Package ‘ridigbio’

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Title Interface to the iDigBio Data API
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Description An interface to iDigBio's search API that allows downloading specimen records. Searches are returned as a data.frame. Other functions such as the metadata end points return lists of information. iDigBio is a US project focused on digitizing and serving museum specimen collections on the web. See <https://www.idigbio.org> for information on iDigBio.

URL https://github.com/iDigBio/ridigbio

BugReports https://github.com/iDigBio/ridigbio/issues

Depends R (>= 3.0.1)

Imports stats, plyr, httr, jsonlite

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LazyData true

Suggests testthat

Repository CRAN

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build_field_lists

Build fields and fields_exclude for queries.

Description

Given the desired fields to be returned, intelligently add an exclusion for the data array if warranted and handle the "all" keyword. And do so without setting both fields and fields_exclude due to fact that the API will return wrong results if are passed. This is still possible if the user deliberately sets both. Not exported.

Usage

build_field_lists(fields, type)

Arguments

fields character vector of fields user wants returned
type type of records to get fields for

Value

list list with fields key for df fields and query key for parameters to be merged with the query sent
idig_build_attrib

**Description**
Function to build attribution dataframe from a query to the iDigBio API

**Usage**
```
idig_build_attrib(dat)
```

**Arguments**
- `dat` dataframe generated by `idig_search` method

**Details**
This function differs from the attribution metadata that is attached to the dataframe returned by the `idig_search_*` methods. It summarizes the record sets used by records in the dataframe, not the record sets that have records that match the query sent to iDigBio. This is useful if only part of the records for a query are downloaded, for example with the limit and offset parameters.

Exported.

**Value**
a data frame

**Author(s)**
Kevin Love

idig_check

**Description**
Checks for HTTP error codes and JSON errors.

**Usage**
```
idig_check(req)
```

**Arguments**
- `req` the returned request
**Details**

Part 1 of the error checking process. This part handles HTTP error codes and then calls part 2 which handles JSON errors in the responses. Not exported.

**Value**

nothing. Stops if HTTP code is >= 400

**Author(s)**

Francois Michonneau

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**idig_check_error**

*Check is the request returned an error.*

---

**Description**

Checks for error messages that can be returned by the API in JSON.

**Usage**

`idig_check_error(req)`

**Arguments**

`req` the returned request

**Details**

Part 2 of the error checking process. Checks the JSON response for error messages and stops if any are found. Not exported.

**Value**

nothing. Stops if request contains an error.

**Author(s)**

Francois Michonneau
**idig_count_media**  
*Count media endpoint*

**Description**
Count media records matching a query.

**Usage**
```
idig_count_media(rq = FALSE, mq = FALSE, ...)
```

**Arguments**
- `rq`: iDigBio record query in nested list format
- `mq`: iDigBio media query in nested list format
- `...`: additional parameters

**Details**
Quickly return a count of the media records matching the query(s) provided.

**Value**
count of media records matching the query(s)

**Author(s)**
Matthew Collins

---

**idig_count_records**  
*Count record endpoint*

**Description**
Count specimen records matching a query.

**Usage**
```
idig_count_records(rq = FALSE, ...)
```

**Arguments**
- `rq`: iDigBio record query in nested list format
- `...`: additional parameters
Details

Quickly return a count of the specimen records matching the query(s) provided.

Value

count of specimen records matching the query(s)

Author(s)

Matthew Collins

---

**idig_GET**

(internal GET request)

Description

Internal function for GET requests.

Usage

```r
idig_GET(path, ...)
```

Arguments

- `path` endpoint
- `...` additional arguments to be passed to `httr::GET`

Details

Generates a GET request and performs the checks on what is returned. Not exported.

Value

the request (as a list)

Author(s)

Francois Michonneau
### idig_meta_fields

**Description**

List of fields in iDigBio.

**Usage**

```
idig_meta_fields(type = "records", subset = FALSE, ...)
```

**Arguments**

- **type**  
  string type of fields to return, defaults to "records"
- **subset**  
  set of fields to return, "indexed", "raw", or unset for all
- **...**  
  additional parameters

**Details**

Return a list of media or specimen fields that are contained in iDigBio.

**Value**

list of fields of the requested type

**Author(s)**

Matthew Collins

---

### idig_parse

**Description**

Parses output of successful query to return a list.

**Usage**

```
idig_parse(req)
```

**Arguments**

- **req**  
  the returned request
Details
Not exported.

Value
a list

Author(s)
Francois Michonneau

Description
Internal function for POST requests.

Usage
idig_POST(path, body, ...)

Arguments
path endpoint
body a list of parameters for the endpoint
... additional arguments to be passed to httr::POST

Details
Generates a POST request and performs the checks on what is returned. Not exported.

Value
the request (as a list)

Author(s)
Francois Michonneau
idig_search

Basic searching of iDigBio records

Description

Base function to query the iDigBio API

Usage

idig_search(type = "records", mq = FALSE, rq = FALSE, fields = FALSE,
max_items = 1e+05, limit = 0, offset = 0, sort = FALSE, ...)

Arguments

type string type of records to query, defaults to "records"

mq iDigBio media query in nested list format

rq iDigBio record query in nested list format

fields vector of fields that will be contained in the data.frame

max_items CURRENTLY IGNORED, SEE ISSUE #33 maximum number of results allowed to be retrieved (fail-safe)

limit maximum number of results returned

offset number of results to skip before returning results

sort vector of fields to use for sorting, UUID is always appended to make paging safe

... additional parameters

Details

This function is wrapped for media and specimen record searches. Please consider using idig_search_media or idig_search_records instead as they supply nice defaults to this function depending on the type of records desired.

Fuller documentation of parameters is in the idig_search_records function’s help.

Exported to facilitate wrapping this package in other packages.

Value

a data frame

Author(s)

Francois Michonneau
Examples

## not run:
# Ten media records related to genus Acer specimens
idig_search(type="media", rq=list(genus="acer"), limit=10)

## end(not run)

---

dig_search_media | Searching of iDigBio media records

Description

Function to query the iDigBio API for media records

Usage

idig_search_media(mq = FALSE, rq = FALSE, fields = FALSE,
max_items = 1e+05, limit = 0, offset = 0, sort = FALSE, ...)

Arguments

mq | iDigBio media query in nested list format
rq | iDigBio record query in nested list format
fields | vector of fields that will be contained in the data.frame, defaults to "all" which is all indexed fields
max_items | maximum number of results allowed to be retrieved (fail-safe)
limit | maximum number of results returned
offset | number of results to skip before returning results
sort | vector of fields to use for sorting, UUID is always appended to make paging safe
... | additional parameters

Details

Also see idig_search_records for the full examples of all the parameters related to searching iDigBio.

Wraps idig_search to provide defaults specific to searching media records. Using this function instead of idig_search directly is recommended. Record queries and media queries objects are allowed (rq and mq parameters) and media records returned will match the requirements of both.

This function defaults to returning all indexed media record fields.

Value

a data frame
Author(s)
Matthew Collins

Examples

```r
## Not run:
# Searching for media using a query on related specimen information - first
# 10 media records with image URIs related to a specimen in the genus Acer:
df <- idig_search_media(rq=list(genus="acer"),
    mq=list("data.ac:accessURI"="list("type"="exists"),
    fields=c("uuid","data.ac:accessURI"), limit=10)

## End(Not run)
```

**idig_search_records**  
Searching of iDigBio records

**Description**

Function to query the iDigBio API for specimen records

**Usage**

```r
idig_search_records(rq, fields = FALSE, max_items = 1e+05, limit = 0,
    offset = 0, sort = FALSE, ...)
```

**Arguments**

- `rq`  
  iDigBio record query in nested list format
- `fields`  
  vector of fields that will be contained in the data.frame, limited set returned by default, use "all" to get all indexed fields
- `max_items`  
  maximum number of results allowed to be retrieved (fail-safe)
- `limit`  
  maximum number of results returned
- `offset`  
  number of results to skip before returning results
- `sort`  
  vector of fields to use for sorting, UUID is always appended to make paging safe
- `...`  
  additional parameters

**Details**

Wraps `idig_search` to provide defaults specific to searching specimen records. Using this function instead of `idig_search` directly is recommended.

Queries need to be specified as a nested list structure that will serialize to an iDigBio query object’s JSON as expected by the iDigBio API:  

As an example, the first sample query looks like this in JSON in the API documentation:
To rewrite this in R for use as the rq parameter to idig_search_records or idig_search_media, it would look like this:

```r
rq <- list("scientificname"=list("type"="exists"),
            "family"="asteraceae" )
```

An example of a more complex JSON query with nested structures:

```json
{
  "geopoint": {
    "type": "geo_bounding_box",
    "top_left": {
      "lat": 19.23,
      "lon": -130
    },
    "bottom_right": {
      "lat": -45.1119,
      "lon": 179.99999
    }
  }
}
```

To rewrite this in R for use as the rq parameter, use nested calls to the list() function:

```r
rq <- list(geopoint=list(
  type="geo_bounding_box",
  top_left=list(lat=19.23, lon=-130),
  bottom_right=list(lat=-45.1119, lon=179.99999)
  )
)
```

See the Examples section below for more samples of simpler and more complex queries. Please refer to the API documentation for the full functionality available in queries.

All matching results are returned up to the max_items cap (default 100,000). If more results are wanted, a higher max_items can be passed as an option. This API loads records 5,000 at a time using HTTP so performance with large sets of data is not very good. Expect result sets over 50,000 records to take tens of minutes. You can use the idig_count_records or idig_count_media functions to find out how many records a query will return; these are fast.

The iDigBio API will only return 5,000 records at a time but this function will automatically page through the results and return them all. Limit and offset are available if manual paging of results is
needed though the max_items cap still applies. The item count comes from the results header not the count of actual records in the limit/offset window.

Return is a data.frame containing the requested fields (or the default fields). The columns in the data frame are untyped and no factors are pre-built. Attribution and other metadata is attached to the dataframe in the data.frame's attributes. (I.e. attributes(df))

Value

a data frame

Author(s)

Matthew Collins

Examples

```r
# Simple example of retrieving records in a genus:
idig_search_records(rq=list(genus="acer"), limit=10)

# This complex query shows that booleans passed to the API are represented
# as strings in R, fields used in the query don't have to be returned, and
# the syntax for accessing raw data fields:
idig_search_records(rq=list("hasImage"="true", genus="acer"),
                       fields=c("uuid", "data.dwc:verbatimLatitude"), limit=100)

# Searching inside a raw data field for a string, note that raw data fields
# are searched as full text, indexed fields are searched with exact matches:
idig_search_records(rq=list("data.dwc:dynamicProperties"="parasite"),
                       fields=c("uuid", "data.dwc:dynamicProperties"), limit=100)

# Retrieving a data.frame for use with MaxEnt. Notice geopoint is expanded
# to two columns in the data.frame: geopoint.lat and geopoint.lon:
df <- idig_search_records(rq=list(genus="acer", geopoint=list(type="exists")),
                          fields=c("uuid", "geopoint"), limit=10)
write.csv(df[cbind("uuid", "geopoint.lon", "geopoint.lat")],
          file="acer_occurrences.csv", row.names=FALSE)
```

## End(Not run)

---

**idig_top_media**

*Top media endpoint*

**Description**

Top media records summaries.
idig_top_records

Usage

```r
idig_top_media(rq = FALSE, mq = FALSE, top_fields = FALSE, count = 0, 
...)
```

Arguments

- **rq**: iDigBio record query in nested list format
- **mq**: iDigBio media query in nested list format
- **top_fields**: vector of field names to summarize by
- **count**: maximum number of results to return, capped at 1000
- **...**: additional parameters

Details

Summarize the count of media records in iDigBio according to unique values in the fields passed. This operates similarly to a SELECT DISTINCT count(field_name) query in SQL. When multiple fields are passed, the summaries are nested eg fields=c("country", "genus") would result in counting the top 10 genera in each of the top 10 countries for a total of 100 counts.

Value

nested list of field values with counts of media records

Author(s)

Matthew Collins

---

idig_top_records]

Top records endpoint

Description

Top specimen records summaries.

Usage

```r
idig_top_records(rq = FALSE, top_fields = FALSE, count = 0, ...)
```

Arguments

- **rq**: iDigBio record query in nested list format
- **top_fields**: vector of field names to summarize by
- **count**: maximum number of results to return, capped at 1000
- **...**: additional parameters
Details

Summarize the count of specimen records in iDigBio according to unique values in the fields passed. This operates similarly to a SELECT DISTINCT count(field_name) query in SQL. When multiple fields are passed, the summaries are nested eg fields=c("country", "genus") would result in counting the top 10 genera in each of the top 10 countries for a total of 100 counts.

Value

nested list of field values with counts of specimen records

Author(s)

Matthew Collins

---

idig_url  

base URL

Description

Return base URL for the API calls.

Usage

idig_url(dev = FALSE)

Arguments

dev  Should be the beta version of the API be used?

Details

Defaults to use beta URL, Not exported.

Value

string for the URL

Author(s)

Francois Michonneau
idig_validate validate fields

**Description**
Stub function for validating parameters.

**Usage**

```R
idig_validate(inputs)
```

**Arguments**

- `inputs` list of inputs to validate

**Details**
Takes list of inputs named by validation rule eg "number":[2, 3] and returns a vector of strings with any validation errors. If the vector is 0 length, everything is valid. Not exported.

**Value**
boolean

**Author(s)**
Matthew Collins

idig_version API version

**Description**
Return the version number to use for the API calls.

**Usage**

```R
idig_version(version = "v2")
```

**Arguments**

- `version` optional argument giving the version of the API to use

**Details**
The current default is "v2". Not exported.
idig_view_media

**Value**

string for the version to use

**Author(s)**

Francois Michonneau

---

idig_view_media view media endpoint

**Description**

View individual media records.

**Usage**

idig_view_media(uuid, ...)

**Arguments**

<table>
<thead>
<tr>
<th>uuid</th>
<th>uuid of media record</th>
</tr>
</thead>
<tbody>
<tr>
<td>...</td>
<td>additional parameters</td>
</tr>
</tbody>
</table>

**Details**

View all information about a specific media record.

**Value**

nested list of data

**Author(s)**

Matthew Collins
idig_view_records  
view specimen endpoint

Description

View individual specimen records.

Usage

idig_view_records(uuid, ...)

Arguments

uuid      uuid of specimen record
...       additional parameters

Details

View all information about a specific specimen record.

Value

nested list of data

Author(s)

Matthew Collins

ridigbio

Retrieve data from the iDigBio specimen data repository.

Description

Retrieve data from the iDigBio specimen data repository.

About

ridigbio provides an interface to the iDigBio data API described here: https://www.idigbio.org/wiki/index.php/IDigBio_API. With this package you can retrieve specimen and media records from the iDigBio data repository. The iDigBio portal https://portal.idigbio.org/ uses the same API so you should be able to retrieve the same information as shown in the portal.

iDigBio contains nearly 30 million data records on museum specimens held at United States institutions. It also holds nearly 5 million images of these specimens.
Getting Started

The main function is `idig_search_records` and reviewing its documentation first with `?idig_search_records` is recommended.

Limitations

This package does not yet provide an interface to the mapping or the download APIs.

Citing

To cite the ridigbio package in your work, please use the following format:


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