Package ‘neotoma’

January 5, 2019

Type Package

Title Access to the Neotoma Paleoecological Database Through R

Version 1.7.4

Date 2019-01-03

Encoding UTF-8

Author Simon J. Goring [aut, cre],
      Gavin L. Simpson [aut],
      Jeremiah P. Marsicek [ctb],
      Karthik Ram [aut],
      Luke Sosalla [ctb]

Maintainer Simon J. Goring <goring@wisc.edu>

Description Access paleoecological datasets from the Neotoma Paleoecological Database using the published API (<http://api.neotomadb.org/>). The functions in this package access various pre-built API functions and attempt to return the results from Neotoma in a usable format for researchers and the public.

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URL https://github.com/ropensci/neotoma

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

Depends R (>= 2.10)

Imports plyr, reshape2, jsonlite, httr, xml2, analogue, leaflet, dplyr

Suggests testthat, knitr

RoxygenNote 6.1.1

NeedsCompilation no

Repository CRAN

Date/Publication 2019-01-05 01:10:03 UTC
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**Description**

Extracts age information from objects and returns them in a useful format.

**Usage**

```r
ages(obj, ...)
```

## S3 method for class 'download'
```
ages(obj, ...)
```

## S3 method for class 'download_list'
```
ages(obj, ...)
```

**Arguments**

- **obj**
  - an R object from which counts are to be extracted.
- **...**
  - arguments passed to other methods.

**Details**

Methods are available for "download" and "download_list" objects.
bind

Value
Either a data frame of ages or a list of such objects.

Author(s)
Simon Goring

Examples
```r
## Not run:
ostracodes <- get_dataset(datasettype = 'ostracode')

ostro.dl <- get_download(ostracodes)
ostro.ages <- ages(ostro.dl)

## End(Not run)
```

bind Function to bind objects together into a longer object.

Description
From multiple download*s, dataset*s or sites, join them together into a single object.

Usage
```r
bind(x, ...)
```

Arguments

**x**
An object returned by one of the get_* commands for download, site or dataset.

**...**
other objects of the same class.

Details
To support further synthesis and analysis compile_download works to transform a list returned by get_download into a large data frame with columns for site and sample attributes and also with the associated assemblage data at each sample depth. This function also does the same for single sites.

Value
This command returns a larger list.

Author(s)
Simon J. Goring <simon.j.goring@gmail.com>
browse

References

Neotoma Project Website: http://www.neotomadb.org

Examples

## Not run:
# Search for sites with "Thuja" pollen that are older than 8kyr BP and
# that are on the west coast of North America:
t8kyr.poa <- get_dataset(taxonname="Thuja*",
  loc=c(-150, 20, -100, 60), ageyoung = 8000)
t8kyr.canis <- get_dataset(taxonname="Canis*",
  loc=c(-150, 20, -100, 60), ageyoung = 8000)

t8kyr.co_site <- bind(t8kyr.poa, t8kyr.canis)
plot(t8kyr.co_site)

###
# We want to look at four different dataset types across a forest-prairie
# boundary:
dataset_types <- c("ostracode surface sample",
  "water chemistry",
  "diatom surface sample",
  "pollen surface sample")

dataset_lists <- lapply(dataset_types,
  function(x) {
    get_dataset(datasettype=x,
      loc = c(-100,43,-92,48))
  })

# Using do.call here to make sure that I don't have to split the list out.
new_datasets <- do.call(bind, dataset_lists)

# And voila!
plot(new_datasets)

## End(Not run)

---

**browse**

Open a browser window to display a Neotoma dataset within the Neotoma Explorer

---

**Description**

Using a download or dataset object, open up a browser window in the users default browser. Passing a download_list or dataset_list will open Neotoma Explorer with the first object and return a warning.
Using a numeric value, download, download_list, dataset or dataset_list object, open up a browser window in the users default browser. Very large objects

Usage

browse(x)

Arguments

x A numeric value, download, download_list, dataset or dataset_list object.

Value

Returns a NULL value, opens a browser.

Author(s)

Simon J. Goring <simon.j.goring@gmail.com>

References

Neotoma Project Website: http://www.neotomadb.org API Reference: http://api.neotomadb.org/doc/resources/sites

Examples

```r
## Not run:
# Where are the XRF data?

xrf.data <- get_dataset(datasettype='X-ray fluorescence (XRF)')
browse(xrf.data)

## End(Not run)
```

---

**browse.dataset**  
Open a browser window to display a Neotoma dataset within the Neotoma Explorer

**Description**

Using a numeric value, download, download_list, dataset or dataset_list object, open up a browser window in the users default browser. Very large objects

Usage

```r
## S3 method for class 'dataset'
browse(x)
```
**browse.dataset_list**

**Arguments**

- **x**  
  A dataset object.

---

**Description**

Using a numeric value, `download`, `download_list`, `dataset` or `dataset_list` object, open up a browser window in the user's default browser. Very large objects.

**Usage**

```r
## S3 method for class 'dataset_list'
browse(x)
```

**Arguments**

- **x**  
  A `dataset_list` object.

---

**browse.default**

**Open a browser window to display a Neotoma dataset within the Neotoma Explorer**

---

**Description**

Using a numeric value, `download`, `download_list`, `dataset` or `dataset_list` object, open up a browser window in the user's default browser. Very large objects.

**Usage**

```r
## Default S3 method:
browse(x)
```

**Arguments**

- **x**  
  A numeric value with the dataset ID.
browse.download  

Open a browser window to display a Neotoma dataset within the Neotoma Explorer

Description

Using a numeric value, download, download_list, dataset or dataset_list object, open up a browser window in the users default browser. Very large objects

Usage

```r
## S3 method for class 'download'
browse(x)
```

Arguments

- `x`: A download object.

browse.download_list

Open a browser window to display a Neotoma dataset within the Neotoma Explorer

Description

Using a numeric value, download, download_list, dataset or dataset_list object, open up a browser window in the users default browser. Very large objects

Usage

```r
## S3 method for class 'download_list'
browse(x)
```

Arguments

- `x`: A download_list object.
compile_downloads

**Description**

Function to convert multiple downloads into a single large table.

From the assemblage data for multiple cores, return a single data.frame with columns for site metadata and assemblage data.

To support further synthesis and analysis compile_download works to transform a list returned by get_download into a large data frame with columns for site and sample attributes and also with the associated assemblage data at each sample depth. This function also does the same for single sites.

**Usage**

```r
compile_downloads(downloads)
```

**Arguments**

- `downloads` A download_list as returned by `get_download`, or multiple downloads joined in a list.

**Value**

This command returns a data frame.

**Author(s)**

Simon J. Goring <simon.j.goring@gmail.com>

**References**

Neotoma Project Website: http://www.neotomadb.org


API Reference: http://api.neotomadb.org/doc/resources/contacts
Examples

```r
## Not run:
# Search for sites with "Thuja" pollen that are older than 8kyr BP and
# that are on the west coast of North America:
t8kyr.datasets <- get_dataset(taxonname='Thuja*',
  loc=c(-150, 20, -100, 60),
  ageyoung = 8000)

# Returns 3 records (as of 04/04/2013), get dataset for the first record,
# Gold Lake Bog.
thuja.sites <- get_download(t8kyr.datasets)
gold.p25 <- compile_taxa(thuja.sites, 'P25')

all.gold <- compile_downloads(gold.p25)
pollen.sums <- rowSums(all.gold[,1:ncol(all.gold)], na.rm=TRUE)

plot(x = all.gold$age,
  y = all.gold$Cupressaceae.Taxaceae / pollen.sums,
  col = all.gold$site.name,
  pch = 19)
```

## End(Not run)

### compile_taxa

Function to convert assemblage taxa to standardized lists.

**Description**

From the assemblage data for the core return assemblage data with the assemblage taxa Currently implemented only for pollen data.

**Usage**

```r
compile_taxa(object, list.name, alt.table = NULL, cf = TRUE,
  type = TRUE)
```

**Arguments**

- `object`  A pollen object returned by `get_download`.
- `list.name`  The taxon compilation list, one of a set of lists from the literature (e.g., "P25", "WhitmoreFull"). More detail in section Details.
- `alt.table`  A user provided table formatted with at least two columns, one called 'taxon' and the other named as in `list.name`.
- `cf`  Should taxa listed as *cf*s (*e.g.*, *cf*. *Gilia*) be considered highly re-solved?
type Should taxa listed as types (*e.g.*, *Iva annua*-type) be considered highly resolved?

Details

The data object uses the smaller pollen subset. As this package develops we will add the capacity to summarize data output from the translation. Currently we can return only subsets that have been defined in the literature. These lists include:

- "P25" This list is derived from Gavin et al., (2003), and includes 25 pollen taxa.
- "W564" This list is derived from Williams and Shuman (2008).
- "WhitmoreFull" This is the full list associated with the Whitmore et al., (2005) North American Modern Pollen Database.
- "WhitmoreSmall" As above, but taxa for which both fully resolved and undifferentiated exist these taxa are summed.

Value

This command returns a list object with the same structure as the parent pollen object returned by `get_download`, or a matrix (or data frame) depending on whether object is one or the other. Any pollen taxon not included in the major taxa defined in the pollen gets returned as 'Other'.

Author(s)

Simon J. Goring <simon.j.goring@gmail.com>

References

Neotoma Project Website: http://www.neotomadb.org


API Reference: http://api.neotomadb.org/doc/resources/contacts

Examples

```r
## Not run:
# Search for sites with "Thuja" pollen that are older than 8kyr BP and
# that are on the west coast of North America:
t8kyr.datasets <- get_dataset(taxonname='Thuja*', loc=c(-150, 20, -100, 60), ageyoung = 8000)

# Returns 3 records (as of 04/04/2013), get dataset for the first record, Gold Lake Bog.
```
counts

GOLDKBG <- get_download(t8kyr.datasets[[1]])
gold.p25 <- compile_taxa(GOLDKBG, 'P25')

## End(Not run)

<table>
<thead>
<tr>
<th>counts</th>
<th>Access proxy count data</th>
</tr>
</thead>
</table>

**Description**

Extract pollen or other proxy counts from data objects and returns them in a useful format.

**Usage**

```r
counts(obj, ...)
```

## S3 method for class 'download'
```r
counts(obj, ...)
```

## S3 method for class 'download_list'
```r
counts(obj, ...)
```

**Arguments**

- `obj` an R object from which counts are to be extracted.
- `...` arguments passed to other methods.

**Details**

Methods are available for "download" and "download_list" objects.

**Value**

Either a data frame of counts or a list of such objects.

**Author(s)**

Gavin Simpson
**depths**

*Extracts the depth values from a ‘download’ object*

**Examples**

```r
## Not run:
marion <- get_site('Marion Lake')
louise <- get_site('Louise Pond')
western.sites <- rbind(marion, louise)
western.data <- get_dataset(western.sites)

western.dl <- get_download(western.data)
western.cnt <- counts(western.dl)
sapply(western.cnt, dim)
marion.cnt <- counts(western.dl[[1]])
dim(marion.cnt)

## End(Not run)
```

**Description**

Using a `download` object, return the sample depths (if available).

Using a numeric value, `download`, `download_list`, `dataset` or `dataset_list` object, open up a browser window in the users default browser. Very large objects

**Usage**

```r
depths(obj, ...)
```

**Arguments**

- **obj**  
  A download object.

- **...**  
  arguments passed to other methods.

**Value**

Retruns a vector of depths.
Author(s)
Simon J. Goring <simon.j.goring@gmail.com>

References
Neotoma Project Website: http://www.neotomadb.org API Reference: http://api.neotomadb.org/doc/resources/sites

Examples
```r
## Not run:
# Provide a vector of depths to generate a new age model:
# The dataset id 684 is for Devils Lake, a record published by Louis Maher Jr.

pollen.data <- get_download(684)
pollen.chron <- get_chroncontrol(pollen.data)[[1]]

age_sds <- pollen.chron$chron.control$age - focal$chron.control$age.young,
get_curves <- ifelse(regexpr("Radiocarbon",
    pollen.chron$chron.control$control.type) > -1,
    'intcal13', 'normal')

new_chron <- Bchron::Bchronology(ages = pollen.chron$chron.control$age,
    ageSds = age_sds
    positions = pollen.chron$chron.control$depth,
    calCurves = ,
    predictPositions = depths(pollen.data))
```

```
## End(Not run)
```

download

A class for download objects.

Description
A download is an object with the full record for a single dataset.
A download is an object with the full record for a single dataset.

Details
TO DO
TO DO

Author(s)
Simon Goring
Simon Goring
get_chroncontrol

Function to return chronological control tables used to build age models.

Description

Using the dataset ID, return all records associated with the data. At present, only returns the dataset in an unparsed format, not as a data table. This function will only download one dataset at a time.

Usage

get_chroncontrol(x, chronology = 1, verbose = TRUE, add = FALSE)

Arguments

x A single numeric chronology ID, a vector of numeric dataset IDs as returned by get_dataset or a download or download_list object.
chronology When download objects have more than associated chronology, which chronology do you want? Default is 1.
verbose logical, should messages on API call be printed?
add logical, should this chron control be added to the download object?

Value

This command returns either an object of class "try-error" containing the error returned from the Neotoma API call, or a full data object containing all the relevant information required to build either the default or prior chronology for a core. When download or download_list objects are passes, the user can add the chroncontrol to the download object explicitly, in which case the function will return a download with chroncontrol embedded.

This is a list comprising the following items:

chron.control A table describing the collection, including dataset information, PI data compatible with get_contact and site data compatible with get_site.
meta Dataset information for the core, primarily the age-depth model and chronology. In cases where multiple age models exist for a single record the most recent chronology is provided here.

If Neotoma returns empty content, either the control table or the associated metadata (which happens in approximately 25

Author(s)

Simon J. Goring <simon.j.goring@gmail.com>
get_chroncontrol.dataset

Function to return chronological control tables from a dataset.

Description

Using a dataset, return the default chron-control table.

Usage

```
## S3 method for class 'dataset'
get_chroncontrol(x, chronology = 1, verbose = TRUE, add = FALSE)
```

Arguments

- **x**: A dataset.
- **chronology**: When download objects have more than associated chronology, which chronology do you want? Default is 1.
- **verbose**: logical; should messages on API call be printed?
- **add**: Should the chroncontrol be added to the download object (only accepts FALSE)
get_chroncontrol.dataset_list

Function to return chronological control tables from a dataset_list.

Description
Using a dataset_list, return the default chron-control table.

Usage
```r
## S3 method for class 'dataset_list'
get_chroncontrol(x, chronology = 1,
                   verbose = TRUE, add = FALSE)
```

Arguments
- `x`: A dataset_list object.
- `chronology`: When download objects have more than associated chronology, which chronology do you want? Default is 1.
- `verbose`: logical; should messages on API call be printed?
- `add`: Should the chroncontrol be added to the download object (only accepts FALSE)

get_chroncontrol.default

Function to return chronological control tables from a chronologic ID.

Description
Using the chronology ID, return the chron control table as a data.frame.

Usage
```r
## Default S3 method:
get_chroncontrol(x, chronology = 1, verbose = TRUE,
                  add = FALSE)
```

Arguments
- `x`: A single numeric chronology ID or a vector of numeric chronology IDs as returned by get_datasets.
- `chronology`: For download methods, which chronology controls should be used?
- `verbose`: logical; should messages on API call be printed?
- `add`: logical, should this chron control be added to the download object?
get_chroncontrol.download

Function to return chronological control tables from a download object.

Description

Using a download, return the default chron-control table as a data.frame.

Usage

```r
## S3 method for class 'download'
get_chroncontrol(x, chronology = 1, verbose = TRUE, 
                  add = FALSE)
```

Arguments

- `x`: A single download object.
- `chronology`: For download methods, which chronology controls should be used?
- `verbose`: logical; should messages on API call be printed?
- `add`: Should the chroncontrol be added to the download object (default FALSE)

get_chroncontrol.download_list

Function to return chronological control tables from a download_list object.

Description

Using a download_list, return the default chron-control table as a data.frame.

Usage

```r
## S3 method for class 'download_list'
get_chroncontrol(x, chronology = 1, verbose = TRUE, 
                  add = FALSE)
```

Arguments

- `x`: A download_list object.
- `chronology`: When download objects have more than associated chronology, which chronology do you want? Default is 1.
- `verbose`: logical; should messages on API call be printed?
- `add`: Should the chroncontrol be added to the download object (default FALSE)
**get_closest**  

*Find the closest dataset records to a site, dataset or long/lat pair in Neotoma*

### Description

Passing in a download object the function outputs a Bacon or Clam formatted file to a user defined destination for age modelling with existing age-depth modeling software.

### Usage

```r
get_closest(x, n, buffer, ...)
```

### Arguments

- `x`  
  A vector long/lat pair, or a dataset, site or download.

- `n`  
  The number of records to return.

- `buffer`  
  The size of the buffer for dataset search (in kilometers)

- `...`  
  Optional arguments to pass into `get_dataset`.

### Value

This command returns a dataset or dataset_list, or NULL if no records exist within the bounding box.

### Author(s)

Simon J. Goring <simon.j.goring@gmail.com>, Andria Dawson <andria.dawson@gmail.com>

### References

Neotoma Project Website: http://www.neotomadb.org  
API Reference: http://api.neotomadb.org/doc/resources/contacts

### Examples

```r
## Not run:  
# The point of pulling chronology tables is to re-build or examine the chronological  
# information that was used to build the age-depth model for the core.  
# Find the closest records to Madison, WI:  
get_closest(x = c(-89.4012, 43.0731), n = 10, buffer = 5, datasettype = "pollen")

## End(Not run)
```
get_contact

Get contact information.

Description

A function to obtain contact information for data contributors from the Neotoma Paleoecological Database.

Usage

get_contact(contactid, contactname, contactstatus, familyname)

Arguments

- **contactid**: Contact ID is a numerical value associated with the Neotoma Contact table’s numerical Contact ID.
- **contactname**: A character string indicating the data contributors’ project, organization or personal name. May be a partial string and can include wildcards.
- **contactstatus**: The current status of the contact. Possible values include: active, deceased, defunct, extant, inactive, retired, unknown.
- **familyname**: A character string. Full or partial string indicating the contact’s last name.

Value

The function takes parameters defined by the user and returns a list of contact information supplied by the Neotoma Paleoecological Database. The user may define all or none of the possible fields. The function contains data checks for each defined parameter.

The function returns either a single item of class "try-error" describing the reason for failure (either mis-defined parameters or an error from the Neotoma API), or a table of contacts, with rows corresponding to the number of individual contacts returned by the Neotoma API. Each row entry includes the following parameters:

- **contact.name**: Full name of the person, last name first (e.g. "Simpson, George Gaylord") or name of organization or project (e.g. "Great Plains Flora Association").
- **contact.status**: Current status of the person, organization, or project. Field links to the Contact-Statuses lookup table.
- **family.name**: Family or surname name of a person.
- **leading.initials**: Leading initials for given or forenames without spaces (e.g. "G.G.").
- **given.names**: Given or forenames of a person (e.g. "George Gaylord"). Initials with spaces are used if full given names are not known (e.g. "G. G").
- **suffix**: Suffix of a person’s name (e.g. "Jr.", "III").
- **title**: A person’s title (e.g. "Dr.", "Prof.", "Prof. Dr").
**get_dataset**

get_dataset

- **phone**: Telephone number.
- **fax**: Fax number.
- **email**: Email address.
- **url**: Universal Resource Locator, an Internet World Wide Web address.
- **address**: Full mailing address.
- **notes**: Free form notes or comments about the person, organization, or project.
- **contact.id**: Unique database record identifier for the contact.
- **alias.id**: The ContactID of a person's current name. If the AliasID is different from the ContactID, the ContactID refers to the person's former name.

**Author(s)**

Simon J. Goring <simon.j.goring@gmail.com>

**References**


**Examples**

```r
# Not run:
# To find all data contributors who are active:
active.cont <- get_contact(contactstatus = 'active')

# To find all data contributors who have the last name "Smith"
smith.cont <- get_contact(familyname = 'Smith')

# End(Not run)
```

---

**get_dataset**

*Obtain dataset information from the Neotoma Paleoecological Database or an existing object.*

**Description**

A function to access the Neotoma API and return datasets corresponding to the parameters defined by the user.

**Usage**

```r
get_dataset(x, datasettype, piid, altmin, altmax, loc, gpid, taxonids, taxonname, ageold, ageyoung, ageof, subdate)
```
get_dataset

Arguments

x  An optional value, either a numeric site ID or object of class download, download_list or site.

datasettype  A character string corresponding to one of the allowed dataset types in the Neotoma Database. Allowed types include: "geochronologic", "loss-on-ignition", "pollen", "plant macrofossils", "vertebrate fauna", "mollusks", and "pollen surface sample".

piid  Numeric value for the Principle Investigator's ID number.

altmin  Numeric value indicating the minimum altitude for the site (can be used alone or with altmax).

altmax  Numeric value indicating the maximum altitude for the site (can be used alone or with altmin).

loc  A numeric vector c(lonW, latS, lonE, latN) representing the bounding box within which to search for sites. The convention here is to use negative values for longitudes west of Greenwich or longitudes south of the equator.

gpid  A character string or numeric value, must correspond to a valid geopolitical identity in the Neotoma Database. Use get.tables('GeoPoliticalUnits') for a list of acceptable values, or link here: http://api.neotomadb.org/apdx/geopol.htm

taxonids  A numeric identifier for the taxon. See get_table and use get_tables('Taxa') for a list of acceptable values.

taxonname  A character string corresponding to a valid taxon identity in the Neotoma Database. See get_table and use get_table('Taxa') for a list of acceptable values.

ageold  The oldest date acceptable for the search (in years before present).

ageyoung  The youngest date acceptable for the search.

ageof  If a taxon ID or taxon name is defined this parameter must be set to "taxon", otherwise it may refer to "sample", in which case the age bounds are for any samples within datasets or "dataset" if you want only datasets that are within the bounds of ageold and ageyoung.

subdate  Date of dataset submission, either YYYY-MM-DD or MM-DD-YYYY.

Value

More details on the use of these parameters can be obtained from http://api.neotomadb.org/doc/resources/datasets.

A list of class ‘dataset_list’, with each item corresponding to an individual record. Searches that return no items will result in a NULL value being returned. Otherwise each list item (each dataset record) includes the following components:

dataset.id  Unique database record identifier for the dataset.

dataset.name  Name of the dataset; not commonly used.

CollUnitHandle  Code name of the Collection Unit with which the dataset is associated. This code may be up to 10 characters. Data are frequently distributed by Collection Unit, and the Handle is used for file names.
CollUnitID  Unique database record identifier for the collection unit.
CollType   The collection type. Types include cores, sections, excavations, and animal middens.
DatasetType The dataset type, such as: geochronologic, loss-on-ignition, pollen, plant macrofossils, vertebrate fauna, etc.
AgeOldest The oldest of all sample ages (in calendar years before present) in the dataset.
AgeYoungest The youngest of all sample ages (in calendar years before present) in the dataset.
SubDates   An array of objects that describe dataset submission events. If multiple submissions occurred then this is a table.
DatasetPIs An array of objects that describe Principal Investigators associated with a dataset.
Site      An object describing the site where the dataset samples were taken.

Author(s)
Simon J. Goring <simon.j.goring@gmail.com>

References

Examples

```r
## Not run:
# Search for sites with "Thuja" pollen that are older than 8kyr BP and
# that are on the west coast of North America:
t8ktyr.datasets <- get_dataset(taxonname='Thuja',
  loc=c(-150, 20, -100, 60),
  ageyoung = 8000)

# Search for vertebrate fossils in Canada (gpid: 756) within the last 2kyr.
gpids <- get_table(table.name='GeoPoliticalUnits')
canID <- gpids[which(gpids$GeoPoliticalName == 'Canada'),1]
v2kyr.datasets <- get_dataset(datasettype='vertebrate fauna',
  gpid=canID,
  ageold = 2000)
```

Description

A function to access the Neotoma API and return datasets corresponding to the parameters defined by the user.
Usage

## Default 53 method:

```r
get_dataset(x, datasettype, piid, altmin, altmax, loc,
            gpid, taxonids, taxonname, ageold, ageyoung, ageof, subdate)
```

Arguments

- **x**: A numeric value corresponding to the site ID.
- **datasettype**: A character string corresponding to one of the allowed dataset types in the Neotoma Database. Allowed types include: "geochronologic", "loss-on-ignition", "pollen", "plant macrofossils", "vertebrate fauna", "mollusks", and "pollen surface sample".
- **piid**: Numeric value for the Principle Investigator’s ID number.
- **altmin**: Numeric value indicating the minimum altitude for the site (can be used alone or with altmax).
- **altmax**: Numeric value indicating the maximum altitude for the site (can be used alone or with altmin).
- **loc**: A numeric vector `c(lonW, latS, lonE, latN)` representing the bounding box within which to search for sites. The convention here is to use negative values for longitudes west of Greenwich or longitudes south of the equator.
- **gpid**: A character string or numeric value, must correspond to a valid geopolitical identity in the Neotoma Database. Use `get.tables('GeoPoliticalUnits')` for a list of acceptable values, or link here: [http://api.neotomadb.org/apdx/geopol.htm](http://api.neotomadb.org/apdx/geopol.htm)
- **taxonids**: A numeric identifier for the taxon. See `get_table` and use `get_tables('Taxa')` for a list of acceptable values.
- **taxonname**: A character string corresponding to a valid taxon identity in the Neotoma Database. See `get_table` and use `get_table('Taxa')` for a list of acceptable values.
- **ageold**: The oldest date acceptable for the search (in years before present).
- **ageyoung**: The youngest date acceptable for the search.
- **ageof**: If a taxon ID or taxon name is defined this parameter must be set to "taxon", otherwise it may refer to "sample", in which case the age bounds are for any samples within datasets or "dataset" if you want only datasets that are within the bounds of ageold and ageyoung.
- **subdate**: Date of dataset submission, either YYYY-MM-DD or MM-DD-YYYY.

---

**get_dataset.download**  
*Obtain dataset information from an existing download object.*

Description

A function to access a dataset within a download object.
Usage

```r
## S3 method for class 'download'
get_dataset(x, ...)
```

Arguments

- `x`: An object of class `download`.
- `...`: Objects passed from the generic. Not used in the call.

Description

Obtain dataset information from a `download_list`.

Usage

```r
## S3 method for class 'download_list'
get_dataset(x, ...)
```

Arguments

- `x`: An object of class `download_list`.
- `...`: Objects passed from the generic. Not used in the call.

Description

A function to return datasets corresponding to the objects within a `download_list`.

Usage

```r
## S3 method for class 'geochronologic'
get_dataset(x, ...)
```

Arguments

- `x`: An object of class `geochronologic`.
- `...`: Objects passed from the generic. Not used in the call.

Description

Obtain dataset information from an object of class `geochronologic`.

Usage

```r
## S3 method for class 'geochronologic'
get_dataset(x, ...)
```

Arguments

- `x`: An object of class `geochronologic`.
- `...`: Objects passed from the generic. Not used in the call.
get_dataset.geochronologic_list

Obtain dataset information from an object of class geochronologic_list.

Description
A function to access the Neotoma API and return datasets corresponding to the parameters defined by the user.

Usage
```r
## S3 method for class 'geochronologic_list'
get_dataset(x, ...)
```

Arguments
- `x` An object of class geochronologic_list.
- `...` objects passed from the generic. Not used in the call.

get_dataset.integer

Obtain dataset information from a vector of dataset IDs.

Description
A function to access the Neotoma API and return datasets corresponding to the parameters defined by the user.

Usage
```r
## S3 method for class 'integer'
get_dataset(x = NULL, ...)
```

Arguments
- `x` A single numeric dataset id, or a numeric vector.
- `...` objects passed from the generic. Not used in the call.
**get_dataset.numeric**  
*Obtain dataset information from a vector of dataset IDs.*

**Description**

A function to access the Neotoma API and return datasets corresponding to the parameters defined by the user.

**Usage**

```r
## S3 method for class 'numeric'
get_dataset(x = NULL, ...)
```

**Arguments**

- `x`  
  A single numeric dataset id, or a numeric vector.

- `...`  
  Objects passed from the generic. Not used in the call.

---

**get_dataset.site**  
*Obtain dataset information from an existing site object.*

**Description**

A function to access the Neotoma API and return datasets corresponding to the parameters defined by the user.

**Usage**

```r
## S3 method for class 'site'
get_dataset(x, ...)
```

**Arguments**

- `x`  
  An object of class site.

- `...`  
  Objects passed from the generic. Not used in the call.
get_download

Function to return full download records using sites, datasets, or dataset IDs.

Description

Using the dataset ID, site object or dataset object, return all records associated with the data as a download_list.

Usage

get_download(x, verbose = TRUE)

Arguments

x
A single numeric dataset ID or a vector of numeric dataset IDs as returned by get_datasets, or a site, dataset, or dataset_list.

verbose
logical; should messages on API call be printed?

Value

This command returns either object of class "try-error" (see try) defined by the error returned from the Neotoma API call, or an object of class download_list, containing a set of download objects, each with relevant assemblage information and metadata: The download object is a list of lists and data frames that describe an assemblage, the constituent taxa, the chronology, site and PIs who contributed the data. The following are important components:

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dataset</td>
<td>A table describing the collection, including dataset information, PI data compatible with get_contact and site data compatible with get_site.</td>
</tr>
<tr>
<td>sample.meta</td>
<td>Dataset information for the core, primarily the age-depth model and chronology. In cases where multiple age models exist for a single record the most recent chronology is provided here.</td>
</tr>
<tr>
<td>taxon.list</td>
<td>The list of taxa contained within the dataset, unordered, including information that can be used in get_taxa</td>
</tr>
<tr>
<td>counts</td>
<td>The assemblage data for the dataset, arranged with each successive depth in rows and the taxa as columns. All taxa are described in taxon.list, the chronology is in sample.data</td>
</tr>
<tr>
<td>lab.data</td>
<td>A data frame of laboratory data, such as exotic pollen spike, amount of sample counted, charcoal counts, etc.</td>
</tr>
<tr>
<td>chronologies</td>
<td>A list of existing chronologies. If only a single chronology exists for a record then this is the same as the age-model in sample.meta.</td>
</tr>
</tbody>
</table>
get_download

Note

The function returns a warning in cases where single taxa are defined by multiple taphonomic characteristics, for example grains that are identified separately as crumpled and torn in the same sample and sums these values within a sample. In the case that a geochronology dataset is passed to get_download the function returns a message and a NULL object (that is later excised). Use get_geochron for these objects. The chronologies can be augmented using the function get_chroncontrol, where the individual chronology objects in chronologies will consist of a table equivalent to sample.meta and a chroncontrol object.

Author(s)

Simon J. Goring <simon.j.goring@gmail.com>

References


Examples

```r
# Not run:
# Search for sites with "Pseudotsuga" pollen that are older than 8kyr BP and
# that are roughly within western British Columbia:
t8kyr.datasets <- get_dataset(taxonname='Picea*', loc=c(-90, 41, -89, 44),
ageold = 20000, ageyoung=10000)

# Returns 20 records (as of 04/04/2013), get the dataset for all records:
pollen.records <- get_download(t8kyr.datasets)

# Standardize the taxonomies for the different records using the WS64 taxonomy.
compiled.sites <- compile_taxa(pollen.records, list.name='WS64')

# Extract the Pseudotsuga curves for the sites:
generate.curve <- function(x, taxa) {
  if (taxa %in% colnames(x$counts)) {
    count <- x$counts[,taxa]/rowSums(x$counts, na.rm=TRUE)
  } else {
    count <- rep(0, nrow(x$count))
  }
data.frame(site = x$dataset$site.data$site.name, age = x$sample.meta$age,
             count = count)
}

curves <- do.call(rbind.data.frame, lapply(compiled.sites, generate.curve, taxa = 'Larix/Pseudotsuga'))

# For illustration, remove the sites with no Pseudotsuga occurrence:
curves <- curves[curves$count > 0, ]

smooth.curve <- predict(loess(sqrt(count)-age, data=curves),
                         data.frame(age=seq(20000, 0, by = -100)))
```

plot(sqrt(count) ~ age, data = curves,
    ylab = '% Pseudotsuga/Larix', xlab='Calibrated Years BP', pch=19,
    col=rgb(0.1, 0.1, 0.1), xlim=c(0, 20000))
lines(seq(20000, 0, by = -100), smooth.curve, lwd=2, lty=2, col=2)

# This figure shows us an apparent peak in Larix/Pseudotsuga pollen in the
# early-Holocene that lends support to a warmer, drier early-Holocene in
# western North America.

## End(Not run)

---

**get_download.dataset**  
*Function to return full download records using a dataset.*

**Description**  
Using a dataset, return all records associated with the data as a download_list.

**Usage**  
```r
## S3 method for class 'dataset'
get_download(x, verbose = TRUE)
```

**Arguments**  
- `x`: An object of class dataset.
- `verbose`: logical; should messages on API call be printed?

---

**get_download.dataset_list**  
*Function to return full download records using a dataset_list.*

**Description**  
Using a dataset_list, return all records associated with the data as a download_list.

**Usage**  
```r
## S3 method for class 'dataset_list'
get_download(x, verbose = TRUE)
```

**Arguments**  
- `x`: An object of class dataset_list.
- `verbose`: logical; should messages on API call be printed?
get_download.default  

Function to return full download records using numeric dataset IDs.

Description

Using the dataset ID, return all records associated with the data as a download_list.

Usage

```r
## Default S3 method:
get_download(x, verbose = TRUE)
```

Arguments

- `x` A single numeric dataset ID or a vector of numeric dataset IDs as returned by `get_datasets`.
- `verbose` logical; should messages on API call be printed?

get_download.site  

Function to return full download records using a site.

Description

Using a site, return all records associated with the data as a download_list.

Usage

```r
## S3 method for class 'site'
get_download(x, verbose = TRUE)
```

Arguments

- `x` An object of class site.
- `verbose` logical; should messages on API call be printed?
get_geochron

Function to return geochronological data from records.

Description

Using the dataset ID, return all geochronological data associated with the dataID. At present, only returns the dataset in an unparsed format, not as a data table. This function will only download one dataset at a time.

Usage

get_geochron(x, verbose = TRUE)

Arguments

x          A numeric dataset ID or a vector of numeric dataset IDs, or an object of class of class site, dataset, dataset_list, download or download_list for which geochrons are required.
verbose    logical; should messages on API call be printed?

Value

This command returns either an object of class "try-error" (see try) defined by the error returned from the Neotoma API call, or a geochronologic object, which is a list with two components, a dataset and a geochronology table, a data.frame with the following components:

- sample.id A unique identifier for the geochronological unit.
- age.type String. The age type, one of calendar years, radiocarbon years, etc.
- age Dated age of the material.
- e.older The older error limit of the age value. Commonly 1 standard deviation.
- e.young The younger error limit of the age value.
- delta13C The measured or assumed delta13C value for radiocarbon dates, if provided.
- material.dated A table describing the collection, including dataset information, PI data compatible with get_contact and site data compatible with get_site.
- geo.chron.type Text string, type of geochronological analysis, i.e., Radiocarbon dating, luminescence.
- notes Text string
- infinite Boolean, does the dated material return an "infinite" date?

A full data object containing all the relevant geochronological data available for a dataset.

Author(s)

Simon J. Goring <simon.j.goring@gmail.com>
get_publication

References


Examples

```r
## Not run:
# Search for the sites around Marion Lake, BC. I want to find sites within
# about 1km.
marion <- get_site(sitename = "Marion Lake")

marion_close <- get_closest(marion, n = 10, buffer = 1)

# Returns 116 records (as of 13/07/2015). These are the pollen records though,
# we want the sites:
geochron.records <- get_geochron(marion_close)

# We want to extract all the radiocarbon ages from the records:
get_ages <- function(x){
  any.ages <- try(x[[2]]$age[[2]]$age.type == 'Radiocarbon years BP'))
  if(class(any.ages) == 'try-error') output <- NA
  if(!class(any.ages) == 'try-error') output <- unlist(any.ages)
  output
}

radio.chron <- unlist(sapply(get_ages))

hist(radio.chron[radio.chron<40000], breaks=seq(0, 25000, by = 1000),
     main = 'Radiocarbon dates for Pseudotsuga records',
     xlab = 'Radiocarbon date (14C years before 1950)')

## End(Not run)
```

---

get_publication

A function to get publications for sites or datasets in the Neotoma Database using the API.

Description

The function takes the parameters, defined by the user, and returns a table with publication information from the Neotoma Paleoecological Database.

Usage

get_publication(x, contactid, datasetid, author, pubtype, year, search)
### Arguments

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>x</code></td>
<td>Numeric Publication ID value, either from <code>get_dataset</code> or known.</td>
</tr>
<tr>
<td><code>contactid</code></td>
<td>Numeric Contact ID value, either from <code>get_dataset</code> or <code>get_contact</code>.</td>
</tr>
<tr>
<td><code>datasetid</code></td>
<td>Numeric Dataset ID, known or from <code>get_dataset</code>.</td>
</tr>
<tr>
<td><code>author</code></td>
<td>Character string for full or partial author's name. Can include wildcards such as 'Smit*' for all names beginning with 'Smit'.</td>
</tr>
<tr>
<td><code>pubtype</code></td>
<td>Character string, one of eleven allowable types, see <code>get_table</code>. For a list of allowed types run <code>get_table(&quot;PublicationTypes&quot;)</code>.</td>
</tr>
<tr>
<td><code>year</code></td>
<td>Numeric publication year.</td>
</tr>
<tr>
<td><code>search</code></td>
<td>A character string to search for within the article citation.</td>
</tr>
</tbody>
</table>

### Value

A list is returned with two data frame components:

- `meta` A single row with Publication ID, type, year of publication and full citation.
- `Authors dataNframe` of author names, order and IDs, can be of variable length.

### Author(s)

Simon J. Goring <simon.j.goring@gmail.com>

### References


### Examples

```r
# Not run:
# To find all publications from 1998:
year.cont <- get_publication(year = 1998)

# To find all data contributors who have the last name "Smith"
smith.cont <- get_publication(author = 'Smith')
```

## Description

A function to get publications for datasets in the Neotoma Database using the API.

The function takes a dataset and returns a table with publication information from the Neotoma Paleoecological Database.
Usage

```r
## S3 method for class 'dataset'
get_publication(x, ...)
```

Arguments

- `x` an object of class dataset.
- `...` objects passed from the generic. Not used in the call.

---

**get_publication.dataset_list**

*A function to get publications for dataset_lists in the Neotoma Database using the API.*

---

Description

The function takes a `dataset_list` and returns a table with publication information from the Neotoma Paleoecological Database.

Usage

```r
## S3 method for class 'dataset_list'
get_publication(x, ...)
```

Arguments

- `x` an object of class `dataset_list`.
- `...` objects passed from the generic. Not used in the call.

---

**get_publication.default**

*A function to get publications for sites or datasets in the Neotoma Database using the API.*

---

Description

The function takes the parameters, defined by the user, and returns a table with publication information from the Neotoma Paleoecological Database.

Usage

```r
## Default S3 method:
get_publication(x, contactid, datasetid, author, pubtype, year, search)
```
get_publication.download

A function to get publications for downloads in the Neotoma Database using the API.

Description

The function takes a download and returns a table with publication information from the Neotoma Paleoecological Database.

Usage

```r
## S3 method for class 'download'
get_publication(x, ...)
```

Arguments

- **x**: an object of class download.
- **...**: objects passed from the generic. Not used in the call.

get_publication.download_list

A function to get publications for datasets in the Neotoma Database using the API.

Description

The function takes a download_list and returns a table with publication information from the Neotoma Paleoecological Database.

Arguments

- **x**: Numeric Publication ID value, either from get_dataset or known.
- **contactid**: Numeric Contact ID value, either from get_dataset or get_contact
- **datasetid**: Numeric Dataset ID, known or from get_dataset
- **author**: Character string for full or partial author’s name. Can include wildcards such as ’Smit*’ for all names beginning with ’Smit’.
- **pubtype**: Character string, one of eleven allowable types, see get_table. For a list of allowed types run get_table("PublicationTypes").
- **year**: Numeric publication year.
- **search**: A character string to search for within the article citation.
Usage

```r
# S3 method for class 'download_list'
get_publication(x, ...)
```

Arguments

- `x`: an object of class `download_list`.
- `...`: objects passed from the generic. Not used in the call.

---

**get_site**

Return Site Information.

Description

Return site information from the Neotoma Paleoecological Database.

`get_site` returns site information from the Neotoma Paleoecological Database based on parameters defined by the user.

Usage

```r
get_site(sitename, altmin, altmax, loc, gpid, ...)
```

Arguments

- `sitename`: character string representing the full or partial site name, or an object of class `dataset`, `dataset_list`, `download` or `download_list`.
- `altmin`: Minimum site altitude (in m).
- `altmax`: Maximum site altitude (in m).
- `loc`: A numeric vector c(lonW, latS, lonE, latN) representing the bounding box within which to search for sites. The convention here is to use negative values for longitudes west of Greenwich or longitudes south of the equator.
- `gpid`: A character string or numeric value, must correspond to a valid geopolitical identity in the Neotoma Database. Use `get.tables('GeoPoliticalUnits')` for a list of acceptable values, or link here: http://api.neotomadb.org/apdx/geopol.htm
- `...`: Optional additional arguments

Value

A data frame:

- `siteid`: Unique database record identifier for the site.
- `sitename`: Name of the site.
- `long`: Mean longitude, in decimal degrees, for a site (-180 to 180).
- `lat`: Mean latitude, in decimal degrees, for a site (-90 to 90).
get_site.dataset

Return Site Information from a numeric list of site ids.

description

Elevation in meters.

Free form description of a site, including such information as physiography and vegetation around the site.

long_acc

If the site is described by a bounding box this is the box width.

lat_acc

If the site is described by a bounding box this is the box height.

Author(s)

Simon J. Goring <simon.j.goring@gmail.com>

References

Neotoma Project Website: http://www.neotomadb.org API Reference: http://api.neotomadb.org/doc/resources/sites

Examples

```r
## Not run:
# What is the distribution of site elevations in Neotoma?
all.sites <- get_site()  # takes a bit of time.

plot(density(all.sites$elev, from = 0, na.rm=TRUE),
     main = 'Altitudinal Distribution of Neotoma Sites', xlab = 'Altitude (m)', log='x')

# Get site information from a dataset:
mw.datasets <- get_dataset(loc = c(-140, 50, -110, 65),
                          datatypem='pollen',
                          taxonname='Pinus*')

mw.sites <- get_site(mw.datasets)

## End(Not run)
```

description

Return site information from the Neotoma Paleoecological Database.

Usage

```r
## S3 method for class 'dataset'
get_site(sitename, ...)
```

Arguments

- `sitename` An object of class `dataset`.
- `...` Arguments passed from the generic method, not used.
get_site.dataset_list  
*Return Site Information from a dataset_list*

---

**Description**

Return site information from the Neotoma Paleoeological Database.

**Usage**

```r
## S3 method for class 'dataset_list'
get_site(sitename, ...)
```

**Arguments**

- `sitename`: An object of class `dataset_list`.
- `...`: Arguments passed from the generic method, not used.

---

get_site.default  
*Return Site Information.*

---

**Description**

Return site information from the Neotoma Paleoeological Database.

**Usage**

```r
## Default S3 method:
get_site(sitename, ...)
```

**Arguments**

- `sitename`: A character string representing the full or partial site name.
- `...`: Arguments passed from the generic method, not used.
get_site.download

Return Site Information from a download

Description

Return site information from the Neotoma Paleoecological Database.

Usage

```r
## S3 method for class 'download'
get_site(sitename, ...)
```

Arguments

- `sitename`: An object of class `download`.
- `...`: Arguments passed from the generic method, not used.

get_site.download_list

Return Site Information from a download_list

Description

Return site information from the Neotoma Paleoecological Database.

Usage

```r
## S3 method for class 'download_list'
get_site(sitename, ...)
```

Arguments

- `sitename`: An object of class `download_list`.
- `...`: Arguments passed from the generic method, not used.
get_site.geochronologic

Return Site Information from a geochronologic

Description

Return site information from the Neotoma Paleoecological Database.

Usage

## S3 method for class 'geochronologic'
get_site(sitename, ...)

Arguments

sitename An object of class geochronologic.
...
Arguments passed from the generic method, not used.

get_site.geochronologic_list

Return Site Information from a geochronologic_list

Description

Return site information from the Neotoma Paleoecological Database.

Usage

## S3 method for class 'geochronologic_list'
get_site(sitename, ...)

Arguments

sitename An object of class geochronologic_list.
...
Arguments passed from the generic method, not used.
get_site.integer  

Description

Return site information from the Neotoma Paleoecological Database.

Usage

## S3 method for class 'integer'
get_site(sitename, ...)

Arguments

sitename  

A integer or vector of integers.

...  
Arguments passed from the generic method, not used.

get_site.numeric  

Description

Return site information from the Neotoma Paleoecological Database.

Usage

## S3 method for class 'numeric'
get_site(sitename = NULL, ...)

Arguments

sitename  

A numeric value or vector of numeric elements.

...  
Arguments passed from the generic method, not used.
**get_table**  

*Get Neotoma value tables.*

**Description**

Get Neotoma value tables.

**Usage**

```r
get_table(table.name = NULL)
```

**Arguments**

- `table.name` Call one of the available tables in the Neotoma Database. A full listing of tables can be found here: [http://api.neotomadb.org/doc/resources/dbtables](http://api.neotomadb.org/doc/resources/dbtables). By default it returns all objects in the table.

**Details**

A table of values corresponding to the parameter of interest.

**Author(s)**

Simon J. Goring <simon.j.goring@gmail.com>

**References**


**Examples**

```r
## Not run:
taxon.table <- get_table('Taxa')

# Get the frequency of a random taxon in Neotoma.
tax_sample <- sample(nrow(taxon.table), 1)
cat("The taxon",
taxon.table$TaxonName[tax_sample],
"occurs in Neotoma",
length(get_dataset(taxonname = taxon.table$TaxonName[tax_sample])),
"times.")

## End(Not run)
```
get_taxa

Get taxon information from Neotoma.

Description
Get taxon information from Neotoma.

Usage
get_taxa(taxonid, taxonname, status, taxagroup, ecolgroup)

Arguments

- taxonid: Numeric taxon identifier used in Neotoma
- taxonname: A character string representing the full or partial name of taxa of interest.
- status: The current status of the taxon, one of 'extinct', 'extant', 'all'.
- taxagroup: The taxonomic grouping for the taxa. See http://api.neotomadb.org/doc/resources/taxa for the list of approved groupings.
- ecolgroup: The ecological group of the taxon. More detailed than taxagroup, can be obtained using get_table("EcolGroupTypes").

Value
Returns a data frame with the following components:

- TaxonID: Unique database record identifier for a taxon
- TaxonCode: Shorthand notation for a taxon identification
- TaxonName: Name of the taxon
- Author: Author(s) of the name. Used almost exclusively with beetle taxa
- Extinct: True if extinct; false if extant
- TaxaGroup: Code for taxa group to which taxon belongs
- EcolGroups: Array of ecological group codes to which the taxon belongs
- HigherTaxonID: TaxonID of the next higher taxonomic rank
- PublicationID: Publication identification number
- Notes: Free-form notes or comments about the taxon

Author(s)
Simon J. Goring <simon.j.goring@gmail.com>

References
Examples

```r
## Not run:
## Return all species taxa with "Abies" in name - note wildcard
taxa <- get_taxa(taxonname = "Abies")

## End(Not run)
```

---

gp.table

A list of all the geopolitical entities in the Neotoma database.

Description

A list of geopolitical entities with associated numeric ID values.

Usage

```r
gp.table
```

Format

```r
a data.frame object
```

Author(s)

Simon J. Goring <simon.j.goring@gmail.com>

Source

The Neotoma database.

---

plot_leaflet

Leaflet plots for neotoma data.

Description

A plotting function to provide interactive data investigation using the leaflet tools. This package requires a connection to the internet for proper functioning.

Usage

```r
plot_leaflet(x, providerTiles = "Stamen.TerrainBackground", ...)
```
pollen.equiv

Arguments

x

A neotoma data object

providerTiles

Default "Stamen.TerrainBackground", a character string indicating the tile background to be used for plotting.

... Other terms to be passed to the function.

Value

A leaflet object

pollen.equiv

A table to convert the pollen taxa identified by investigators to standardized lists.

Description

A list of standardized (published) taxonomies from the literature to help standardize taxonomies for synthesis work.

Usage

translate.table

Format

a data.frame object

Details

Taxon conversion table (readable).

Author(s)

Simon J. Goring <simon.j.goring@gmail.com>; Jeremiah Marsicek
read.tilia

Read proxy data from Tilia TLX files

Description
Read proxy data from a Tilia TLX format file.

Usage
read.tilia(file)

Arguments
file a string representing a Tilia TLX format file.

Value
Return a 'download' object.

Author(s)
Simon J. Goring <simon.j.goring@gmail.com>

Examples
### Not run:
crystal <- read.tilia('crystal.tlx')

### End(Not run)

---

read_bacon

Function to read in defined Bacon outputs.

Description
Reads in Bacon output and formats it for inclusion in a download object.

Usage
read_bacon(x, path = ".", add = FALSE, chron_name = "Bacon",
as_default = TRUE, download = NULL, sections = NULL,
age_field = "median", interp = TRUE)
Arguments

- x: A folder path that contains a Bacon age file.
- path: The location of the Cores folder.
- add: Should the results be added to an existing download? Defaults to FALSE.
- chron_name: The name for the chronology if the Bacon file is being added to a download.
- as_default: Should the chronology become the default?
- download: The target download if add is TRUE.
- sections: If there are multiple Bacon runs in a folder, identify the file by the number of sections in the run.
- age_field: Should the age be assigned to the "median" or the "wmean"?
- interp: If the depths don’t match up, should we interpolate from the Bacon output? (default TRUE)

Details

The function expects that you are in a working directory containing a "Cores" which would then contain output files from Bacon runs. The output can either be added to an existing record (for example, replacing the default age model returned by Neotoma), or it can be loaded on its own. If the depths for the loaded file do not match with the depths in the ‘download’ ‘sample.meta’ then the user can use the ‘interp’ parameter to interpolate between depths. This method uses linear interpolation.

Examples

```r
## Not run:
# Download the record for Lake O' Pines:
lake_o_dl <- get_download(15925)

# This assumes that you have Bacon installed in a folder and have
# set it to your working directory.
write_agefile(lake_o_dl[[1]], path = ".", chronology = 1,
  corename = "LAKEPINES", cal.prog = 'Bacon')

source("Bacon.R")

# These defaults just help the core run quickly, they're not
# necessarily good parameters.

Bacon("LAKEPINES", acc.mean = 10,
  thick = 50, depths.file = TRUE,
  suggest = FALSE, ask = FALSE)

lake_o_dl <- read_bacon("LAKEPINES", add = TRUE,
  download = download, sections = 17)

## End(Not run)
```
Stratiplot.download  
*Palaeoecological stratigraphic diagrams*

**Description**

Draws palaeoecological diagrams from a download object. Allows control of variable type (using the `tran` function from the analogue package), and taxonomic grouping.

**Usage**

```r
## S3 method for class 'download'
Stratiplot(x, yaxis = "age", method = "none",
          group = NULL, ...)
```

**Arguments**

- `x`: A download object.
- `yaxis`: One of the columns in `sample.meta`, including `depth`, `age`, `age.younger`, or `age.older`, default `age`.
- `method`: An option for axis transformation using `tran` from the analogue package. "none" by default.
- `group`: An ecological group from the taxon table.
- `...`: variables to be passed to `stratiplot`.

**Details**

A wrapper for the analogue package’s `Stratiplot` function. Allowing the user to plot a stratigraphic diagram directly from a download object.

**Value**

A `trellis` object.

**Examples**

```r
## Not run:
lake_o_dl <- get_download(15925)
Stratiplot(lake_o_dl[[1]])

## End(Not run)
```
Stratiplot.download_list

Palaeoecological stratigraphic diagrams

Description

Draws paleoecological diagrams from a download_list object. Allows control of variable type (using the tran function from the analogue package), and taxonomic grouping. This function only works for download_list objects that contain a single object.

Usage

## S3 method for class 'download_list'
Stratiplot(x, yaxis = "age", method = "none", group = NULL, ...)

Arguments

- **x**: A download_list object.
- **yaxis**: One of the columns in sample.meta, including depth, age, age.younger, or age.older, default age.
- **method**: An option for axis transformation using tran from the analogue package. "none" by default.
- **group**: An ecological group from the taxon table.
- **...**: Variables to be passed to Stratiplot.

Details

A wrapper for the analogue package's Stratiplot function. Allowing the user to plot a stratigraphic diagram directly from a download object.

Value

A trellis object.

Examples

```r
## Not run:
lake_o_dl <- get_download(15925)
# This works:
Stratiplot(lake_o_dl)

lakes_o_nw <- get_download(get_site(sitename = "Lake B"))
# This fails:
# Stratiplot(lake_o_nw)
```

## End(Not run)
Description

Extracts taxa from download objects and returns them in a useful format.

Usage

taxa(obj, ...)

## S3 method for class 'download'
taxa(obj, ...)  

## S3 method for class 'download_list'
taxa(obj, collapse = TRUE, hierarchy = FALSE, ...)

Arguments

obj       an R object from which counts are to be extracted.
...        arguments passed to other methods.
collapse  should the results be returned as a list, one for each site (FALSE), or a single dataframe of all taxa? Default is TRUE
hierarchy Should the taxonomic hierarchy be included?

Details

Methods are available for "download" and "download_list" objects.

Value

Either a data frame of taxa or a list of such objects.

Author(s)

Simon Goring

Examples

## Not run:
ostracodes <- get_dataset(datasettype = 'ostracode')

ostro.dl <- get_download(ostracodes)
ostro.taxa <- taxa(ostro.dl)

## End(Not run)
### taxon.list  
**Neotoma taxon list**

**Description**
The taxonomy table for datasets in neotoma, as would be returned by `get_table`.

**Usage**
taxon.list

**Format**
a `data.frame` object

**Author(s)**
Simon J. Goring <simon.j.goring@gmail.com>

**Source**
The Neotoma database.

### translate.table

**A table to convert the original taxa to standardized lists.**

**Description**
A list of standardized (published) taxonomies from the literature to help standardize taxonomies for synthesis work.

**Usage**
translate.table

**Format**
a `data.frame` object

**Author(s)**
Simon J. Goring <simon.j.goring@gmail.com>

**Source**
The Neotoma database.
**write_agefile**  

*Write age control file to disk formatted for either Bacon or Clam*

**Description**

Passing in a download object the function outputs a Bacon or Clam formatted file to a user defined destination for age modelling with existing age-depth modeling software.

**Usage**

```r
write_agefile(download, chronology = 1, path, corename,
  cal.prog = "Bacon")
```

**Arguments**

- **download**: A single site returned by `get_download`.
- **chronology**: Default is 1, the default chronology for the core. If a core has more than one chronology the user can define a different set of chronological controls.
- **path**: The location of the 'Cores' folder & working directory for Bacon. Do not include "Cores" in the path name.
- **corename**: The intended handle for the core, to be used in writing to file.
- **cal.prog**: The method intended to build the age model, either 'Bacon' or 'Clam'.

**Value**

This command returns a file in location `path/Cores` containing all the relevant information required to build either the default or prior chronology for a core.

**Author(s)**

Simon J. Goring <simon.j.goring@gmail.com>

**References**


**Examples**

```r
## Not run:
# Find a particular record:

three_pines <- get_download(get_dataset(get_site("Three Pines Bog"),
  datasettype = "pollen"))

# You will need to edit the 'path' argument here to point to a directory that
# contains a 'Cores' directory.
```
write_agefile(download = three_pines[[1]],
    path = "./inst",
    corename = "THREEPINES",
    cal.prog = "Bacon")

## End(Not run)
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