Package ‘rbison’

June 8, 2020

Title Interface to the 'USGS' 'BISON' API

Description Interface to the 'USGS' 'BISON' (<https://bison.usgs.gov/>)

API, a 'database' for species occurrence data. Data comes from
species in the United States from participating data providers. You can get
data via 'taxonomic' and location based queries. A simple function
is provided to help visualize data.

Version 1.0.0

License MIT + file LICENSE

URL https://github.com/ropensci/rbison (devel)

https://docs.ropensci.org/rbison (docs)

BugReports https://github.com/ropensci/rbison/issues

LazyData true

VignetteBuilder knitr

Encoding UTF-8

Language en-US

Imports plyr, crul (>= 0.7.4), ggplot2, mapproj, grid, sp, dplyr (>=

0.5.0), jsonlite (>= 1.1), data.table, tibble

Suggests knitr, rmarkdown, testthat, taxize, vcr

RoxygenNote 7.1.0

X-schema.org-applicationCategory DataAccess

X-schema.org-keywords species, occurrences, biodiversity, maps, USGS,

BISON

X-schema.org-isPartOf “https://ropensci.org”

NeedsCompilation no

Author Scott Chamberlain [aut, cre] (<https://orcid.org/0000-0003-1444-9135>),

rOpenSci [fnd] (https://ropensci.org)

Maintainer Scott Chamberlain <myrmecocystus@gmail.com>

Repository CRAN

Date/Publication 2020-06-08 18:30:03 UTC
rbison-package

R topics documented:

rbison-package ................................................. 2
all_states ....................................................... 3
bison ............................................................. 3
bisonmap .......................................................... 6
bison_datause ..................................................... 7
bison_providers ................................................... 8
bison_solr .......................................................... 9
bison_stats ........................................................ 14
bison_tax .......................................................... 15
fips ................................................................. 16
is.bison ............................................................ 17
is.bison_solr ...................................................... 17

Index 18

rbison-package  rbison is an interface to the USGS Bison API.

Description

To get started, see the vignette vignette(package="rbison")

Details

See https://bison.usgs.gov/doc/api.jsp for API docs for the BISON API.

To cite rbison, do citation(package='rbison')

Use the following format to cite data retrieved from BISON:


For example:


Base URL for the BISON API: https://bison.usgs.gov

Author(s)

Scott Chamberlain <myrmecocystus@gmail.com>
all_states

---

Data for a states map

---

bison

Search for and collect data from the USGS Bison API.

---

**Description**

Search for and collect data from the USGS Bison API.

**Usage**

```r
bison(
  species = NULL,
  type = "scientific_name",
  tsn = NULL,
  start = 0,
  count = 25,
  countyFips = NULL,
  county = NULL,
  state = NULL,
  aoi = NULL,
  aoibbox = NULL,
  params = NULL,
  ...
)
```

**Arguments**

- `species` (character): A species name.
- `type` (character): Type, one of `scientific_name` or `common_name`.
- `tsn` (numeric): Specifies the TSN to query by. If you supply a tsn it doesn’t make sense to supply a species name as well. Example: 162003.
- `start` (numeric): Record to start at. Default: 0. See "Pagination" in Details.
- `count` (numeric): Number of records to return. Default: 25. See "Pagination" in Details.
- `countyFips` (character): Specifies the county fips code to geographically constrain the search to one county. Character must be supplied as a number starting with zero may lose the zero. Eg: "49015".
county (character) County name. As codes are a pain in the ass, you can put in the county name here instead of specifying a countyFips entry, and bison will attempt to look up the countyFips code. (character)

state (character) Specifies the state name to geographically constrain the search. Example: Tennessee.

aoi (character) Specifies a WKT (Well-Known Text) polygon to geographically constrain the search. Eg.: c(-111.06 38.84, -110.80 39.377, -110.20 39.17, -110.20 38.90, -110.63 38.67, -111.06 38.84), which calls up the occurrences within the specified area. Check out the Wikipedia page here http://en.wikipedia.org/wiki/Well-known_text for an in depth look at the options, terminology, etc.

aobbox (character) Specifies a four-sided bounding box to geographically constrain the search (using format: minx,miny,maxx,maxy). The coordinates are Spherical Mercator with a datum of WGS84. Example: -111.31,38.81,-110.57,39.21

params (character) String of parameters, one of providerID, resourceID, basisOfRecord, catalogNumber, year, computedStateFips, hierarchy_homonym_string, TSNs, recordedBy, occurrenceID, collectorNumber, provider, ownerInstitutionCollectionCode, eventDate, providedScientificName, scientificName, ITISscientificName, providedCommonName, ITIScommonName, kingdom, ITISTsn, centroid, higherGeographyID, computedCountyFips, providedCounty, calculatedCounty, stateProvince, calculatedState, countryCode. See examples.

Further args passed on to crul::HttpClient(). See examples.

Pagination

bison() paginates internally for you on the count parameter, so that for example, if you request 2000 records, then we’ll do two requests to get all those records. If you request for example 50 records, then we just do one request.

References

https://bison.usgs.gov/#opensearch

See Also

bison_solr() bison_tax()

Examples

```r
## Not run:
bison(species="Bison bison", count=50)

# lots of results
res <- bison(species="Bison bison", count=2000)
res$summary
NROW(res$points)

out <- bison(species="Helianthus annuus", count=300)
out$summary # see summary
```
bison

out$counties # see county data
out$states # see state data
bisonmap(out, tomap = "points")
bisonmap(out, tomap = "county")
bisonmap(out, tomap = "state")

# Search for a common name
bison(species="Tufted Titmouse", type="common_name")

# Constrain search to a certain county, 49015 is Emery County in Utah
bison(species="Helianthus annuus", countyFips = "49015")

# Constrain search to a certain county, specifying county name instead of
# code
bison(species="Helianthus annuus", county = "Los Angeles")
bison(species="Helianthus annuus", county = "Los")

# Constrain search to a certain aoi, which turns out to be Emery County,
# Utah as well
bison(species="Helianthus annuus",
aoi = "POLYGON((-111.06360117772908 38.84001566645886,
-110.80542246679359 39.377077711107983,
-110.2017441992392 39.17722368276862,
-110.20666758398464 38.908844075244811,
-110.63513438085685 38.67724220095734,
-111.06360117772908 38.84001566645886))")

# Constrain search to a certain aoibbox, which, you guessed it, is also
# Emery Co., Utah
bison(species="Helianthus annuus", aoibbox = '-111.31,38.81,-110.57,39.21')

# Taxonomic serial number
bison(tsn = 162003)

## If you don't have tsn's, search for a taxonomic serial number
library('taxize')
poa_tsn <- get_tsn('Poa annua')
bison(tsn = poa_tsn)

# Curl debugging and some of these examples aren't
# that useful, but are given for demonstration purposes
## get curl verbose output to see what's going on with your request
bison(tsn = 162003, count=1, verbose = TRUE)
## set a timeout so that the call stops after time x, compare 1st to 2nd call
# bison(tsn=162003, count=1, timeout_ms = 1)
## set cookies
bison(tsn=162003, count=1, cookie = "a=1;b=2")
## user agent and verbose
bison(tsn=162003, count=1, useragent = "rbison",
   verbose = TRUE)

# Params - the params function accepts a number of search terms
## Find the provider with ID 318.
bison(params='providerID:("318")')
## Find all resources with id of '318,1902' OR '318,9151', with values separated by spaces.
```r
bison(params='resourceID:("318,1902" "318,9151")')
```
## Criterion may be combined using the semicolon (';') character, which translates to a logical AND operator. Note that field names and values are case sensitive.
```r
bison(params='providerID:("408" "432");resourceID:("14027")')
```
## Search by basisOfRecord, for specimen types in this case
```r
bison(params='basisOfRecord:(specimen)')
```
## Search by computedStateFips, 01 for Alabama
```r
bison(params='computedStateFips:01')
```
## Search by ITIStsn
```r
bison(params='ITIStsn:162003')
```
## Search by countryCode
```r
bison(params='countryCode:US')
```
## Search by ITIScommonName
```r
bison(params='ITIScommonName:"Canada goose"')
```
## End(Not run)

---

**bisonmap**

*Make map to visualize BISON data.*

### Description

Make map to visualize BISON data.

### Usage

```r
bisonmap(
  input = NULL,
  tomap = "points",
  geom = geom_point,
  jitter = NULL,
  customize = NULL
)
```

```r
# S3 method for class 'bison'
bisonmap(
  input = NULL,
  tomap = "points",
  geom = geom_point,
  jitter = NULL,
  customize = NULL
)
```

```r
# S3 method for class 'bison_solr'
bisonmap(
  input = NULL,
  tomap = "points",
  geom = geom_point,
  jitter = NULL,
  customize = NULL
)
```
input = NULL,
tomap = "points",
geom = geom_point,
jitter = NULL,
customize = NULL
)

Arguments

input Input bison object.
tomap One of points (occurrences), county (counts by county), or state (counts by state). For bison class objects, we stop with message if you pass in data from bison() that doesn’t include data by each state. For bison_solr class objects, we only allow points.
geom geom_point or geom_jitter, not quoted.
jitter jitter position, see ggplot2 help.
customize Pass in more to the plot.

Value

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

Examples

## Not run:
# Using function bison
library("ggplot2")
out <- bison(species="Accipiter", type="scientific_name", count=300)
bisonmap(input=out)
bisonmap(input=out, geom=geom_jitter, jitter=position_jitter(width = 0.3, height = 0.3))

# Using function bison_solr
out <- bison_solr(scientificName=’Ursus americanus’, rows=200)
bisonmap(out)

## End(Not run)

bison_datause

Get BISON data use agreement details and examples for how to cite data.

Description

Get BISON data use agreement details and examples for how to cite data.
Usage

bison_datause()

bison_citation()

References

https://bison.usgs.gov/doc/api.jsp#data

bison_providers  Get information about BISON data providers.

Description

Get information about BISON data providers.

Usage

bison_providers(details = FALSE, provider_no = NULL, ...)

Arguments

details  (logical) If TRUE, returns a list of data.frame’s for each provider, including their resource details. If FALSE (default), only coarse grained data returned.

provider_no  (numeric) Provider number. If this parameter is provided, details is forced to be FALSE

...  Further args passed on to crul::HttpClient() See examples in bison()

Value

A data.frame or list of data.frame’s

Examples

## Not run:
head(bison_providers())
head(bison_providers(provider_no=131))
out <- bison_providers(details=TRUE)
out$National_Herbarium_of_New_South_Wales

## End(Not run)
bison_solr

Search for and collect occurrence data from the USGS Bison API using their solr endpoint.

Description

This fnx is somewhat similar to bison(), but interacts with the SOLR interface https://bison.usgs.gov/#solr instead of the OpenSearch interface https://bison.usgs.gov/#opensearch, which bison() uses.

Usage

bison_solr(
  decimalLatitude = NULL,
  decimalLongitude = NULL,
  year = NULL,
  providerID = NULL,
  resourceID = NULL,
  pointPath = NULL,
  basisOfRecord = NULL,
  eventDate = NULL,
  computedCountyFips = NULL,
  computedStateFips = NULL,
  scientificName = NULL,
  hierarchy_homonym_string = NULL,
  TSNs = NULL,
  recordedBy = NULL,
  occurrenceID = NULL,
  catalogNumber = NULL,
  ITIScommonName = NULL,
  kingdom = NULL,
  collectorNumber = NULL,
  provider = NULL,
  ownerInstitutionCollectionCode = NULL,
  providedScientificName = NULL,
  ITISscientificName = NULL,
  providedCommonName = NULL,
  ITIStsn = NULL,
  centroid = NULL,
  higherGeographyID = NULL,
  providedCounty = NULL,
  calculatedCounty = NULL,
  stateProvince = NULL,
  calculatedState = NULL,
  countryCode = NULL,
  callopts = list(),
  verbose = TRUE,
Arguments

decimalLatitude
Geographic coordinate that specifies the north-south position of a location on
the Earth surface.

decimalLongitude
Geographic coordinate that specifies the east-west position of a location on the
Earth surface.

year
The year the collection was taken.

providerID
(character) Unique identifier assigned by GBIF.

resourceID
(character) A unique identifier that is a concatenation of the provider identifier
and the resource ID separated by a comma.

pointPath
A dynamic field that contains the location in longitude and latitude followed
by the basis of record and an optional Geo (Spatial) precision. Geo (Spatial)
precision is an added descriptor when the record is a county centroid.

basisOfRecord
One of these enumerated values: Observation, Germplasm, Fossil, Specimen,
Literature, Unknown, or Living.

eventDate
The date when the occurrence was recorded. Dates need to be of the form
YYYY-MM-DD

computedCountyFips
County FIPS code conforming to standard FIPS 6-4 but with leading zeros re-
moved. See fips dataset for codes

computedStateFips
The normalized state FIPS code. See fips dataset for codes

scientificName
The species scientific name that is searchable in a case insensitive way.

hierarchy_homonym_string
Hierarchy of the accepted or valid species name starting at kingdom. If the
name is a taxonomic homonym more than one string is provided separated by
';'.

TSNs
Accepted or valid name is provided. If the name is a taxonomic homonym more
than one TSN is provided.

recordedBy
Individual responsible for the scientific record.

occurrenceID
Non-persistent unique identifier.

catalogNumber
Unique key for every record (occurrence/row) within a dataset that is not ma-
nipulated nor changed (nor generated, if not provided) during the data ingest.

ITIScommonName
Common name(s) from ITIS, e.g. "Canada goose"

kingdom
Kingdom name, from GBIF raw occurrence or BISON provider.

collectorNumber
An identifier given to the occurrence at the time it was recorded, such as a spec-
imen collector's number. e.g., "SIM030022".

provider
Non-persistent unique identifier.
ownerInstitutionCollectionCode
   Name for the dataset, format = OwnerInstitution-Collection. / e.g., "USGS
   NAWQA BioData - Fish Occurrence Records"

providedScientificName
   Full scientific name as provided in the dataset, with authorship and date inform-
   ation if known.

ITISscientificName
   Scientific name from join on ITIS table, calculated e.g., "Bison bison"

providedCommonName
   A list (concatenated and separated) of the available vernacular species names. / e.g., "common shrew, Masked Shrew"

ITIStsn
   Phase II: ITIS TSN corresponding to clean_provided_scientific_name. May be invalid,unaccepted. Calculated. e.g., "3250", "05713"

centroid
   Text string indicating that provided lat/lon point represents a polygon centroid. Text provides description of the centroid.

higherGeographyID
   5-digit numeric text string geographic code for the state-county combination
   provided by data provider. / e.g., "13029"

providedCounty
   Full county, parish, or organized borough name, as provided in the dataset. If
   provided, Verbatim State is required. Is not changed during data ingest. / e.g.,
   "Fairfax"

calculatedCounty
   Full county, parish, or organized borough name of the occurrence calculated. / e.g.,
   "Fairfax"

stateProvince
   Full name of state or territory of the occurrence, as provided in the dataset.

calculatedState
   U.S. State or territory name calculated. e.g., "Puerto Rico"

countryCode
   The geographic location of the specific occurrence, expressed through a con-
   strained vocabulary of countries using 2-letter ISO country code.

callopts
   Further args passed on to \texttt{crul::HttpClient()} for HTTP debugging/inspecting. In \texttt{bison}, \texttt{bison_providers}, and \texttt{bison_stats}, \ldots is used instead of callopts, but \ldots is used here to pass additional Solr params.

verbose
   Print message with url (TRUE, default).

... Additional SOLR query arguments. See details.

Details

Named parameters in this function are combined with \texttt{AND} and passed on to \texttt{q SOLR} parameter. Of course parameters can be more flexibly combined - let us know if you want that flexibility and we can think about that.

Value

An object of class \texttt{bison_solr} - which is a list with slots for number of records found (\texttt{num_found}), records, highlight, or facets.
SOLR search parameters passed on via . . .

- fl: Fields to return in the query
- rows: Number of records to return
- start: Record number to start at (an offset)
- sort: Field to sort by, see examples
- facet: Facet or not, logical
- facet.field: Fields to facet by

To do pagination, use rows and start together.

You can also use highlighting in solr search, but I’m not sure I see a use case for it with BISON data, though you can do it with this function.

For a tutorial see here http://lucene.apache.org/solr/3_6_2/doc-files/tutorial.html

Range searches

If you pass a vector of length 2 to a parameter we construct a range query for you. For example, c(4, 5) turns into [4 TO 5]. The [] syntax means the search is inclusive, meaning 4 to 5, including 4 and 5. Let us know if you think you need more flexible searching. That is, doing exclusive {} or mixed searches ({} or {}). Range searches can only be done with variables that are numeric/integer or dates or strings that can be coerced to dates. Dates need to be of the form YYYY-MM-DD

References

https://bison.usgs.gov/#solr

See Also

bison_tax(), bison()

The USGS BISON Solr installation version as of 2014-10-14 was 4.4.

Examples

```r
# Not run:
x = bison_solr(scientificName = 'Ursus americanus')

bison_solr(scientificName = 'Ursus americanus', computedStateFips = '02', fl = 'scientificName')

x <- bison_solr(scientificName = 'Ursus americanus', computedStateFips = '02', rows = 50)
x$points$computedStateFips
head(x$points)

bison_solr(ITISscientificName = 'Ursus americanus', rows = 50)

bison_solr(providerID = 220)

# pagination
bison_solr(scientificName = 'Ursus americanus', rows = 10)
```
bison_solr(scientificName = 'Ursus americanus', rows = 10, start = 10)

# combining parameters
x <- bison_solr(eventDate = c('2008-01-01', '2010-12-31'),
    ITISscientificName="Helianthus annuus", rows = 100)
head(x$points)
sort(x$points$eventDate)

# range queries
## range search with providerID
bison_solr(providerID = c(220, 221))
## date range search
x <- bison_solr(eventDate = c('2010-08-08', '2010-08-21'))
sort(x$points$eventDate)
## TSN range search
x <- bison_solr(TSNs = c(174773, 174775), rows = 100)
sort(x$points$TSN)
## can't do range searches with character strings (that are not dates)
# bison_solr(kingdom = c("Animalia", "Plantae"))

# more examples
bison_solr(TSNs = 174773)
bison_solr(occurrenceID = 576630651)
bison_solr(catalogNumber = 'OBS101299944')
bison_solr(ITIScommonName = "Canada goose")
bison_solr(kingdom = "Animalia")
bison_solr(kingdom = "Plantae")

# Mapping
out <- bison_solr(scientificName='Ursus americanus', rows=200)
bisonmap(out)

out <- bison_solr(scientificName='Helianthus annuus', rows=800)
bisonmap(out)

# Using additional solr fields
## Faceting
bison_solr(scientificName='Helianthus annuus', rows=0, facet='true',
    facet.field='computedStateFips')

## Highlighting
bison_solr(scientificName='Helianthus annuus', rows=10, hl='true',
    hl.fl='scientificName')

## Use of hierarchy_homonym_string
bison_solr(hierarchy_homonym_string = '-202423-914154-914156-158852-')
## -- This is a bit unwieldy, but you can find this string in the output
## of a call, like this
x <- bison_solr(scientificName='Ursus americanus', rows=1)
string <- x$points$hierarchy_homonym_string
bison_solr(hierarchy_homonym_string = string)

# The pointPath parameter
bison_stats(pointPath = '-110.0,45.0/specimen')

# Curl options
bison_solr(scientificName='Ursus americanus', callopts=list(verbos e = TRUE))

## End(Not run)

---

**bison_stats**  
*Get statistics about BISON downloads.*

**Description**  
Get statistics about BISON downloads.

**Usage**  
`bison_stats(what = "stats", ...)`

**Arguments**  
- **what** (character) One of stats (default), search, download, or wms. See Details.
- **...** Further args passed on to `curl::HttpClient()`. See examples in `bison`

**Details**  
For the `what` parameter:
- **stats** - Retrieve all data provider accumulated statistics.
- **search** - Retrieve data provider statistics for BISON searches.
- **download** - Retrieve data provider statistics for data downloads from BISON.
- **wms** - Retrieve data provider statistics for BISON OGC WMS tile requests.

**Value**  
A list of data frame’s with names of the list the different data sources

**Examples**  
## Not run:
```r
out <- bison_stats()
out <- bison_stats(what='wms')
out$Arctos
out$Harvard_University_Herbaria
out$ZooKeys
```

## End(Not run)
**bison_tax**

Search for and collect taxonomic name data from the USGS Bison API using solr

---

**Description**

See the SOLR documentation here [http://lucene.apache.org/solr/](http://lucene.apache.org/solr/) for other parameters you can use.

The following two methods are possible, as far as I know you can only use one at a time:

- **vernacularName** The species specific common names that is searchable in a case insensitive way.
- **scientificName** The species scientific name that is associated with a common name that is searchable in a case insensitive way.

**Usage**

```r
bison_tax(
  query,
  method = "vernacularName",
  exact = FALSE,
  parsed = TRUE,
  callopts = list(),
  ...
)
```

**Arguments**

- **query** Name to search for. Required.
- **method** The field to query by. See description below for details.
- **exact** Exact matching or not. See examples. Defaults to FALSE.
- **parsed** If TRUE (default) creates data.frame of names data output. Otherwise, a list.
- **callopts** Further args passed on to `crul::HttpClient()` for HTTP debugging/inspecting. In `bison`, `bison_providers`, and `bison_stats`, ... is used instead of callopts, but ... is used here to pass additional Solr params.
- **...** Further solr arguments passed in to the query. See examples below.

**Value**

A list.

**See Also**

`bison_solr()`, `bison()`
Examples

```r
## Not run:
# All taxa
bison_tax("*:*")

# Some example calls
bison_tax(query="*bear")
bison_tax(query="Helianthus", method="scientificName")

# Exact argument, here nothing found with latter call as 'bear'
# doesn't exist, which makes sense
bison_tax(query="*bear", exact=FALSE)
bison_tax(query="*bear", exact=TRUE)

# Using solr arguments (not all Solr arguments work)
## Return a certain number of rows
bison_tax(query="*bear", method="vernacularName", rows=3)
## Return certain fields
bison_tax(query="*bear", method="vernacularName", fl='vernacularName')

# Curl options
bison_tax(query='*dolphin', callopts=list(verbos = TRUE))
```

## End(Not run)

---

### fips

*Fips codes for states and counties*

**Description**

See https://www.census.gov/geo/reference/codes/cou.html and https://www.census.gov/geo/reference/ansi_statetables.html for more information on FIPS codes

**Format**

A data frame with 3142 rows and 4 variables:

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>state</td>
<td>State name</td>
</tr>
<tr>
<td>county</td>
<td>County name</td>
</tr>
<tr>
<td>fips_state</td>
<td>State FIPS code</td>
</tr>
<tr>
<td>fips_county</td>
<td>County FIPS code</td>
</tr>
</tbody>
</table>
is.bison

Check if object is of class bison

Description

Check if object is of class bison

Usage

is.bison(x)

Arguments

x  input

is.bison_solr

Check if object is of class bison_solr

Description

Check if object is of class bison_solr

Usage

is.bison_solr(x)

Arguments

x  input
Index

*Topic **data**
  - all_states, 3
  - fips, 16
*Topic **package**
  - rbison-package, 2

all_states, 3

bison, 3
bison(), 7–9, 12, 15
bison_citation (bison_datause), 7
bison_datause, 7
bison_providers, 8
bison_solr, 9
bison_solr(), 4, 15
bison_stats, 14
bison_tax, 15
bison_tax(), 4, 12
bisonmap, 6

crul::HttpClient(), 4, 8, 11, 14, 15

fips, 10, 16

is.bison, 17
is.bison_solr, 17

rbison (rbison-package), 2
rbison-package, 2