Package: naturecounts (via r-universe)

October 11, 2024

```
Title Access and download data on plant and animal populations from
     NatureCounts
Version 0.4.1
Description Access and download data on plant and animal populations
     from various databases through NatureCounts, a service managed
     by Bird Studies Canada.
License GPL-3
Encoding UTF-8
LazyData true
Depends R (>= 3.1.2)
Imports askpass (>= 1.1), DBI (>= 1.0.0), dbplyr (>= 1.4.4), dplyr (>=
     1.1.0), ggplot2 (>= 3.4.0), httr (>= 1.4.0), isonlite (>= 1.6),
     lubridate (>= 1.7.4), magrittr, memoise (>= 2.0.1), purrr (>=
     0.3.2), rlang (>= 0.1.2), RSQLite (>= 2.1.1), stringi (>=
     1.2.4), stringr (>= 1.3.1), tibblify (>= 0.3.0), tidyr (>=
     0.8.2), tidyselect (>= 1.0.0), units (>= 0.8.1), wk (>= 0.7.1)
Suggests ggspatial, here, knitr, mapview, rnaturalearth, roxygen2,
     rmarkdown, sf (>= 1.0-9), spelling, testthat, vdiffr
Language en-US
Roxygen list(markdown = TRUE)
RoxygenNote 7.3.1
URL https://github.com/BirdsCanada/naturecounts,
     https://naturecounts.ca,
     https://birdscanada.github.io/naturecounts/
VignetteBuilder knitr
Config/testthat/edition 3
Repository https://birdscanada.r-universe.dev
RemoteUrl https://github.com/birdscanada/naturecounts
RemoteRef HEAD
RemoteSha ec11e608201d7dbd2bd9e7a878a8c248b435e4f4
```

Type Package

2 naturecounts-package

Contents

natur	recounts-package	Access and a tureCounts	downlo	ad data o	on plant and	animal populo	ations from Na	-
Index								28
	search_species_code			• • • •				26
	search_species							
	search_region							
	nc_requests							
	nc_remove_cache .							
	nc_query_table							21
	nc_permissions							20
	nc_metadata							19
	nc_data_dl							16
	nc_count							13
	meta							11
	map_canada							10
	hofi							10
	grid_canada							9
	format_zero_fill							7
	format_dates							7
	cosewic_ranges							4
	cosewic_plot							
	bcch							
	naturecounts-packag	e						2

Description

naturecounts is an R package for accessing and downloading data on plant and animal populations from various databases through NatureCounts, a service managed by Bird Studies Canada.

Details

See the vignettes (vignettes (package = "naturecounts"), or at https://birdscanada.github.io/naturecounts for getting started!

Author(s)

Maintainer: Denis Lepage <dlepage@birdscanada.org> Authors:

• Steffi LaZerte <steffi@steffi.ca>

bcch 3

See Also

Useful links:

```
• https://github.com/BirdsCanada/naturecounts
```

• https://naturecounts.ca

• https://birdscanada.github.io/naturecounts/

bcch

Example black-capped chickadee data

Description

An example of black-capped chickadee data downloaded from NatureCounts

Usage

bcch

Format

A data frame with 160 rows and 57 variables:

cosewic_plot

Plot COSEWIC IAO and EOO

Description

Creates a plot of COSEWIC ranges for illustration and checking.

Usage

```
cosewic_plot(
  ranges,
  points = NULL,
  grid = NULL,
  map = NULL,
  species = "species_id",
  title = ""
)
```

4 cosewic_ranges

Arguments

ranges	List. Output of cosewic_ranges() with spatial = TRUE.
points	Data frame. Optional naturecounts data used to compute ranges. Raw data points will be added to the plot if provided.
grid	sf data frame. Optional grid over which to summarize IAO values (useful for species with many points over a broad distribution).
map	sf data frame. Optional base map over which to plot the values.
species	Character. Name of the column containing species identification.
title	Character. Optional title to add to the map. Can be a named by species vector to supply different titles for different species.

Value

ggplot2 map

Examples

cosewic_ranges

Calculate COSEWIC IAO and EOO

Description

The COSEWIC Index of Area of Occupancy (IAO; also called Area of Occupancy, AOO by the IUCN) and Extent of Occurrence (EOO; IUCN as well) are metrics used to support status assessments for potentially endangered species.

cosewic_ranges 5

Usage

```
cosewic_ranges(
  df_db,
  record = "record_id",
  coord_lon = "longitude",
  coord_lat = "latitude",
  species = "species_id",
  iao_grid_size_km = 2,
  eoo_p = 0.95,
  filter_unique = FALSE,
  spatial = TRUE
)
```

Arguments

df_db Either data frame or a connection to database with naturecounts table.

record Character. Name of the column containing record identification.

coord_lon Character. Name of the column containing longitude.

coord_lat Character. Name of the column containing latitude.

species Character. Name of the column containing species identification.

iao_grid_size_km

Numeric. Size of grid (km) to use when calculating IAO. Default is COSEWIC

requirement (2). Use caution if changing.

eoo_p Numeric. The percentile to calculate the convex hull over. Defaults to 0.95 for

a 95% convex hull to ensure outlier points do not artificially inflate the EOO. Note that for a final COSEWIC report, this may not be appropriate. Set to 1 to

include all points.

filter_unique Logical. Whether to filter observations to unique locations. Use this only if

there are too many data points to work with. This changes the nature of what an observation is, and also may bias EOO calculations if using less than 100% of

points (see eoo_p).

spatial Logical. Whether to return sf spatial objects showing calculations. If FALSE (the

default) returns a data frame with the IAO and EOO values. If TRUE returns a

list of objects with both the values and the spatial grid/polygons.

Details

Note that the while the IUCN calls this metric AOO, in COSEWIC, AOO is actually a different measure, the *biological* area of occupancy. See the Distribution section in 'Instructions for preparing COSEWIC status reports' for more details.

By default the EOO is calculated only using the inner 95% of points (based on distance to the centroid). This is to ensure that a first-pass of the EOO does not reject a species from consideration if there are any outlier observations. However, for a final COSEWIC assessment report, it is likely better to carefully explore the data to ensure there are no outliers and then use the full data set (i.e. set $eoo_p = 1$).

6 cosewic_ranges

The IAO is calculated by first assessing large grids (10x large than the specified size). Only then are smaller grids created within large grid cells containing observations. This speeds up the process by avoiding the creation of grids in areas where there are no observations. This means that the plots and spatial objects may not have grids over large areas lacking observations. See examples.

Details on how IAO and EOO are calculated and used

- COSEWIC Guidelines for use of the Index of Area of Occupancy in COSEWIC Assessments
- COSEWIC Instructions for preparing COSEWIC status reports
- COSEWIC Table 2 COSEWIC quantitative criteria and guidelines for the status assessment of Wildlife Species

Value

Summarized data frame (ranges) or list containing ranges, a summarized data frame, and spatial, a list of two spatial data frames.

ranges contains columns

- n_records_total Total number of records used to create ranges
- min_record Minimum number of records within IAO cells
- max_record Maximum number of records within IAO cells
- median_record Median number of records within IAO cells
- grid_size_km IAO cell size (area is this squared)
- n_occupied Number of IAO cells with at least one record
- iao IAO value (grid_size_km^2 * n_occupied)
- eoo_pXX EOO area calculated with a convex hull at percentile eoo_p (e.g., 95%)

spatial contains spatial data frames

- iao_sf Polygons of the IAO grids with the n_records per cell
- eoo_sf Polygon of the Convex Hull at percentile eoo_p

```
# Using the included, test data on black-capped chickadees
bcch # look at the data

r <- cosewic_ranges(bcch)
r <- cosewic_ranges(bcch, spatial = FALSE)

# Calculate for multiple species
mult <- rbind(bcch, hofi)
r <- cosewic_ranges(mult)
r <- cosewic_ranges(mult, spatial = FALSE)</pre>
```

format_dates 7

format_dates

Add date and day-of-year field/columns to data

Description

Creates and adds columns date and doy (day-of-year) to the data source (either data frame or database table naturecounts).

Usage

```
format_dates(df_db, overwrite = FALSE)
```

Arguments

df_db Either data frame or a connection to database with naturecounts table. Must

have fields/columns of survey_year, survey_month, survey_day

overwrite Logical. Overwrite existing columns date and/or doy?

Value

If df_dbwas a data frame, return a data frame with new columns date and doy. Otherwise return database connection.

Examples

```
bcch_with_dates <- format_dates(bcch)</pre>
```

format_zero_fill

Zero-fill data

Description

Zero-fill the species presence data by adding zero observation counts (absences) data to an existing nature counts dataset.

Usage

```
format_zero_fill(
  df_db,
  by = "SamplingEventIdentifier",
  species = "all",
  fill = "ObservationCount",
  extra_species = NULL,
  extra_event = NULL,
  warn = TRUE,
  verbose = TRUE
)
```

8 format_zero_fill

Arguments

df_db	Either data frame or a connection to database with naturecounts table (a data frame is returned).
by	Character vector. By default, "SamplingEventIdentifier" or a vector of specific column names to fill by (see details)
species	Character vector. Either "all", for species in the data, or a vector of species ID codes to fill in.
fill	Character. The column name to fill in. Defaults to "ObservationCount".
extra_species	Character vector. Extra columns/fields uniquely associated with species_id to keep in the data (all columns not in by, species, fill, extra_species, or extra_event will be omitted from the result).
extra_event	Character vector. Extra columns/fields uniquely associated with the Sampling Event (the field defined by by) to keep in the data (all columns not in by, species, fill, extra_species, or extra_event) will be omitted from the result).
warn	Logical. If TRUE, stop zero-filling if >100 species and >1000 unique sampling events. If FALSE, ignore and proceed.
verbose	Logical. Show messages?

Details

by refers to the combination of columns which are used to detect missing values. By default SamplingEventIdentifier is used. Otherwise users can specify their own combination of columns.

If species is supplied, all records will be used to determine observation events, but only records (zero-filled or otherwise) which correspond to a species in species will be returned (all others will be omitted). Note that records where species_id is NA (generally for 0 counts for presence/absence), will be converted to a list of 0's for the individual species.

Value

Data frame

grid_canada 9

grid_canada

Create grid across Canada

Description

Create grid across Canada

Usage

```
grid_canada(cell_size = 200, buffer = 500)
```

Arguments

cell_size

Numeric. Size of grid (km) to use when creating grid. If using this grid as input

to cosewic_ranges(), should use default COSEWIC grid size of 2.

buffer

Numeric. Extra buffer (km) to add around the outline of Canada before calcu-

lating grid.

Value

sf data frame with polygon grid

```
gc <- grid_canada(200)
gc_buff <- grid_canada(200, buffer = 0)
# Plot to illustrate</pre>
```

map_canada

```
library(ggplot2)
ggplot() +
  geom_sf(data = map_canada()) +
  geom_sf(data = gc, fill = NA) +
  labs(caption = "200km buffer")

ggplot() +
  geom_sf(data = map_canada()) +
  geom_sf(data = gc_buff, fill = NA) +
  labs(caption = "No buffer")
```

hofi

Example house finch data

Description

An example of house finch data downloaded from NatureCounts

Usage

hofi

Format

A data frame with 19 rows and 57 variables:

map_canada

Map of Canada

Description

Wrapper around rnaturalearth::ne_countries() to creates a simple features basic map of Canada with CRS 3347 (Statistics Canada Lambert).

Usage

```
map_canada()
```

Value

Sf data frame

meta 11

Examples

```
map_canada()
plot(map_canada())
library(ggplot2)
ggplot(data = map_canada()) + geom_sf()
```

meta

Metadata

Description

These functions return metadata codes, names, descriptions, and information associated with the data downloaded from NatureCounts.

Usage

```
meta_country_codes()
meta_statprov_codes()
meta_subnational2_codes()
meta_iba_codes()
meta_bcr_codes()
meta_bcr_codes()
meta_utm_squares()
meta_species_authority()
meta_species_codes()
meta_species_taxonomy()
meta_collections()
meta_breeding_codes()
meta_project_protocols()
meta_projects()
meta_protocol_types()
```

12 meta

```
meta_bmde_versions()
meta_bmde_fields(version = "minimum")
```

Arguments

version

Character. BMDE version for which to return fields. NULL returns all versions

Details

Some of these metadata are stored locally and can be updated with the nc_metadata() function. Others are downloaded as requested.

Value

Data frame

Functions

- meta_country_codes(): Country codes
- meta_statprov_codes(): State/Province codes
- meta_subnational2_codes(): Subnational2 codes
- meta_iba_codes(): Important Bird Area (IBA) codes
- meta_bcr_codes(): Bird Conservation Region (BCR) codes
- meta_utm_squares(): UTM Square codes
- meta_species_authority(): Species taxonomic authorities
- meta_species_codes(): Alpha-numeric codes for avian species
- meta_species_taxonomy(): Codes and taxonomic information for all species
- meta_collections(): Collections names and descriptions
- meta_breeding_codes(): Breeding codes and descriptions
- meta_project_protocols(): Project protocols
- meta_projects(): Projects ids, names, websites, and descriptions
- meta_protocol_types(): Protocol types and descriptions
- meta_bmde_versions(): Names and descriptions of the available versions of BMDE (Bird Monitoring Data Exchange). These refer to sets of fields/columns which can be downloaded for a given group of data. See nc_data_dl() for more details.
- meta_bmde_fields(): Fields/columns associated with a particular BMDE (Bird Monitoring Data Exchange) version. See meta_bmde_versions() for the different versions available, meta_collections() for which version is used by which project, and nc_data_dl() for more details on downloading data with a given set of fields/columns.

nc_count 13

Examples

```
# Return fields/columns in the 'minimum' version
meta_bmde_fields()

# Retrun fields/columns in the 'core' version
meta_bmde_fields(version = "core")

# Return all possible fields
meta_bmde_fields(version = "extended")
```

nc_count

Download information about NatureCounts collections

Description

Download the number of records available for different collections filtered by location (if provided). If authorization is provided, the collections are filtered to only those available to the user (unless using show = "all"). Without authorization all collections are returned.

Usage

```
nc_count(
  collections = NULL,
  project_ids = NULL,
  species = NULL,
  doy = NULL,
  region = NULL,
  site_type = NULL,
  show = "available",
  username = NULL,
  timeout = 120,
  verbose = TRUE
)
```

Arguments

collections	Character vector. The collection codes from which to download data. NULL (default) downloads data from all available collections
project_ids	Character/Numeric vector. The project ids from which to download data. First the collections associated with a project_id are determined, and then data is downloaded for each collection. If both collections and project_ids are supplied, they are combined.
species	Numeric vector. Numeric species ids (see details)
years	Numeric vector. The start/end years of data to download. Can use NA for either start or end, or a single value to return data from a single year.

14 nc_count

Character/Numeric vector. The start/end day-of-year to download (1-366 or doy dates that can be converted to day of year). Can use NA for either start or end region List. Named list with *one* of the following options: country, statprov, subnational2, iba, bcr, utm_squares, bbox. See details Character vector. The type of site to return (e.g., IBA). site_type Character. Either "all" or "available". "all" returns counts from all data sources. show "available" only returns counts for data available for the username provided. If no username is provided, defaults to "all". Character vector. Username for http://naturecounts.ca. If provided, the username

user will be prompted for a password. If left NULL, only public collections will

be returned.

timeout Numeric. Number of seconds before connecting to the server times out.

verbose Logical. Show messages?

Details

The akn_level column describes the level of data access for that collection (see descriptions online). The access column describes the accessibility of a collection for a given username (or no access if no username supplied). See the section on Access and request_ids for more details.

Value

Data frame

NatureCounts account

All public data is available with a username/password (sign up for a free NatureCounts account). However, to access private/semi-public projects/collections you must request access. See the Access and request_ids section for more information.

Species ids (species)

Numeric species id codes can determined from the functions search_species() or search_species_code(). See also the article on species codes for more information.

Day of Year (doy)

The format for day of year (doy) is fairly flexible and can be a whole number between 1 and 366 or anything recognized by lubridate-package's ymd() function. However, it must have the order of year, month, day. Note that year is ignored when converting to day of year, except that it will result in a 1 day offset for leap years.

Regions (region)

Regions are defined by codes reflecting the country, state/province, subnational (level 2), Important Bird Areas (IBA), and Bird Conservation Regions (BCR) (see search_region() for codes). They can also be defined by providing specific UTM squares to download or a bounding box area which specifies the min/max longitude and min/max latitude (bbox). See the article on regional filters for more information.

nc_count 15

Access and request_ids

Access to a data collection is either available as "full" or "by request". Use nc_count(username = "USER", show = "all"), to see the accessibility of collections.

"Full" access means that data can be immediately requested directly through the naturecounts R package. "By request" means that a request must be submitted online and approved before the data can be downloaded through naturecounts.

This means that there are two types of data requests: ones made through this naturecounts R package (API requests) and those made through the online Web Request Form (Web requests). Every request (from either method) generates a request_id which identifies the filter set and collections requested. Details of all of requests can be reviewed with the nc_requests() function.

To download data with "full" access, users can either specify filters, or if they are repeating a download, can use the request_id from nc_requests().

Otherwise, if the user doesn't have "full" access, they must supply an approved request_id to the nc_data_dl() function (e.g., nc_data_dl(request_id = 152000, username = "USER")). Use nc_requests() to see request_ids, filters, and approval status.

Requests for "full" access to additional collections can be made online through the Web Request Form by checking the "Full access?" box in Step 2 of the form.

See Also

```
nc_requests()
```

```
# Count all publicly available records:

nc_count()

# Count publicly available records for Manitoba, Canada

nc_count(region = list(statprov = "MB"))

# Count all records for all collections user "sample" has access to

## Not run:

nc_count(username = "sample")

## End(Not run)

# Count records with house finches in Ontario
search_species("house finch")

nc_count(species = 20350, region = list(statprov = "ON"), username = "sample")

# Count all records available in the Christmas Bird Count and Breeding Bird

# Survey collections (regardless of user permissions)

nc_count(collections = c("CBC", "BBS"), show = "all", username = "sample")
```

nc_data_dl

nc_data_dl

Download NatureCounts data records

Description

Download data records from various collections filtered by various options. In order to ease the load on the server, note that only **three** of collections/project_ids, species, years, doy, region, and site_type can be used in any one request. See the vignette for filtering your data after download for more options: vignette("filtering_data", package = "naturecounts").

Usage

```
nc_data_dl(
  collections = NULL,
  project_ids = NULL,
  species = NULL,
  years = NULL,
  doy = NULL,
  region = NULL,
  site_type = NULL,
  fields_set = "minimum",
  fields = NULL,
  username,
  info = NULL,
  request_id = NULL,
  sql_db = NULL,
  warn = TRUE,
  timeout = 120,
  verbose = TRUE
)
```

Arguments

collections	Character vector. The collection codes from which to download data. NULL (default) downloads data from all available collections
project_ids	Character/Numeric vector. The project ids from which to download data. First the collections associated with a project_id are determined, and then data is downloaded for each collection. If both collections and project_ids are supplied, they are combined.
species	Numeric vector. Numeric species ids (see details)
years	Numeric vector. The start/end years of data to download. Can use NA for either start or end, or a single value to return data from a single year.
doy	Character/Numeric vector. The start/end day-of-year to download (1-366 or dates that can be converted to day of year). Can use NA for either start or end
region	List. Named list with <i>one</i> of the following options: country, statprov, subnational2, iba, bcr, utm_squares, bbox. See details

nc_data_dl 17

site_type Character vector. The type of site to return (e.g., IBA). fields_set Character. Set of fields/columns to download. See details. fields Character vector. If fields set = custom, which fields/columns to download. See details username Character vector. Username for http://naturecounts.ca. If provided, the user will be prompted for a password. If left NULL, only public collections will be returned. info Character vector. Short description of reason for the download. E.g., "COSEWIC report", "Impact Assessment Study", "School project", etc. This kind of information helps NatureCounts.ca justify the utility of the database. Required unless resuming/re-downloaded with a request_id. Numeric. Specific request id to check or download. request_id Character vector. Name and location of SQLite database to either create or add sql_db Logical. Interactive warning if request more than 1,000,000 records to downwarn load. timeout Numeric. Number of seconds before connecting to the server times out.

Value

Data frame or connection to SQLite database

Logical. Show messages?

NatureCounts account

verbose

All public data is available with a username/password (sign up for a free NatureCounts account). However, to access private/semi-public projects/collections you must request access. See the Access and request_ids section for more information.

Species ids (species)

Numeric species id codes can determined from the functions search_species() or search_species_code(). See also the article on species codes for more information.

Day of Year (doy)

The format for day of year (doy) is fairly flexible and can be a whole number between 1 and 366 or anything recognized by lubridate-package's ymd() function. However, it must have the order of year, month, day. Note that year is ignored when converting to day of year, except that it will result in a 1 day offset for leap years.

Regions (region)

Regions are defined by codes reflecting the country, state/province, subnational (level 2), Important Bird Areas (IBA), and Bird Conservation Regions (BCR) (see search_region() for codes). They can also be defined by providing specific UTM squares to download or a bounding box area which specifies the min/max longitude and min/max latitude (bbox). See the article on regional filters for more information.

18 nc_data_dl

Data Fields/Columns (fields_set and fields)

By default data is downloaded with the minimum set of fields/columns. However, for more advanced applications, users may wish to specify which fields/columns to return. The Bird Monitoring Data Exchange (BMDE) schema keeps track of variables used to augment observation data. There are different versions reflecting different collections of variables which can be specified for download in one of four ways:

- fields_set can be a specific shorthand reflecting a BMDE version: core, extended or minimum (default). See meta_bmde_versions() to see which BMDE version the shorthand refers to.
- 2. fields_set can be default which uses the default BMDE version for a particular collection (note that if you download more than one collection, the field sets will expand to cover all fields/columns in the combined collections)
- 3. fields_set can be the exact BMDE version. See meta_bmde_versions() for options.
- 4. fields_set can be custom and the fields argument can be a character vector specifying the exact fields/columns to return. See meta_bmde_fields()) for potential fields values.

Note that in all cases there are a set of fields/columns that are *always* returned, no matter what fields_set is used.

Access and request_ids

Access to a data collection is either available as "full" or "by request". Use nc_count(username = "USER", show = "all"), to see the accessibility of collections.

"Full" access means that data can be immediately requested directly through the naturecounts R package. "By request" means that a request must be submitted online and approved before the data can be downloaded through naturecounts.

This means that there are two types of data requests: ones made through this naturecounts R package (API requests) and those made through the online Web Request Form (Web requests). Every request (from either method) generates a request_id which identifies the filter set and collections requested. Details of all of requests can be reviewed with the nc_requests() function.

To download data with "full" access, users can either specify filters, or if they are repeating a download, can use the request_id from nc_requests().

Otherwise, if the user doesn't have "full" access, they must supply an approved request_id to the nc_data_dl() function (e.g., nc_data_dl(request_id = 152000, username = "USER")). Use nc_requests() to see request_ids, filters, and approval status.

Requests for "full" access to additional collections can be made online through the Web Request Form by checking the "Full access?" box in Step 2 of the form.

nc_metadata 19

```
info = "nc_example")
# Black-capped Chickadees (BCCH) in SAMPLE2 collection in 2013
search_species("black-capped chickadee") # Find the species_id
bcch <- nc_data_dl(collection = "SAMPLE2", species = 14280, year = 2013,</pre>
                   username = "sample", info = "nc_example")
# All BCCH observations since 2015 accessible to user "sample"
bcch <- nc_data_dl(species = 14280, years = c(2015, NA), username = "sample",</pre>
                    info = "nc_example")
# All BCCH observations from mid-July to late October in all years for user "sample"
bcch <- nc_data_dl(species = 14280, doy = c(200, 300), username = "sample",</pre>
                     info = "nc_example")
# All BCCH observations from a specific bounding box for user "sample"
bcch <- nc_data_dl(species = 14280, username = "sample",</pre>
                   region = list(bbox = c(left = -100, bottom = 45,
                                            right = -80, top = 60)),
                    info = "nc_example")
# All American Bittern observations from user "sample"
search_species("american bittern")
bittern <- nc_data_dl(species = 2490, username = "sample", info = "nc_example")</pre>
# Different fields/columns
bittern <- nc_data_dl(species = 2490, fields_set = "core",
                      username = "sample", info = "nc_example")
bittern <- nc_data_dl(species = 2490, fields_set = "custom",</pre>
                      fields = c("Locality", "AllSpeciesReported"),
                      username = "sample", info = "nc_example")
## Not run:
# All collections by request id
# Specific collection by request id
my_data <- nc_data_dl(collections = "ABATLAS1",</pre>
                       request_id = 000000, username = "USER",
                       info = "MY REASON")
## End(Not run)
```

nc_metadata

Update NatureCounts metadata files

Description

Updates the local copies of meta data used by the package.

20 nc_permissions

Usage

```
nc_metadata(force = FALSE, utm = FALSE, verbose = TRUE)
```

Arguments

force Logical. Force update even if the remote version matches local?

utm Logical. Update meta_utm_squares() as well? WARNING: This is a large

and time consuming download!

verbose Logical. Show progress messages?

Examples

nc_metadata()

nc_permissions

Download list of accessible collections

Description

Returns a list of collections accessible by 'username'.

Usage

```
nc_permissions(username = NULL, timeout = 60)
```

Arguments

username Character vector. Username for http://naturecounts.ca. If provided, the

user will be prompted for a password. If left NULL, only public collections will

be returned.

timeout Numeric. Number of seconds before connecting to the server times out.

NatureCounts account

All public data is available with a username/password (sign up for a free NatureCounts account). However, to access private/semi-public projects/collections you must request access. See the Access and request_ids section for more information.

nc_query_table 21

Access and request_ids

Access to a data collection is either available as "full" or "by request". Use nc_count(username = "USER", show = "all"), to see the accessibility of collections.

"Full" access means that data can be immediately requested directly through the naturecounts R package. "By request" means that a request must be submitted online and approved before the data can be downloaded through naturecounts.

This means that there are two types of data requests: ones made through this naturecounts R package (API requests) and those made through the online Web Request Form (Web requests). Every request (from either method) generates a request_id which identifies the filter set and collections requested. Details of all of requests can be reviewed with the nc_requests() function.

To download data with "full" access, users can either specify filters, or if they are repeating a download, can use the request_id from nc_requests().

Otherwise, if the user doesn't have "full" access, they must supply an approved request_id to the nc_data_dl() function (e.g., nc_data_dl(request_id = 152000, username = "USER")). Use nc_requests() to see request_ids, filters, and approval status.

Requests for "full" access to additional collections can be made online through the Web Request Form by checking the "Full access?" box in Step 2 of the form.

Examples

```
nc_permissions()
nc_permissions(username = "sample")
```

nc_query_table

Custom table queries

Description

Generate custom table queries with the table name and filter arguments.

Usage

```
nc_query_table(
  table = NULL,
    ...,
  username = NULL,
  timeout = 120,
  verbose = FALSE
)
```

22 nc_query_table

Arguments

table Character. Table to query (see details)

... Name/value pairs for custom queries/filters (see details)

username Character vector. Username for http://naturecounts.ca. If provided, the

user will be prompted for a password. If left NULL, only public collections will

be returned.

timeout Numeric. Number of seconds before connecting to the server times out.

verbose Logical. Show messages?

Details

```
nc_query_table(username = "sample") for available options
```

Value

data.frame()

```
# What tables are available? What 'filters' do they take? Are any 'required'?
nc_query_table(username = "sample")
# Query the bmdefilter_bad_dates table
d <- nc_query_table(table = "bmde_filter_bad_dates", username = "sample")</pre>
head(d)
# Filter our query
d <- nc_query_table(table = "bmde_filter_bad_dates",</pre>
                     SiteCode = "DMBO", username = "sample")
# Filter our query
d <- nc_query_table(table = "bmde_filter_bad_dates",</pre>
                     species_id = 15770, username = "sample")
# Want more than one species? Either filter after, or combine two queries
# Filter after
library(dplyr)
d <- nc_query_table(table = "bmde_filter_bad_dates", username = "sample")</pre>
d <- filter(d, species_id %in% c(15770, 9750))</pre>
# Combine two queries
d1 <- nc_query_table(table = "bmde_filter_bad_dates",</pre>
                      species_id = 15770, username = "sample")
d2 <- nc_query_table(table = "bmde_filter_bad_dates",</pre>
                      species_id = 9750, username = "sample")
d <- rbind(d1, d2)</pre>
```

nc_remove_cache 23

nc_remove_cache

Remove in-memory cache

Description

All server queries are cached for four hours to reduce server load. You can reset the cache at any time by either restarting your R session or running nc_remove_cache().

Usage

```
nc_remove_cache()
```

Value

TRUE if it worked

Examples

nc_remove_cache()

nc_requests

Check on status of data requests

Description

List pending or completed data requests for a given user.

Usage

```
nc_requests(request_id = NULL, type = "web", username)
```

Arguments

request_id Numeric. Specific request id to check or download.

type Character One of "web", "api", or "all" specifying which types of request to

return (defaults to "web").

username Character vector. Username for http://naturecounts.ca. If provided, the

user will be prompted for a password. If left NULL, only public collections will

be returned.

Value

data frame

24 search_region

Access and request_ids

Access to a data collection is either available as "full" or "by request". Use nc_count(username = "USER", show = "all"), to see the accessibility of collections.

"Full" access means that data can be immediately requested directly through the naturecounts R package. "By request" means that a request must be submitted online and approved before the data can be downloaded through naturecounts.

This means that there are two types of data requests: ones made through this naturecounts R package (API requests) and those made through the online Web Request Form (Web requests). Every request (from either method) generates a request_id which identifies the filter set and collections requested. Details of all of requests can be reviewed with the nc_requests() function.

To download data with "full" access, users can either specify filters, or if they are repeating a download, can use the request_id from nc_requests().

Otherwise, if the user doesn't have "full" access, they must supply an approved request_id to the nc_data_dl() function (e.g., nc_data_dl(request_id = 152000, username = "USER")). Use nc_requests() to see request_ids, filters, and approval status.

Requests for "full" access to additional collections can be made online through the Web Request Form by checking the "Full access?" box in Step 2 of the form.

Examples

```
nc_requests(username = "sample")
nc_requests(request_id = 152446, username = "sample")
```

search_region

Find country, state/province, subnational2, IBA, or BCR codes

Description

Search for the correct codes to identify countries, states/provinces, subnational2 areas, Important Bird Areas (IBA), or Bird Conservation Regions (BCR). These are then used in the nc_data_dl() and nc_count() functions.

Usage

```
search_region(name = NULL, type = "country")
```

Arguments

name Character. The location name to search for

type Character. One of "country", "statprov", "subnational2", "iba", or "bcr". The

type of information to return.

Details

```
region_search() is deprecated in favour of search_region()
```

search_species 25

Value

A data frame with the relevant codes and other information

Examples

```
search_region("Mexico", type = "country") # MX
search_region("Yucatan", type = "statprov") # Yucatán
search_region("Alberta", type = "statprov") # AB
search_region("Edmonton", type = "subnational2") # CA.AB.11
search_region("Brandon", type = "subnational2") # CA.MB.07
search_region("hays reservoir", type = "iba") # AB075
search_region("rainforest", type = "bcr") # 5

# Show all codes
search_region(type = "country")
search_region(type = "statprov")
search_region(type = "subnational2")
search_region(type = "iba")
search_region(type = "bcr")

# Using the codes
nc_count(region = list(statprov = "AB"), years = 2010)
```

search_species

Find species codes

Description

Find species id codes by searching for scientific, English and French species names.

Usage

```
search_species(name = NULL, show = "names", authority = NULL)
```

Arguments

name	Character. The species name to search for	
show	Character. Either "all" or "names" (default). Whether to return all taxonomic information or only a subset with species names	
authority	Character. If not NULL (default), return the alphanumeric code associated with avian species for this taxonomic authority.	

26 search_species_code

Details

```
species_search() is deprecated in favour of search_species()
```

Value

Data frame of species ids and taxonomic information

Examples

```
# Show all ids
search_species()
search_species("chickadee")
search_species("black-capped chickadee")

# Add alphanumeric code for BSCDATA authority
search_species("black-capped chickadee", authority = "BSCDATA")

# Show all taxonomic information
search_species("black-capped chickadee", show = "all")

# Using the codes
nc_count(species = 14280)
```

search_species_code

Search for bird species id codes by alphanumeric codes

Description

This is an advanced function for returning all Bird-related species id codes based on the various alphanumeric codes used by different authorities.

Usage

```
search_species_code(code = NULL, authority = "BSCDATA", results = "all")
```

Arguments

code Vector. Character or numeric code indicating a species for a given authority.

Character. The authority to compare codes against (defaults to "BSCDATA")

results Character. "all" returns codes for all related species (including subspecies and

main species). "exact" returns only the code for exact species indicated by the

code.

Details

```
species_code_search() is deprecated in favour of search_species_code()
Species ids returned reflect both species and sub-species levels.
```

search_species_code 27

Value

A data frame of numeric species id codes and names

```
# Show all ids
search_species_code()
# Get all species ids for house finches
search_species_code("HOFI")
# Get all species ids for Dark-eyed Juncos
search_species_code("DEJU")
# Get all species ids related to Yellow-rumped Warbler (Myrtle)
# NOTE! This includes Audubon's and the main, Yellow-rumped Warbler species
search_species_code("MYWA")
# Get ONLY specific id related to Yellow-rumped Warbler (Myrtle)
search_species_code("MYWA", results = "exact")
# Use the Christmas Bird Count authority
search_species_code(11609, authority = "CBC")
# Look in more than one authority (note that the code only needs to match on
# of the authorities)
search_species_code("MYWA", authority = c("BCMA", "CBC"))
```

Index

* datasets	nc_count, 13, 24
bcch, 3	nc_data_dl, 16, 24
hofi, 10	nc_data_dl(), <i>12</i> , <i>15</i> , <i>18</i> , <i>21</i> , <i>24</i>
	nc_metadata, 19
bcch, 3	nc_metadata(), 12
	nc_permissions, 20
cosewic_plot, 3	nc_query_table, 21
cosewic_ranges, 4	nc_remove_cache, 23
Connect datas 7	nc_requests, 23
format_dates, 7	nc_requests(), 15, 18, 21, 24
<pre>format_zero_fill, 7</pre>	
grid_canada,9	region_search(search_region), 24
	search_region, 24
hofi, 10	search_region(), <i>14</i> , <i>17</i>
man panada 10	search_species, 25
map_canada, 10	search_species(), <i>14</i> , <i>17</i>
meta, 11	search_species_code, 26
meta_bcr_codes (meta), 11	search_species_code(), 14, 17
meta_bmde_fields (meta), 11	species_code_search
meta_bmde_fields(), 18	(search_species_code), 26
meta_bmde_versions (meta), 11	species_search (search_species), 25
meta_bmde_versions(), 12, 18	, – , – , , , , , , , , , , , , , , , ,
meta_breeding_codes (meta), 11	ymd, <i>14</i> , <i>17</i>
meta_collections (meta), 11	
meta_collections(), 12	
meta_country_codes (meta), 11	
meta_iba_codes (meta), 11	
meta_project_protocols (meta), 11	
meta_projects (meta), 11	
meta_protocol_types (meta), 11	
meta_species_authority (meta), 11	
meta_species_codes (meta), 11	
meta_species_taxonomy (meta), 11	
meta_statprov_codes (meta), 11	
meta_subnational2_codes (meta), 11	
meta_utm_squares (meta), 11	
meta_utm_squares(), 20	
naturecounts (naturecounts-package), 2	
naturecounts-package, 2	