

Package ‘rcol’

February 5, 2021

Type Package

Title Catalogue of Life Client

Description Client for the Catalogue of Life ('CoL') (<<https://www.catalogueoflife.org/>>); based on the new 'CoL' service, not the old one. Catalogue of Life is a database of taxonomic names. Includes functions for each of the API methods, including searching for names, and more.

Version 0.1.0

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URL <https://docs.ropensci.org/rcol/> (docs),
<https://github.com/ropensci/rcol>

BugReports <https://github.com/ropensci/rcol/issues>

LazyData true

Encoding UTF-8

Language en-US

Imports crul, tibble, jsonlite, data.table, glue

Suggests testthat, vcr

RoxygenNote 7.1.1

X-schema.org-applicationCategory Biology

X-schema.org-keywords biology, science, API, web, api-client,
taxonomy, species

X-schema.org-isPartOf <https://ropensci.org>

NeedsCompilation no

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Repository CRAN

Date/Publication 2021-02-05 08:30:03 UTC

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Description

Catalogue of Life (CoL) Client

Note

CoL API docs: <https://api.catalogueoflife.org/>

Author(s)

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cp_children	<i>Children</i>
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Description

Children

Usage

```
cp_children(dataset_key, taxon_id, ...)
```

Arguments

dataset_key	(character/integer/numeric) dataset identifier
taxon_id	(character/integer/numeric) taxon identifier
...	curl options passed on to crul::verb-GET

Value

list with two slots

- result (data.frame/tibble): results, a zero row data.frame if no results found
- meta (data.frame/tibble): number of results found

Examples

```
chk <- function(x) {
  z <- tryCatch(crul::ok(x), error = function(e) e)
  if (inherits(z, "error")) FALSE else z
}
if (chk("https://api.catalogueoflife.org/version")) {
  z <- cp_children(dataset_key=1000, taxon_id='1')
  z
  z$result
  if (NROW(z$result) > 0) {
    z$result$scientificName
    z$result$created
  }
}
```

cp_classification	<i>Classification</i>
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Description

Classification

Usage

```
cp_classification(dataset_key, taxon_id, ...)
```

Arguments

dataset_key (character/integer/numeric) dataset identifier
taxon_id (character/integer/numeric) taxon identifier
... curl options passed on to [crul::verb-GET](#)

Value

a data.frame/tibble with results, a zero row data.frame if no results found

Examples

```

if (cp_up("/dataset/1000/taxon/10/classification")) {
  cp_classification(dataset_key=1000, taxon_id=10)
}
## Not run:
cp_classification(dataset_key=1000, taxon_id=20)
cp_classification(dataset_key=3,
  taxon_id="6565450e-1cf2-4dc2-acbb-db728e42e635")

## End(Not run)

```

cp_datasets

Datasets

Description

Datasets

Usage

```
cp_datasets(q = NULL, start = 0, limit = 10, ...)
```

```
cp_dataset(dataset_keys, ...)
```

Arguments

q	(character) main query string. optional
start	(integer) requested number of offset records. Default: 0
limit	(integer) requested number of maximum records to be returned. Default: 10; max: 1000
...	curl options passed on to verb-GET
dataset_keys	(character) one or more dataset keys. required

Details

for `cp_dataset()`, separate http requests are made for each dataset key. unfortunately, the output of `cp_dataset()` is a list for each dataset key because the nested structure of the data is hard to rectangularize

Value

list with two slots

- `result` (data.frame/tibble): results, a zero row data.frame if no results found
- `meta` (data.frame/tibble): number of results found

Examples

```

if (cp_up("/dataset")) {
  cp_datasets(limit = 1)
}
## Not run:
cp_datasets(q = "life")
cp_dataset(dataset_keys = 1000)
cp_dataset(dataset_keys = c(3, 1000, 1014))

## End(Not run)

```

cp_ds

*Datasets API route catch all method***Description**

Datasets API route catch all method

Usage

```
cp_ds(route, ..., .list = list())
```

Arguments

route	(character) an API route. the /dataset route part is added internally; so just include the route following that. required.
...	named parameters, passed on to <code>glue::glue()</code> . required. param names must match must match names given in the route. For example, if you have route = <code>\\{key\\}/name\\{id\\}</code> , then you need to pass in a key and an id parameter. The names in the route (here, key and id) don't have to match the names in the API route you are trying to use - they just need to match the named parameters you pass in. Having said that, it may be easier to remember what you're doing if you match the names to the route parts.
.list	a named list. instead of passing in named parameters through ..., you can pre-prepare a named list and give to this parameter

Details

There are A LOT of datasets API routes. Instead of making an R function for each route, we have R functions for some of the "more important" routes, then `cp_ds()` will allow you to make requests to the remainder of the datasets API routes.

Value

output varies depending on the route requested, but output will always be a named list. when no results found, an error message will be returned

Not supported dataset routes

Some dataset routes do not return JSON so we don't support those. Thus far, the only route we don't support is /dataset/{key}/logo

Examples

```
## Not run:
cp_ds(route = "{key}/tree", key = "1000")
cp_ds(route = "{key}/tree", key = "1014")
cp_ds(route = "{key}/name/{id}", key = 1005, id = 100003)

# pass a named list to the .list parameter
args <- list(key = 1005, id = 100003)
cp_ds("{key}/name/{id}", .list = args)

## End(Not run)
```

cp_importer

Importer metrics

Description

Importer metrics

Usage

```
cp_importer(
  dataset_key = NULL,
  state = NULL,
  running = FALSE,
  start = 0,
  limit = 10,
  ...
)
```

Arguments

dataset_key	(character) a dataset key to filter by. optional
state	(character) filter listed import metrics by their state, e.g. the last failed import. one of: downloading, processing, inserting, unchanged, finished, canceled, failed. optional
running	(logical) if only a list of running imports should be returned. default: FALSE. optional
start	(integer) requested number of offset records. Default: 0
limit	(integer) requested number of maximum records to be returned. Default: 10; max: 1000
...	curl options passed on to verb-GET

Value

a named list, with slots `offset` (integer), `limit` (integer), `total` (integer), `result` (list), `empty` (boolean), and `last` (boolean). The `result` slot is a list itself, with any number of results as named lists

Examples

```
if (cp_up("/importer?limit=1")) {
  cp_importer(limit = 1)
}
```

cp_name_match	<i>Name Matching</i>
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Description

Match name against the name index

Usage

```
cp_name_match(
  q = NULL,
  rank = NULL,
  code = NULL,
  trusted = NULL,
  ver_bose = NULL,
  start = 0,
  limit = 10,
  ...
)
```

Arguments

<code>q</code>	(character) scientific name to match
<code>rank</code>	(character) rank to restrict matches to. one of: domain, realm, subrealm, superkingdom, kingdom, subkingdom, infrakingdom, superphylum, phylum, subphylum, infraphylum, superclass, class, subclass, infraclass, parvclass, superdivision, division, subdivision, infradivision, superlegion, legion, sublegion, infralegion, supercohort, cohort, subcohort, infracohort, gigaorder, magnorder, grandorder, mirorder, superorder, order, nanorder, hypoorder, minorder, suborder, infraorder, parvorder, megafamily, grandfamily, superfamily, epifamily, family, subfamily, infrafamily, supertribe, tribe, subtribe, infratribe, suprageneric_name, genus, subgenus, infragenus, supersection, section, subsection, superseries, series, subseries, infrageneric_name, species_aggregate, species, infraspecific_name, grex, subspecies, cultivar_group, convariety, infrasubspecific_name, proles, natio, aberration, morph, variety, subvariety, form, subform, pathovar, biovar, chemovar, morphovar, phagovar, serovar, chemoform, forma_specialis, cultivar, strain, other, unranked

code	(character) nomenclatural code to restrict matches to. one of: bacterial, botanical, cultivars, viral, zoological, phytosociological
trusted	(logical) if TRUE, unmatched name will be inserted into the names index. default: FALSE
ver_bose	(logical) if TRUE, list alternatively considered name matches. default: FALSE
start	(integer) requested number of offset records. Default: 0
limit	(integer) requested number of maximum records to be returned. Default: 10; max: 1000
...	curl options passed on to verb-GET

Details

Matches by the canonical name, it's authorship and rank. Authorship matching is somewhat loose, but name matching is quite strict and only allows for a few common misspellings frequently found in epithets (silent h, gender suffix, double letters, i/y), but not in uninomials. Suprageneric ranks are all considered to be the same, otherwise a different rank results in a different match.

Value

a named list, with slots name (list), type (character), alternatives (data.frame), and nameKey (integer)

Examples

```
if (cp_up("/name/matching?q=Apis")) {
  cp_name_match(q="Apis")
}
## Not run:
cp_name_match(q="Agapostemon")
cp_name_match(q="Apis mellifera")
cp_name_match(q="Apis mellifer") # no fuzzy match apparently

## End(Not run)
```

cp_nu_search

Name Usage: Search

Description

Name Usage: Search

Usage

```

cp_nu_search(
  q = NULL,
  dataset_key = NULL,
  min_rank = NULL,
  max_rank = NULL,
  content = NULL,
  highlight = NULL,
  reverse = NULL,
  fuzzy = NULL,
  type = NULL,
  nomstatus = NULL,
  status = NULL,
  issue = NULL,
  published_in = NULL,
  facet = NULL,
  sortBy = NULL,
  start = 0,
  limit = 10,
  ...
)

```

Arguments

q	(character) vector of one or more scientific names
dataset_key	(character) dataset key
min_rank, max_rank	(character) filter by rank. one of: domain, superkingdom, kingdom, subkingdom, infrakingdom, superphylum, phylum, subphylum, infraphylum, superclass, class, subclass, infraclass, parvclass, superlegion, legion, sublegion, infralegion, supercohort, cohort, subcohort, infracohort, magnorder, superorder, grandorder, order, suborder, infraorder, parvorder, superfamily, family, subfamily, infrafamily, supertribe, tribe, subtribe, infratribe, suprageneric name, genus, subgenus, infragenus, supersection, section, subsection, superseries, series, subseries, infragenic name, species aggregate, species, infraspecific name, grex, subspecies, cultivar group, convariety, infrasubspecific name, proles, natio, aberration, morph, variety, subvariety, form, subform, pathovar, biovar, chemovar, morphovar, phago-var, serovar, chemoform, forma specialis, cultivar, strain, other, unranked
content	(character) one of: 'scientific_name' or 'authorship'
highlight	(logical) TRUE or FALSE. default: NULL
reverse	(logical) TRUE or FALSE. default: NULL
fuzzy	(logical) TRUE or FALSE. default: NULL
type	(character) one of: 'prefix', 'whole_words', 'exact'
nomstatus	(character) filter by nomenclatural status. one of: ok, unavailable, illegitimate, variant, conserved, rejected, doubtful, unevaluated

status	(character) filter by taxonomic status. one of: accepted, doubtful, ambiguous synonym
issue	(character) filter by issue found
published_in	(character) reference id to filter names by
facet	(character) request a facet to be returned. one of: dataset_key, rank, nom_status, status, issue, type, field. facet limit default: 50
sortBy	(character) one of: "relevance", "name", "taxonomic", "index_name_id", or "native"
start	(integer) requested number of offset records. Default: 0
limit	(integer) requested number of maximum records to be returned. Default: 10; max: 1000
...	curl options passed on to verb-GET

Value

list with two slots

- result (data.frame/tibble): results, a zero row data.frame if no results found
- meta (data.frame/tibble): number of results found

Examples

```
if (cp_up("/nameusage/search?q=Apis")) {
  cp_nu_search(q="Apis", limit = 1)
}
## Not run:
cp_nu_search(q="Agapostemon")
cp_nu_search(q="Agapostemon", dataset_key = 3)
cp_nu_search(q="Agapostemon", min_rank = "genus")
cp_nu_search(q="Agapostemon", nomstatus = "doubtful")
cp_nu_search(q="Agapostemon", status = "accepted")
cp_nu_search(q="Bombus", facet = "rank")
cp_nu_search(q="Agapostemon", dataset_key = 3, hasField="uninomial")

x <- cp_nu_search(q="Poa")
x
x$result
x$result$usage
x$result$usage$name

## End(Not run)
```

cp_nu_suggest	<i>Name Usage: Suggest</i>
---------------	----------------------------

Description

Name Usage: Suggest

Usage

```
cp_nu_suggest(
  q,
  dataset_key,
  fuzzy = FALSE,
  min_rank = NULL,
  max_rank = NULL,
  sort = NULL,
  reverse = FALSE,
  accepted = FALSE,
  limit = 10,
  ...
)
```

Arguments

q	(character) main query string. required
dataset_key	(character) a dataset key. required
fuzzy	(logical) Whether or not to do fuzzy search (default: FALSE)
min_rank, max_rank	(character) See rank options in cp_name_match()
sort	(character) one of name, taxonomic, index_name_id, native, relevance
reverse	(logical) reverse order i assume (default: FALSE)
accepted	(logical) limit to accepted names (default: FALSE)
limit	(integer) requested number of maximum records to be returned. Default: 10; max: 1000
...	curl options passed on to crul::verb-GET

Value

a data.frame/tibble of results. a zero row data.frame if no results

Examples

```
if (cp_up("/dataset/3/nameusage/suggest?q=Apis")) {
  cp_nu_suggest(q="Apis", 3)
}
```

cp_parser

Name Parser

Description

Name Parser

Usage

```
cp_parser(names, ...)
```

Arguments

names (character) one or more scientific names to parse
... curl options passed on to [crul::verb-POST](#)

Value

tibble, with one row for each parsed name

Examples

```
## Not run:  
cp_parser(names = "Apis mellifera")  
cp_parser(names = c("Apis mellifera", "Homo sapiens var. sapiens"))  
  
## End(Not run)
```

cp_vocab

CoL Vocabularies

Description

CoL Vocabularies

Usage

```
cp_vocab(vocab, ...)
```

Arguments

vocab (character) a vocabulary name
... curl options passed on to [crul::verb-GET](#)

Value

character vector of words

Examples

```
## Not run:  
cp_vocab("rank")  
cp_vocab("datasetorigin")  
cp_vocab("datasettype")  
cp_vocab("matchtype")  
cp_vocab("taxonomicstatus")  
  
## End(Not run)
```

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