Package ‘rgbif’

March 9, 2015

Title Interface to the Global Biodiversity Information Facility API

Description  A programmatic interface to the Web Service methods provided by the Global Biodiversity Information Facility (GBIF). GBIF is a database of species occurrence records from sources all over the globe. rgbif includes functions for searching for taxonomic names, retrieving information on data providers, getting species occurrence records, and getting counts of occurrence records.

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Date 2015-03-09

URL https://github/ropensci/rgbif

BugReports https://github/ropensci/rgbif/issues

LazyData true

VignetteBuilder knitr

Imports XML, plyr, ggplot2, maps, httr, rgeos, sp, data.table,
    stringr, whisker, magrittr, jsonlite (>= 0.9.12)

Suggests roxygen2, testthat, knitr, reshape2

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R topics documented:

<table>
<thead>
<tr>
<th>Package/Function</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>rgbif-package</td>
<td>2</td>
</tr>
<tr>
<td>check_wkt</td>
<td>3</td>
</tr>
<tr>
<td>count_facet</td>
<td>4</td>
</tr>
<tr>
<td>datasets</td>
<td>5</td>
</tr>
<tr>
<td>dataset_metrics</td>
<td>6</td>
</tr>
<tr>
<td>dataset_search</td>
<td>7</td>
</tr>
<tr>
<td>dataset_suggest</td>
<td>9</td>
</tr>
<tr>
<td>elevation</td>
<td>11</td>
</tr>
<tr>
<td>gbifmap</td>
<td>12</td>
</tr>
<tr>
<td>gbif_bbox2wkt</td>
<td>14</td>
</tr>
<tr>
<td>gbif_issues</td>
<td>15</td>
</tr>
<tr>
<td>gbif_names</td>
<td>15</td>
</tr>
<tr>
<td>gbif_photos</td>
<td>16</td>
</tr>
<tr>
<td>installations</td>
<td>17</td>
</tr>
<tr>
<td>isocodes</td>
<td>18</td>
</tr>
<tr>
<td>name_backbone</td>
<td>19</td>
</tr>
<tr>
<td>name_lookup</td>
<td>20</td>
</tr>
<tr>
<td>name_suggest</td>
<td>24</td>
</tr>
<tr>
<td>name_usage</td>
<td>25</td>
</tr>
<tr>
<td>networks</td>
<td>27</td>
</tr>
<tr>
<td>nodes</td>
<td>29</td>
</tr>
<tr>
<td>occ_count</td>
<td>31</td>
</tr>
<tr>
<td>occ_fields</td>
<td>32</td>
</tr>
<tr>
<td>occ_get</td>
<td>33</td>
</tr>
<tr>
<td>occ_issues</td>
<td>34</td>
</tr>
<tr>
<td>occ_issues_lookup</td>
<td>36</td>
</tr>
<tr>
<td>occ_metadata</td>
<td>36</td>
</tr>
<tr>
<td>occ_search</td>
<td>37</td>
</tr>
<tr>
<td>organizations</td>
<td>48</td>
</tr>
<tr>
<td>parsenames</td>
<td>49</td>
</tr>
<tr>
<td>rgbif-defunct</td>
<td>50</td>
</tr>
<tr>
<td>rgb_country_codes</td>
<td>50</td>
</tr>
<tr>
<td>taxrank</td>
<td>51</td>
</tr>
<tr>
<td>typestatus</td>
<td>51</td>
</tr>
</tbody>
</table>

Index 52

---

**rgbif-package**

*Interface to the Global Biodiversity Information Facility API.*

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**Description**

*rgbif: A programmatic interface to the Web Service methods provided by the Global Biodiversity Information Facility.*
About

This package gives you access to data from GBIF http://www.gbif.org/ via their API.

A note about the old GBIF API

The old GBIF API was at http://data.gbif.org/tutorial/services, but is now defunct - that is, not available anymore. We used to have functions that worked with the old API, but those functions are now not available anymore because GBIF made the old API defunct.

Documentation for the GBIF API

- summary http://www.gbif.org/developer/summary - Summary of the GBIF API
- registry http://www.gbif.org/developer/registry - Metadata on datasets, and contributing organizations
- species names http://www.gbif.org/developer/species - Species names and metadata
- occurrences http://www.gbif.org/developer/occurrence - Occurrences
- maps http://www.gbif.org/developer/maps - Maps - these APIs are not implemented in rgbif, and are meant more for intergration with web based maps.

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---

check_wkt

Check input WKT

Description

Check input WKT

Usage

check_wkt(wkt = NULL)

Arguments

wkt A Well Known Text object
count_facet

Facetted count occurrence search.

Description
Facetted count occurrence search.

Usage
count_facet(keys = NULL, by = "country", countries = 10, removezeros = FALSE)

Arguments
keys (numeric) GBIF keys, a vector.
by (character) One of georeferenced, basisOfRecord, country, or publishingCountry.
countries (numeric) Number of countries to facet on, or a vector of country names
removezeros (logical) Default is FALSE

Examples
## Not run:
# Select number of countries to facet on
count_facet(by = "country", countries = 3, removezeros = TRUE)
# Or, pass in country names
count_facet(by = c("AR"), removezeros = TRUE)


```r
keys <- sapply(spplist, function(x) name_backbone(x, rank="species")$usageKey)
count_facet(keys, by='country', countries=3, removezeros = TRUE)
count_facet(keys, by='country', countries=3, removezeros = FALSE)
count_facet(keys, by='country', countries=20, removezeros = TRUE)

# Pass in country names instead
countries <- isocodes$code[1:10]
count_facet(by='country', countries=countries, removezeros = TRUE)

# get occurrences by georeferenced state
# across all records
count_facet(by='georeferenced')

## by keys
out <- count_facet(keys, by='georeferenced')
library("reshape2")
dcast(out, .id ~ georeferenced)

## by basisOfRecord
count_facet(by="basisOfRecord")

## End(Not run)
```

**datasets**

Search for datasets and dataset metadata.

### Description

Search for datasets and dataset metadata.

### Usage

```r
datasets(data = "all", type = NULL, uuid = NULL, query = NULL, id = NULL, limit = 100, start = NULL, …)
```

### Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>data</td>
<td>The type of data to get. Default is all data.</td>
</tr>
<tr>
<td>type</td>
<td>Type of dataset, options include OCCURRENCE, etc.</td>
</tr>
<tr>
<td>uuid</td>
<td>UUID of the data node provider. This must be specified if data is anything other than 'all'.</td>
</tr>
<tr>
<td>query</td>
<td>Query term(s). Only used when data='all'</td>
</tr>
<tr>
<td>id</td>
<td>A metadata document id.</td>
</tr>
<tr>
<td>limit</td>
<td>Number of records to return. Default: 100. Maximum: 1000.</td>
</tr>
<tr>
<td>start</td>
<td>Record number to start at. Default: 0. Use in combination with limit to page through results.</td>
</tr>
</tbody>
</table>
Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.

Value
A list.

References
http://www.gbif.org/developer/registry#datasets

Examples
```r
## Not run:
datasets(limit=5)
datasets(type="OCCURRENCE")
datasets(uuid="a6998220-7e3a-485d-9cd6-73076bd85657")
datasets(data='contact', uuid="a6998220-7e3a-485d-9cd6-73076bd85657")
datasets(data='metadata', uuid="a6998220-7e3a-485d-9cd6-73076bd85657")
datasets(data='metadata', uuid="a6998220-7e3a-485d-9cd6-73076bd85657", id=598)
datasets(data=c('deleted','duplicate'))
datasets(data=c('deleted','duplicate'), limit=1)

# httr options
library(httr)
res <- datasets(data=c('deleted','duplicate'), config=progress())

## End(Not run)
```

Description
Get details on a GBIF dataset.

Usage
dataset_metrics(uuid, ...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>uuid</td>
<td>(character) One or more dataset UUIDs. See examples.</td>
</tr>
<tr>
<td>...</td>
<td>Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.</td>
</tr>
</tbody>
</table>

References
http://www.gbif.org/developer/registry#datasetMetrics
dataset_search

Examples

```r
## Not run:
dataset_metrics(uuid='3f8a1297-3259-4700-91fc-acc4170b27ce')
dataset_metrics(uuid='66dd0960-2d7d-46ee-a491-87b9adcf7b1')
dataset_metrics(uuid=c('3f8a1297-3259-4700-91fc-acc4170b27ce',
  '66dd0960-2d7d-46ee-a491-87b9adcf7b1'))

library("httr")
dataset_metrics(uuid='66dd0960-2d7d-46ee-a491-87b9adcf7b1', config=verbose())

## End(Not run)
```

dataset_search  

Search datasets in GBIF.

Description

This function does not search occurrence data, only metadata on the datasets that contain occurrence data.

Usage

```r
dataset_search(query = NULL, country = NULL, type = NULL,
  keyword = NULL, owningOrg = NULL, publishingOrg = NULL,
  hostingOrg = NULL, publishingCountry = NULL, decade = NULL,
  facet = NULL, facetMincount = NULL, facetMultiselect = NULL,
  limit = 100, start = NULL, pretty = FALSE, return = "all", ...)
```

Arguments

- **query**: Query term(s) for full text search. The value for this parameter can be a simple word or a phrase. Wildcards can be added to the simple word parameters only, e.g. `q=*puma*`
- **country**: NOT YET IMPLEMENTED. Filters by country as given in `isocodes$gbif_name`, e.g. `country=CANADA`.
- **type**: Type of dataset, options include OCCURRENCE, etc.
- **keyword**: Keyword to search by. Datasets can be tagged by keywords, which you can search on. The search is done on the merged collection of tags, the dataset keywordCollections and temporalCoverages.
- **owningOrg**: Owning organization. A uuid string. See `organizations`
- **publishingOrg**: Publishing organization. A uuid string. See `organizations`
- **hostingOrg**: Hosting organization. A uuid string. See `organizations`
- **publishingCountry**: Publishing country. See options at `isocodes$gbif_name`
dataset_search

**decade**
Decade, e.g., 1980. Filters datasets by their temporal coverage broken down to decades. Decades are given as a full year, e.g. 1880, 1960, 2000, etc, and will return datasets wholly contained in the decade as well as those that cover the entire decade or more. Facet by decade to get the break down, e.g. /search?facet=DECADE&facet_only=true (see example below)

**facet**
A list of facet names used to retrieve the 100 most frequent values for a field. Allowed facets are: datasetKey, highertaxonKey, rank, status, extinct, habitat, and nameType. Additionally threat and nomenclaturalStatus are legal values but not yet implemented, so data will not yet be returned for them.

**facetMincount**
Used in combination with the facet parameter. Set facetMincount=# to exclude facets with a count less than #, e.g. http://bit.ly/1bMdByP only shows the type value 'ACCEPTED' because the other statuses have counts less than 7,000,000

**facetMultiselect**
Used in combination with the facet parameter. Set facetMultiselect=true to still return counts for values that are not currently filtered, e.g. http://bit.ly/19YLXPO still shows all status values even though status is being filtered by status=ACCEPTED

**limit**
Number of records to return. Default: 100. Maximum: 1000.

**start**
Record number to start at. Default: 0. Use in combination with limit to page through results.

**pretty**
Print informative metadata using cat. Not easy to manipulate output though.

**return**
What to return. One of meta, descriptions, data, facets, or all (Default).

... Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.

**Value**
A data.frame, list, or message printed to console (using pretty=TRUE).

**References**
http://www.gbif.org/developer/registry#datasetSearch

**Examples**

```
## Not run:
# Gets all datasets of type "OCCURRENCE".
dataset_search(type="OCCURRENCE")

# Gets all datasets tagged with keyword "france".
dataset_search(keyword="france")

# Fulltext search for all datasets having the word "amsterdam" somewhere in
# its metadata (title, description, etc).
dataset_search(query="amsterdam")

# Limited search
dataset_search(type="OCCURRENCE", limit=2)
dataset_search(type="OCCURRENCE", limit=2, start=10)
```
dataset_suggest

Suggest datasets in GBIF.

Description
Search that returns up to 20 matching datasets. Results are ordered by relevance.

Usage
dataset_suggest(query = NULL, country = NULL, type = NULL,
subtype = NULL, keyword = NULL, owningOrg = NULL,
publishingOrg = NULL, hostingOrg = NULL, publishingCountry = NULL,
decade = NULL, continent = NULL, limit = 100, start = NULL,
pretty = FALSE, description = FALSE, ...)

Arguments

query        Query term(s) for full text search. The value for this parameter can be a simple
            word or a phrase. Wildcards can be added to the simple word parameters only,
e.g. q=*puma*
country     NOT YET IMPLEMENTED. Filters by country as given in isocodes$gbif_name,
e.g. country=CANADA.
dataset_suggest

type
Type of dataset, options include OCCURRENCE, etc.

subtype
NOT YET IMPLEMENTED. Will allow filtering of datasets by their dataset subtypes, DC or EML.

keyword
Keyword to search by. Datasets can be tagged by keywords, which you can search on. The search is done on the merged collection of tags, the dataset keywordCollections and temporalCoverages.

owningOrg
Owning organization. A uuid string. See organizations

publishingOrg
Publishing organization. A uuid string. See organizations

hostingOrg
Hosting organization. A uuid string. See organizations

publishingCountry
Publishing country. See options at isocodes$gbif_name

decade
Decade, e.g., 1980. Filters datasets by their temporal coverage broken down to decades. Decades are given as a full year, e.g. 1880, 1960, 2000, etc, and will return datasets wholly contained in the decade as well as those that cover the entire decade or more. Facet by decade to get the break down, e.g. /search?facet=DECADE&facet_only=true (see example below)

continent
Not yet implemented, but will eventually allow filtering datasets by their continent(s) as given in our Continent enum.

limit
Number of records to return. Default: 100. Maximum: 1000.

start
Record number to start at. Default: 0. Use in combination with limit to page through results.

pretty
Print informative metadata using cat. Not easy to manipulate output though.

description
Return descriptions only (TRUE) or all data (FALSE, default)

... Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.

Value
A data.frame, list, or message printed to console (using pretty=TRUE).

References
httpZOOwwwNgbifNorgOdeveloperOregistryCdatasetsearch

Examples

## Not run:
# Suggest datasets of type "OCCURRENCE".
dataset_suggest(query="Amazon", type="OCCURRENCE")

# Suggest datasets tagged with keyword "france".
dataset_suggest(keyword="france")

# Suggest datasets owned by the organization with key # "07f617d0-c688-11d8-bf62-b8a03c50a862" (UK NBN).
head(dataset_suggest(owningOrg="07f617d0-c688-11d8-bf62-b8a03c50a862"))
# Fulltext search for all datasets having the word "amsterdam" somewhere in
# its metadata (title, description, etc).
head(dataset_suggest(query="amsterdam"))

# Limited search
dataset_suggest(type="OCCURRENCE", limit=2)
dataset_suggest(type="OCCURRENCE", limit=2, start=10)

# Return just descriptions
dataset_suggest(type="OCCURRENCE", description=TRUE)

# Return metadata in a more human readable way (hard to manipulate though)
dataset_suggest(type="OCCURRENCE", pretty=TRUE)

# Search by country code. Lookup isocodes first, and use US for United States
isocodes[agrep("UNITED", isocodes$gbif_name),]
head(dataset_suggest(country="US"))

# Search by decade
head(dataset_suggest(decade=1980))

# httr options
library('httr')
dataset_suggest(type="OCCURRENCE", limit=2, config=verbose())

## End(Not run)

elevation

Get elevation for lat/long points from a data.frame or list of points.

Description

Get elevation for lat/long points from a data.frame or list of points.

Usage

elevation(input = NULL, latitude = NULL, longitude = NULL,
latlong = NULL, ...)

Arguments

input A data.frame of lat/long data. There must be columns decimalLatitude and decimalLongitude.
latitude A vector of latitude’s. Must be the same length as the longitude vector.
longitude A vector of longitude’s. Must be the same length as the latitude vector.
latlong A vector of lat/long pairs. See examples.
... Further named parameters, such as query, path, etc, passed on to modify_url
within GET call. Unnamed parameters will be combined with config.
gbifmap

Make a map to visualize GBIF occurrence data.

Description

Make a map to visualize GBIF occurrence data.

Usage

```r
gbifmap(input = NULL, mapdatabase = "world", region = ".", geom = geom_point, jitter = NULL, customize = NULL)
```

Arguments

- **input**: Either a single data.frame or a list of data.frame’s (e.g., from different species). The data.frame has to have, in addition to any other columns, columns named exactly "decimalLatitude" and "decimalLongitude".

Value

A new column named elevation in the supplied data.frame or a vector with elevation of each location in meters.

References

Uses the Google Elevation API at the following link https://developers.google.com/maps/documentation/elevation/

Examples

```r
## Not run:
key <- name_suggest('Puma concolor')$key[1]
dat <- occ_search(taxonKey=key, return='data', limit=300, hasCoordinate=TRUE)
head(elevation(dat))

# Pass in a vector of lat's and a vector of long's
elevation(latitude=dat$decimalLatitude, longitude=dat$decimalLongitude)

# Pass in lat/long pairs in a single vector
pairs <- list(c(31.8496,-110.576060), c(29.15503,-103.59828))
elevation(latlong=pairs)

# Pass on options to httr
library('httr')
pairs <- list(c(31.8496,-110.576060), c(29.15503,-103.59828))
elevation(latlong=pairs, config=verbose())

## End(Not run)
```
gbifmap takes care of cleaning up the data.frame (removing NA's, etc.) returned from rgbif functions, and creating the map. This function gives a simple map of your data. You can look at the code behind the function itself if you want to build on it to make a map according to your specifications.

Note that this function removes values that are impossible on the globe, and those rows that have both lat and long as NA or zeros.

Value

Map (using ggplot2 package) of points on a map or tiles on a map.

Examples

```r
## Not run:
# Make a map of Puma concolor occurrences
key <- name_backbone(name='Puma concolor')$speciesKey
dat <- occ_search(taxonKey=key, return='data', limit=100)
gbifmap(input=dat)

# Plot more Puma concolor occurrences
dat <- occ_search(taxonKey=key, return='data', limit=1200)
nrow(dat)
gbifmap(input=dat)

# More than 1 species
library("plyr")
splist <- c('Cyanocitta stelleri', 'Junco hyemalis', 'Aix sponsa')
keys <- sapply(splist, function(x) name_backbone(x)$speciesKey, USE.NAMES=FALSE)
dat <- occ_search(taxonKey=keys, return='data', limit=50)
gbifmap(ldply(dat))

## End(Not run)
```
gbif_bbox2wkt

Convert a bounding box to a Well Known Text polygon, and a WKT to a bounding box

Description
Convert a bounding box to a Well Known Text polygon, and a WKT to a bounding box

Usage

gbif_bbox2wkt(minx = NA, miny = NA, maxx = NA, maxy = NA, bbox = NULL)

gbif_wkt2bbox(wkt = NULL)

Arguments

minx Minimum x value, or the most western longitude
miny Minimum y value, or the most southern latitude
maxx Maximum x value, or the most eastern longitude
maxy Maximum y value, or the most northern latitude
bbox A vector of length 4, with the elements: minx, miny, maxx, maxy
wkt A Well Known Text object.

Value

gbif_bbox2wkt returns an object of class character, a Well Known Text string of the form 'POLYGON((minx miny, maxx miny, maxx maxy, minx maxy, minx miny))'.
gbif_wkt2bbox returns a numeric vector of length 4, like c(minx, miny, maxx, maxy).

Examples

## Not run
# Convert a bounding box to a WKT
library("rgeos")

## Pass in a vector of length 4 with all values
mm <- gbif_bbox2wkt(bbox=c(38.4,-125.0,40.9,-121.8))
plot(readWKT(mm))

## Or pass in each value separately
mm <- gbif_bbox2wkt(minx=38.4, miny=-125.0, maxx=40.9, maxy=-121.8)
plot(readWKT(mm))

# Convert a WKT object to a bounding box
wkt <- "POLYGON((38.4 -125,40.9 -125,40.9 -121.8,38.4 -121.8,38.4 -125))"
gbif_wkt2bbox(wkt)

## End(Not run)
gbif_issues

Table of GBIF issues, with codes used in data output, full issue name, and descriptions.

Description

Table has the following fields:

Usage
gbff_issues()

Details

- issue. Full name of the issue.
- description. Description of the issue.

Source


---

gbff_names

View highlighted terms in name results from GBIF.

Description

View highlighted terms in name results from GBIF.

Usage
gbff_names(input, output = NULL, browse = TRUE)

Arguments

input Input output from occ_search
output Output folder path. If not given uses temporary folder.
browse (logical) Browse output (default: TRUE)
### Examples

```r
## Not run:
gbif_names(name_lookup(query='snake', hl=TRUE), browse=FALSE)

(out <- name_lookup(query='canada', hl=TRUE, limit=5))
gbif_names(out)
gbif_names(name_lookup(query='snake', hl=TRUE))
gbif_names(name_lookup(query='bird', hl=TRUE))

# or not highlight
gbif_names(name_lookup(query='bird', limit=200))

## End(Not run)
```

---

**gbif_photos**

*View photos from GBIF.*

---

**Description**

View photos from GBIF.

**Usage**

```r
gbif_photos(input, output = NULL, which = "table", browse = TRUE)
```

**Arguments**

- **input**: Input output from occ_search
- **output**: Output folder path. If not given uses temporary folder.
- **which**: One of map or table (default).
- **browse**: (logical) Browse output (default: TRUE)

**Details**

The max number of photos you can see when which="map" is ~160, so cycle through if you have more than that.

**Examples**

```r
## Not run:
(res <- occ_search(mediatype = 'StillImage', return = "media"))
gbif_photos(res)
gbif_photos(res, which='map')

res <- occ_search(scientificName = "Aves", mediatype = 'StillImage', return = "media", limit=150)
gbif_photos(res)
gbif_photos(res, output = "~/barfoo")

## End(Not run)
```
installations

Installations metadata.

Description

Installations metadata.

Usage

installations(data = "all", uuid = NULL, query = NULL, identifier = NULL, identifierType = NULL, limit = 100, start = NULL, ...)

Arguments

data The type of data to get. Default is all data. If not 'all', then one or more of 'contact', 'endpoint', 'dataset', 'comment', 'deleted', 'nonPublishing'.

uuid UUID of the data node provider. This must be specified if data is anything other than 'all'.

query Query nodes. Only used when data='all'. Ignored otherwise.

identifier The value for this parameter can be a simple string or integer, e.g. identifier=120. This parameter doesn’t seem to work right now.

identifierType Used in combination with the identifier parameter to filter identifiers by identifier type. See details. This parameter doesn’t seem to work right now.

limit Number of records to return. Default: 100. Maximum: 1000.

start Record number to start at. Default: 0. Use in combination with limit to page through results.

... Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.

Details

identifierType options:

- DOI No description.
- FTP No description.
- GBIF_NODE Identifies the node (e.g: 'DK' for Denmark, 'sp2000' for Species 2000).
- GBIF_PARTICIPANT Participant identifier from the GBIF IMS Filemaker system.
- GBIF_PORTAL Indicates the identifier originated from an auto_increment column in the portal.data_provider or portal.data_resource table respectively.
- HANDLER No description.
- LSID Reference controlled by a separate system, used for example by DOI.
- SOURCE_ID No description.
• UNKNOWN No description.
• URI No description.
• URL No description.
• UUID No description.

References

http://www.gbif.org/developer/registry#installations

Examples

```r
## Not run:
installations(limit=5)
installations(query="france")
installations(uuid="b77901f9-d9b0-47fa-94e0-dd96450aa2b4")
installations(data='contact', uuid="b77901f9-d9b0-47fa-94e0-dd96450aa2b4")
installations(data='endpoint', uuid="b77901f9-d9b0-47fa-94e0-dd96450aa2b4")
installations(data='dataset', uuid="b77901f9-d9b0-47fa-94e0-dd96450aa2b4")
installations(data='deleted')
installations(data='deleted', limit=2)
installations(data=c('deleted','nonPublishing'), limit=2)
installations(identifierType='DOI', limit=2)

# Pass on options to httr
library('httr')
res <- installations(data='deleted', config=progress())

## End(Not run)
```

<table>
<thead>
<tr>
<th>isocodes</th>
<th>Table of country two character ISO codes, and GBIF names</th>
</tr>
</thead>
</table>

Description

• code. Two character ISO country code.
• name. Name of country.
• gbif_name. Name of country used by GBIF - this is the name you want to use when searching by country in this package.

Lookup-table for 2 character country ISO codes
name_backbone

Lookup names in the GBIF backbone taxonomy.

Description

Lookup names in the GBIF backbone taxonomy.

Usage

name_backbone(name, rank = NULL, kingdom = NULL, phylum = NULL, class = NULL, order = NULL, family = NULL, genus = NULL, strict = FALSE, verbose = FALSE, start = NULL, limit = 100, ...)

Arguments

name (character) Full scientific name potentially with authorship (required)
rank (character) The rank given as our rank enum. (optional)
kingdom (character) If provided default matching will also try to match against this if no direct match is found for the name alone. (optional)
phylum (character) If provided default matching will also try to match against this if no direct match is found for the name alone. (optional)
class (character) If provided default matching will also try to match against this if no direct match is found for the name alone. (optional)
order (character) If provided default matching will also try to match against this if no direct match is found for the name alone. (optional)
family (character) If provided default matching will also try to match against this if no direct match is found for the name alone. (optional)
genus (character) If provided default matching will also try to match against this if no direct match is found for the name alone. (optional)
strict (logical) If TRUE it (fuzzy) matches only the given name, but never a taxon in the upper classification (optional)
verbose (logical) If TRUE show alternative matches considered which had been rejected.
start Record number to start at. Default: 0. Use in combination with limit to page through results.
limit Number of records to return. Default: 100. Maximum: 1000.
...

Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.

Details

If you don’t get a match GBIF gives back a list of length 3 with slots synonym, confidence, and matchType='NONE'.
Value

A list for a single taxon with many slots (with verbose=FALSE - default), or a list of length two, first element for the suggested taxon match, and a data.frame with alternative name suggestions resulting from fuzzy matching (with verbose=TRUE).

References

http://www.gbif.org/developer/species#searching

Examples

```r
## Not run:
name_backbone(name="Helianthus annuus", kingdom="plants")
name_backbone(name="Helianthus", rank="genus", kingdom="plants")
name_backbone(name="Poa", rank="genus", family="Poaceae")

# Verbose - gives back alternatives
name_backbone(name="Helianthus annuus", kingdom="plants", verbose=TRUE)

# Strictness
name_backbone(name="Poa", kingdom="plants", verbose=TRUE, strict=FALSE)
name_backbone(name="Helianthus annuus", kingdom="plants", verbose=TRUE, strict=TRUE)

# Non-existent name - returns list of length 3 stating no match
name_backbone(name="Aso")
name_backbone(name="Oenante")

# Pass on httr options
library('httr')
name_backbone(name="Oenante", config=timeout(1))

## End(Not run)
```

name_lookup

Lookup names in all taxonomies in GBIF.

Description

This service uses fuzzy lookup so that you can put in partial names and you should get back those things that match. See examples below.

Faceting: If facet=FALSE or left to the default (NULL), no faceting is done. And therefore, all parameters with facet in their name are ignored (facetOnly, facetMincount, facetMultiselect).

Usage

```r
name_lookup(query = NULL, rank = NULL, higherTaxonKey = NULL, 
status = NULL, isExtinct = NULL, habitat = NULL, nameType = NULL, 
datasetKey = NULL, nomenclaturalStatus = NULL, limit = 100,
```
Arguments

query
Query term(s) for full text search.

rank
CLASS, CULTIVAR, CULTIVAR_GROUP, DOMAIN, FAMILY, FORM, GENUS, INFORMAL, INFRAGENERIC_NAME, INFRAORDER, INFRASPECIFIC_NAME, INFRASUBSPECIFIC_NAME, KINGDOM, ORDER, PHYLUM, SECTION, SERIES, SPECIES, STRAIN, SUBCLASS, SUBFAMILY, SUBFORM, SUBGENUS, SUBKINGDOM, SUBORDER, SUBPHYLUM, SUBSECTION, SUBSERIES, SUBSPECIES, SUBTRIBE, SUBVARIETY, SUPERCLASS, SUPERFAMILY, SUPERORDER, SUPERPHYLUM, SUPRAGENERIC_NAME, TRIBE, UNRANKED, VARIETY

higherTaxonKey
Filters by any of the higher Linnean rank keys. Note this is within the respective checklist and not searching nub keys across all checklists.

status
Filters by the taxonomic status as one of:
- ACCEPTED
- DETERMINATION_SYNONYM Used for unknown child taxa referred to via spec, ssp, ...
- DOUBTFUL Treating as accepted, but doubtful whether this is correct.
- HETEROTYPIC_SYNONYM More specific subclass of SYNONYM.
- HOMOTYPIC_SYNONYM More specific subclass of SYNONYM.
- INTERMEDIATE_RANK_SYNONYM Used in nub only.
- MISAPPLIED More specific subclass of SYNONYM.
- PROPARTE_SYNONYM More specific subclass of SYNONYM.
- SYNONYM A general synonym, the exact type is unknown.

isExtinct
(logical) Filters by extinction status (e.g. isExtinct=TRUE)

habitat
(logical) Filters by the habitat, though currently only as boolean marine or not-marinie (i.e. TRUE=marine, FALSE=not-marine)

nameType
Filters by the name type as one of:
- BLACKLISTED surely not a scientific name.
- CANDIDATUS Candidatus is a component of the taxonomic name for a bacterium that cannot be maintained in a Bacteriology Culture Collection.
- CULTIVAR a cultivated plant name.
- DOUBTFUL doubtful whether this is a scientific name at all.
- HYBRID a hybrid formula (not a hybrid name).
- INFORMAL a scientific name with some informal addition like "cf." or indetermined like Abies spec.
- SCINAME a scientific name which is not well formed.
- VIRUS a virus name.
- WELLFORMED a well formed scientific name according to present nomenclatural rules.
datasetKey Filters by the dataset’s key (a uuid)
nomenclaturalStatus Not yet implemented, but will eventually allow for filtering by a nomenclatural status enum
limit Number of records to return. Maximum: 1000.
start Record number to start at.
facet A list of facet names used to retrieve the 100 most frequent values for a field. Allowed facets are: datasetKey, higherTaxonKey, rank, status, isExtinct, habitat, and nameType. Additionally threat and nomenclaturalStatus are legal values but not yet implemented, so data will not yet be returned for them.
facetMincount Used in combination with the facet parameter. Set facetMincount=# to exclude facets with a count less than #, e.g. http://bit.ly/1bMdByP only shows the type value 'ACCEPTED' because the other statuses have counts less than 7,000,000
facetMultiselect (logical) Used in combination with the facet parameter. Set facetMultiselect=TRUE to still return counts for values that are not currently filtered, e.g. http://bit.ly/19YLXPO still shows all status values even though status is being filtered by status=ACCEPTED
type Type of name. One of occurrence, checklist, or metadata.
hl (logical) Set hl=TRUE to highlight terms matching the query when in fulltext search fields. The highlight will be an emphasis tag of class 'gbifH1' e.g. query='plant', hl=TRUE. Fulltext search fields include: title, keyword, country, publishing country, publishing organization title, hosting organization title, and description. One additional full text field is searched which includes information from metadata documents, but the text of this field is not returned in the response.
verbose (logical) If TRUE, all data is returned as a list for each element. If FALSE (default) a subset of the data that is thought to be most essential is organized into a data.frame.
return One of data, meta, facets, names, or all. If data, a data.frame with the data. facets returns the facets, if facets=TRUE, or empty list if facets=FALSE. meta returns the metadata for the entire call. names returns the vernacular (common) names for each taxon. all gives all data back in a list. Each element is NULL if there is no contents in that element. hierarchies and names slots are named by the GBIF key, which matches the first column of the data.frame in the data slot. So if you wanted to combine those somehow, you could easily do so using the key.

Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.

Value
A list of length three. The first element is metadata. The second is either a data.frame (verbose=FALSE, default) or a list (verbose=TRUE), and the third element is the facet data.

References
http://www.gbif.org/developer/species#searching
Examples

```r
## Not run:
# Look up names like mammalia
name_lookup(query='mammalia')

# Paging
name_lookup(query='mammalia', limit=1)
name_lookup(query='mammalia', limit=1, start=2)

# large requests, use start parameter
first <- name_lookup(query='mammalia', limit=1000)
second <- name_lookup(query='mammalia', limit=1000, start=1000)
tail(first$data)
head(second$data)
first$meta
second$meta

# Get all data and parse it, removing descriptions which can be quite long
out <- name_lookup('Helianthus annuus', rank='species', verbose=TRUE)
lapply(out$data, function(x) x!names(x) %in% c("descriptions","descriptionsSerialized"))

# Search for a genus, returning just data
name_lookup(query='Cnaemidophorus', rank='genus', return="data")

# Just metadata
name_lookup(query='Cnaemidophorus', rank='genus', return="meta")

# Just hierarchies
name_lookup(query='Cnaemidophorus', rank='genus', return="hierarchy")

# Just vernacular (common) names
name_lookup(query='Cnaemidophorus', rank='genus', return="names")

# Fuzzy searching
name_lookup(query='Cnaemidophor', rank='genus')

# Limit records to certain number
name_lookup('Helianthus annuus', rank='species', limit=2)

# Using faceting
name_lookup(facet='status', limit=0, facetMinCount='70000')
name_lookup(facet=c('status', 'higherTaxonKey'), limit=0, facetMinCount='700000')
name_lookup(facet='nameType', limit=0)
name_lookup(facet='habitat', limit=0)
name_lookup(facet='datasetKey', limit=0)
name_lookup(facet='rank', limit=0)
name_lookup(facet='isExtinct', limit=0)
name_lookup(isExtinct=TRUE, limit=0)

# text highlighting
```
## name_suggest

### Description

A quick and simple autocomplete service that returns up to 20 name usages by doing prefix matching against the scientific name. Results are ordered by relevance.

### Usage

```r
name_suggest(q = NULL, datasetKey = NULL, rank = NULL, fields = NULL, start = NULL, limit = 100, ...)
```

### Arguments

- **q** (character, required) Simple search parameter. The value for this parameter can be a simple word or a phrase. Wildcards can be added to the simple word parameters only, e.g. `q=*puma*`
- **datasetKey** (character) Filters by the checklist dataset key (a uuid, see examples)
- **rank** (character) A taxonomic rank. One of class, cultivar, cultivar_group, domain, family, form, genus, informal, infrageneric_name, infraorder, infraspecific_name, infrasubspecific_name, kingdom, order, phylum, section, series, species, strain, subclass, subfamily, subform, subgenus, subkingdom, suborder, subphylum, subsection, subseries, subspecies, subtribe, subvariety, superclass, superfAMILY, superorder, superphylum, suprageneric_name, tribe, unranked, or variety.
- **fields** (character) Fields to return in output data.frame (simply prunes columns off)
- **start** Record number to start at. Default: 0. Use in combination with `limit` to page through results.
name_usage

Usage

name_usage(key = NULL, name = NULL, data = "all", language = NULL, datasetKey = NULL, uuid = NULL, sourceId = NULL, rank = NULL, shortname = NULL, start = NULL, limit = 100, return = "all", ...)
Arguments

- `key` (numeric) A GBIF key for a taxon
- `name` (character) Filters by a case insensitive, canonical namestring, e.g. 'Puma con-color'
- `data` (character) Specify an option to select what data is returned. See Description below.
- `language` (character) Language, default is english
- `datasetKey` (character) Filters by the dataset’s key (a uuid)
- `uuid` (character) A uuid for a dataset. Should give exact same results as datasetKey.
- `sourceId` (numeric) Filters by the source identifier. Not used right now.
- `rank` (character) Taxonomic rank. Filters by taxonomic rank as one of: CLASS, CULTIVAR, CULTIVAR_GROUP, DOMAIN, FAMILY, FORM, GENUS, INFORMAL, INFRAGENERIC_NAME, INFRAORDER, INFRASPECIFIC_NAME, INFRASUBSPECIFIC_NAME, KINGDOM, ORDER, PHYLUM, SECTION, SERIES, SPECIES, STRAIN, SUBCLASS, SUBFAMILY, SUBFORM, SUBGENUS, SUBKINGDOM, SUBORDER, SUBPHYLUM, SUBSECTION, SUBSERIES, SUBSPECIES, SUBTRIBE, SUBVARIETY, SUPERCLASS, SUPER FAMILY, SUPERORDER, SUPERPHYLUM, SUPRAGENERIC_NAME, TRIBE, UNRANKED, VARIETY
- `shortname` (character) A short name..need more info on this?
- `start` Record number to start at. Default: 0. Use in combination with `limit` to page through results.
- `limit` Number of records to return. Default: 100. Maximum: 1000.
- `return` One of data, meta, or all. If data, a data.frame with the data. meta returns the metadata for the entire call. all gives all data back in a list.
- ... Further named parameters, such as query, path, etc, passed on to `modify_url` within `GET` call. Unnamed parameters will be combined with `config`.

Details

This service uses fuzzy lookup so that you can put in partial names and you should get back those things that match. See examples below.

This function is different from `name_lookup` in that that function searches for names. This function encompasses a bunch of API endpoints, most of which require that you already have a taxon key, but there is one endpoint that allows name searches (see examples below).

Note that `data="verbatim"` hasn’t been working.

Options for the data parameter are: 'all', 'verbatim', 'name', 'parents', 'children', 'related', 'synonyms', 'descriptions', 'distributions', 'images', 'references', 'speciesProfiles', 'vernacularNames', 'typeSpecimens', 'root'

This function used to be vectorized with respect to the data parameter, where you could pass in multiple values and the function internally loops over each option making separate requests. This has been removed. You can still loop over many options for the data parameter, just use an `lapply` family function, or a for loop, etc.
Value

A list of length two. The first element is metadata. The second is a data.frame

References

http://www.gbif.org/developer/species#nameUsages

Examples

```r
## Not run:
# A single name usage
name_usage(key=1)

# Name usage for a taxonomic name
name_usage(name='Puma', rank="GENUS")

# All name usages
name_usage()

# References for a name usage
name_usage(key=3119195, data='references')

# Species profiles, descriptions
name_usage(key=3119195, data='speciesProfiles')
name_usage(key=3119195, data='descriptions')
name_usage(key=2435099, data='children')

# Vernacular names for a name usage
name_usage(key=3119195, data='vernacularNames')

# Limit number of results returned
name_usage(key=3119195, data='vernacularNames', limit=3)

# Search for names by dataset with datasetKey parameter
name_usage(datasetKey="d7dddbf4-2cf0-4f39-9b2a-bb099caae36c")

# Search for a particular language
name_usage(key=3119195, language="FRENCH", data='vernacularNames')

# Pass on httr options
## here, print progress, notice the progress bar
library('httr')
res <- name_usage(name='Puma concolor', limit=300, config=progress())

## End(Not run)
```
Description

Networks metadata.

Usage

```
networks(data = "all", uuid = NULL, query = NULL, identifier = NULL, identifierType = NULL, limit = 100, start = NULL, ...)
```

Arguments

- **data**: The type of data to get. Default is all data.
- **uuid**: UUID of the data network provider. This must be specified if data is anything other than 'all'.
- **query**: Query nodes. Only used when data='all'. Ignored otherwise.
- **identifier**: The value for this parameter can be a simple string or integer, e.g. identifier=120. This parameter doesn’t seem to work right now.
- **identifierType**: Used in combination with the identifier parameter to filter identifiers by identifier type. See details. This parameter doesn’t seem to work right now.
- **limit**: Number of records to return. Default: 100. Maximum: 1000.
- **start**: Record number to start at. Default: 0. Use in combination with limit to page through results.
- **...**: Further named parameters, such as query, path, etc, passed on to `modify_url` within `GET` call. Unnamed parameters will be combined with `config`.

Details

**identifierType** options:

- **DOI**: No description.
- **FTP**: No description.
- **GBIF_NODE**: Identifies the node (e.g: 'DK' for Denmark, 'sp2000' for Species 2000).
- **GBIF_PARTICIPANT**: Participant identifier from the GBIF IMS Filemaker system.
- **GBIF_PORTAL**: Indicates the identifier originated from an auto_increment column in the portal.data_provider or portal.data_resource table respectively.
- **HANDLER**: No description.
- **LSID**: Reference controlled by a separate system, used for example by DOI.
- **SOURCE_ID**: No description.
- **UNKNOWN**: No description.
- **URI**: No description.
- **URL**: No description.
- **UUID**: No description.
References

http://www.gbif.org/developer/registry#networks

Examples

```r
## Not run:
networks(limit=5)
networks(uuid='16ab5405-6c94-4189-ac71-16ca3b753df7')
networks(data='endpoint', uuid='16ab5405-6c94-4189-ac71-16ca3b753df7')

# Pass on options to httr
library('httr')
res <- networks(limit=5, config=progress())
## End(Not run)
```

---

**nodes**

<table>
<thead>
<tr>
<th>nodes</th>
<th>Nodes metadata.</th>
</tr>
</thead>
</table>

**Description**

Nodes metadata.

**Usage**

```r
nodes(data = "all", uuid = NULL, query = NULL, identifier = NULL,
      identifierType = NULL, limit = 100, start = NULL, isocode = NULL, ...)
```

**Arguments**

- **data**
  - The type of data to get. Default is all data.
- **uuid**
  - UUID of the data node provider. This must be specified if data is anything other than ’all’.
- **query**
  - Query nodes. Only used when data=’all’
- **identifier**
  - The value for this parameter can be a simple string or integer, e.g. identifier=120. This parameter doesn’t seem to work right now.
- **identifierType**
  - Used in combination with the identifier parameter to filter identifiers by identifier type. See details. This parameter doesn’t seem to work right now.
- **limit**
  - Number of records to return. Default: 100. Maximum: 1000.
- **start**
  - Record number to start at. Default: 0. Use in combination with limit to page through results.
- **isocode**
  - A 2 letter country code. Only used if data=’country’.
- **...**
  - Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.
Details

identifierType options:

- **DOI** No description.
- **FTP** No description.
- **GBIF_NODE** Identifies the node (e.g. 'DK' for Denmark, 'sp2000' for Species 2000).
- **GBIF_PARTICIPANT** Participant identifier from the GBIF IMS Filemaker system.
- **GBIF_PORTAL** Indicates the identifier originated from an auto_increment column in the portal.data_provider or portal.data_resource table respectively.
- **HANDLER** No description.
- **LSID** Reference controlled by a separate system, used for example by DOI.
- **SOURCE_ID** No description.
- **UNKNOWN** No description.
- **URI** No description.
- **URL** No description.
- **UUID** No description.

References

http://www.gbif.org/developer/registry\#nodes

Examples

```r
## Not run:
res <- res[!sapply(res, length) == 0]
ldply(res)
```

library('plyr')
```r
res <- res[, 'identifier']$data
```

library('httr')
occ_count

Get number of occurrence records.

Description
Get number of occurrence records.

Usage

```r
occ_count(taxonKey = NULL, georeferenced = NULL, basisOfRecord = NULL,
datasetKey = NULL, date = NULL, catalogNumber = NULL, country = NULL,
hostCountry = NULL, year = NULL, from = 2000, to = 2012,
type = "count", publishingCountry = "US", nubKey = NULL,
protocol = NULL, ...)
```

Arguments

taxonKey Species key
georeferenced Return only occurrence records with lat/long data (TRUE) or all records (FALSE, default).
basisOfRecord Basis of record
datasetKey Dataset key
date Collection date
catalogNumber Catalog number. PARAMETER GONE.
country Country data was collected in, two letter abbreviation. See http://countrycode.org/ for abbreviations.
hostCountry Country that hosted the data. PARAMETER GONE.
year Year data were collected in
from Year to start at
to Year to end at
type One of count (default), schema, basis_of_record, countries, or year.
publishingCountry Publishing country, two letter ISO country code
nubKey Species key. PARAMETER NAME CHANGED TO taxonKey.
protocol Protocol. E.g., ’DWC_ARCHIVE’
... Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.
Value

A single numeric value, or a list of numerics.

References

http://www.gbif.org/developer/occurrence#metrics

Examples

```r
## Not run:
occ_count(basisOfRecord='OBSERVATION')
occ_count(georeferenced=TRUE)
occ_count(country='DE')
occ_count(country='CA', georeferenced=TRUE, basisOfRecord='OBSERVATION')
occ_count(datasetKey='9e7ea10e-0bf8-4087-bb61-dfe4f29e0f17')
occ_count(year=2012)
occ_count(taxonKey=2435099)
occ_count(taxonKey=2435099, georeferenced=TRUE)
occ_count(protocol='DWC_ARCHIVE')

# Just schema
occ_count(type='schema')

# Counts by basisOfRecord types
occ_count(type='basisOfRecord')

# Counts by countries. publishingCountry must be supplied (default to US)
occ_count(type='countries')

# Counts by year. from and to years have to be supplied, default to 2000 and 2012
occ_count(type='year', from=2000, to=2012)

# Counts by publishingCountry, must supply a country (default to US)
occ_count(type='publishingCountry')
occ_count(type='publishingCountry', country='BZ')

# Pass on options to httr
library('httr')
res <- occ_count(type='year', from=2000, to=2012, config=progress())
res

## End(Not run)
```

---

**occ_fields**

Vector of fields in the output for the function **occ_search**

Description

These fields can be specified in the fields parameter in the **occ_search** function.
occ_get  

Get data for specific GBIF occurrences.

Description

Get data for specific GBIF occurrences.

Usage

occ_get(key = NULL, return = "all", verbatim = FALSE, 
fields = "minimal", ...)

Arguments

key Occurrence key
return One of data, hier, meta, or all. If 'data', a data.frame with the data. 'hier' returns 
the classifications in a list for each record. meta returns the metadata for the 
entire call. 'all' gives all data back in a list. Ignored if verbatim=TRUE.
verbatim Return verbatim object (TRUE) or cleaned up object (FALSE, default).
fields (character) Default ('minimal') will return just taxon name, key, latitude, and 
longitude. 'all' returns all fields. Or specify each field you want returned by 
name, e.g. fields = c('name','decimalLatitude','altitude').
... Further named parameters, such as query, path, etc, passed on to modify_url 
within GET call. Unnamed parameters will be combined with config.

Value

A data.frame or list of data.frame's.

References

httpZOOwwwNgbifNorgOdeveloperOoccurrenceCoccurrence

Examples

## Not run:
occ_get(key=766766824, return='data')
occ_get(key=766766824, 'hier')
occ_get(key=766766824, 'all')

# many occurrences
occ_get(key=c(101010,240713150,855998194,49819470), return='data')

# Verbatim data
occ_get(key=766766824, verbatim=TRUE)
occ_get(key=766766824, fields='all', verbatim=TRUE)
occ_get(key=766766824, fields=c('scientificName','lastCrawled','county'), verbatim=TRUE)
occ_issues

Parse and examine further GBIF issues on a dataset.

Description

Parse and examine further GBIF issues on a dataset.

Usage

occ_issues(.data, ..., mutate = NULL)

Arguments

.data Output from a call to occ_search
... Named parameters to only get back (e.g., cdround), or to remove (e.g., -cdround).
mutate (character) One of:

* split Split issues into new columns.
* split.expand Split into new columns, and expand issue names.
* expand Expand issue abbreviated codes into descriptive names.

For split and split.expand, values in cells become y ("yes") or n ("no")

Details

See also the vignette Cleaning data using GBIF issues.

Note that you can also query based on issues, e.g., occ_search(taxonKey=1, issue='DEPTH_UNLIKELY'). However, I imagine it’s more likely that you want to search for occurrences based on a taxonomic name, or geographic area, not based on issues, so it makes sense to pull data down, then clean as needed using this function.

This function only affects the data element in the gbif class that is returned from a call to occ_search. Maybe in a future version we will remove the associated records from the hierarchy and media elements as they are remove from the data element.
occ_issues

References


Examples

## Not run:
## what do issues mean, can print whole table, or search for matches
head(gbif_issues())
gbif_issues()[ gbif_issues()$code %in% c('cround','cudc','gass84','txmathi'), ]

# compare out data to after occ_issues use
(out <- occ_search(limit=100))
out %>% occ_issues(cudc)

## End(Not run)

## Not run:
# Parsing output by issue
(res <- occ_search(geometry='POLYGON((30.1 10.1, 10 20, 20 40, 40 40, 30.1 10.1))', limit = 50))

## or parse issues in various ways
### include only rows with gass84 issue
gg <- res %>% occ_issues(gass84)
NROW(res$data)
NROW(gg$data)
head(res$data)[,c(1:5)]
head(gg$data)[,c(1:5)]

### remove data rows with certain issue classes
res %>% occ_issues(-gass84, -mdatunl)

### split issues into separate columns
res %>% occ_issues(mutate = "split")
res %>% occ_issues(-gass84, -mdatunl, mutate = "split")
res %>% occ_issues(gass84, mutate = "split")

### expand issues to more descriptive names
res %>% occ_issues(mutate = "expand")

### split and expand
res %>% occ_issues(mutate = "split_expand")

### split, expand, and remove an issue class
res %>% occ_issues(-gass84, mutate = "split_expand")

## Or you can use occ_issues without %>%
occ_issues(res, -gass84, mutate = "split_expand")

## End(Not run)
occ_issues_lookup  
Lookup occurrence issue definitions and short codes

Description

Lookup occurrence issue definitions and short codes

Usage

occ_issues_lookup(issue = NULL, code = NULL)

Arguments

issue  Full name of issue, e.g. CONTINENT_COUNTRY_MISMATCH
code  an issue short code, e.g. ccm

Examples

occ_issues_lookup(issue = 'CONTINENT_COUNTRY_MISMATCH')
occ_issues_lookup(issue = 'MULTIMEDIA_DATE_INVALID')
occ_issues_lookup(issue = 'ZERO_COORDINATE')
occ_issues_lookup(code = 'cdiv')

occ_metadata  
Search for catalog numbers, collection codes, collector names, and institution codes.

Description

Search for catalog numbers, collection codes, collector names, and institution codes.

Usage

occ_metadata(type = "catalogNumber", q = NULL, limit = 5, pretty = TRUE, ...)
occ_search

Search for GBIF occurrences.

Description

Search for GBIF occurrences.

Usage

occ_search(taxonKey = NULL, scientificName = NULL, country = NULL, publishingCountry = NULL, hasCoordinate = NULL, typeStatus = NULL, recordNumber = NULL, lastInterpreted = NULL, continent = NULL, geometry = NULL, collectorName = NULL, basisOfRecord = NULL, datasetKey = NULL, eventDate = NULL, catalogNumber = NULL, year = NULL, month = NULL, decimallatitude = NULL, decimallongitude = NULL, elevation = NULL, depth = NULL, institutionCode = NULL, collectionCode = NULL, spatialIssues = NULL, issue = NULL, search = NULL, mediatype = NULL, limit = 500,
start = 0, fields = "all", return = "all", ...)

## S3 method for class 'gbif'
print(x, ..., n = 10)

### Arguments

taxonKey A taxon key from the GBIF backbone. All included and synonym taxa are included in the search, so a search for aves with taxononKey=212 (i.e. /occurrence/search?taxonKey=212) will match all birds, no matter which species. You can pass many keys by passing occ_search in a call to an lapply-family function (see last example below).

scientificName A scientific name from the GBIF backbone. All included and synonym taxa are included in the search.

country The 2-letter country code (as per ISO-3166-1) of the country in which the occurrence was recorded. See here [http://en.wikipedia.org/wiki/ISO_3166-1_alpha-2](http://en.wikipedia.org/wiki/ISO_3166-1_alpha-2)

publishingCountry The 2-letter country code (as per ISO-3166-1) of the country in which the occurrence was recorded.

hasCoordinate (logical) Return only occurrence records with lat/long data (TRUE) or all records (FALSE, default).

typeStatus Type status of the specimen. One of many options. See ?typestatus

recordNumber Number recorded by collector of the data, different from GBIF record number. See [http://rs.tdwg.org/dwc/terms/#recordNumber](http://rs.tdwg.org/dwc/terms/#recordNumber) for more info

lastInterpreted Date the record was last modified in GBIF, in ISO 8601 format: yyyy, yyyy-MM, yyyy-MM-dd, or MM-dd. Supports range queries, smaller,larger (e.g., '1990,1991', whereas '1991,1990' wouldn’t work)

continent Continent. One of africa, antarctica, asia, europe, north_america (North America includes the Caribbean and reaches down and includes Panama), oceania, or south_america

geometry Searches for occurrences inside a polygon described in Well Known Text (WKT) format. A WKT shape written as either POINT, LINESTRING, LINEARRING or POLYGON. Example of a polygon: ((30.1 10.1, 20, 20 40, 40 40, 30.1 10.1)) would be queried as [http://bit.ly/1BzNwDq](http://bit.ly/1BzNwDq).

 collectorName The person who recorded the occurrence.

basisOfRecord Basis of record, as defined in our BasisOfRecord enum here [http://gbif.github.io/gbif-api/apidocs/org/gbif/api/vocabulary/BasisOfRecord.html](http://gbif.github.io/gbif-api/apidocs/org/gbif/api/vocabulary/BasisOfRecord.html) Acceptable values are:

- FOSSIL_SPECIMEN An occurrence record describing a fossilized specimen.
- HUMAN_OBSERVATION An occurrence record describing an observation made by one or more people.
- LITERATURE An occurrence record based on literature alone.
- LIVING_SPECIMEN An occurrence record describing a living specimen, e.g.
- MACHINE_OBSERVATION An occurrence record describing an observation made by a machine.
- OBSERVATION An occurrence record describing an observation.
- PRESERVED_SPECIMEN An occurrence record describing a preserved specimen.
- UNKNOWN Unknown basis for the record.

dataSetKey  The occurrence dataset key (a uuid)
eventDate   Occurrence date in ISO 8601 format: yyyy, yyyy-MM, yyyy-MM-dd, or MM-dd. Supports range queries, smaller, larger (e.g., ’1990,1991’, whereas ’1991,1990’ wouldn’t work)
catalogNumber  An identifier of any form assigned by the source within a physical collection or digital dataset for the record which may not unique, but should be fairly unique in combination with the institution and collection code.
year  The 4 digit year. A year of 98 will be interpreted as AD 98. Supports range queries, smaller, larger (e.g., ’1990,1991’, whereas ’1991,1990’ wouldn’t work)
month  The month of the year, starting with 1 for January. Supports range queries, smaller, larger (e.g., ’1,2’, whereas ’2,1’ wouldn’t work)
decimalLatitude  Latitude in decimals between -90 and 90 based on WGS 84. Supports range queries, smaller, larger (e.g., ’25,30’, whereas ’30,25’ wouldn’t work)
decimalLongitude  Longitude in decimals between -180 and 180 based on WGS 84. Supports range queries (e.g., ’-0.4,-0.2’, whereas ’-0.2,-0.4’ wouldn’t work).
elevation  Elevation in meters above sea level. Supports range queries, smaller, larger (e.g., ’5,30’, whereas ’30,5’ wouldn’t work)
depth  Depth in meters relative to elevation. For example 10 meters below a lake surface with given elevation. Supports range queries, smaller, larger (e.g., ’5,30’, whereas ’30,5’ wouldn’t work)
institutionCode  An identifier of any form assigned by the source to identify the institution the record belongs to. Not guaranteed to be unique.
collectionCode  An identifier of any form assigned by the source to identify the physical collection or digital dataset uniquely within the text of an institution.
spatialIssues  (logical) Includes/excludes occurrence records which contain spatial issues (as determined in our record interpretation), i.e. spatialIssues=TRUE returns only those records with spatial issues while spatialIssues=FALSE includes only records without spatial issues. The absence of this parameter returns any record with or without spatial issues.
issue  (character) One or more of many possible issues with each occurrence record. See Details. Issues passed to this parameter filter results by the issue.
search  Query terms. The value for this parameter can be a simple word or a phrase.
mediatype Media type. Default is NULL, so no filtering on mediatype. Options: NULL, 'MovingImage', 'Sound', and 'StillImage'.

limit Number of records to return. Default: 500. Note that the per request maximum is 300, but since we set it at 500 for the function, we do two requests to get you the 500 records (if there are that many). Note that there is a hard maximum of 200,000, which is calculated as the limit+start, so start=199,000 and limit=2000 won't work.

start Record number to start at. Use in combination with limit to page through results. Note that we do the paging internally for you, but you can manually set the start parameter.

fields (character) Default (‘all’) returns all fields. ’minimal’ returns just taxon name, key, latitude, and longitude. Or specify each field you want returned by name, e.g. fields = c(‘name’,’latitude’,’elevation’).

return One of data, hier, meta, or all. If data, a data.frame with the data. hier returns the classifications in a list for each record. meta returns the metadata for the entire call. all gives all data back in a list.

Details

Note that you can pass in a vector to one of taxonkey, datasetKey, and catalogNumber parameters in a function call, but not a vector >1 of the three parameters at the same time.

Hierarchies: hierarchies are returned with each occurrence object. There is no option no to return them from the API. However, within the occ_search function you can select whether to return just hierarchies, just data, all of data and hierarchies and metadata, or just metadata. If all hierarchies are the same we just return one for you.

Data: By default only three data fields are returned: name (the species name), decimallatitude, and decimallongitude. Set parameter minimal=FALSE if you want more data.

Nerds: You can pass parameters not defined in this function into the call to the GBIF API to control things about the call itself using . . . . See an example below that passes in the verbose function to get details on the http call.

Scientific names vs. taxon keys: In the previous GBIF API and the version of rgbif that wrapped that API, you could search the equivalent of this function with a species name, which was convenient. However, names are messy right. So it sorts make sense to sort out the species key numbers you want exactly, and then get your occurrence data with this function. GBIF has added a parameter scientificName to allow searches by scientific names in this function - which includes synonym taxa. Note: that if you do use the scientificName parameter, we will check internally that it’s not a synonym of an accepted name, and if it is, we’ll search on the accepted name. If you want to force searching by a synonym do so by finding the GBIF identifier first with any name_*. functions, then pass that ID to the taxonkey parameter.

WKT: Examples of valid WKT objects:

- 'POLYGON((30.1 10.1, 10 20, 20 60, 60 60, 30.1 10.1))'
Range queries: A range query is as it sounds - you query on a range of values defined by a lower and upper limit. Do a range query by specifying the lower and upper limit in a vector like depth = '50,100'. It would be more R like to specify the range in a vector like c(50,100), but that sort of syntax allows you to do many searches, one for each element in the vector - thus range queries have to differ. The following parameters support range queries.

- decimalLatitude
- decimalLongitude
- depth
- elevation
- eventDate
- lastInterpreted
- month
- year


- BASIS_OF_RECORD_INVALID The given basis of record is impossible to interpret or seriously different from the recommended vocabulary.
- CONTINENT_COUNTRY_MISMATCH The interpreted continent and country do not match up.
- CONTINENT_DERIVED_FROMCOORDINATES The interpreted continent is based on the coordinates, not the verbatim string information.
- CONTINENT_INVALID Uninterpretable continent values found.
- COORDINATEINVALID Coordinate value given in some form but GBIF is unable to interpret it.
- COORDINATE_OUT_OF_RANGE Coordinate has invalid lat/lon values out of their decimal max range.
- COORDINATE_REPROJECTED The original coordinate was successfully reprojected from a different geodetic datum to WGS84.
- COORDINATE_REPROJECTION_FAILED The given decimal latitude and longitude could not be reprojected to WGS84 based on the provided datum.
- COORDINATE_REPROJECTION_SUSPICIOUS Indicates successful coordinate reprojection according to provided datum, but which results in a datum shift larger than 0.1 decimal degrees.
- COORDINATE_ROUNDED Original coordinate modified by rounding to 5 decimals.
- COUNTRY_COORDINATE_MISMATCH The interpreted occurrence coordinates fall outside of the indicated country.
- COUNTRY_DERIVED_FROMCOORDINATES The interpreted country is based on the coordinates, not the verbatim string information.
• COUNTRY_INVALID Uninterpretable country values found.
• COUNTRY_MISMATCH Interpreted country for dwc:country and dwc:countryCode contradict each other.
• DEPTH_MIN_MAX_SWAPPED Set if supplied min>max
• DEPTH_NON_NUMERIC Set if depth is a non numeric value
• DEPTH_NOT_METRIC Set if supplied depth is not given in the metric system, for example using feet instead of meters
• DEPTH_UNLIKELY Set if depth is larger than 11.000m or negative.
• ELEVATION_MIN_MAX_SWAPPED Set if supplied min > max elevation
• ELEVATION_NON_NUMERIC Set if elevation is a non numeric value
• ELEVATION_NOT_METRIC Set if supplied elevation is not given in the metric system, for example using feet instead of meters
• ELEVATION_UNLIKELY Set if elevation is above the troposphere (17km) or below 11km (Mariana Trench).
• GEODETIC_DATUM_ASSUMED_WGS84 Indicating that the interpreted coordinates assume they are based on WGS84 datum as the datum was either not indicated or interpretable.
• GEODETIC_DATUM_INVALID The geodetic datum given could not be interpreted.
• IDENTIFIED_DATE_INVALID The date given for dwc:dateIdentified is invalid and can't be interpreted at all.
• IDENTIFIED_DATE_UNLIKELY The date given for dwc:dateIdentified is in the future or before Linnean times (1700).
• MODIFIED_DATE_INVALID A (partial) invalid date is given for dc:modified, such as a non existing date, invalid zero month, etc.
• MODIFIED_DATE_UNLIKELY The date given for dc:modified is in the future or predates unix time (1970).
• MULTIMEDIA_DATE_INVALID An invalid date is given for dc:created of a multimedia object.
• MULTIMEDIA_URI_INVALID An invalid uri is given for a multimedia object.
• PRESUMED_NEGATED_LATITUDE Latitude appears to be negated, e.g. 32.3 instead of -32.3
• PRESUMED_NEGATED_LONGITUDE Longitude appears to be negated, e.g. 32.3 instead of -32.3
• PRESUMED_SWAPPED_COORDINATE Latitude and longitude appear to be swapped.
• RECORDED_DATE_INVALID A (partial) invalid date is given, such as a non existing date, invalid zero month, etc.
• RECORDED_DATE_MISMATCH The recording date specified as the eventDate string and the individual year, month, day are contradicting.
• RECORDED_DATE_UNLIKELY The recording date is highly unlikely, falling either into the future or represents a very old date before 1600 that predates modern taxonomy.
• REFERENCES_URI_INVALID An invalid uri is given for dc:references.
• TAXON_MATCH_FUZZY Matching to the taxonomic backbone can only be done using a fuzzy, non exact match.

• TAXON_MATCH_HIGHER_RANK Matching to the taxonomic backbone can only be done on a higher rank and not the scientific name.

• TAXON_MATCH_NONE Matching to the taxonomic backbone cannot be done cause there was no match at all or several matches with too little information to keep them apart (homonyms).

• TYPE_STATUS_INVALID The given type status is impossible to interpret or seriously different from the recommended vocabulary.

• ZERO_COORDINATE Coordinate is the exact 0/0 coordinate, often indicating a bad null coordinate.

Value
A data.frame or list

References
http://www.gbif.org/developer/occurrence#search

Examples
```r
## Not run:
# Search by species name, using \code{name_backbone} first to get key
(key <- name_suggest(q='Helianthus annuus', rank='species')$key[1])
occ_search(taxonKey=key, limit=2)

# Return 20 results, this is the default by the way
occ_search(taxonKey=key, limit=20)

# Return just metadata for the search
occ_search(taxonKey=key, limit=100, return='meta')

# Instead of getting a taxon key first, you can search for a name directly
# However, note that using this approach (with \code{scientificName="..."})
# you are getting synonyms too. The results for using \code{scientificName} and
# \code{taxonKey} parameters are the same in this case, but I wouldn't be surprised if for some
# names they return different results
occ_search(scientificName = 'Ursus americanus')
key <- name_backbone(name = 'Ursus americanus', rank='species')$usageKey
occ_search(taxonKey = key)

# Search by dataset key
occ_search(datasetKey='7b5d6a48-f762-11e1-a439-00145eb45e9a', return='data', limit=20)

# Search by catalog number
occ_search(catalogNumber="49366", limit=20)
occ_search(catalogNumber=c("49366","Bird.27847588"), limit=20)

# Get all data, not just lat/long and name
occ_search(taxonKey=key, fields='all', limit=20)
```
# Or get specific fields. Note that this isn't done on GBIF's side of things. This
# is done in R, but before you get the return object, so other fields are garbage
# collected
occ_search(taxonKey=key, fields=c('name', 'basisOfRecord', 'protocol'), limit=20)

# Use paging parameters (limit and start) to page. Note the different results
# for the two queries below.
occ_search(datasetKey='7b5d6a48-f762-11e1-a439-00145eb45e9a', start=10, limit=5,
       return="data")
occ_search(datasetKey='7b5d6a48-f762-11e1-a439-00145eb45e9a', start=20, limit=5,
       return="data")

# Many dataset keys
occ_search(datasetKey=c("50c9509d-22c7-4a22-a47d-8c48425ef4a7",
                     "7b5d6a48-f762-11e1-a439-00145eb45e9a"), limit=20)

# Occurrence data: lat/long data, and associated metadata with occurrences
# If return='data' the output is a data.frame of all data together
# for easy manipulation
occ_search(taxonKey=key, return='data', limit=20)

# Taxonomic hierarchy data
# If return='meta' the output is a list of the hierarch for each record
occ_search(taxonKey=key, return='hier', limit=10)

# Search by collector name
occ_search(collectorName="smith", limit=20)

# Many collector names
occ_search(collectorName=c("smith","bj stacey"), limit=20)

# Pass in curl options for extra fun
library('httr')
occ_search(taxonKey=key, limit=20, return='hier', config=verbose())
occ_search(taxonKey=key, limit=20, return='hier', config=progress())
occ_search(taxonKey=key, limit=20, return='hier', config=timeout(1))

# Search for many species
splist <- c('Cyanocitta stelleri', 'Junco hyemalis', 'Aix sponsa')
keys <- sapply(splist, function(x) name_suggest(x)$key[1], USE NAMES = FALSE)
occ_search(taxonKey=keys, limit=5, return='data')

# Search using a synonym name
# Note that you'll see a message printing out that the accepted name will be used
occ_search(scientificName = 'Pulsatilla patens', fields = c('name', 'scientificName'), limit=5)

# Search on latitidue and longitude
occ_search(search="kingfisher", decimalLatitude=50, decimalLongitude=-10)

# Search on a bounding box
# in well known text format
occ_search(geometry='POLYGON((30.1 10.1, 10 20, 20 40, 40 40, 30.1 10.1))', limit=20)
key <- name_suggest(q='Aesculus hippocastanum')$key[1]
occ_search(taxonKey=key, geometry='POLYGON((30.1 10.1, 10 20, 20 40, 40 40, 30.1 10.1))', limit=20)
## or using bounding box, converted to WKT internally
occ_search(accuracy=c(-125.0,38.4,-121.8,40.9), limit=20)
## Visualize a WKT area
library('rgeos')
plot(readWKT('POLYGON((30.1 10.1, 10 20, 20 40, 40 40, 30.1 10.1))'))

# Search on country
occ_search(country='US', fields=c('name','country'), limit=20)
occ_search(country='FR', fields=c('name','country'), limit=20)
occ_search(country='DE', fields=c('name','country'), limit=20)
occ_search(country=c('US','DE'), fields=c('name','country'), limit=20)

# Get only occurrences with lat/long data
occ_search(taxonKey=key, hasCoordinate=TRUE, limit=20)

# Get only occurrences that were recorded as living specimens
occ_search(taxonKey=key, basisOfRecord='LIVING_SPECIMEN', hasCoordinate=TRUE, limit=20)

# Get occurrences for a particular eventDate
occ_search(taxonKey=key, eventDate='2013', limit=20)
occ_search(taxonKey=key, year='2013', limit=20)
occ_search(taxonKey=key, month='6', limit=20)

# Get occurrences based on depth
key <- name_backbone(name='Salmo salar', kingdom='animals')$speciesKey
occ_search(taxonKey=key, depth='5', limit=20)

# Get occurrences based on elevation
key <- name_backbone(name='Puma concolor', kingdom='animals')$speciesKey
occ_search(taxonKey=key, elevation=50, hasCoordinate=TRUE, limit=20)

# Get occurrences based on institutionCode
occ_search(institutionCode='TLMF', limit=20)
occ_search(institutionCode=c('TLMF','ArtDatabanken'), limit=20)

# Get occurrences based on collectionCode
occ_search(collectionCode='Floristic Databases MV - Higher Plants', limit=20)
occ_search(collectionCode=c('Floristic Databases MV - Higher Plants','Artport'))

# Get only those occurrences with spatial issues
occ_search(taxonKey=key, spatialIssues=TRUE, limit=20)

# Search using a query string
occ_search(search='kingfisher', limit=20)

# Range queries
## See Detail for parameters that support range queries
occ_search(depth='50,100') # this is a range depth, with lower/upper limits in character string
occ_search(depth=c(50,100)) # this is not a range search, but does two searches for each depth
## Range search with year
```
occ_search(year='1999:2000', limit=20)
```

## Range search with latitude
```
occ_search(decimallatitude='29.59,29.6')
```

# Search by specimen type status
## Look for possible values of the typeStatus parameter looking at the typestatus dataset
```
occ_search(typestatus = 'allotype', fields = c('name','typeStatus'))
```

# Search by specimen record number
## This is the record number of the person/group that submitted the data, not GBIF's numbers
## You can see that many different groups have record number 1, so not super helpful
```
occ_search(recordNumber = 1, fields = c('name','recordNumber','recordedBy'))
```

# Search by last time interpreted: Date the record was last modified in GBIF
## The lastInterpreted parameter accepts ISO 8601 format dates, including
## yyyy, yyyy-MM, yyyy-MM-dd, or MM-dd. Range queries are accepted for lastInterpreted
```
occ_search(lastInterpreted = '2014-04-02', fields = c('name','lastInterpreted'))
```

# Search by continent
## One of africa, antarctica, asia, europe, north_america, oceania, or south_america
```
occ_search(continent = 'south_america', return = 'meta')
occ_search(continent = 'africa', return = 'meta')
occ_search(continent = 'oceania', return = 'meta')
occ_search(continent = 'antarctica', return = 'meta')
```

# Search for occurrences with images
```
occ_search(mediatype = 'StillImage', return='media')
occ_search(mediatype = 'MovingImage', return='media')
occ_search(mediatype = 'Sound', return='media')
```

# Query based on issues - see Details for options
## one issue
```
occ_search(taxonKey=1, issue='DEPTH_UNLIKELY', fields = c('name','key','decimallatitude','decimallongitude','depth'))
```
## two issues
```
occ_search(taxonKey=1, issue=c('DEPTH_UNLIKELY','COORDINATE_ROUNDED'))
```

# Show all records in the Arizona State Lichen Collection that cant be matched to the GBIF
## backbone properly:
```
occ_search(datasetKey='84c0e1a0-f762-11e1-a439-00145eb45e9a',
           issue=c('TAXON_MATCH_NONE','TAXON_MATCH_HIGHER_RANK'))
```

# Parsing output by issue
```
(res <- occ_search( geometry='POLYGON((30.1 10.1, 10 20, 20 40, 40 40, 30.1 10.1))', limit = 50))
```

## what do issues mean, can print whole table, or search for matches
```
head(gbif_issues())
gbif_issues()[ gbif_issues()$code %in% c('cdround','cudc','gass84','txmathi'), ]
```

## or parse issues in various ways
## remove data rows with certain issue classes
```
library('magrittr')
res %>% occ_issues(-gass84, -mdatunl)
```
### split issues into separate columns
res %>% occ_issues(mutate = "split")
### expand issues to more descriptive names
res %>% occ_issues(mutate = "expand")
### split and expand
res %>% occ_issues(mutate = "split\_expand")
### split, expand, and remove an issue class
res %>% occ_issues(~gass84, mutate = "split\_expand")

# If you try multiple values for two different parameters you are wacked on the hand
# occ_search(taxonKey=c(2482598,2492010), collectorName=c("smith","BJ Stacey"))

# Get a lot of data, here 1500 records for Helianthus annuus
out <- occ_search(taxonKey=key, limit=1500, return="data")
nrow(out)

# If you pass in an invalid polygon you get hopefully informative errors

### the WKT string is fine, but GBIF says bad polygon
wkt <- 'POLYGON((-178.59375 64.8325898321493,-165.9375 59.24622380285539,
-147.3046875 59.065977905449806,-130.78125 51.0448476446178,-125.859375 36.70806354647625,
-112.1484375 23.367471307359686,-105.1171875 16.09332085359257,-86.8359375 9.23767076398516,
-82.96875 2.9485268155666175,-82.6171875 -14.812060061226388,-74.8828125 -18.84911862023985,
-77.34375 -47.661687803329166,-84.375 -49.975955187343295,174.7265625 -50.649460483096114,
179.296875 -42.19189902447192,-176.8359375 -35.634976650677295,176.8359375 -31.835565983656227,
163.4765625 -6.528187613695323,152.578125 1.894796132058301,135.703125 4.782353722559447,
127.96875 15.077427674847987,127.96875 23.68980451429606,139.921875 32.06861069132688,
149.410625 42.65416193033991,159.2578125 48.3106811803533,168.3984375 57.81980436633165,
178.2421875 59.95776046458139,-179.6484375 61.1670863140347,-178.59375 64.83258989321493))'

# occ_search( geometry = gsub("\n", ", wkt))

### unable to parse due to last number pair needing two numbers, not one
# wkt <- 'POLYGON((-178.5 64.8,-165.9 59.2,-147.3 59.0,-130.7 51.0,-125.8))'
# occ_search( geometry = wkt)

### unable to parse due to unclosed string
# wkt <- 'POLYGON((-178.5 64.8,-165.9 59.2,-147.3 59.0,-130.7 51.0))'
# occ_search( geometry = wkt)

### another of the same
# wkt <- 'POLYGON((-178.5 64.8,-165.9 59.2,-147.3 59.0,-130.7 51.0,-125.8 36.7))'
# occ_search( geometry = wkt)

### returns no results
# wkt <- 'LINESTRING(3 4,10 50,20 25)'
# occ_search( geometry = wkt)

### Apparently a point is allowed, but haven't successfully retrieved data, so returns nothing
# wkt <- 'POINT(45 -122)'
# occ_search( geometry = wkt)

# End(Not run)
Description

Organizations metadata.

Usage

organizations(data = "all", uuid = NULL, query = NULL, limit = 100,
         start = NULL, ...)

Arguments

data   The type of data to get. Default is all data.
uuid   UUID of the data node provider. This must be specified if data is anything other than 'all'.
query  Query nodes. Only used when data='all'
limit  Number of records to return. Default: 100. Maximum: 1000.
start  Record number to start at. Default: 0. Use in combination with limit to page through results.
...    Further named parameters, such as query, path, etc, passed on to modify_url within GET call. Unnamed parameters will be combined with config.

Value

A list of length one or two. If uuid is NULL, then a data.frame with call metadata, and a data.frame, but if uuid given, then a list.

References

http://www.gbif.org/developer/registry#organizations

Examples

## Not run:
organizations(limit=5)
organizations(query="france")
organizations(uuid="4b4b2111-ee51-45f5-bf5e-f535f4a1c9dc")
organizations(data='contact', uuid="4b4b2111-ee51-45f5-bf5e-f535f4a1c9dc")
organizations(data='pending')
organizations(data=c('contact','endpoint'), uuid="4b4b2111-ee51-45f5-bf5e-f535f4a1c9dc")

# Pass on options to httr
library('httr')
res <- organizations(query="spain", config=progress())

## End(Not run)
parsenames

Parse taxon names using the GBIF name parser.

Description

Parse taxon names using the GBIF name parser.

Usage

parsenames(scientificname, ...)

Arguments

- **scientificname**: A character vector of scientific names.
- **...**: Further named parameters, such as `query`, `path`, etc, passed on to `modify_url` within `GET` call. Unnamed parameters will be combined with `config`.

Value

A `data.frame` containing fields extracted from parsed taxon names. Fields returned are the union of fields extracted from all species names in `scientificname`.

Author(s)

John Baumgartner (johnbb@student.unimelb.edu.au)

References

[http://www.gbif.org/developer/species#parser](http://www.gbif.org/developer/species#parser)

Examples

```r
## Not run:
parsenames(scientificname='x Agropogon littoralis')
parsenames(c('Arrhenatherum elatius var. elatius', 'Secale cereale subsp. cereale', 'Secale cereale ssp. cereale', 'Vanessa atalanta (Linnaeus, 1758)'))

# Pass on options to httr
library('httr')
res <- parsenames(c('Arrhenatherum elatius var. elatius', 'Secale cereale subsp. cereale', 'Secale cereale ssp. cereale', 'Vanessa atalanta (Linnaeus, 1758)'), config=progress())

## End(Not run)
```
Defunct functions in rgbif

Description

- `density_spplist`: service no longer provided
- `densitylist`: service no longer provided
- `gbifdata`: service no longer provided
- `gbifmap_dens`: service no longer provided
- `gbifmap_list`: service no longer provided
- `occurrencedensity`: service no longer provided
- `providers`: service no longer provided
- `resources`: service no longer provided
- `taxoncount`: service no longer provided
- `taxonget`: service no longer provided
- `taxonsearch`: service no longer provided
- `stylegeojson`: moving this functionality to spocc package, will be removed soon
- `togejson`: moving this functionality to spocc package, will be removed soon
- `gist`: moving this functionality to spocc package, will be removed soon

Details

The above functions have been removed. See https://github.com/ropensci/rgbif and poke around the code if you want to find the old functions in previous versions of the package, or email Scott at <myrmecocystus@gmail.com>

rgb_country_codes

Look up 2 character ISO country codes

Description

Look up 2 character ISO country codes

Usage

`rgb_country_codes(country_name, fuzzy = FALSE, ...)`

Arguments

- `country_name`: Name of country to look up
- `fuzzy`: If TRUE, uses agrep to do fuzzy search on names.
- `...`: Further arguments passed on to agrep or grep
Examples

rgb_country_codes(country_name="United")

---

**taxrank**

*Get the possible values to be used for (taxonomic) rank arguments in GBIF API methods.*

Description

Get the possible values to be used for (taxonomic) rank arguments in GBIF API methods.

Usage

taxrank()

Examples

```r
## Not run:
taxrank()
## End(Not run)
```

---

**typestatus**

*Type status options for GBIF searching*

Description

- name. Name of type.
- description. Description of the type.
Index

*Topic data
  isocodes, 18
  occ_fields, 32
  typestatus, 51

  cat, 8, 10
  check_wkt, 3
  config, 6, 8, 10, 11, 17, 19, 22, 25, 26, 28, 29, 31, 33, 36, 40, 48, 49
  count_facet, 4
  dataset_metrics, 6
  dataset_search, 7
  dataset_suggest, 9
  datasets, 5
  density_spplist, 50
  densitylist, 50
  elevation, 11
  gbif_bbox2wkt, 14
  gbif_issues, 15
  gbif_names, 15
  gbif_photos, 16
  gbif_wkt2bbox (gbif_bbox2wkt), 14
  gbifdata, 50
  gbifmap, 12
  gbifmap_dens, 50
  gbifmap_list, 50
  GET, 6, 8, 10, 11, 17, 19, 22, 25, 26, 28, 29, 31, 33, 36, 40, 48, 49
  gist, 50
  installations, 17
  isocodes, 18
  modify_url, 6, 8, 10, 11, 17, 19, 22, 25, 26, 28, 29, 31, 33, 36, 40, 48, 49
  name_backbone, 19
  name_lookup, 20
  name_suggest, 24
  name_usage, 25
  networks, 27
  nodes, 29
  occ_count, 31
  occ_fields, 32
  occ_get, 33
  occ_issues, 34
  occ_issues_lookup, 36
  occ_metadata, 36
  occ_search, 34, 37
  occurrence_density, 50
  organizations, 7, 10, 48
  parsenames, 49
  print.gbif (occ_search), 37
  providers, 50
  resources, 50
  rgb_country_codes, 50
  rgb (rgbif-package), 2
  rgbif-defunct, 50
  rgbif-package, 2
  stylegeojson, 50
  taxoncount, 50
  taxonget, 50
  taxonsearch, 50
  taxrank, 51
  togeojson, 50
  typestatus, 51