_match_names
	context_name name of the taxonomic context to be searched (length-one char- acter vector or NULL). Must match (case sensitive) one of the values returned by tnrs_contexts. Default to "All life".
	do_approximate_matching A logical indicating whether or not to perform approximate string (a.k.a. "fuzzy") matching. Using FALSE will greatly improve speed. Default, however, is TRUE.
	ids A vector of ids to use for identifying names. These will be assigned to each name in the names array. If ids is provided, then ids and names must be identical in length.

include_suppressed Ordinarily, some quasi-taxa, such as incertae sedis buckets and other non-OTUs, are suppressed from TNRS results. If this parameter is true, these quasi-taxa are allowed as possible TNRS results.

Value

A datelifeTNRS object, which is a list of three elements:

\$cleaned_names A character vector of names provided as input.
\$tnrs_names A character vector of taxon names processed with TNRS.
\$ott_ids A numeric vector of Open Tree of Life Taxonomy (OTT) ids.

get_valid_children	Extract valid children from given taxonomic name(s) or Open Tree of
	Life Taxonomic identifiers (OTT ids) from a taxonomic source.

Description

Extract valid children from given taxonomic name(s) or Open Tree of Life Taxonomic identifiers (OTT ids) from a taxonomic source.

Usage

```
get_valid_children(input = NULL, ott_ids = NULL, reference_taxonomy = "ncbi")
```

Arguments

input	Optional. A character vector of names or a datelifeQuery object.	
ott_ids	If not NULL, it takes this argument and ignores input. A numeric vector of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().	
reference_taxonomy		
	A character vector with the desired taxonomic sources. Options are "ncbi", "gbif" or "irmng". Any other value will retrieve data from all taxonomic sources.	

Details

GBIF and other taxonomies contain deprecated taxa that are not marked as such in the Open Tree of Life Taxonomy. We are relying mainly in the NCBI taxonomy for now.

Value

A named list containing valid taxonomic children of given taxonomic name(s).

The function defaults to "ncbi".

input_process

Examples

```
# genus Dictyophyllidites with ott id = 6003921 has only extinct children
# in cases like this the same name will be returned
```

```
tti <- rotl::taxonomy_taxon_info(6003921, include_children = TRUE)
gvc <- get_valid_children(ott_ids = 6003921)</pre>
```

More examples:

get_valid_children(ott_ids = 769681) # Psilotopsida
get_valid_children(ott_ids = 56601) # Marchantiophyta

input_process	Process a phylo object or a character string to determine if it's correct
	newick

Description

Process a phylo object or a character string to determine if it's correct newick

Usage

input_process(input)

Arguments

input	Taxon names as one of the following:
	A character vector of taxon names With taxon names as a single comma sep-
	arated starting or concatenated with c().
	A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo
	object, OR as a newick character string.

Value

A phylo object or NA if input is not a tree.

is_datelife_query Check if input is a datelifeQuery object

Description

is_datelife_query checks for two things to be TRUE or FALSE. First, that input is of class datelifeQuery. Second, that input is a list that contains at least two elements of a datelifeQuery object:

cleaned_names A character vector of taxon names.

phy Either NA or a phylo object.

Usage

is_datelife_query(input)

Arguments

input An object to be checked as an object with essential properties of a 'datelife-Query' object.

Details

If the object has the correct format but it has a class different than datelifeQuery, the class is not modified.

Value

Is determined by the second condition.

is_datelife_result_empty

Check if we obtained an empty search with the given taxon name(s).

Description

Check if we obtained an empty search with the given taxon name(s).

Usage

```
is_datelife_result_empty(datelife_result, use_tnrs = FALSE)
```

Arguments

datelife_result		
	A datelifeResult object, usually an output of get_datelife_result().	
use_tnrs	Whether to use Open Tree of Life's Taxonomic Name Resolution Service (TNRS) to process input taxon names. Default to TRUE, it corrects misspellings and taxonomic name variations with tnrs_match(), a wrapper of rot1::tnrs_match_names().	

Value

Boolean. If TRUE, no chronograms were found for the given taxon name(s). If FALSE, the chronogram search was successful. is_good_chronogram Check if a tree is a valid chronogram.

Description

Check if a tree is a valid chronogram.

Usage

is_good_chronogram(phy)

Arguments

phy A phylo object.

Value

TRUE if it is a valid tree.

is_n_overlap	Function for computing n-overlap for two vectors of names (ie.,
	phy1\$tip.label, phy2\$tip.label) and seeing if they have n overlap

Description

This function implements definition 2.8 for n-overlap from Ané et al. (2009) doi:10.1007/s00026-0090017x.

Usage

is_n_overlap(names_1, names_2, n = 2)

Arguments

names_1	First vector of names
names_2	Second vector of names
n	Degree of overlap required

Value

Boolean for whether the degree of overlap was met or not.

References

Ané, C., Eulenstein, O., Piaggio-Talice, R., & Sanderson, M. J. (2009). "Groves of phylogenetic trees". Annals of Combinatorics, 13(2), 139-167, doi:10.1007/s000260090017x.

make_all_associations Find all authors and where they have deposited their trees

Description

Find all authors and where they have deposited their trees

Usage

```
make_all_associations(outputfile = "depositorcache.RData")
```

Arguments

outputfile Path including file name. NULL to prevent saving.

Value

a data.frame of "person" and "urls".

make_bladj_tree	Use the BLADJ algorithm to get a chronogram from a tree topology
	for which you have age data for some of its nodes.

Description

The function takes a tree topology and uses the BLADJ algorithm implemented with phylocomr::ph_bladj() to assign node ages and branch lengths, given a set of fixed node ages and respective node names.

Usage

make_bladj_tree(tree = NULL, nodenames = NULL, nodeages = NULL)

Arguments

tree	A tree either as a newick character string or as a phylo object.
nodenames	A character vector with names of nodes in tree with known ages
nodeages	A numeric vector with the actual ages of named nodes

Details

Input tree can be dated or not, \$edge.length is ignored. Ages given in nodeages are fixed on their corresponding nodes given in nodenames.

Value

A phylo object.

make_bold_otol_tree

Use genetic data from the Barcode of Life Database (BOLD) to reconstruct branch lengths on a tree.

Description

make_bold_otol_tree takes taxon names from a tree topology or a vector of names to search for genetic markers in the Barcode of Life Database (BOLD), create an alignment, and reconstruct branch lengths on a tree topology with Maximum Likelihood.

Usage

```
make_bold_otol_tree(
    input = c("Rhea americana", "Struthio camelus", "Gallus gallus"),
    marker = "COI",
    otol_version = "v3",
    chronogram = TRUE,
    doML = FALSE,
    aligner = "muscle",
    ...
)
```

Arguments

input	One of the following:
	A character vector With taxon names as a single comma separated starting or concatenated with c().
	A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.
	A datelifeQuery object An output from make_datelife_query().
marker	A character vector indicating the gene from BOLD system to be used for branch length estimation.
otol_version	Version of Open Tree of Life to use
chronogram	Default to TRUE, branch lengths returned are estimated with ape::chronoMPL(). If FALSE, branch lengths returned are estimated with phangorn::acctran() and represent relative substitution rates.
doML	Default to FALSE. If TRUE, it does a ML branch length optimization with phangorn::optim.pml().
aligner	A character vector indicating whether to use MAFFT or MUSCLE to align BOLD sequences. It is not case sensitive. Default to MUSCLE, supported using the msa package from Bioconductor, which needs to be installed using BiocManager::install().
	Arguments passed on to get_otol_synthetic_tree
	<pre>resolve Defaults to TRUE. Whether to resolve the tree at random or not. ott_ids If not NULL, it takes this argument and ignores input. A numeric vec- tor of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().</pre>

Details

If input is a phylo object or a newick string, it is used as backbone topology. If input is a character vector of taxon names, an induced synthetic OpenTree subtree is used as backbone.

Value

A phylo object. If there are enough BOLD sequences available for the input taxon names, the function returns a tree with branch lengths proportional to relative substitution rate. If not enough BOLD sequences are available for the input taxon names, the function returns the topology given as input, or a synthetic Open Tree of Life for the taxon names given in input, obtained with get_otol_synthetic_tree().

make_contributor_cache

Create a cache from Open Tree of Life

Description

Create a cache from Open Tree of Life

Usage

make_contributor_cache(outputfile = "contributorcache.RData")

Arguments

outputfile Path including file name

Value

List containing author and curator results

make_datelife_query Go from taxon names to a datelifeQuery object

Description

Go from taxon names to a datelifeQuery object

Usage

```
make_datelife_query(
    input = c("Rhea americana", "Pterocnemia pennata", "Struthio camelus"),
    use_tnrs = TRUE,
    get_spp_from_taxon = FALSE,
    reference_taxonomy = "ott"
)
```

60

input	Taxon names as one of the following:	
	A character vector of taxon names With taxon names as a single comma separated starting or concatenated with c().	
	A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.	
use_tnrs	Whether to use Open Tree of Life's Taxonomic Name Resolution Service (TNRS) to process input taxon names. Default to TRUE, it corrects misspellings and taxonomic name variations with tnrs_match(), a wrapper of rotl::tnrs_match_names().	
get_spp_from_taxon		
	Whether to search ages for all species belonging to a given taxon or not. De- fault to FALSE. If TRUE, it must have same length as input. If input is a newick string with some clades it will be converted to a phylo object, and the order of get_spp_from_taxon will match phy\$tip.label.	
reference_taxonomy		
	A character vector specifying the reference taxonomy to use for TNRS. Options are "ott", "ncbi", "gbif" or "irmng". The function defaults to "ott".	

Details

It processes phylo objects and newick character string inputs with input_process(). If input is a multiPhylo object, only the first phylo element will be used. Similarly, if an input newick character string has multiple trees, only the first one will be used.

Value

A datelifeQuery object, which is a list of three elements:

\$phy A phylo object or NA, if input is not a tree.

\$cleaned_names A character vector of cleaned taxon names.

\$ott_ids A numeric vector of OTT ids if use_tnrs = TRUE, or NULL if use_tnrs = FALSE.

make_datelife_query2 Go from taxon names to a datelifeQuery object

Description

Go from taxon names to a datelifeQuery object

Usage

```
make_datelife_query2(
    input = c("Rhea americana", "Pterocnemia pennata", "Struthio camelus"),
    get_spp_from_taxon = FALSE,
    reference_taxonomy = "ott",
    ...
)
```

input	Taxon names as one of the following:	
	A character vector of taxon names With taxon names as a single comma separated starting or concatenated with c().	
	A phylogenetic tree with taxon names as tip labels As a phylo or multiPhylo object, OR as a newick character string.	
<pre>get_spp_from_ta</pre>	axon	
	Whether to search ages for all species belonging to a given taxon or not. De- fault to FALSE. If TRUE, it must have same length as input. If input is a newick string with some clades it will be converted to a phylo object, and the order of get_spp_from_taxon will match phy\$tip.label.	
reference_taxonomy		
	A character vector specifying the reference taxonomy to use for TNRS. Options are "ott", "ncbi", "gbif" or "irmng". The function defaults to "ott".	
	Arguments passed on to rotl::tnrs_match_names	
	context_name name of the taxonomic context to be searched (length-one char- acter vector or NULL). Must match (case sensitive) one of the values returned by tnrs_contexts. Default to "All life".	
	do_approximate_matching A logical indicating whether or not to perform approximate string (a.k.a. "fuzzy") matching. Using FALSE will greatly improve speed. Default, however, is TRUE.	
	ids A vector of ids to use for identifying names. These will be assigned to each name in the names array. If ids is provided, then ids and names must be identical in length.	
	include_suppressed Ordinarily, some quasi-taxa, such as incertae sedis buck- ets and other non-OTUs, are suppressed from TNRS results. If this param- eter is true, these quasi-taxa are allowed as possible TNRS results.	

Details

It processes phylo objects and newick character string inputs with input_process(). If input is a multiPhylo object, only the first phylo element will be used. Similarly, if an input newick character string has multiple trees, only the first one will be used.

Value

A datelifeQuery object, which is a list of four elements:

\$input_names A character vector of input taxon names.

\$tnrs_names A character vector of taxon names processed with TNRS.

\$ott_ids A numeric vector of OTT ids.

\$phy A phylo object or NA, if input is not a tree.

make_mrbayes_runfile

Make a mrBayes run block file with a constraint topology and a set of node calibrations and missing taxa

Description

Make a mrBayes run block file with a constraint topology and a set of node calibrations and missing taxa

Usage

```
make_mrbayes_runfile(
  constraint = NULL,
  taxa = NULL,
  ncalibration = NULL,
  missing_taxa = NULL,
  age_distribution = "fixed",
  root_calibration = FALSE,
  mrbayes_output_file = "mrbayes_run.nexus"
)
```

Arguments

constraint	The constraint tree: a phylo object or a newick character string, with or without branch lengths.	
taxa	A character vector with taxon names to be maintained in tree	
ncalibration	The node calibrations: a phylo object with branch lengths proportional to time; in this case all nodes from nealibration will be used as calibration points. Alter- natively, a list with two elements: the first is a character vector with node names from phy to calibrate; the second is a numeric vector with the corresponding ages to use as calibrations.	
missing_taxa	A tree, a data frame or a vector enlisting all missing taxa you want to include.	
	A tree Either as a phylo object or as a newick character string. It contains all taxa that you want at the end, both missing and non missing. This tree will be used as a hard constraint.	
	A data.frame It contains two columns named "taxon" and "clade". The first one contains a character vector of missing taxon names. The second one contains a character or numeric vector of nodes from a constraint tree to which each taxon will be assigned.	
	A character vector It contains the names of the missing taxa. They will be added at random to the constraint tree.	
age_distribution		
	A character string specifying the type of calibration. Only "fixed" and "uniform" are implemented for now.	
	fixed The age given in ncalibration will be used as fixed age.	

lognormal The age given in nealibration will be used as mean age. The stan-
dard deviation can be provided. # still need to add this option. By default,
a 95 CI sd is used.
uniform The age given in ncalibration will be used as mean age. Where min_age
$= 0.9 *$ mean age, and max_age $= 1.1 *$ mean age.
root_calibration
Used to set a calibration at the root or not. Default to FALSE. Only relevant if ncalibration is specified.
<pre>mrbayes_output_file</pre>
A character vector specifying the name of mrBayes run file and outputs (can specify directory too).

Value

A MrBayes block run file in nexus format.

<pre>make_mrbayes_tree</pre>	Take a constraint tree and use mrBayes to get node ages and branch
	lengths given a set of node calibrations without any data.

Description

Take a constraint tree and use mrBayes to get node ages and branch lengths given a set of node calibrations without any data.

Usage

```
make_mrbayes_tree(
   constraint = NULL,
   taxa = NULL,
   ncalibration = NULL,
   missing_taxa = NULL,
   age_distribution = "fixed",
   root_calibration = FALSE,
   mrbayes_output_file = "mrbayes_run.nexus"
)
```

Arguments

constraint	The constraint tree: a phylo object or a newick character string, with or without branch lengths.
taxa	A character vector with taxon names to be maintained in tree
ncalibration	The node calibrations: a phylo object with branch lengths proportional to time; in this case all nodes from ncalibration will be used as calibration points. Alter- natively, a list with two elements: the first is a character vector with node names from phy to calibrate; the second is a numeric vector with the corresponding ages to use as calibrations.

missing_taxa A tree, a data frame or a vector enlisting all missing taxa you want to include.

- A tree Either as a phylo object or as a newick character string. It contains all taxa that you want at the end, both missing and non missing. This tree will be used as a hard constraint.
- A data.frame It contains two columns named "taxon" and "clade". The first one contains a character vector of missing taxon names. The second one contains a character or numeric vector of nodes from a constraint tree to which each taxon will be assigned.
- A character vector It contains the names of the missing taxa. They will be added at random to the constraint tree.

age_distribution

A character string specifying the type of calibration. Only "fixed" and "uniform" are implemented for now.

fixed The age given in nealibration will be used as fixed age.

- **lognormal** The age given in ncalibration will be used as mean age. The standard deviation can be provided. # still need to add this option. By default, a 95 CI sd is used.
- **uniform** The age given in ncalibration will be used as mean age. Where min_age = 0.9 * mean age, and max_age = 1.1 * mean age.

root_calibration

Used to set a calibration at the root or not. Default to FALSE. Only relevant if ncalibration is specified.

mrbayes_output_file

A character vector specifying the name of mrBayes run file and outputs (can specify directory too).

Value

A phylo object with branch lengths proportional to time. It saves all mrBayes outputs in the working directory.

make_otol_associations

Associate Open Tree of Life authors with studies

Description

Associate Open Tree of Life authors with studies

Usage

```
make_otol_associations()
```

Value

data.frame with author last name, author first and other names, and comma delimited URLs for OToL studies

make_overlap_table Create an overlap table

Description

Create an overlap table

Usage

```
make_overlap_table(results_table)
```

Arguments

results_table An "author.results" or "curator.results" data.frame

Value

A data.frame with information on curators and what clades they've worked on

make_sdm	Make a Super Distance Matrix (SDM) from a list of good matrices
	<pre>obtained with get_goodmatrices()</pre>

Description

Make a Super Distance Matrix (SDM) from a list of good matrices obtained with get_goodmatrices()

Usage

make_sdm(unpadded.matrices, weighting = "flat")

Arguments

unpadded.matr	ices
	A list of patristic matrices, a datelifeResult object.
weighting	A character vector indicating how much weight to give to each tree in input during the SDM analysis. Options are:
	weighting = "flat" All trees have equal weighting.
	weighting = "taxa" Weight is proportional to number of taxa.
	weighting = "inverse" Weight is proportional to 1 / number of taxa.
	Defaults to weighting = "flat".

Value

A matrix.

make_treebase_associations

Associate TreeBase authors with studies

Description

Associate TreeBase authors with studies

Usage

make_treebase_associations()

Value

data.frame with author last name, author first and other names, and comma delimited URLs for TreeBase studies

make_treebase_cache Create a cache from TreeBase

Description

Create a cache from TreeBase

Usage

```
make_treebase_cache(outputfile = "treebasecache.RData")
```

Arguments

outputfile Path including file name

Value

List containing author and curator results

map_nodes_ott

Description

Add Open Tree of Life Taxonomy to tree nodes.

Usage

```
map_nodes_ott(tree)
```

Arguments

tree A tree either as a newick character string or as a phylo object.

Value

A phylo object with "nodelabels".

Examples

Not run: # This is a flag for package development. You are welcome to run the example.

```
# Load the Open Tree chronograms database cached in datelife:
utils::data(opentree_chronograms)
```

Get the small chronograms (i.e., chronograms with less that ten tips) to generate a pretty plot: small <- opentree_chronograms\$trees[unlist(sapply(opentree_chronograms\$trees, ape::Ntip)) < 10]</pre>

```
# Now, map the Open Tree taxonomy to the nodes of the first tree
phy <- map_nodes_ott(tree = small[[1]])
# and plot it:
# plot_phylo_all(phy)
library(ape)
plot(phy)
nodelabels(phy$node.label)
```

End(Not run) #end dontrun

match_all_calibrations

Match calibrations to nodes of a given tree

Description

match_all_calibrations searches a given tree for the most recent common ancestor (mrca) of all taxon name pairs in a datelifeCalibration. It uses phytools::findMRCA().

Usage

match_all_calibrations(phy, calibrations)

Arguments

phy	A phylo object.
calibrations	A calibrations object, an output of extract_calibrations_phylo().

Details

The function takes pairs of taxon names in a secondary calibrations data frame, and looks for them in the vector of tip labels of the tree. If both are present, then it gets the node that represents the most recent common ancestor (mrca) for that pair of taxa in the tree. Nodes of input phy can be named or not.

Value

A list of two elements:

- **phy** A phylo object with nodes renamed with tree_add_nodelabels().
- matched_calibrations A matchedCalibrations object, which is the input calibrations object with two additional columns storing results from the mrca search with phytools::findMRCA(): \$mrca_node_number and \$mrca_node_name.

matrices_to_table Go from a list of patristic distance matrix to a table of node ages

Description

Go from a list of patristic distance matrix to a table of node ages

Usage

matrices_to_table(matrices)

matrices A names list of patristic distance matrices. Names correspond to the study reference.

Value

A single data.frame of "taxonA", "taxonB", and "age".

matrix_to_table Go from a patristic distance matrix to a node ages table

Description

Go from a patristic distance matrix to a node ages table

Usage

```
matrix_to_table(matrix, reference)
```

Arguments

matrix	A patristic distance matrix.
reference	A character vector with the study reference from where the ages come from.

Value

A data.frame of "taxonA", "taxonB", and "age".

message_multiphylo *Message for a* multiPhylo *input*

Description

Message for a multiPhylo input

Usage

```
message_multiphylo()
```

Value

A relevant message as a character string.

missing_taxa_check Checks that missing_taxa argument is ok to be used by make_mrbayes_runfile inside tree_add_dates functions.

Description

Checks that missing_taxa argument is ok to be used by make_mrbayes_runfile inside tree_add_dates functions.

Usage

```
missing_taxa_check(missing_taxa = NULL, dated_tree = NULL)
```

Arguments

missing_taxa	A tree, a data frame or a vector enlisting all missing taxa you want to include.
	A tree Either as a phylo object or as a newick character string. It contains all taxa that you want at the end, both missing and non missing. This tree will be used as a hard constraint.
	A data.frame It contains two columns named "taxon" and "clade". The first one contains a character vector of missing taxon names. The second one contains a character or numeric vector of nodes from a constraint tree to which each taxon will be assigned.
	A character vector It contains the names of the missing taxa. They will be added at random to the constraint tree.
dated_tree	a tree (newick or phylo) with branch lengths proportional to absolute time

Value

A phylo object, a newick character string or a dataframe with taxonomic assignations

mrca_calibrations	Identify nodes of a tree topology that are most recent common ancestor
	(mrca) of taxon pairs from a calibrations object

Description

mrca_calibrations get nodes of a tree topology given in phy that correspond to the most recent common ancestor (mrca) of taxon pairs given in calibrations. It uses phytools::findMRCA() to get mrca nodes.

Usage

mrca_calibrations(phy, calibrations)

phy	A phylo object.
calibrations	A calibrations object, an output of extract_calibrations_phylo().

Details

The function takes pairs of taxon names in a calibrations data frame, and looks for them in the vector of tip labels of the tree. If both are present, then it gets the node that represents the most recent common ancestor (mrca) for that pair of taxa in the tree. Nodes of input phy can be named or not. They will be renamed.

Value

A list of two elements:

- **matched_phy** A phylo object with nodes renamed to match results of the mrca search. Nodes are renamed using tree_add_nodelabels().
- matched_calibrations A matchedCalibrations object, which is the input calibrations object with two additional columns storing results from the mrca search with phytools::findMRCA(): \$mrca_node_number and \$mrca_node_name.

opentree_chronograms Chronogram database

Description

Now storing >200 chronograms from Open Tree of Life

Usage

opentree_chronograms

Format

A list of four elements, containing data from Open Tree of Life chronograms

- **authors** A list of lists of author names of the original studies that published chronograms in the Open Tree of Life database.
- **curators** A list of lists of curator names that uploaded chronograms to the Open Tree of Life database.

studies A list of study identifiers.

trees A multiPhylo object storing the chronograms from Open Tree of Life database.

- update A character vector indicating the time when the database object was last updated.
- **version** A character vector indicating the datelife utils::packageVersion() when the database was last updated.

Details

Generated with devtools::install_github("ropensci/rotl", ref = devtools::github_pull("137")) remotes::install_github("ROpenS opentree_chronograms <- get_opentree_chronograms() opentree_chronograms\$update <- Sys.time() opentree_chronograms\$version <- '2022.01.28' usethis::use_data(opentree_chronograms, overwrite = T, compress = "xz") and updated with update_datelife_cache()

Source

http://opentreeoflife.org

patristic_matrix_array_congruify

patristic_matrix_array_congruify is used for patristic_matrix_array_subset_both and patristic_matrix_array_congruify.

Description

patristic_matrix_array_congruify is used for patristic_matrix_array_subset_both and patristic_matrix_array_congruify.

Usage

```
patristic_matrix_array_congruify(
   patristic_matrix_array,
   taxa,
   phy = NULL,
   dating_method = "PATHd8"
)
```

Arguments

patristic_matrix_array		
	A patristic matrix array, rownames and colnames must be taxa.	
taxa	Vector of taxon names to get a subset for.	
phy	A user tree to congruify as phylo object (ape).	
dating_method	The method used for tree dating.	

Value

A patristic matrix with ages for the target taxa.

patristic_matrix_array_phylo_congruify

Congruify a patristic matrix array from a given phylo object.

Description

Congruify a patristic matrix array from a given phylo object.

Usage

```
patristic_matrix_array_phylo_congruify(
   patristic_matrix,
   target_tree,
   dating_method = "PATHd8",
   attempt_fix = TRUE
)
```

Arguments

patristic_matrix

	A patristic matrix, rownames and colnames must be taxa.
target_tree	A phylo object. Use this in case you want a specific backbone for the output tree.
dating_method	The method used for tree dating.
attempt_fix	Default to TRUE. If congruification results in NA branch lengths, it will attempt to fix them.

Value

A matrix.

Description

Split a patristic matrix array Used inside: patristic_matrix_array_congruify

Usage

patristic_matrix_array_split(patristic_matrix_array)

patristic_matrix_array_subset

Arguments

patristic_matrix_array

A patristic matrix array, rownames and colnames must be taxa.

Value

A patristic matrix 3d array.

patristic_matrix_array_subset
 Subset a patristic matrix array

Description

Subset a patristic matrix array

Usage

```
patristic_matrix_array_subset(patristic_matrix_array, taxa, phy4 = NULL)
```

Arguments

patristic_matrix_array		
	A patristic matrix array, rownames and colnames must be taxa.	
taxa	Vector of taxon names to get a subset for.	
phy4	A user tree to congruify in phylo4 format (phylobase).	

Value

A list with a patristic matrix array and a \$problem if any.

 ${\tt patristic_matrix_array_subset_both}$

Are all desired taxa in the patristic matrix array?

Description

patristic_matrix_array_subset_both is used inside get_subset_array_dispatch().

Usage

```
patristic_matrix_array_subset_both(
   patristic_matrix_array,
   taxa,
   phy = NULL,
   phy4 = NULL,
   dating_method = "PATHd8"
)
```

Arguments

patristic_matrix_array		
	A patristic matrix array, rownames and colnames must be taxa.	
taxa	Vector of taxon names to get a subset for.	
phy	A user tree to congruify as phylo object (ape).	
phy4	A user tree to congruify in phylo4 format (phylobase).	
dating_method	The method used for tree dating.	

Value

A patristic matrix with ages for the target taxa.

Description

patristic_matrix_list_to_array us ised inside summarize_datelife_result(), patristic_matrix_array_congrui

Usage

```
patristic_matrix_list_to_array(patristic_matrix_list, pad = TRUE)
```

Arguments

patristic_matr:	ix_list
	List of patristic matrices
pad	If TRUE, pad missing entries

Value

A 3d array of patristic matrices

Description

Get time of MRCA from patristic matrix. Used in datelife_result_MRCA().

Usage

```
patristic_matrix_MRCA(patristic_matrix, na_rm = TRUE)
```

Arguments

patristic_matr	ix
	A patristic matrix (aka a datelifeResult object of length 1)
na_rm	If TRUE, it drops rows containing NAs from the datelifeResult patristic matrix;
	if FALSE, it returns NA where there are missing entries.

Value

The depth of the MRCA as a numeric vector.

```
patristic_matrix_name_order_test
```

Test the name order of a patristic matrix so that row and column labels are in alphabetical order.

Description

patristic_matrix_name_order_test is only used in patristic_matrix_list_to_array().

Usage

```
patristic_matrix_name_order_test(
   patristic_matrix,
   standard.rownames,
   standard.colnames
)
```

Arguments

```
patristic_matrix
A patristic matrix, rownames and colnames must be taxa.
standard.rownames
A character vector of row names.
standard.colnames
A character vector of column names.
```

Value

Boolean.

```
patristic_matrix_name_reorder
```

Reorder a matrix so that row and column labels are in alphabetical order.

Description

patristic_matrix_name_reorder is only used in: patristic_matrix_pad().

Usage

patristic_matrix_name_reorder(patristic_matrix)

Arguments

patristic_matrix

A patristic matrix, rownames and colnames must be taxa.

Value

A patristic matrix with row and column names for taxa in alphabetical order.

patristic_matrix_pad Fill in empty cells in a patristic matrix for missing taxa.

Description

```
Used in: patristic_matrix_list_to_array().
```

Usage

```
patristic_matrix_pad(patristic_matrix, all_taxa)
```

Arguments

patristic_matrix

A patristic matrix, rownames and colnames must be taxa.

all_taxa A vector of names of all taxa you want, including ones not in the patristic matrix.

Value

A patristic matrix, with NA for entries between taxa where at least one was not in the original patristic matrix.

Description

patristic_matrix_taxa_all_matching is used inside: results_list_process().

Usage

```
patristic_matrix_taxa_all_matching(patristic_matrix, taxa)
```

Arguments

patristic_matrix		
	A patristic matrix, rownames and colnames must be taxa.	
taxa	Vector of taxon names to get a subset for.	

Value

A Boolean.

```
patristic_matrix_to_newick
```

Convert patristic matrix to a newick string. Used inside: summarize_datelife_result.

Description

Convert patristic matrix to a newick string. Used inside: summarize_datelife_result.

Usage

```
patristic_matrix_to_newick(patristic_matrix)
```

Arguments

```
patristic_matrix
```

A patristic matrix

Value

A newick string

```
patristic_matrix_to_phylo
```

Convert a patristic matrix to a phylo object.

Description

Function patristic_matrix_to_phylo is used inside summarize_datelife_result().

Usage

```
patristic_matrix_to_phylo(
   patristic_matrix,
   clustering_method = "nj",
   fix_negative_brlen = TRUE,
   fixing_method = 0,
   ultrametric = TRUE,
   variance_matrix = NULL
)
```

Arguments

patristic_matrix A patristic matrix clustering_method A character vector indicating the method to construct the tree. Options are: **nj** Neighbor-Joining method applied with ape::nj(). upgma Unweighted Pair Group Method with Arithmetic Mean method applied with phangorn::upgma(). **bionj** An improved version of the Neighbor-Joining method applied with ape::bionj(). triangle Triangles method applied with ape::triangMtd() **mvr** Minimum Variance Reduction method applied with ape::mvr(). fix_negative_brlen Boolean indicating whether to fix negative branch lengths in resulting tree or not. Default to TRUE. fixing_method A character vector specifying the method to fix branch lengths: "bladj", "mrbayes" or a number to be assigned to all branches meeting fixing_criterion Boolean indicating whether to force ultrametric or not. ultrametric variance_matrix A variance matrix from a datelifeResult object, usually an output from datelife_result_variance_ Only used if clustering_method = "mvr".

Details

We might add the option to insert a function as clustering_method in the future. Before, we had hard-coded the function to try Neighbor-Joining (NJ) first; if it errors, it will try UPGMA. Now, it uses NJ for a "phylo_all" summary, and we are using our own algorithm to get a tree from a summary matrix.

Value

A rooted phylo object.

```
patristic_matrix_unpad
```

Function to remove missing taxa from a datelifeResult object.

Description

Used in datelife_result_sdm_phylo().

Usage

patristic_matrix_unpad(patristic_matrix)

Arguments

patristic_matrix

A patristic matrix with row and column names for taxa

Value

patristic_matrix for all_taxa

phylo_check	Checks if phy is a phylo object and/or a chronogram.	
-------------	------------------------------------------------------	--

Description

Checks if phy is a phylo object and/or a chronogram.

Usage

phylo_check(phy = NULL, brlen = FALSE, dated = FALSE)

Arguments

phy	A phylo object.
brlen	Boolean. If TRUE it checks if phylo object has branch lengths.
dated	Boolean. If TRUE it checks if phylo object is ultrametric.

Value

Nothing

phylo_congruify

Description

Congruify a reference tree and a target tree given as phylo objects.

Usage

```
phylo_congruify(
  reference_tree,
  target_tree,
  dating_method = "PATHd8",
  attempt_fix = TRUE
)
```

Arguments

reference_tree A phylo object.

target_tree	A phylo object. Use this in case you want a specific backbone for the output tree.
dating_method	The method used for tree dating.
attempt_fix	Default to TRUE. If congruification results in NA branch lengths, it will attempt to fix them.

Value

A matrix.

phylo_generate_uncertainty Generate uncertainty in branch lengths using a lognormal.

Description

Generate uncertainty in branch lengths using a lognormal.

Usage

```
phylo_generate_uncertainty(
   phy,
   size = 100,
   uncertainty_method = "other",
   age_distribution = "uniform",
   age_sd = NULL,
   age_var = 0.1,
   age_scale = 0,
   alpha = 0.025,
   rescale = TRUE
)
```

Arguments

phy	A phylo object.
size	A numeric vector indicating the number of samples to be generated.
uncertainty_met	thod
	A character vector specifying the method to generate uncertainty. mrbayes is default.
age_distributio	on
	A character string specifying the type of calibration. Only "fixed" and "uniform" are implemented for now.
	fixed The age given in nealibration will be used as fixed age.
	lognormal The age given in ncalibration will be used as mean age. The stan- dard deviation can be provided. # still need to add this option. By default, a 95 CI sd is used.
	uniform The age given in ncalibration will be used as mean age. Where min_age = $0.9 *$ mean age, and max_age = $1.1 *$ mean age.
age_sd	The standard deviation around the age to use for generating the uncertainty. If not a numeric value, var will be used to calculate it.
age_var	The variance to calculate age_sd and generate uncertainty.
age_scale	How to scale sd by the depth of the node. If 0, same sd for all. If not, older nodes have more uncertainty
alpha	The significance level on uncertainty to generate. By default 0.025
rescale	Boolean. If true, observed age will be rescaled each round.

Details

If you want to change the size of sampled trees you do not need to run mrbayes again. Just use sample_trees("mrbayes_trees_file_directory", size = new_size) and you will get a multiPhylo object with a new tree sample.

Value

A phylo or multiPhylo object with the same topology as phy but different branch lengths

Examples

```
## Not run:
# Generate uncertainty over feline species SDM chronogram.
# Load the data:
data(felid_sdm)
# By default, generates a sample of 100 trees with var = 0.1:
unc <- phylo_generate_uncertainty(felid_sdm$phy)</pre>
length(unc)
# Make an LTT plot:
max_age <- max(sapply(unc, ape::branching.times))</pre>
ape::ltt.plot(phy = unc[[1]], xlim = c(-max_age, 0), col = "#cce5ff50")
for (i in 2:100) {
  ape::ltt.lines(phy = unc[[i]], col = "#cce5ff50")
}
ape::ltt.lines(felid_sdm$phy, col = "red")
title(c("fake uncertainty", "in Felidae SDM chronogram"))
## End(Not run) # end dontrun
```

```
phylo_get_node_numbers
```

Gets node numbers from any phylogeny

Description

Gets node numbers from any phylogeny

Usage

```
phylo_get_node_numbers(phy)
```

Arguments

phy A phylo object.

Value

A numeric vector with node numbers

phylo_get_subset_array

Get a subset array from a phylo object

Description

Get a subset array from a phylo object

Usage

```
phylo_get_subset_array(
   reference_tree,
   taxa,
   phy4 = NULL,
   dating_method = "PATHd8"
)
```

Arguments

reference_tree	A phylo object.
taxa	Vector of taxon names to get a subset for.
phy4	A user tree to congruify in phylo4 format (phylobase).
dating_method	The method used for tree dating.

Value

A list with a patristic matrix array and a \$problem if any.

phylo_get_subset_array_congruify Get a congruified subset array from a phylo object

Description

Get a congruified subset array from a phylo object

Usage

```
phylo_get_subset_array_congruify(
  reference_tree,
  taxa,
  phy = NULL,
  dating_method = "PATHd8"
)
```

Arguments

reference_tree	A phylo object.
taxa	Vector of taxon names to get a subset for.
phy	A user tree to congruify as phylo object (ape).
dating_method	The method used for tree dating.

Value

A list with a patristic matrix array and a \$problem if any.

phylo_has_brlen Check if a tree has branch lengths

Description

Check if a tree has branch lengths

Usage

phylo_has_brlen(phy)

Arguments

phy A phylo object.

Value

A TRUE or FALSE

phylo_prune_missing_taxa

Prune missing taxa from a phylo object Used inside phylo_get_subset_array and phylo_get_subset_array_congruify.

Description

Prune missing taxa from a phylo object Used inside phylo_get_subset_array and phylo_get_subset_array_congruify.

Usage

phylo_prune_missing_taxa(phy, taxa)

phylo_subset_both

Arguments

phy	A user tree to congruify as phylo object (ape).
taxa	Vector of taxon names to get a subset for.

Value

A phylo object.

phylo_subset_both *Subset a reference and a target tree given as* phylo *objects.*

Description

Subset a reference and a target tree given as phylo objects.

Usage

```
phylo_subset_both(
  reference_tree,
  taxa,
  phy = NULL,
  phy4 = NULL,
  dating_method = "PATHd8"
)
```

Arguments

reference_tree	A phylo object.
taxa	Vector of taxon names to get a subset for.
phy	A user tree to congruify as phylo object (ape).
phy4	A user tree to congruify in phylo4 format (phylobase).
dating_method	The method used for tree dating.

Value

A list with a patristic matrix array and a \$problem if any.

phylo_tiplabel_space_to_underscore

Convert spaces to underscores in trees.

Description

phylo_tiplabel_space_to_underscore is used in: make_mrbayes_runfile(), tree_get_singleton_outgroup(), congruify_and_check(), patristic_matrix_array_phylo_congruify().

Usage

phylo_tiplabel_space_to_underscore(phy)

Arguments

phy A phylo object.

Value

A phylo object.

phylo_tiplabel_underscore_to_space Convert underscores to spaces in trees.

Description

phylo_tiplabel_underscore_to_space is used inside patristic_matrix_array_phylo_congruify(), congruify_and_check().

Usage

phylo_tiplabel_underscore_to_space(phy)

Arguments

phy A phylo object.

Value

A phylo object.

phylo_to_patristic_matrix

Get a patristic matrix from a phylo *object.*

Description

Get a patristic matrix from a phylo object.

Usage

```
phylo_to_patristic_matrix(phy, test = TRUE, tol = 0.01, option = 2)
```

Arguments

phy	A phylo object.
test	Default to TRUE. Whether to test if phy has branch lengths and is ultrametric or
	not.
tol	branching time in reference above which secondary constraints will be applied
	to target
option	an integer (1 or 2; see details).

Value

A patristic matrix.

pick_grove	Pick a grove in the case of multiple groves in a set of trees.
------------	----------------------------------------------------------------

Description

Pick a grove in the case of multiple groves in a set of trees.

Usage

```
pick_grove(grove_list, criterion = "taxa", datelife_result)
```

Arguments

grove_list	A list of vectors of tree indices. Each element is a grove.
criterion	Defaults to criterion = "taxa". Used for chronogram summarizing, i.e., ob-
	taining a single summary chronogram from a group of input chronograms. For
	summarizing approaches that return a single summary tree from a group of phy-
	logenetic trees, it is necessary that the latter form a grove, roughly, a sufficiently
	overlapping set of taxa between trees, see Ané et al. (2009) doi:10.1007/s00026-
	0090017x. In rare cases, a group of trees can have multiple groves. This ar-
	gument indicates whether to get the grove with the most trees (criterion =
	"trees") or the most taxa (criterion = "taxa").

```
datelife_result
A datelifeResult object. Only needed for criterion = "taxa".
```

Value

A numeric vector of the elements of the picked grove.

plant_bold_otol_tree Some plants chronogram

Description

Some plants chronogram

Usage

plant_bold_otol_tree

Format

A phylo object with 6 tips and 5 internal nodes

edge Integer vector with edge (branch) numberstip.label Character vector with species names of plantsNnode Integer vector with the number of nodesnode.label Character vector with node namesedge.length Numeric vector with edge (branch) lengths

Details

Generated with make_bold_otol_tree(input = "((Zea mays,Oryza sativa),((Arabidopsis thaliana,(Glycine max,Medicago sativa)),Solanum lycopersicum)Pentapetalae);") usethis::use_data(plant_bold_otol_tree)

Author(s)

 $Luna \ L. \ Sanchez-Reyes < \texttt{lsanche7@utk.edu} >$

Brian O'Meara <bomeara@utk.edu>

Source

http://opentreeoflife.org
http://www.boldsystems.org

problems

Description

Problematic chronograms from Open Tree of Life.

Usage

problems

Format

A list of trees with unmapped taxa

Details

Before we developed tools to clean and map tip labels for our cached trees we found some trees that were stored with unmapped tip labels we extracted them and saved them to be used for testing functions. Generated with problems <- opentree_chronograms\$trees[sapply(sapply(opentree_chronograms\$trees, "[", "tip.label"), function(x) any(grepl("not.mapped", x)))] usethis::use_data(problems) opentree_chronograms object from commit https://github.com/phylotastic/datelife/tree/be894448f6fc437241cd0916fab45e84ac3e09c6

[", "tip.label"), function(x) any(grepl("not.mapped", x)))]: R:%22,%20%22tip.label%22),%20function(x)%20any(grepl(%22

Source

http://opentreeoflife.org

recover_mrcaott Get an mrcaott tag from an OpenTree induced synthetic tree and get its name and ott id

Description

Get an mrcaott tag from an OpenTree induced synthetic tree and get its name and ott id

Usage

```
recover_mrcaott(tag)
```

Arguments

tag A character vector with the mrca tag

Value

A numeric vector with ott id from original taxon named with the corresponding ott name

relevant_curators_tabulate

Return the relevant curators for a set of studies.

Description

Return the relevant curators for a set of studies.

Usage

```
relevant_curators_tabulate(results.index, cache = "opentree_chronograms")
```

Arguments

results.index	A vector from datelife_result_study_index() with the indices of the relevant studies.
cache	The cached chronogram database.

Value

A vector with counts of each curator, with names equal to curator names.

results_list_process Take results_list and process it.

Description

results_list_process is used inside: get_datelife_result()

Usage

```
results_list_process(results_list, taxa = NULL, partial = FALSE)
```

Arguments

results_list	A list returned from using get_subset_array_dispatch() on open tree_chronograms trees
taxa	Vector of taxon names to get a subset for.
partial	If TRUE, return matrices that have only partial matches.

Value

A list with the patristic.matrices that are not NA.

Description

Core function to generate results

Usage

```
run(
    input = c("Rhea americana", "Pterocnemia pennata", "Struthio camelus"),
    format = "citations",
    partial = "yes",
    plot.width = 600,
    plot.height = 600,
    use_tnrs = "no",
    opentree_chronograms = NULL
)
```

Arguments

input	A newick string or vector of taxa
format	The output format
partial	How to deal with trees that have a subset of taxa in the query
plot.width	Width in pixels for output plot
plot.height	Height in pixels for output plot
use_tnrs	Whether to use OpenTree's TNRS for the input
opentree_chronograms	
	The list of lists containing the input trees and other info

Value

results in the desired format

run_mrbayes Runs MrBayes from R

Description

Runs MrBayes from R

Usage

run_mrbayes(mrbayes_output_file = NULL)

run

run

Arguments

mrbayes_output_file

A character vector specifying the name of mrBayes run file and outputs (can specify directory too).

Value

A phylo object with the consensus tree. MrBayes output files are stored in the working directory.

sample_trees	Sample trees from a file containing multiple trees.	Usually from a
	bayesian analysis output trees file.	

Description

Sample trees from a file containing multiple trees. Usually from a bayesian analysis output trees file.

Usage

```
sample_trees(trees_file, trees_object = NULL, burnin = 0.25, size = 100)
```

Arguments

trees_file	A character vector indicating the name and directory of file with trees to sample.
trees_object	An R object containing a list of trees already read into R from a tree file from a bayesian analysis output.
burnin	A numeric vector indicating the burnin fraction. It should be a number between 0 and 1. Default to 0.25
size	A numeric vector indicating the number of samples to be generated.

Value

A multiPhylo object with a random sample of trees.

some_ants_datelife_result

datelifeResult object of some ants

Description

datelifeResult object of some ants

Usage

some_ants_datelife_result

Format

A list of one element, containing a named patristic matrix

Details

Generated with: some_ants_input <- "(Aulacopone_relicta,(((Myrmecia_gulosa,(Aneuretus_simoni,Dolichoderus_mariae)), some_ants_datelife_query <- make_datelife_query(input = some_ants_input) some_ants_datelife_result <- get_datelife_result(input = some_ants_datelife_query) usethis::use_data(some_ants_datelife_result)

Source

http://opentreeoflife.org

subset2_search	A list with datelifeQuery and datelifeResult objects from a search of
	taxon names from subset2_taxa

Description

A list with datelifeQuery and datelifeResult objects from a search of taxon names from subset2_taxa

Usage

subset2_search

Format

A list with two named elements. datelifeResult object with 24 patristic matrices

datelife_query A datelifeQuery object using names_subset 2 as input.

datelife_result A datelifeResult object resulting from a search of names in datelifeQuery

Details

Generated with: datelife_query <- make_datelife_query(subset2_taxa) datelife_result <- get_datelife_result(datelife_query) subset2_search <- list(query = datelife_query, result = datelife_result) usethis::use_data(subset2_search, overwrite = TRUE)

subset2_taxa

Long list of >2.7k virus, bacteria, plant and animal taxon names

Description

Long list of >2.7k virus, bacteria, plant and animal taxon names

Usage

subset2_taxa

Format

A character vector of length 2778

Details

Generated with: subset2_taxa <- rphylotastic::url_get_scientific_names("https://github.com/phylotastic/rphylotastic/blob/mausethis::use_data(subset2_taxa)

Source

https://github.com/phylotastic/rphylotastic/tree/master/tests/testthat

Description

Function summarize_congruifiedCalibrations returns a table of summary statistics for each node in congruified_calibrations argument.

Usage

summarize_congruifiedCalibrations(congruified_calibrations, age_column)

Arguments

congruified_calibrations		
	$A \ congruified {\tt Calibrations} \ object, \ output \ of \ congruify_and_mrca_multiPhylo().$	
age_column	A character string indicating the name of the column to be summarized.	

Value

A data.frame of summarized ages.

summarize_datelife_result

Summarize a datelifeResult object.

Description

Get different types of summaries from a datelifeResult object, an output from get_datelife_result(). This allows rapid processing of data. If you need a list of chronograms from your datelifeResult object, this is the function you are looking for.

Usage

```
summarize_datelife_result(
  datelife_result = NULL,
  datelife_query = NULL,
  summary_format = "phylo_all",
  na_rm = TRUE,
  summary_print = c("citations", "taxa"),
  taxon_summary = c("none", "summary", "matrix"),
  criterion = "taxa"
)
```

Arguments

datelife_result	
	A datelifeResult object, usually an output of get_datelife_result().
datelife_query	A datelifeQuery object, usually an output of make_datelife_query().
summary_format A character vector of length one, indicating the output format for result. DateLife search. Available output formats are:	
	"citations" A character vector of references where chronograms with some or all of the target taxa are published (source chronograms).
	"mrca" A named numeric vector of most recent common ancestor (mrca) ages of target taxa defined in input, obtained from the source chronograms. Names of mrca vector are equal to citations.
	"newick_all" A named character vector of newick strings corresponding to tar- get chronograms derived from source chronograms. Names of newick_all vector are equal to citations.

- "newick_sdm" Only if multiple source chronograms are available. A character vector with a single newick string corresponding to a target chronogram obtained with SDM supertree method (Criscuolo et al. 2006).
- "newick_median" Only if multiple source chronograms are available. A character vector with a single newick string corresponding to a target chronogram from the median of all source chronograms.
- "phylo_sdm" Only if multiple source chronograms are available. A phylo object with a single target chronogram obtained with SDM supertree method (Criscuolo et al. 2006).
- "**phylo_median**" Only if multiple source chronograms are available. A phylo object with a single target chronogram obtained from source chronograms with median method.
- "**phylo_all**" A named list of phylo objects corresponding to each target chronogram obtained from available source chronograms. Names of phylo_all list correspond to citations.
- "**phylo_biggest**" The chronogram with the most taxa. In the case of a tie, the chronogram with clade age closest to the median age of the equally large trees is returned.
- "html" A character vector with an html string that can be saved and then opened in any web browser. It contains a 4 column table with data on target taxa: mrca, number of taxa, citations of source chronogram and newick target chronogram.
- "data_frame" A 4 column data.frame with data on target taxa: mrca, number of taxa, citations of source chronograms and newick string.
- na_rm If TRUE, it drops rows containing NAs from the datelifeResult patristic matrix; if FALSE, it returns NA where there are missing entries.
- summary_print A character vector specifying the type of summary information to be printed to screen. Options are:

"citations" Prints references of chronograms where target taxa are found.

- "taxa" Prints a summary of the number of chronograms where each target taxon is found.
- "**none**" Nothing is printed to screen.

Defaults to c("citations", "taxa"), which displays both.

- taxon_summary A character vector specifying if data on target taxa missing in source chronograms should be added to the output as a "summary" or as a presence/absence "matrix". Default to "none", no information on taxon_summary added to the output.
- criterion Defaults to criterion = "taxa". Used for chronogram summarizing, i.e., obtaining a single summary chronogram from a group of input chronograms. For summarizing approaches that return a single summary tree from a group of phylogenetic trees, it is necessary that the latter form a grove, roughly, a sufficiently overlapping set of taxa between trees, see Ané et al. (2009) doi:10.1007/s00026-0090017x. In rare cases, a group of trees can have multiple groves. This argument indicates whether to get the grove with the most trees (criterion = "trees") or the most taxa (criterion = "taxa").

Value

The output is determined by the argument summary_format:

- If summary_format = "citations" The function returns a character vector of references.
- If summary_format = "mrca" The function returns a named numeric vector of most recent common ancestor (mrca) ages.
- If summary_format = "phylo_[all, sdm, median, or biggest]" The function returns output chronograms as phylo or multiPhylo objects.
- If summary_format = "html" or "data_frame" The function returns a 4 column table with data on mrca ages, number of taxa, references, and output chronograms as newick strings.

References

Ané, C., Eulenstein, O., Piaggio-Talice, R., & Sanderson, M. J. (2009). "Groves of phylogenetic trees". Annals of Combinatorics, 13(2), 139-167, doi:10.1007/s000260090017x.

summarize_fossil_range

Summarize taxon age from PBDB to just a single min and max age

Description

This uses the Paleobiology Database's API to gather information on the ages for all specimens of a taxon. It will also look for all descendants of the taxon. It fixes name misspellings if possible. It is basically a wrapper for get_fossil_range.

Usage

summarize_fossil_range(taxon, recent = FALSE, assume_recent_if_missing = TRUE)

Arguments

taxon	The scientific name of the taxon you want the range of occurrences of
recent	If TRUE, forces the minimum age to be zero
assume_recent_i	f_missing
	If TRUE, any taxon missing from pbdb is assumed to be recent

Value

a single row data.frame of max_ma and min_ma for the specimens, with rowname equal to taxon input

```
summarize_summary_matrix
```

Gets all ages per taxon pair from a distance matrix Internal function used in summary_matrix_to_phylo_all().

Description

Gets all ages per taxon pair from a distance matrix Internal function used in summary_matrix_to_phylo_all().

Usage

```
summarize_summary_matrix(summ_matrix)
```

Arguments

summ_matrix Any summary patristic distance matrix, such as the ones obtained with datelife_result_sdm_matrix()
or datelife_result_median_matrix().

Value

A data.frame of pairwise ages, with row number equal to the combinatory of column names (or row names), estimated as ncol(summ_matrix)^2 - sum(1:(ncol(summ_matrix)-1)).

summary.datelifeResult

Summarize a datelifeResult object.

Description

Summarize a datelifeResult object.

Usage

```
## S3 method for class 'datelifeResult'
summary(object, datelife_query, na_rm = TRUE, ...)
```

Arguments

object	An object of class datelifeResult, usually an output of get_datelife_result().
datelife_query	A datelifeQuery object, usually an output of make_datelife_query().
na_rm	Default to TRUE, whether to include partial matches or not.
	Further arguments passed to or from other methods.

Value

A named list of 11 elements:

- "citations" A character vector of references where chronograms with some or all of the target taxa are published (source chronograms).
- "mrca" A named numeric vector of most recent common ancestor (mrca) ages of target taxa defined in input, obtained from the source chronograms. Names of mrca vector are equal to citations.
- "newick_all" A named character vector of newick strings corresponding to target chronograms derived from source chronograms. Names of newick_all vector are equal to citations.
- "**newick_sdm**" Only if multiple source chronograms are available. A character vector with a single newick string corresponding to a target chronogram obtained with SDM supertree method (Criscuolo et al. 2006).
- "**newick_median**" Only if multiple source chronograms are available. A character vector with a single newick string corresponding to a target chronogram from the median of all source chronograms.
- "**phylo_sdm**" Only if multiple source chronograms are available. A phylo object with a single target chronogram obtained with SDM supertree method (Criscuolo et al. 2006).
- "phylo_median" Only if multiple source chronograms are available. A phylo object with a single target chronogram obtained from source chronograms with median method.
- "**phylo_all**" A named list of phylo objects corresponding to each target chronogram obtained from available source chronograms. Names of phylo_all list correspond to citations.
- "**phylo_biggest**" The chronogram with the most taxa. In the case of a tie, the chronogram with clade age closest to the median age of the equally large trees is returned.
- "html" A character vector with an html string that can be saved and then opened in any web browser. It contains a 4 column table with data on target taxa: mrca, number of taxa, citations of source chronogram and newick target chronogram.
- "data_frame" A 4 column data.frame with data on target taxa: mrca, number of taxa, citations of source chronograms and newick string.

summary.matchedCalibrations

SummarizeamatchedCalibrationsobjectsummary.matchedCalibrationsgetsthenodeagedistributionfrom amatchedCalibrationsobject.

Description

Summarize a matchedCalibrations object summary.matchedCalibrations gets the node age distribution from a matchedCalibrations object.

Usage

```
## S3 method for class 'matchedCalibrations'
summary(object, ...)
```

Arguments

object	$A \ {\tt matchedCalibrations\ object,\ usually\ an\ element\ of\ the\ output\ of\ {\tt match_all_calibrations\ }}).$
	Further arguments passed to or from other methods.

Details

Columns in_phy\$mrca_node_name and in_phy\$reference are factors.

Value

A summaryMatchedCalibrations object, which is a list of two matchedCalibrations objects:

- **not_in_phy** A data.frame subset of input matchedCalibrations object containing taxon name pairs that were not present in the given tree. NULL if all input taxon names are found in the given tree.
- **in_phy** A data.frame subset of input matchedCalibrations object containing all taxon name pairs that were present in the given tree.

summary_matrix_to_phylo

Go from a summary matrix to an ultrametric phylo *object.*

Description

Go from a summary matrix to an ultrametric phylo object.

Usage

```
summary_matrix_to_phylo(
   summ_matrix,
   datelife_query = NULL,
   target_tree = NULL,
   total_distance = TRUE,
   use = "mean",
   ...
)
```

Arguments

summ_matrix	Any summary patristic distance matrix, such as the ones obtained with datelife_result_sdm_matrix() or datelife_result_median_matrix().
datelife_query	A datelifeQuery object, usually an output of make_datelife_query().
target_tree	A phylo object. Use this in case you want a specific backbone for the output tree.

total_distance	Whether the input summ_matrix stores total age distance (from tip to tip) or distance from node to tip. Default to TRUE, divides the matrix in half, if FALSE it will take it as is.
use	A character vector indicating what type of age to use for summary tree. One of the following:
	"mean" It will use the mean() of the node ages in summ_matrix.
	"median" It uses the stats::median() age of node ages in summ_matrix.
	"min" It will use the min() age from node ages in summ_matrix.
	"max" Choose this if you wanna be conservative; it will use the max() age from node ages in summ_matrix.
	"midpoint" It will use the mean of minimum age and maximum age.
	Arguments passed on to <pre>summary_matrix_to_phylo_all</pre>

Details

It can take a regular patristic distance matrix, but there are simpler methods for that implemented in patristic_matrix_to_phylo().

Value

An ultrametric phylo object.

summary_matrix_to_phylo_all

Get minimum, median, mean, midpoint, and maximum summary chronograms from a summary matrix of a datelifeResult object.

Description

Get minimum, median, mean, midpoint, and maximum summary chronograms from a summary matrix of a datelifeResult object.

Usage

```
summary_matrix_to_phylo_all(
  summ_matrix,
  datelife_query = NULL,
  target_tree = NULL,
  total_distance = TRUE,
  ...
)
```

Arguments

summ_matrix	Any summary patristic distance matrix, such as the ones obtained with datelife_result_sdm_matrix() or datelife_result_median_matrix().
datelife_query	A datelifeQuery object, usually an output of make_datelife_query().
target_tree	A phylo object. Use this in case you want a specific backbone for the output tree.
total_distance	Whether the input summ_matrix stores total age distance (from tip to tip) or distance from node to tip. Default to TRUE, divides the matrix in half, if FALSE it will take it as is.
	Arguments passed on to get_otol_synthetic_tree
	otol_version Version of Open Tree of Life to use
	resolve Defaults to TRUE. Whether to resolve the tree at random or not.
	input Optional. A character vector of names or a datelifeQuery object.
	<pre>ott_ids If not NULL, it takes this argument and ignores input. A numeric vec- tor of ott ids obtained with rotl::taxonomy_taxon_info() or rotl::tnrs_match_names() or tnrs_match().</pre>

Details

With this function users can choose the minimum, mean or maximum ages from the summary matrix as calibration points to get a single summary chronogram. Users get all three summary chronograms in a multiPhylo object.

Value

A multiPhylo object of length 5. It contains min, mean, median, midpoint, and max summary chronograms.

summary_patristic_matrix_array
Summarize patristic matrix array (by default, median). Used inside:
 summarize_datelife_result.

Description

Summarize patristic matrix array (by default, median). Used inside: summarize_datelife_result.

Usage

```
summary_patristic_matrix_array(patristic_matrix_array, fn = stats::median)
```

Arguments

patristic_matrix_array	
	3D array of patristic matrices
fn	The function to use to summarize

threebirds_dr

Value

A 2d array with the median (or max, or mean, etc) of the input array

threebirds_dr	datelifeResult object of three birds "Rhea americana", "Pterocne-
	mia pennata", and "Struthio camelus"

Description

datelifeResult object of three birds "Rhea americana", "Pterocnemia pennata", and "Struthio camelus"

Usage

threebirds_dr

Format

A list of 9 named patristic matrix

Details

Generated with: threebirds_dr <- get_datelife_result(input=c("Rhea americana", "Pterocnemia pennata", "Struthio camelus"), partial = TRUE, use_tnrs = FALSE, approximate_match = TRUE, cache = "opentree_chronograms") use_data(threebirds_dr)

Source

http://opentreeoflife.org

tnrs_match	Taxon name resolution service (tnrs) applied to a vector of names by
	batches

Description

Taxon name resolution service (tnrs) applied to a vector of names by batches

Usage

```
tnrs_match(input, reference_taxonomy, tip, ...)
## Default S3 method:
tnrs_match(input, reference_taxonomy = "ott", ...)
## S3 method for class 'phylo'
tnrs_match(input, reference_taxonomy = "ott", tip = NULL, ...)
```

Arguments

input	A character vector of taxon names, or a phylo object with tip names, to be
	matched to taxonomy.
reference_taxor	nomy
	A character vector specifying the reference taxonomy to use for TNRS. Options are "ott", "ncbi", "gbif" or "irmng". The function defaults to "ott".
tip	A vector of mode numeric or character specifying the tips to match. If left empty all tips will be matched.
	Arguments passed on to rotl::tnrs_match_names
	<pre>context_name name of the taxonomic context to be searched (length-one char- acter vector or NULL). Must match (case sensitive) one of the values returned by tnrs_contexts. Default to "All life".</pre>
	do_approximate_matching A logical indicating whether or not to perform approximate string (a.k.a. "fuzzy") matching. Using FALSE will greatly improve speed. Default, however, is TRUE.
	ids A vector of ids to use for identifying names. These will be assigned to each name in the names array. If ids is provided, then ids and names must be identical in length.
	include_suppressed Ordinarily, some quasi-taxa, such as incertae sedis buck- ets and other non-OTUs, are suppressed from TNRS results. If this param- eter is true, these quasi-taxa are allowed as possible TNRS results.

Details

There is no limit to the number of names that can be queried and matched.

The output will preserve all elements from original input phylo object and will add

phy\$mapped A character vector indicating the state of mapping of phy\$tip.labels:

original Thrs matching was not attempted. Original labeling is preserved.

ott Matching was manually made by a curator in Open Tree of Life.

- **tnrs** Tnrs matching was attempted and successful with no approximate matching. Original label is replaced by the matched name.
- **approximated** Tnrs matching was attempted and successful but with approximate matching. Original labeling is preserved.
- unmatched Tnrs matching was attempted and unsuccessful. Original labeling is preserved.

phy\$original.tip.label A character vector preserving all original labels.

phy\$ott_ids A numeric vector with ott id numbers of matched tips. Unmatched and original tips will be NaN.

if tips are duplicated, thrs will only be run once (avoiding increases in function running time) but the result will be applied to all duplicated tip labels

Value

An object of class data frame or phylo, with the added class match_names.

NULL

NULL

treebase_cache

Examples

```
tnrs_match(input = c("Mus"))
tnrs_match(input = c("Mus", "Mus musculus"))
tnrs_match(input = c("Mus", "Echinus", "Hommo", "Mus"))
```

```
treebase_cache
```

Information on contributors, authors, study ids and clades from studies with chronograms in Open tree of Life

Description

Information on contributors, authors, study ids and clades from studies with chronograms in Open tree of Life

Usage

treebase_cache

Format

A list of five data sets

tb.author.pretty A dataframe with two elements: author names and number of studies in TreeBase authored by each

tb.author.results A dataframe with two elements: author names and study identifiers

Details

Generated with make_treebase_cache()

Source

TreeBASE database, no longer available online https://en.wikipedia.org/wiki/TreeBASE

tree_add_dates	Add missing taxa to a dated tree and fabricate node ages for these
	missing taxa.

Description

This function adds missing taxa to a chronogram given in dated_tree. It is still work in progress.

Usage

```
tree_add_dates(
    dated_tree = NULL,
    missing_taxa = NULL,
    dating_method = "mrbayes",
    adding_criterion = "random",
    mrbayes_output_file = "mrbayes_tree_add_dates.nexus"
)
```

Arguments

dated_tree	a tree (newick or phylo) with branch lengths proportional to absolute time
missing_taxa	A tree, a data frame or a vector enlisting all missing taxa you want to include.
	A tree Either as a phylo object or as a newick character string. It contains all taxa that you want at the end, both missing and non missing. This tree will be used as a hard constraint.
	A data.frame It contains two columns named "taxon" and "clade". The first one contains a character vector of missing taxon names. The second one contains a character or numeric vector of nodes from a constraint tree to which each taxon will be assigned.
	A character vector It contains the names of the missing taxa. They will be added at random to the constraint tree.
dating_method	The method used for tree dating, options are "mrbayes" and "bladj".
adding_criteri	on
	Only used when dating_method = "mrbayes". A character vector to specify how missing_taxa should be added to dated_tree. Choose one of:
	adding_method = "random" missing_taxa will be added at random to dated_tree.
	adding_method = ''taxonomy'' taxa will be added to dated_tree following a dataframe with taxonomic assignations given in missing_taxa argument. If no dataframe is given, OpenTree's reference taxonomy will be used.
	adding_method = "tree" taxa will be added to dated_tree following a tree given in missing_taxa argument. If no tree is given, OpenTree's synthetic tree will be used.
<pre>mrbayes_output</pre>	_file
	A character vector specifying the name of mrBayes run file and outputs (can specify directory too).
Value	

A phylo object.

tree_add_nodelabels Adds labels to nodes with no assigned label

Description

Adds labels to nodes with no assigned label

Usage

```
tree_add_nodelabels(tree = NULL, node_prefix = "n", node_index = "node_number")
```

Arguments

tree	A tree either as a newick character string or as a phylo object.
node_prefix	Character vector. If length 1, it will be used to name all nodes with no labels, fol- lowed by a number which can be the node_number or consecutive, as specified in node_index.
node_index	Character vector. Choose between "from_1" and "node_number" as numeric index for node labels. It will use consecutive numbers from 1 to total node number in the first case and phylo node numbers in the second case (i.e, from Ntip + 1).

Value

A phylo object

tree_add_outgroup	Function to add an outgroup to any phylogeny, in phylo or newick
	format

Description

Function to add an outgroup to any phylogeny, in phylo or newick format

Usage

```
tree_add_outgroup(tree = NULL, outgroup = "outgroup")
```

Arguments

tree	A tree either as a newick character string or as a phylo object.
outgroup	A character vector with the name of the outgroup. If it has length>1, only first element will be used.

Value

A phylo object with no root edge.

tree_check

Checks if a tree is a phylo class object otherwise it uses input_process. Additionally it can check if tree is a chronogram with phylo_check

Description

Checks if a tree is a phylo class object otherwise it uses input_process. Additionally it can check if tree is a chronogram with phylo_check

Usage

tree_check(tree = NULL, ...)

Arguments

tree	A tree either as a newick character string or as a phylo object.
	Arguments passed on to phylo_check
	brlen Boolean. If TRUE it checks if phylo object has branch lengths.
	dated Boolean. If TRUE it checks if phylo object is ultrametric.

Value

If tree is correctly formatted, it returns a phylo object.

<pre>tree_fix_brlen</pre>	Take a tree with branch lengths and fix negative or zero length
	branches.

Description

Take a tree with branch lengths and fix negative or zero length branches.

Usage

```
tree_fix_brlen(
   tree = NULL,
   fixing_criterion = "negative",
   fixing_method = 0,
   ultrametric = TRUE
)
```

Arguments

tree	A tree either as a newick character string or as a phylo object.
fixing_criterion	
	A character vector specifying the type of branch length to be fixed: "negative" or "zero" (the number 0 is also allowed).
fixing_method	A character vector specifying the method to fix branch lengths: "bladj", "mr- bayes" or a number to be assigned to all branches meeting fixing_criterion
ultrametric	Boolean indicating whether to force ultrametric or not.

Value

A phylo object with no negative or zero branch lengths.

tree_from_taxonomy Gets a taxonomic tree from a vector of taxa

Description

This uses the taxize package's wrapper of the Global Names Resolver to get taxonomic paths for the vector of taxa you pass in. Sources is a vector of source labels in order (though it works best if everything uses the same taxonomy, so we recommend doing just one source). You can see options by doing taxize::gnr_datasources(). Our default is Catalogue of Life. The output is a phylo object (typically with many singleton nodes if collapse_singles is FALSE: nodes with only one descendant (like "Homo" having "Homo sapiens" as its only descendant) but these singletons typically have node.labels

Usage

```
tree_from_taxonomy(
   taxa,
   sources = "Catalogue of Life",
   collapse_singles = TRUE
)
```

Arguments

taxa	Vector of taxon names	
sources	Vector of names of preferred sources; see taxize::gnr_datasources(). Currently supports 100 taxonomic resources, see details.	
collapse_singles		
	If true, collapses singleton nodes	

Value

A list containing a phylo object with resolved names and a vector with unresolved names

Examples

Not run: # This is a flag for package development. You are welcome to run the example.

```
taxa <- c(
    "Homo sapiens", "Ursus arctos", "Pan paniscus", "Tyrannosaurus rex",
    "Ginkgo biloba", "Vulcan", "Klingon"
)
results <- tree_from_taxonomy(taxa)
print(results$unresolved) # The taxa that do not match
ape::plot.phylo(results$phy) # may generate warnings due to problems with singletons
ape::plot.phylo(ape::collapse.singles(results$phy), show.node.label = TRUE)
# got rid of singles, but this also removes a lot of the node.labels
## End(Not run) # end dontrun</pre>
```

tree_get_node_data Get node numbers, node names, descendant tip numbers and labels of nodes from any tree, and node ages from dated trees.

Description

Get node numbers, node names, descendant tip numbers and labels of nodes from any tree, and node ages from dated trees.

Usage

```
tree_get_node_data(
   tree = NULL,
   nodes = NULL,
   node_data = c("node_number", "node_label", "node_age", "descendant_tips_number",
        "descendant_tips_label")
)
```

Arguments

tree	A tree either as a newick character string or as a phylo object.
nodes	Numeric vector with node numbers from which you want to obtain data. Default to NULL: obtain data for all nodes in the tree.
node_data	A character vector containing one or all from: "node_number", "node_label", "node_age", "descendant_tips_number", "descendant_tips_label"

Value

A list

tree_get_singleton_outgroup

Identify the presence of a single lineage outgroup in a phylogeny

Description

Identify the presence of a single lineage outgroup in a phylogeny

Usage

```
tree_get_singleton_outgroup(tree = NULL)
```

Arguments

tree A tree either as a newick character string or as a phylo object.

Value

A character vector with the name of the single lineage outgroup. Returns NA if there is none.

tree_node_tips To get tip numbers descending from any given node of a tree

Description

To get tip numbers descending from any given node of a tree

Usage

tree_node_tips(tree = NULL, node = NULL, curr = NULL)

Arguments

tree	a phylogenetic tree as an object of class "phylo".
node	an integer specifying a node number in the tree.
curr	the set of previously stored node numbers - used in recursive function calls.

Value

A numeric vector with tip numbers descending from a node

update_all_cached

Description

This includes opentree chronograms, contributors, treebase and curators For speed, datelife caches chronograms and other information. Running this (within the checked out version of datelife) will refresh these. Then git commit and git push them back

Usage

update_all_cached()

Value

None

update_datelife_cache Create an updated OpenTree chronograms database object

Description

The function calls get_opentree_chronograms() to update the OpenTree chronograms database cached in datelife. It has the option to write the updated object as an .Rdata file, that will be independent of the opentree_chronograms data object that you can load with data("opentree_chronograms", package = "datelife").

Usage

```
update_datelife_cache(
  write = TRUE,
  updated_name = "opentree_chronograms_updated",
  file_path = file.path(tempdir()),
   ...
)
```

Arguments

write	Defaults to TRUE, it saves an .Rdata file named indicated by argument name, con- taining available chronograms from Open Tree of Life. Saves to path indicated by argument path.
updated_name	Used if write = TRUE. Defaults to "opentree_chronograms_updated". A char- acter vector of length one indicating the name to assign to both the updated OpenTree chronogram database object and the ".Rdata" file. For example, if name = "my_database", the function will assign the updated chronogram database to an object named my_database and will write it to a file named "my_database.Rdata" in the path indicated by argument file_path.

file_path	Used if write = TRUE. A character vector of length 1 indicating the path to write the updated database ".Rdata" file to, excluding file name. Defaults to temporary directory obtained with base::tempdir() and formatted with base::file.path().
	Arguments passed on to get_opentree_chronograms
	max_tree_count Default to "all", it gets all available chronograms. For testing purposes, a numeric value indicating the max number of trees to be cached.

Value

A list of 4 elements:

- **authors** A list of lists of author names of the original studies that published chronograms currently stored in the Open Tree of Life database.
- **curators** A list of lists of curator names that uploaded chronograms to the Open Tree of Life database.
- **studies** A list of study identifiers from original studies that published chronograms currently stored in the Open Tree of Life database.
- trees A multiPhylo object storing the chronograms from Open Tree of Life database.
- update A character vector indicating the time when the database object was last updated.
- version A character vector indicating the datelife package version when the object was last updated.

use_all_calibrations Date a given tree topology using a given set of congruified calibrations or ages

Description

use_all_calibrations generates one or multiple chronograms (i.e., phylogenetic trees with branch lengths proportional to time) by dating a tree topology given in phy, and secondary calibrations given in calibrations, using the algorithm specified in the argument dating_method.

Usage

```
use_all_calibrations(
   phy = NULL,
   calibrations = NULL,
   each = FALSE,
   dating_method = "bladj",
   ...
)
```

Arguments

phy	A phylo object to use as tree topology.
calibrations	A calibrations object, an output of get_all_calibrations().
each	Boolean, default to FALSE: all calibrations are returned in the same data.frame. If TRUE, calibrations from each chronogram are returned in separate data frames.
dating_method	Tree dating algorithm to use. Options are "bladj" or "pathd8" (Webb et al., 2008, doi:10.1093/bioinformatics/btn358; Britton et al., 2007, doi:10.1080/10635150701613783).
	Arguments passed on to use_calibrations
	type The type of age to use as calibration. Options are "median", "mean", "min", or "max".

Details

If phy has no branch lengths, dating_method is ignores, and the function applies secondary calibrations to date the tree with the BLADJ algorithm. See make_bladj_tree() and use_calibrations_bladj(). If phy has branch lengths, the function can use the PATHd8 algorithm. See use_calibrations_pathd8().

Value

A phylo or multiPhylo object with branch lengths proportional to time.

More

The output object stores the used calibrations and dating_method as attributes(output)\$datelife_calibrations and attributes(output)\$dating_method.

References

Webb, C. O., Ackerly, D. D., & Kembel, S. W. (2008). "Phylocom: software for the analysis of phylogenetic community structure and trait evolution". Bioinformatics, 24(18), doi:10.1093/bioinformatics/btn358.

Britton, T., Anderson, C. L., Jacquet, D., Lundqvist, S., & Bremer, K. (2007). "Estimating divergence times in large phylogenetic trees". Systematic biology, 56(5), 741-752. doi:10.1080/10635150701613783.

use_calibrations Date a given tree topology using a combined set of given calibrations

Description

use_calibrations combines all given calibrations and uses them as constraints to perform a dating analysis on a given tree topology, using BLADJ if it has no branch lengths, or PATHd8 if the given tree topology has initial branch lengths.

use_calibrations_bladj

Usage

```
use_calibrations(
  phy = NULL,
  calibrations = NULL,
  dating_method = "bladj",
  type = "median",
   ...
)
```

Arguments

phy	A phylo object to use as tree topology.
calibrations	A calibrations object, an output of <pre>get_all_calibrations().</pre>
dating_method	Tree dating algorithm to use. Options are "bladj" or "pathd8" (Webb et al., 2008, doi:10.1093/bioinformatics/btn358; Britton et al., 2007, doi:10.1080/10635150701613783).
type	The type of age to use as calibration. Options are "median", "mean", "min", or "max".
	Arguments passed on to use_calibrations_pathd8
	expand How much to expand by each step to get consistent calibrations. Should be between 0 and 1.
	giveup How many expansions to try before giving up

Details

If phy has no branch lengths, dating_method is ignores, and the function applies secondary calibrations to date the tree with the BLADJ algorithm. See make_bladj_tree() and use_calibrations_bladj(). If phy has branch lengths, the function can use the PATHd8 algorithm. See use_calibrations_pathd8().

Value

A phylo object with branch lengths proportional to time.

More

The output object stores the used calibrations and dating_method as attributes(output)\$datelife_calibrations and attributes(output)\$dating_method.

use_calibrations_bladj

Use calibrations to date a topology with the BLADJ algorithm.

Description

The function use_calibrations_bladj prepares the input for BLADJ and calls make_bladj_tree().

Usage

```
use_calibrations_bladj(phy = NULL, calibrations, type = "median", root_age)
```

Arguments

phy	A phylo object with or without branch lengths.
calibrations	A data.frame of secondary calibrations for any pair of taxon names in phy, usually obtained with get_all_calibrations().
type	The type of age to use as calibration. Options are "median", "mean", "min", or "max".
root_age	Numeric specifying the age of the root. Only used if there are no ages for the root node in calibrations argument. If missing, NULL, or not numeric, the value of the oldest calibration plus one unit of the mean differences across calibrations, will be used as root calibration. If there is one single age point provided as calibrations, the root age will be set to 10% more than the age of the single calibration.

Details

The BLADJ algorithm is part of the Phylocom software, presented in Webb et al. (2008) doi:10.1093/ bioinformatics/btn358.

Value

A chronogram: a phylo object with branch lengths proportional to time.

References

Webb, C. O., Ackerly, D. D., & Kembel, S. W. (2008). "Phylocom: software for the analysis of phylogenetic community structure and trait evolution". Bioinformatics, 24(18), doi:10.1093/bioinformatics/btn358.

use_calibrations_bladj.matchedCalibrations Use calibrations to date a topology with the BLADJ algorithm.

Description

The function prepares the input for BLADJ and calls make_bladj_tree()

Usage

```
use_calibrations_bladj.matchedCalibrations(
   calibrations,
   type = "mean",
   root_age = NULL
)
```

Arguments

calibrations	A data.frame of secondary calibrations for any pair of taxon names in phy, usually obtained with get_all_calibrations().
type	The type of age to use as calibration. Options are "median", "mean", "min", or "max".
root_age	Numeric specifying an age for the root, provided by the user. Only used if there are no time calibrations for the root node in the chronograms database. If NULL or not numeric, the maximum calibration age plus one unit of the sd (calculated with stats::sd()) of all node ages available for the tree will be used as root calibration. If there is only one calibration available for the whole tree, the root node age will be proportional to 1.1 of the age of that calibration.

Details

The BLADJ algorithm is part of the Phylocom software, presented in Webb et al. (2008) doi:10.1093/ bioinformatics/btn358.

Value

A phylo object with branch lengths proportional to time.

References

Webb, C. O., Ackerly, D. D., & Kembel, S. W. (2008). "Phylocom: software for the analysis of phylogenetic community structure and trait evolution". Bioinformatics, 24(18), doi:10.1093/bioinformatics/btn358.

use_calibrations_each Date a given tree topology by using a given list of calibrations independently, to generate multiple hypothesis of time of divergence

Description

use_calibrations_each wraps use_calibrations to take each set of given calibrations and use it independently as constraints for BLADJ or PATHd8 to date a given tree topology.

Usage

```
use_calibrations_each(phy = NULL, calibrations = NULL, ...)
```

Arguments

phy	A phylo object to use as tree topology.
calibrations	A calibrations object, an output of get_all_calibrations().
	Arguments passed on to use_calibrations

- dating_method Tree dating algorithm to use. Options are "bladj" or "pathd8" (Webb et al., 2008, doi:10.1093/bioinformatics/btn358; Britton et al., 2007, doi:10.1080/10635150701613783).
- type The type of age to use as calibration. Options are "median", "mean", "min", or "max".

Details

If phy has no branch lengths, dating_method is ignores, and the function applies secondary calibrations to date the tree with the BLADJ algorithm. See make_bladj_tree() and use_calibrations_bladj(). If phy has branch lengths, the function can use the PATHd8 algorithm. See use_calibrations_pathd8().

Value

A multiPhylo object of trees with branch lengths proportional to time.

More

The output object stores the used calibrations and dating_method as attributes(output)\$datelife_calibrations and attributes(output)\$dating_method.

use_calibrations_pathd8

Date a tree with secondary calibrations using PATHd8

Description

use_calibrations_pathd8 uses secondary calibrations to date a tree with initial branch lengths using PATHd8.

Usage

```
use_calibrations_pathd8(
   phy = NULL,
   calibrations = NULL,
   expand = 0.1,
   giveup = 100
)
```

Arguments

phy	A phylo object with branch lengths.
calibrations	A data.frame of secondary calibrations for any pair of taxon names in phy. usually obtained with get_all_calibrations().
expand	How much to expand by each step to get consistent calibrations. Should be between 0 and 1.
giveup	How many expansions to try before giving up

Details

This function implements the PATHd8 algorithm described in Britton et al. (2007) doi:10.1080/ 10635150701613783, with geiger::PATHd8.phylo(). The function first attempts to use the given calibrations as fixed ages. If that fails (often due to conflict between calibrations), it will expand the range of the minimum age and maximum age and try again. And repeat. If expand = 0, it uses the summarized calibrations. In some cases, it returns edge lengths in relative time (with maximum tree depth = 1) instead of absolute time, as given by calibrations. In this case, the function returns NA. This is an issue from PATHd8.

Value

A phylo object with branch lengths proportional to time.

References

Britton, T., Anderson, C. L., Jacquet, D., Lundqvist, S., & Bremer, K. (2007). "Estimating divergence times in large phylogenetic trees". Systematic biology, 56(5), 741-752. doi:10.1080/10635150701613783.

use_calibrations_treePL

Date a tree with initial branch lengths with treePL.

Description

Date a tree with initial branch lengths with treePL.

Usage

use_calibrations_treePL(phy, calibrations)

Arguments

phy	A phylo object with or without branch lengths.
calibrations	A data.frame of secondary calibrations for any pair of taxon names in phy, usually obtained with get_all_calibrations().
	usually obtained with get_att_cattor attors().

Details

This function uses treePL as described in Smith, S. A., & O'Meara, B. C. (2012). doi:10.1093/ bioinformatics/bts492, with the function treePL.phylo. It attempts to use the calibrations as fixed ages. If that fails (often due to conflict between calibrations), it will expand the range of the minimum age and maximum age and try again. And repeat. If expand = 0, it uses the summarized calibrations. In some cases, it returns edge lengths in relative time (with maximum tree depth = 1) instead of absolute time, as given by calibrations. In this case, the function returns NA. This is an issue from PATHd8.

Value

A phylo object

References

Smith, S. A., & O'Meara, B. C. (2012). "treePL: divergence time estimation using penalized likelihood for large phylogenies". Bioinformatics, 28(20), 2689-2690, doi:10.1093/bioinformatics/bts492.

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