Package: rotl (via r-universe)

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Title Interface to the 'Open Tree of Life' API

Version 3.1.0

Description An interface to the 'Open Tree of Life' API to retrieve phylogenetic trees, information about studies used to assemble the synthetic tree, and utilities to match taxonomic names to 'Open Tree identifiers'. The 'Open Tree of Life' aims at assembling a comprehensive phylogenetic tree for all named species.

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get_study

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Description

Returns the trees associated with a given study

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Usage

```
get_study(
  study_id = NULL,
  object_format = c("phylo", "nexml"),
  file_format,
  file,
  ...
)
```

Arguments

study_id the study ID for the study of interest (character)

object_format the class of the object the query should return (either phylo or nexml). Ignored if file_format is specified.

file_format the format of the file to be generated (newick, nexus, nexml or json).

file the file name where the output of the function will be saved.

additional arguments to customize the API request (see rot1 package documentation).

Details

If file_format is missing, the function returns an object of the class phylo from the ape package (default), or an object of the class nexml from the RNeXML package.

Otherwise file_format can be either newick, nexus, nexml or json, and the function will generate a file of the selected format. In this case, a file name needs to be provided using the argument file. If a file with the same name already exists, it will be silently overwritten.

Value

if file_format is missing, an object of class phylo or nexml, otherwise a logical indicating whether the file was successfully created.

See Also

```
get_study_meta
```

```
## Not run:
that_one_study <- get_study(study_id="pg_719", object_format="phylo")
if (require(RNeXML)) { ## if RNeXML is installed get the object directly
    nexml_study <- get_study(study_id="pg_719", object_format="nexml")
} else { ## otherwise write it to a file
    get_study(study_id="pg_719", file_format="nexml", file=tempfile(fileext=".nexml"))
}

## End(Not run)</pre>
```

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get_study_subtree

Study Subtree

Description

Retrieve subtree from a specific tree in the Open Tree of Life data store

Usage

```
get_study_subtree(
   study_id,
   tree_id,
   subtree_id,
   object_format = c("phylo"),
   tip_label = c("original_label", "ott_id", "ott_taxon_name"),
   file_format,
   file,
   deduplicate = TRUE,
   ...
)
```

Arguments

study_id	the study identifier (character)
tree_id	the tree identifier (character)
subtree_id	either a node id that specifies a subtree or "ingroup" which returns the ingroup for this subtree.
object_format	the class of the object returned by the function (default, and currently only possibility phylo from the ape package)
tip_label	the format of the tip labels. "original_label" (default) returns the original labels as provided in the study, "ott_id" labels are replaced by their ott IDs, "ott_taxon_name" labels are replaced by their Open Tree Taxonomy taxon name.
file_format	character, the file format to use to save the results of the query (possible values, 'newick' or 'nexus').
file	character, the path and file name where the output should be written.
deduplicate	logical (default TRUE). If the tree returned by the study contains duplicated taxon names, should they be made unique? It is normally illegal for NEXUS/Newick tree strings to contain duplicated tip names. This is a workaround to circumvent this requirement. If TRUE, duplicated tip labels will be appended _1, _2, etc.
• • •	additional arguments to customize the API request (see rotl package documentation).

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Examples

```
## Not run:
small_tr <- get_study_subtree(study_id="pg_1144", tree_id="tree5800", subtree_id="node991044")
ingroup <- get_study_subtree(study_id="pg_1144", tree_id="tree5800", subtree_id="ingroup")</pre>
nexus_file <- tempfile(fileext=".nex")</pre>
get_study_subtree(study_id="pg_1144", tree_id="tree5800", subtree_id="ingroup", file=nexus_file,
                  file_format="nexus")
## End(Not run)
```

get_study_tree

Study Tree

Description

Returns a specific tree from within a study

Usage

```
get_study_tree(
  study_id = NULL,
  tree_id = NULL,
  object_format = c("phylo"),
  tip_label = c("original_label", "ott_id", "ott_taxon_name"),
  file_format,
  file.
  deduplicate = TRUE,
)
```

Arguments

study_id the identifier of a study (character) tree_id the identifier of a tree within the study object_format

the class of the object to be returned (default and currently only possible value

phylo from the ape package).

tip_label the format of the tip labels. "original_label" (default) returns the original

> labels as provided in the study, "ott_id" labels are replaced by their ott IDs, "ott_taxon_name" labels are replaced by their Open Tree Taxonomy taxon

name.

the format of the file to be generated (newick default, nexus, or json). file_format

file the file name where the output of the function will be saved.

deduplicate logical (default TRUE). If the tree returned by the study contains duplicated taxon

names, should they be made unique? It is normally illegal for NEXUS/Newick tree strings to contain duplicated tip names. This is a workaround to circumvent this requirement. If TRUE, duplicated tip labels will be appended _1, _2, etc.

get_tree_ids

... additional arguments to customize the API request (see rot1 package documentation).

Value

if file_format is missing, an object of class phylo, otherwise a logical indicating whether the file was successfully created.

Examples

```
## Not run:
    tree <- get_study_tree(study_id="pg_1144", tree_id="tree2324")

## comparison of the first few tip labels depending on the options used
head(get_study_tree(study_id="pg_1144", tree_id="tree2324", tip_label="original_label")$tip.label)
head(get_study_tree(study_id="pg_1144", tree_id="tree2324", tip_label="ott_id")$tip.label)
head(get_study_tree(study_id="pg_1144", tree_id="tree2324", tip_label="ott_taxon_name")$tip.label)
## End(Not run)

get_tree_ids

Study Metadata</pre>
```

Description

Retrieve metadata about a study in the Open Tree of Life datastore.

Usage

```
get_tree_ids(sm)
get_publication(sm)
candidate_for_synth(sm)
get_study_year(sm)
## S3 method for class 'study_meta'
get_tree_ids(sm)
## S3 method for class 'study_meta'
get_publication(sm)
## S3 method for class 'study_meta'
candidate_for_synth(sm)
## S3 method for class 'study_meta'
candidate_for_synth(sm)
## S3 method for class 'study_meta'
get_study_year(sm)
```

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Arguments

```
    sm an object created by get_study_meta
    study_id the study identifier (character)
    additional arguments to customize the API request (see rotl package documentation).
```

Details

get_study_meta returns a long list of attributes for the studies that are contributing to the synthetic tree. To help with the extraction of relevant information from this list, several helper functions exists:

- get_tree_ids The identifiers of the trees associated with the study
- get_publication The citation information of the publication for the study. The DOI (or URL) for the study is available as an attribute to the returned object (i.e., attr(object, "DOI")).
- candidate_for_synth The identifier of the tree(s) from the study used in the synthetic tree. This is a subset of the result of get_tree_ids.

 get_study_year The year of publication of the study.

Value

named-list containing the metadata associated with the study requested

Examples

```
## Not run:
req <- get_study_meta("pg_719")
get_tree_ids(req)
candidate_for_synth(req)
get_publication(req)
get_study_year(req)
## End(Not run)</pre>
```

inspect.match_names

Inspect and Update alternative matches for a name returned by tnrs_match_names

Description

Taxonomic names may have different meanings in different taxonomic contexts, as the same genus name can be applied to animals and plants for instance. Additionally, the meaning of a taxonomic name may have change throughout its history, and may have referred to a different taxon in the past. In such cases, a given names might have multiple matches in the Open Tree Taxonomy. These functions allow users to inspect (and update) alternative meaning of a given name and its current taxonomic status according to the Open Tree Taxonomy.

Usage

```
## S3 method for class 'match_names'
inspect(response, row_number, taxon_name, ott_id, ...)
inspect(response, ...)
## S3 method for class 'match_names'
update(object, row_number, taxon_name, ott_id, new_row_number, new_ott_id, ...)
```

Arguments

an object generated by the tnrs_match_names function response row_number the row number corresponding to the name to inspect taxon_name the taxon name corresponding to the name to inspect the ott id corresponding to the name to inspect ott_id

currently ignored . . .

object an object created by tnrs_match_names

new_row_number the row number in the output of inspect to replace the taxa specified by row_number,

taxon_name, or ott_id.

the ott id of the taxon to replace the taxa specified by row_number, taxon_name, new_ott_id

or ott_id.

Details

To inspect alternative taxonomic meanings of a given name, you need to provide the object resulting from a call to the tnrs_match_names function, as well as one of either the row number corresponding to the name in this object, the name itself (as used in the original query), or the ott_id listed for this

To update one of the name, you also need to provide the row number in which the name to be replaced appear or its ott id.

Value

a data frame

See Also

```
tnrs_match_names
```

```
## Not run:
matched_names <- tnrs_match_names(c("holothuria", "diadema", "boletus"))</pre>
 inspect(matched_names, taxon_name="diadema")
new_matched_names <- update(matched_names, taxon_name="diadema",</pre>
                             new_ott_id = 631176)
new_matched_names
```

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

is_in_tree

Check that OTT ids occur in the Synthetic Tree

Description

Some valid taxonomic names do not occur in the Synthetic Tree. This convenience function allows you to check whether a given Open Tree Taxonomy identifier (OTT id) is in the tree. A taxonomic name may not occur in the synthetic tree because (1) it is an extinct or invalid taxon, or (2) it is part of a group that is not monophyletic in the tree.

Usage

```
is_in_tree(ott_ids, ...)
```

Arguments

ott_ids a vector of Open Tree Taxonomy identifiers
... additional arguments to customize the API request (see rot1 package documentation).

Value

A named logical vector. TRUE indicates that the OTT id is in the synthetic tree, and FALSE that it is not

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list_trees	List trees ids in objects returned by studies_find_studies and studies_find_trees.

Description

list_trees returns all trees associated with a particular study when used on an object returned by studies_find_studies, but only the trees that match the search criteria when used on objects returned by studies_find_trees.

Usage

```
list_trees(matched_studies, ...)
## S3 method for class 'matched_studies'
list_trees(matched_studies, study_id, ...)
```

Arguments

```
matched_studies
an object created by studies_find_trees or studies_find_studies.
... Currently unused
study_id a study_id listed in the object returned by studies_find_trees
```

Value

list_trees returns a list of the tree_ids for each study that match the requested criteria. If a study_id is provided, then only the trees for this study are returned as a vector.

See Also

studies_find_studies and studies_find_trees. The help for these functions have examples demonstrating the use of list_trees.

```
ott_id.match_names ott_id and flags for taxonomic names matched by tnrs_match_names
```

Description

rotl provides a collection of functions that allows users to extract relevant information from an object generated by tnrs_match_names function.

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Usage

```
## S3 method for class 'match_names'
ott_id(tax, row_number, taxon_name, ott_id, ...)
## S3 method for class 'match_names'
flags(tax, row_number, taxon_name, ott_id, ...)
flags(tax, ...)
```

Arguments

tax	an object returned by tnrs_match_names
row_number	the row number corresponding to the name for which to list the synonyms
taxon_name	the taxon name corresponding to the name for which to list the synonyms
ott_id	the ott id corresponding to the name for which to list the synonyms
	currently ignored

Details

These methods optionally accept one of the arguments row_number, taxon_name or ott_id to retrieve the corresponding information for one of the matches in the object returned by the tnrs_match_names function.

If these arguments are not provided, these methods can return information for the matches currently listed in the object returned by tnrs_match_names.

Value

A list of the ott ids or flags for the taxonomic names matched with tnrs_match_names, for either one or all the names.

```
## Not run:
    rsp <- tnrs_match_names(c("Diadema", "Tyrannosaurus"))
    rsp$ott_id  # ott id for match currently in use
    ott_id(rsp)  # similar as above but elements are named

## flags() is useful for instance to determine if a taxon is extinct
    flags(rsp, taxon_name="Tyrannosaurus")

## End(Not run)</pre>
```

source_list

rotl

An Interface to the Open Tree of Life API

Description

The Open Tree of Life is an NSF funded project that is generating an online, comprehensive phylogenetic tree for 1.8 million species. rot1 provides an interface that allows you to query and retrieve the parts of the tree of life that is of interest to you.

Details

rotl provides function to most of the end points the API provides. The documentation of the API is available at: https://github.com/OpenTreeOfLife/opentree/wiki/Open-Tree-of-Life-APIs

Customizing API calls

All functions that use API end points can take 2 arguments to customize the API call and are passed as ... arguments.

- otl_v This argument controls which version of the API your call is using. The default value for this argument is a call to the non-exported function otl_version() which returns the current version of the Open Tree of Life APIs (v2).
- dev_url This argument controls whether to use the development version of the API. By default, dev_url is set to FALSE, using dev_url = TRUE in your function calls will use the development version.

For example, to use the development version of the API, you could use: tnrs_match_names("anas", dev_url=TRUE)

Additional arguments can also be passed to the GET and POST methods.

Acknowledgments

This package was started during the Open Tree of Life Hackathon organized by the OpenTree of Life, the NESCent Hackathon Interoperability Phylogenetic group, and Arbor.

source_list

List of studies used in the Tree of Life

Description

Retrieve the detailed information for the list of studies used in the Tree of Life.

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Usage

```
source_list(tax, ...)
## S3 method for class 'tol_summary'
source_list(tax, ...)
```

Arguments

```
tax a list containing a source_id_map slot.
... additional arguments (currently unused)
```

Details

This function takes the object resulting from tol_about(study_list = TRUE), tol_mrca(), tol_node_info(), and returns a data frame listing the tree_id, study_id and git_sha for the studies currently included in the Tree of Life.

Value

a data frame

strip_ott_ids

Strip OTT ids from tip labels

Description

```
Strip OTT ids from tip labels
```

Usage

```
strip_ott_ids(tip_labels, remove_underscores = FALSE)
```

Arguments

```
tip_labels a character vector containing tip labels (most likely the tip.label element from a tree returned by tol_induced_subtree
```

remove_underscores

logical (defaults to FALSE). If set to TRUE underscores in tip labels are converted to spaces

Value

A character vector containing the contents of tip_labels with any OTT ids removed.

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Examples

```
## Not run:
genera <- c("Perdix", "Setophaga", "Cinclus", "Struthio")
tr <- tol_induced_subtree(ott_ids=c(102710, 285198, 267845, 292466))
tr$tip.label %in% genera
tr$tip.label <- strip_ott_ids(tr$tip.label)
tr$tip.label %in% genera
## End(Not run)</pre>
```

studies_find_studies Find a Study

Description

Return the identifiers of studies that match given properties

Usage

```
studies_find_studies(
  property = NULL,
  value = NULL,
  verbose = FALSE,
  exact = FALSE,
  detailed = TRUE,
  ...
)
```

Arguments

property
value
The property value to be searched on (character)
verbose
Should the output include all metadata (logical default FALSE)
exact
Should exact matching be used? (logical, default FALSE)

detailed
If TRUE (default), the function will return a data frame that summarizes information about the study (see 'Value'). Otherwise, it only returns the study identifiers.

...
additional arguments to customize the API request (see rot1 package documentation).

Value

If detailed=TRUE, the function returns a data frame listing the study id (study_ids), the number of trees associated with this study (n_trees), the tree ids (at most 5) associated with the studies (tree_ids), the tree id that is a candidate for the synthetic tree if any (candidate), the year of

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publication of the study (study_year), the title of the publication for the study (title), and the DOI (Digital Object Identifier) for the study (study_doi).

If detailed=FALSE, the function returns a data frame with a single column containing the study identifiers.

See Also

studies_properties which lists properties against which the studies can be searched. list_trees that returns a list for all tree ids associated with a study.

Examples

studies_find_trees

Find Trees

Description

Return a list of studies for which trees match a given set of properties

Usage

```
studies_find_trees(
  property = NULL,
  value = NULL,
  verbose = FALSE,
  exact = FALSE,
  detailed = TRUE,
  ...
)
```

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Arguments

property	The property to be searched on (character)
value	The property-value to be searched on (character)
verbose	Should the output include all metadata? (logical, default FALSE)
exact	Should exact matching be used for the value? (logical, default FALSE)
detailed	Should a detailed report be provided? If TRUE (default), the output will include metadata about the study that include trees matching the property. Otherwise, only information about the trees will be provided.
• • •	additional arguments to customize the API request (see rot1 package documentation).

Details

The list of possible values to be used as values for the argument property can be found using the function studies_properties.

Value

A data frame that summarizes the trees found (and their associated studies) for the requested criteria. If a study has more than 5 trees, the tree_ids of the first ones will be shown, followed by . . . to indicate that more are present.

If detailed=FALSE, the data frame will include the study ids of the study (study_ids), the number of trees in this study that match the search criteria (n_matched_trees), the tree ids that match the search criteria (match_tree_ids).

If detailed=TRUE, in addition of the fields listed above, the data frame will also contain the total number of trees associated with the study (n_trees), all the tree ids associated with the study (tree_ids), the tree id that is a potential candidate for inclusion in the synthetic tree (if any) (candidate), the year the study was published (study_year), the title of the study (title), the DOI for the study (study_doi).

See Also

studies_properties which lists properties the studies can be searched on. list_trees for listing the trees that match the query.

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```
list_trees(res, study_id = "pg_2769")
## End(Not run)
```

studies_properties

Properties of the Studies

Description

Return the list of study properties that can be used to search studies and trees used in the synthetic tree.

Usage

```
studies_properties(...)
```

Arguments

additional arguments to customize the API request (see rot1 package documentation).

Details

The list returned has 2 elements tree_properties and studies_properties. Each of these elements lists additional arguments to customize the API request properties that can be used to search for trees and studies that are contributing to the synthetic tree. The definitions of these properties are available from https://github.com/OpenTreeOfLife/phylesystem-api/wiki/NexSON

Value

A list of the study properties that can be used to find studies and trees that are contributing to the synthetic tree.

See Also

```
studies_find_trees
```

```
## Not run:
   all_the_properties <- studies_properties()
   unlist(all_the_properties$tree_properties)
## End(Not run)</pre>
```

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study_external_IDs

Get external identifiers for data associated with an Open Tree study

Description

Data associated with studies contributing to the Open Tree synthesis may be available from other databases. In particular, trees and alignments may be available from treebase and DNA sequences and bibliographic information associated with a given study may be available from the NCBI. This function retrieves that information for a given study.

Usage

```
study_external_IDs(study_id)
```

Arguments

study_id

An open tree study ID

Value

A study_external_data object (which inherits from a list) which contains some of the following.

doi, character, the DOI for the paper describing this study

external_data_url, character, a URL to an external data repository (e.g. a treebase entry) if one exists.

pubmed_id character, the unique ID for this study in the NCBI's pubmed database

popset_ids character, vector of IDs for the NCBI's popset database

nucleotide_ids character, vector of IDs for the NCBI's nucleotide database

See Also

```
studies_find_studies (used to discover study IDs)
```

```
## Not run:
flies <- studies_find_studies(property="ot:focalCladeOTTTaxonName", value="Drosophilidae")
study_external_IDs(flies[2,]$study_ids)
## End(Not run)</pre>
```

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```
synonyms.match_names    List the synonyms for a given name
```

Description

When querying the Taxonomic Name Resolution Services for a particular taxonomic name, the API returns as possible matches all names that include the queried name as a possible synonym. This function allows you to explore other synonyms for an accepted name, and allows you to determine why the name you queried is returning an accepted synonym.

Usage

```
## S3 method for class 'match_names'
synonyms(tax, row_number, taxon_name, ott_id, ...)
```

Arguments

tax	a data frame generated by the thrs_match_names function
row_number	the row number corresponding to the name for which to list the synonyms
taxon_name	the taxon name corresponding to the name for which to list the synonyms
ott_id	the ott id corresponding to the name for which to list the synonyms
	currently ignored

Details

To list synonyms for a given taxonomic name, you need to provide the object resulting from a call to the tnrs_match_names function, as well as one of either the row number corresponding to the name in this object, the name itself (as used in the original query), or the ott_id listed for this name. Otherwise, the synonyms for all the currently matched names are returned.

Value

a list whose elements are all synonym names (as vectors of character) for the taxonomic names that match the query (the names of the elements of the list).

```
## Not run:
    echino <- tnrs_match_names(c("Diadema", "Acanthaster", "Fromia"))
    ## These 3 calls are identical
    synonyms(echino, taxon_name="Acanthaster")
    synonyms(echino, row_number=2)
    synonyms(echino, ott_id=337928)
## End(Not run)</pre>
```

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taxonomy_about

Information about the Open Tree Taxonomy

Description

Summary information about the Open Tree Taxonomy (OTT)

Usage

```
taxonomy_about(...)
```

Arguments

... additional arguments to customize the API request (see rot1 package documentation).

Details

Return metadata and information about the taxonomy itself. Currently, the available metadata is fairly sparse, but includes (at least) the version, and the location from which the complete taxonomy source files can be downloaded.

Value

A list with the following properties:

- weburl String. The release page for this version of the taxonomy.
- author String. The author string.
- name String. The name of the taxonomy.
- source String. The full identifying information for this version of the taxonomy.
- version String. The version number of the taxonomy.

```
## Not run:
taxonomy_about()
## End(Not run)
```

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taxonomy_mrca

Taxonomic MRCA

Description

Taxonomic Least Inclusive Common Ancestor (MRCA)

Usage

```
taxonomy_mrca(ott_ids = NULL, ...)
## S3 method for class 'taxon_mrca'
tax_rank(tax, ...)
## S3 method for class 'taxon_mrca'
tax_name(tax, ...)
## S3 method for class 'taxon_mrca'
ott_id(tax, ...)
## S3 method for class 'taxon_mrca'
unique_name(tax, ...)
## S3 method for class 'taxon_mrca'
tax_sources(tax, ...)
## S3 method for class 'taxon_mrca'
flags(tax, ...)
## S3 method for class 'taxon_mrca'
suppressed(tax, ...)
```

Arguments

```
    a vector of ott ids for the taxa whose MRCA is to be found (numeric).
    additional arguments to customize the API request (see rot1 package documentation).
    an object generated by the taxonomy_mrca function
```

Details

Given a set of OTT ids, get the taxon that is the most recent common ancestor (the MRCA) of all the identified taxa.

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Value

- taxonomy_mrca returns a list about the taxonomic information relating to the MRCA for the ott_ids provided.
- tax_rank returns a character vector of the taxonomic rank for the MRCA.
- tax_name returns a character vector the Open Tree Taxonomy name for the MRCA.
- ott_id returns a numeric vector of the ott id for the MRCA.

Examples

```
## Not run:
req <- taxonomy_mrca(ott_ids=c(515698,590452,643717))
tax_rank(req)
tax_name(req)
ott_id(req)
## End(Not run)</pre>
```

taxonomy_subtree

Taxonomy subtree

Description

Given an ott id, return the inclusive taxonomic subtree descended from the specified taxon.

Usage

```
taxonomy_subtree(
  ott_id = NULL,
  output_format = c("taxa", "newick", "phylo", "raw"),
  label_format = NULL,
  file,
  ...
)
```

Arguments

ott_id	The ott id of the taxon of interest.
output_format	the format of the object to be returned. See the 'Return' section.
label_format	Character. Defines the label type; one of "name", "id", or "name_and_id" (the default).
file	the file name where to save the output of the function. Ignored unless output_format is set to "phylo".
• • •	additional arguments to customize the API request (see rotl package documentation).

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Details

If the output of this function is exported to a file, the only possible value for the output_format argument is "newick". If the file provided already exists, it will be silently overwritten.

Value

If the file argument is missing:

- "taxa" a list of the taxa names (species) in slot tip_label, and higher-level taxonomy (e.g., families, genera) in slot edge_label, descending from the taxa corresponding to the ott_id provided.
- "newick" a character vector containing the newick formatted string corresponding to the taxonomic subtree for the ott_id provided.
- "phylo" an object of the class phylo from the ape package.
- "raw" the direct output from the API, i.e., a list with an element named 'newick' that contains the subtree as a newick formatted string.

If a file argument is provided (and output_format is set to "phylo"), a logical indicating whether the file was successfully created.

Examples

```
## Not run:
req <- taxonomy_subtree(ott_id=515698)
plot(taxonomy_subtree(ott_id=515698, output_format="phylo"))
## End(Not run)</pre>
```

taxonomy_taxon_info

Taxon information

Description

Information about taxa.

Usage

```
taxonomy_taxon_info(
  ott_ids,
  include_children = FALSE,
  include_lineage = FALSE,
  include_terminal_descendants = FALSE,
  ...
)

## S3 method for class 'taxon_info'
tax_rank(tax, ...)
```

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```
## S3 method for class 'taxon_info'
tax_name(tax, ...)
## S3 method for class 'taxon_info'
unique_name(tax, ...)
## S3 method for class 'taxon_info'
synonyms(tax, ...)
## S3 method for class 'taxon_info'
ott_id(tax, ...)
## S3 method for class 'taxon_info'
tax_sources(tax, ...)
## S3 method for class 'taxon_info'
is_suppressed(tax, ...)
## S3 method for class 'taxon_info'
is_suppressed(tax, ...)
```

Arguments

ott_ids the ott ids of the taxon of interest (numeric or character containing only numbers)

include_children

whether to include information about all the children of this taxon. Default FALSE.

include_lineage

whether to include information about all the higher level taxa that include the ott_ids. Default FALSE.

include_terminal_descendants

whether to include the list of terminal ott_ids contained in the ott_ids provided.

. . . additional arguments to customize the API request (see rot1 package documentation).

tax an object generated by the taxonomy_taxon_info function

Details

Given a vector of ott ids, taxonomy_taxon_info returns information about the specified taxa.

The functions tax_rank, tax_name, and synonyms can extract this information from an object created by the taxonomy_taxon_info().

Value

taxonomy_taxon_info returns a list detailing information about the taxa. tax_rank and tax_name

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return a vector. synonyms returns a list whose elements are the synonyms for each of the ott_id requested.

See Also

tnrs_match_names to obtain ott_id from a taxonomic name.

Examples

```
## Not run:
req <- taxonomy_taxon_info(ott_id=515698)
tax_rank(req)
tax_name(req)
synonyms(req)
## End(Not run)</pre>
```

taxon_external_IDs

Get external identifiers for data associated with an Open Tree taxon

Description

The Open Tree taxonomy is a synthesis of multiple reference taxonomies. This function retrieves identifiers to external taxonomic records that have contributed the rank, position and definition of a given Open Tree taxon.

Usage

```
taxon_external_IDs(taxon_id)
```

Arguments

taxon_id

An open tree study ID

Value

a data.frame in which each row represents a unique record in an external database. The column "source" provides and abbreviated name for the database, and "id" the unique ID for the record.

See Also

tnrs_matchnames, which can be used to search for taxa by name. taxonomy_taxon, for more information about a given taxon.

```
## Not run:
    gibbon_IDs <- taxon_external_IDs(712902)
## End(Not run)</pre>
```

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tax_lineage

Lineage of a taxon

Description

Extract the lineage information (higher taxonomy) from an object returned by taxonomy_taxon_info.

Usage

```
tax_lineage(tax, ...)
## S3 method for class 'taxon_info'
tax_lineage(tax, ...)
```

Arguments

tax an object created by taxonomy_taxon_info using the argument include_lineage=TRUE.
... additional arguments (currently unused).

Details

The object passed to this function must have been created using the argument include_lineage=TRUE.

Value

A list with one slot per taxon that contains a data frame with 3 columns: the taxonomy rank, the name, and unique name for all taxa included in the lineage of the taxon up to the root of the tree.

tax_rank

Methods for Taxonomy

Description

Methods for dealing with objects containing taxonomic information (Taxonomy, TNRS endpoints)

Usage

```
tax_rank(tax, ...)
ott_id(tax, ...)
synonyms(tax, ...)
tax_sources(tax, ...)
```

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```
is_suppressed(tax, ...)
unique_name(tax, ...)
tax_name(tax, ...)
```

Arguments

```
tax an object returned by taxonomy_taxon_info, taxonomy_mrca, or tnrs_match_names
... additional arguments (see tnrs_match_names)
```

Details

This is the page for the generic methods. See the help pages for taxonomy_taxon_info, taxonomy_mrca, and tnrs_match_names for more information.

tnrs_contexts

TNRS contexts

Description

This function returns a list of pre-defined taxonomic contexts (i.e. clades) which can be used to limit the scope of thrs queries.

Usage

```
tnrs_contexts(...)
```

Arguments

additional arguments to customize the API request (see rot1 package documentation).

Details

Taxonomic contexts are available to limit the scope of TNRS searches. These contexts correspond to uncontested higher taxa such as 'Animals' or 'Land plants'. This service returns a list containing all available taxonomic context names, which may be used as input (via the context_name argument in other functions) to limit the search scope of other services including tnrs_match_names.

Value

Returns invisibly a list for each major clades (e.g., animals, microbes, plants, fungi, life) whose elements contains the possible contexts.

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tnrs_infer_context

Infer the taxonomic context from a list of names

Description

Return a taxonomic context given a list of taxonomic names

Usage

```
tnrs_infer_context(names = NULL, ...)
```

Arguments

names Vector of taxon names.

... additional arguments to customize the API request (see rot1 package documen-

tation).

Details

Find the least inclusive taxonomic context that includes all the unambiguous names in the input set. Unambiguous names are names with exact matches to non-homonym taxa. Ambiguous names (those without exact matches to non-homonym taxa) are indicated in results.

Value

A list including the context name, the context out id and possibly the names in the query that have an ambiguous taxonomic meaning in the query.

Examples

```
## Not run:
res <- tnrs_infer_context(names=c("Stellula calliope", "Struthio camelus"))
## End(Not run)</pre>
```

tnrs_match_names

Match names to the Open Tree Taxonomy

Description

Match taxonomic names to the Open Tree Taxonomy.

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Usage

```
tnrs_match_names(
  names = NULL,
  context_name = "All life",
  do_approximate_matching = TRUE,
  ids = NULL,
  include_suppressed = FALSE,
  ...
)
```

Arguments

names taxon names to be queried. Currently limited to 10,000 names for exact matches

and 2,500 names for approximate matches (character vector)

context_name of the taxonomic context to be searched (length-one character 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.

allowed as possible TNRS results.

. . additional arguments to customize the API request (see rot1 package documen-

tation).

Details

Accepts one or more taxonomic names and returns information about potential matches for these names to known taxa in the Open Tree Taxonomy.

This service uses taxonomic contexts to disambiguate homonyms and misspelled names; a context may be specified using the context_name argument. The default value for context_name is "All life". If no context is specified (i.e., context_name is set to NULL), then the context will be inferred (i.e., the shallowest taxonomic context that contains all unambiguous names in the input). Taxonomic contexts are uncontested higher taxa that have been selected to allow limits to be applied to the scope of TNRS searches (e.g. 'match names only within flowering plants'). Once a context has been identified (either user-specified or inferred), all taxon name matches will performed only against taxa within that context. For a list of available taxonomic contexts, see tnrs_contexts.

A name is considered unambiguous if it is not a synonym and has only one exact match to any taxon name in the entire taxonomy.

When the name search returns multiple matches, the taxon with the highest match score is returned. If the name returned is not the one you intended, you can use the inspect function to check the

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other taxa returned by your search. The Getting Started vignette has more information on how to do this.

Several functions listed in the 'See also' section can be used to inspect and manipulate the object generated by this function.

Value

A data frame summarizing the results of the query. The original query output is appended as an attribute to the returned object (and can be obtained using attr(object, "original_response")).

See Also

inspect.match_names, update.match_names, synonyms.match_names.

Examples

tol_about

Information about the Tree of Life

Description

Basic information about the Open Tree of Life (the synthetic tree)

Usage

```
tol_about(include_source_list = FALSE, ...)
## $3 method for class 'tol_summary'
tax_rank(tax, ...)
## $3 method for class 'tol_summary'
tax_sources(tax, ...)
## $3 method for class 'tol_summary'
unique_name(tax, ...)
## $3 method for class 'tol_summary'
tax_name(tax, ...)
## $3 method for class 'tol_summary'
ott_id(tax, ...)
```

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Arguments

```
    include_source_list
    Logical (default = FALSE). Return an ordered list of source trees.
    additional arguments to customize the API call (see rotl for more information).
    tax an object created with a call to tol_about.
```

Details

Summary information about the current draft tree of life, including information about the list of trees and the taxonomy used to build it. The object returned by tol_about can be passed to the taxonomy methods (tax_name(), tax_rank(), tax_sources(), ott_id), to extract relevant taxonomic information for the root of the synthetic tree.

Value

An invisible list of synthetic tree summary statistics:

- date_created String. The creation date of the tree.
- num_source_studies Integer. The number of studies (publications)used as sources.
- num_source_trees The number of trees used as sources (may be >1 tree per study).
- taxonomy_version The Open Tree Taxonomy version used as a source.
- filtered_flags List. Taxa with these taxonomy flags were not used in construction of the tree.
- root List. Describes the root node:
 - node_id String. The canonical identifier of the node.
 - num_tips Numeric. The number of descendant tips.
 - taxon A list of taxonomic properties:
 - * ott_id Numeric. The OpenTree Taxonomy ID (ott_id).
 - * name String. The taxonomic name of the queried node.
 - * unique_name String. The string that uniquely identifies the taxon in OTT.
 - * rank String. The taxonomic rank of the taxon in OTT.
 - * tax_sources List. A list of identifiers for taxonomic sources, such as other taxonomies, that define taxa judged equivalent to this taxon.
- source_list List. Present only if include_source_list is TRUE. The sourceid ordering is the precedence order for synthesis, with relationships from earlier trees in the list having priority over those from later trees in the list. See source_id_map below for study details.
- source_id_map Named list of lists. Present only if include_source_list is TRUE. Names
 correspond to the 'sourceids' used in source_list above. Source trees will have the following properties:
 - git_sha String. The git SHA identifying a particular source version.
 - tree_id String. The tree id associated with the study id used.
 - study_id String. The study identifier. Will typically include a prefix ("pg_" or "ot_").
- synth_id The unique string for this version of the tree.

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See Also

source_list to explore the list of studies used in the synthetic tree (see example).

Examples

```
## Not run:
res <- tol_about()
tax_sources(res)
ott_id(res)
studies <- source_list(tol_about(include_source_list=TRUE))
## End(Not run)</pre>
```

tol_induced_subtree

Subtree from the Open Tree of Life

Description

Return the induced subtree on the synthetic tree that relates a list of nodes.

Usage

```
tol_induced_subtree(
  ott_ids = NULL,
  node_ids = NULL,
  label_format = NULL,
  file,
  ...
)
```

Arguments

ott_ids	Numeric vector. OTT ids indicating nodes to be used as tips in the induced tree.
node_ids	Character vector. Node ids indicating nodes to be used as tips in the induced tree.
label_format	Character. Defines the label type; one of "name", "id", or "name_and_id" (the default).
file	If specified, the function will write the subtree to a file in newick format.
	additional arguments to customize the API call (see rotl for more information).

Details

Return a tree with tips corresponding to the nodes identified in the input set that is consistent with the topology of the current synthetic tree. This tree is equivalent to the minimal subtree induced on the draft tree by the set of identified nodes.

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Value

If no value is specified to the file argument (default), a phylogenetic tree of class phylo.

Otherwise, the function returns invisibly a logical indicating whether the file was successfully created.

Note that the tree returned when specifying a file name with the file argument is the "raw" Newick string returned by Open Tree of Life. This string contains singleton nodes, and therefore will be different from the tree returned as a phylo object which will not contain these singleton nodes.

Examples

tol_lineage

Node info

Description

Get summary information about a node in the synthetic tree

Usage

```
tol_lineage(tax, ...)

tol_node_info(ott_id = NULL, node_id = NULL, include_lineage = FALSE, ...)

## S3 method for class 'tol_node'
tax_rank(tax, ...)

## S3 method for class 'tol_node'
tax_sources(tax, ...)

## S3 method for class 'tol_node'
unique_name(tax, ...)

## S3 method for class 'tol_node'
tax_name(tax, ...)
```

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```
## S3 method for class 'tol_node'
ott_id(tax, ...)
## S3 method for class 'tol_node'
source_list(tax, ...)
## S3 method for class 'tol_node'
tax_lineage(tax, ...)
## S3 method for class 'tol_node'
tol_lineage(tax, ...)
```

Arguments

tax an object returned by tol_node_info.
... additional arguments to customize the API call (see ?rotl for more information)
ott_id Numeric. The OpenTree taxonomic identifier.
node_id Character. The OpenTree node identifier.
include_lineage

Logical (default = FALSE). Whether to return the lineage of the node from the synthetic tree.

Details

Returns summary information about a node in the graph. The node of interest may be specified using either a node id or an taxon id, but not both. If the specified node or OTT id is not in the graph, an error will be returned.

If the argument include_lineage=TRUE is used, you can use tax_lineage() or tol_lineage to return the taxonomic information or the node information for all the ancestors to this node, down to the root of the tree.

Value

tol_node_info returns an invisible list of summary information about the queried node:

- node_id String. The canonical identifier of the node.
- num_tips Numeric. The number of descendant tips.
- partial_path_of List. The edge below this synthetic tree node is compatible with the edge below each of these input tree nodes (one per tree). Each returned element is reported as sourceid:nodeid.
- query The node id that resolved to this node. This can differ from the node_id field if the query id is not canonical.
- taxon A list of taxonomic properties. Only returned if the queried node is a taxon. Each source has:
 - ott_id Numeric. The OpenTree Taxonomy ID (ottID).
 - name String. The taxonomic name of the queried node.

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- unique_name String. The string that uniquely identifies the taxon in OTT.
- rank String. The taxonomic rank of the taxon in OTT.
- tax_sources List. A list of identifiers for taxonomic sources, such as other taxonomies, that define taxa judged equivalent to this taxon.

The following properties list support/conflict for the node across synthesis source trees. All properties involve sourceid keys and nodeid values (see source_id_map below).

- supported_by List. Input tree nodes (one per tree) that support this synthetic tree node. Each returned element is reported as sourceid:nodeid.
- terminal List. Input tree nodes (one per tree) that are equivalent to this synthetic tree node (via an exact mapping, or the input tree terminal may be the only terminal descended from this synthetic tree node. Each returned element is reported as sourceid:nodeid.
- conflicts_with Named list of lists. Names correspond to sourceid keys. Each list contains input tree node ids (one or more per tree) that conflict with this synthetic node.

tol_lineage and tax_lineage return data frames. tol_lineage indicate for each ancestor its node identifier, the number of tips descending from that node, and whether it corresponds to a taxonomic level.

Examples

```
## Not run:
birds <- tol_node_info(ott_id=81461, include_lineage=TRUE)
source_list(birds)
tax_rank(birds)
ott_id(birds)
tax_lineage(birds)
tol_lineage(birds)
## End(Not run)</pre>
```

tol_mrca

MRCA of taxa from the synthetic tree

Description

Most Recent Common Ancestor for a set of nodes

Usage

```
tol_mrca(ott_ids = NULL, node_ids = NULL, ...)
## S3 method for class 'tol_mrca'
tax_sources(tax, ...)
## S3 method for class 'tol_mrca'
unique_name(tax, ...)
```

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```
## S3 method for class 'tol_mrca'
tax_name(tax, ...)
## S3 method for class 'tol_mrca'
tax_rank(tax, ...)
## S3 method for class 'tol_mrca'
ott_id(tax, ...)
## S3 method for class 'tol_mrca'
source_list(tax, ...)
```

Arguments

ott_ids	Numeric vector. The ott ids for which the MRCA is desired.
node_ids	Character vector. The node ids for which the MRCA is desired.
	additional arguments to customize the API call (see rot1 for more information).
tax	an object returned by tol_mrca().

Details

Get the MRCA of a set of nodes on the current synthetic tree. Accepts any combination of node ids and ott ids as input. Returns information about the most recent common ancestor (MRCA) node as well as the most recent taxonomic ancestor (MRTA) node (the closest taxonomic node to the MRCA node in the synthetic tree; the MRCA and MRTA may be the same node). If they are the same, the taxonomic information will be in the mrca slot, otherwise they will be in the nearest_taxon slot of the list. If any of the specified nodes is not in the synthetic tree an error will be returned.

Taxonomic methods (tax_sources(), ott_id(), unique_name(), ...) are available on the objects returned by tol_mrca(). If the MRCA node is MRTA, the name of the object returned by these methods will start with 'ott', otherwise it will start with 'mrca'.

Value

An invisible list of the MRCA node properties:

- mrca List of node properties.
 - node_id String. The canonical identifier of the node.
 - num tips Numeric. The number of descendant tips.
 - taxon A list of taxonomic properties. Only returned if the queried node is a taxon. (If the node is not a taxon, a nearest_taxon list is returned (see below)).
 - * ott_id Numeric. The OpenTree Taxonomy ID (ottID).
 - * name String. The taxonomic name of the queried node.
 - * unique_name String. The string that uniquely identifies the taxon in OTT.
 - * rank String. The taxonomic rank of the taxon in OTT.
 - * tax_sources List. A list of identifiers for taxonomic sources, such as other taxonomies, that define taxa judged equivalent to this taxon.

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The following properties list support/conflict for the node across synthesis source trees. All properties involve sourceid keys and nodeid values (see source_id_map below) Not all properties are are present for every node.

- partial_path_of List. The edge below this synthetic tree node is compatible with the edge below each of these input tree nodes (one per tree). Each returned element is reported as sourceid:nodeid.
- supported_by List. Input tree nodes (one per tree) that support this synthetic tree node.
 Each returned element is reported as sourceid:nodeid.
- terminal List. Input tree nodes (one per tree) that are equivalent to this synthetic tree node (via an exact mapping, or the input tree terminal may be the only terminal descended from this synthetic tree node. Each returned element is reported as sourceid:nodeid.
- conflicts_with Named list of lists. Names correspond to sourceid keys. Each list contains
 input tree node ids (one or more per tree) that conflict with this synthetic node.
- nearest_taxon A list of taxonomic properties of the nearest rootward taxon node to the MRCA node. Only returned if the MRCA node is a not taxon (otherwise the taxon list above is returned).
 - ott_id Numeric. The OpenTree Taxonomy ID (ottID).
 - name String. The taxonomic name of the queried node.
 - unique_name String. The string that uniquely identifies the taxon in OTT.
 - rank String. The taxonomic rank of the taxon in OTT.
 - tax_sources List. A list of identifiers for taxonomic sources, such as other taxonomies, that define taxa judged equivalent to this taxon.
- source_id_map Named list of lists. Names correspond to the sourceid keys used in the support/conflict properties of the mrca list above. Source trees will have the following properties:
 - git_sha The git SHA identifying a particular source version.
 - tree_id The tree id associated with the study id used.
 - study_id The study identifier. Will typically include a prefix ("pg_" or "ot_").

The only sourceid that does not correspond to a source tree is the taxonomy, which will have the name "ott"+'taxonomy_version', and the value is the ott_id of the taxon in that taxonomy version. "Taxonomy" will only ever appear in supported_by.

```
## Not run:
birds_mrca <- tol_mrca(ott_ids=c(412129, 119214))
ott_id(birds_mrca)
tax_sources(birds_mrca)
## End(Not run)</pre>
```

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tol_subtree	Extract a subtree from the synthetic tree	

Description

Extract a subtree from the synthetic tree from an Open Tree node id.

Usage

```
tol_subtree(ott_id = NULL, node_id = NULL, label_format = NULL, file, ...)
```

Arguments

ott_id	Numeric. The ott id of the node in the tree that should serve as the root of the tree returned.
node_id	Character. The node id of the node in the tree that should serve as the root of the tree returned.
label_format	Character. Defines the label type; one of "name", "id", or "name_and_id" (the default).
file	If specified, the function will write the subtree to a file in newick format.
	additional arguments to customize the API call (see rot1 for more information).

Details

Given a node, return the subtree of the synthetic tree descended from that node. The start node may be specified using either a node id or an ott id, but not both. If the specified node is not in the synthetic tree an error will be returned. There is a size limit of 25000 tips for this method.

Value

If no value is specified to the file argument (default), a phylogenetic tree of class phylo. Otherwise, the function returns invisibly a logical indicating whether the file was successfully created.

```
## Not run:
res <- tol_subtree(ott_id=241841)
## End(Not run)</pre>
```

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