

Package: traits (via r-universe)

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Title Species Trait Data from Around the Web

Version 0.6.0

Description Species trait data from many sources, including sequence data from NCBI (<<https://www.ncbi.nlm.nih.gov/>>), plant traits from BETYdb, and data from EOL Traitbank and BirdLife International.

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URL <https://docs.ropensci.org/traits/>,
<https://github.com/ropensci/traits>

BugReports <https://github.com/ropensci/traits/issues>

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betydb	<i>Search for traits from BETYdb</i>
--------	--------------------------------------

Description

Search for traits from BETYdb
Get details about a single item from a table

Usage

```
betydb_record(  
  id,  
  table,  
  api_version = NULL,  
  betyurl = NULL,  
  fmt = NULL,  
  key = NULL,  
  user = NULL,  
  pwd = NULL,  
  progress = TRUE,  
  ...  
)
```

```
betydb_trait(  
    id,  
    genus = NULL,  
    species = NULL,  
    api_version = NULL,  
    betyurl = NULL,  
    fmt = "json",  
    key = NULL,  
    user = NULL,  
    pwd = NULL,  
    progress = TRUE,  
    ...  
)
```

```
betydb_specie(  
    id,  
    genus = NULL,  
    species = NULL,  
    api_version = NULL,  
    betyurl = NULL,  
    fmt = "json",  
    key = NULL,  
    user = NULL,  
    pwd = NULL,  
    progress = TRUE,  
    ...  
)
```

```
betydb_citation(  
    id,  
    genus = NULL,  
    species = NULL,  
    api_version = NULL,  
    betyurl = NULL,  
    fmt = "json",  
    key = NULL,  
    user = NULL,  
    pwd = NULL,  
    progress = TRUE,  
    ...  
)
```

```
betydb_site(  
    id,  
    api_version = NULL,  
    betyurl = NULL,  
    fmt = "json",  
    key = NULL,
```

```

    user = NULL,
    pwd = NULL,
    progress = TRUE,
    ...
)

betydb_experiment(
  id,
  api_version = NULL,
  betyurl = NULL,
  fmt = "json",
  key = NULL,
  user = NULL,
  pwd = NULL,
  progress = TRUE,
  ...
)

```

Arguments

<code>id</code>	(integer) One or more ids for a species, site, variable, etc.
<code>table</code>	(character) Name of the database table with which this ID is associated.
<code>api_version</code>	(character) Which version of the BETY API should we query? One of "v0" or "beta". Default is <code>options("betydb_api_version")</code> if set, otherwise "v0".
<code>betyurl</code>	(string) URL to target BETYdb. Default is "https://www.betydb.org/". This can be also be set with <code>options("betydb_url")</code> . Note: only https://www.betydb.org/ is officially supported.
<code>fmt</code>	(character) Format to return data in, one of json, xml, csv. Only json currently supported.
<code>key</code>	(character) An API key. Use this or user/pwd combo. Save in your .Rprofile file as <code>options(betydb_key = "your40digitkey")</code> . Optional
<code>user, pwd</code>	(character) A user name and password. Use a user/pwd combo or an API key. Save in your .Rprofile file as <code>options(betydb_user = "yournamehere")</code> and <code>options(betydb_pwd = "yourpasswordhere")</code> . Optional
<code>progress</code>	show progress bar? default: TRUE
<code>...</code>	Curl options passed on to GET . Optional
<code>genus</code>	(character) A genus name. Optional
<code>species</code>	(character) A specific epithet. Optional

Details

BETYdb is focused on bioenergy crops and related ecological data. This package officially supports the main public instance at <https://www.betydb.org/>.

This package queries plant traits, phenotypes, biomass yields, and ecosystem functions. It does not currently interface with the workflow and provenance data that support PECAn Project (pecanproject.org) and TERRA REF (terraref.org) software.

API documentation: <https://pecan.gitbooks.io/betydb-data-access/content/API.html>
API endpoints are here: <https://www.betydb.org/api/docs> This package currently uses the original 'v0' API by default. To use a newer version, set `api_version`. Newer versions of the API will support database inserts.

Value

For `betydb_trait`, `betydb_specie`, `betydb_citation`, `betydb_site`, and `betydb_experiment`, a named list of information about the requested object. For functions that query multiple records, a `data.frame` containing the query results.

Authentication

Defers to use API key first since it's simpler, but if you don't have an API key, you can supply a username and password.

Functions

Singular functions like `betydb_trait` accept an id and additional parameters, and return a list of variable outputs depending on the inputs.

However, plural functions like `betydb_traits` accept query parameters, but not ids, and always return a single `data.frame`.

`betydb_search("Search terms", ...)` is a convenience wrapper that passes all further arguments to `betydb_query(table = "search", search = "Search terms", ...)`. See there for details on possible arguments.

References

API documentation <https://pecan.gitbooks.io/betydb-data-access/content/API.html> and <https://www.betydb.org/api/docs>

See Also

[betydb_query](#)

Examples

```
if (interactive()) {  
  # General Search  
  out <- betydb_search(query = "Switchgrass Yield")  
  if (NROW(out)) {  
    mean_result <- tapply(as.numeric(out$mean), out$id, function(z) mean(z, na.rm = TRUE))  
    sort(mean_result, decreasing = TRUE)  
  }  
  # Get by ID  
  ## Traits  
  betydb_trait(id = 10)  
  ## Species  
  betydb_specie(id = 1)  
  ## Citations
```

```

betydb_citation(id = 1)
## Site information
betydb_site(id = 795)
}

```

betydb_query

Query a BETY table

Description

Query a BETY table

Usage

```

betydb_query(
  ...,
  table = "search",
  key = NULL,
  api_version = NULL,
  betyurl = NULL,
  user = NULL,
  pwd = NULL,
  progress = TRUE
)

betydb_search(
  query = "Maple SLA",
  ...,
  include_unchecked = NULL,
  progress = TRUE
)

```

Arguments

...	(named character) Columns to query, as key="value" pairs. Note that betydb_query passes these along to BETY with no check whether the requested keys exist in the specified table.
table	(character) The name of the database table to query, or "search" (the default) for the traits and yields view
key	(character) An API key. Use this or user/pwd combo. Save in your .Rprofile file as options(betydb_key = "your40digitkey"). Optional
api_version	(character) Which version of the BETY API should we query? One of "v0" or "beta". Default is options("betydb_api_version") if set, otherwise "v0".
betyurl	(string) url to target instance of betydb. Default is options("betydb_url") if set, otherwise "https://www.betydb.org/"

user, pwd	(character) A user name and password. Use a user/pwd combo or an API key. Save in your .Rprofile file as options(betydb_user = "yournamehere") and options(betydb_pwd = "yourpasswordhere"). Optional
progress	show progress bar? default: TRUE
query	(character) A string containing one or more words to be queried across all columns of the "search" table.
include_unchecked	(logical) Include results that have not been quality checked? Applies only to tables with a "checked" column: "search", "traits", "yields". Default is to exclude unchecked values.

Details

Use betydb_query to retrieve records from a table that match on all the column filters specified in '...'. If no filters are specified, retrieves the whole table. In API versions that support it (i.e. not in v0), filter strings beginning with "~" are treated as regular expressions.

Value

A data.frame with attributes containing request metadata, or NULL if the query produced no results

Examples

```
if (interactive()) {
  # literal vs regular expression vs anchored regular expression:
  betydb_query(units = "Mg", table = "variables")
  # NULL
  betydb_query(units = "Mg/ha", table = "variables")[["name"]]
  # $name
  # [1] "a_biomass"                "root_live_biomass"
  # [3] "leaf_dead_biomass_in_Mg_ha" "SDM"

  nrow(betydb_query(genus = "Miscanthus", table = "species"))
  # [1] 10
  unique(betydb_query(genus = "~misc", table = "species", api_version = "beta")[["genus"]])
  # $genus
  # [1] "Platymiscium" "Miscanthus"   "Dermiscellum"

  unique(betydb_query(genus = "^misc", table = "species", api_version = "beta")[["genus"]])
  # $genus
  # [1] "Miscanthus"
}
```

birdlife_habitat	<i>Get bird habitat information from BirdLife/IUCN</i>
------------------	--

Description

Get bird habitat information from BirdLife/IUCN

Usage

```
birdlife_habitat(id)
```

Arguments

id	A single IUCN species ID
----	--------------------------

Value

a data.frame with level 1 and level 2 habitat classes, as well as importance ratings and occurrence type (e.g. breeding or non-breeding). The habitat classification scheme is described at <https://www.iucnredlist.org/resources/classification-schemes>

Author(s)

David J. Harris <harry491@gmail.com>

See Also

Other birdlife: [birdlife_threats\(\)](#)

Examples

```
if (interactive()) {  
  # Setophaga chrysoparia  
  birdlife_habitat(22721692)  
  # Passer domesticus  
  birdlife_habitat(103818789)  
}
```

birdlife_threats	<i>Get bird threat information from BirdLife/IUCN</i>
------------------	---

Description

Get bird threat information from BirdLife/IUCN

Usage

```
birdlife_threats(id)
```

Arguments

id	A single IUCN species ID
----	--------------------------

Value

a data.frame with the species ID and two levels of threat descriptions, plus stresses, timing, scope, severity, and impact associated with each stressor.

Author(s)

David J. Harris <harry491@gmail.com>

See Also

Other birdlife: [birdlife_habitat\(\)](#)

Examples

```
if (interactive()) {  
  # Setophaga chrysoparia  
  birdlife_threats(22721692)  
  # Aburria aburri  
  birdlife_threats(22678440)  
}
```

leda	<i>Access LEDA trait data</i>
------	-------------------------------

Description

Access LEDA trait data

Usage

```
leda(trait = "age_first_flowering", ...)
```

Arguments

trait	(character) Trait to get. See Details.
...	Curl options passed on to crul::verb-GET

Details

For parameter trait, one of age_first_flowering, branching, buds_seasonality, buds_vertical_dist, canopy_height, dispersal_type, leaf_distribution, ldmc_geo, leaf_mass, leaf_size, morphology_disperal, growth_form, life_span, releasing_height, seed_longevity, seed_mass, seed_number, seed_shape, shoot_growth_form, snp, ssd, tv, or clonal_growth_organs

The following are not supported as they are too much of a pain to parse: buoyancy, seed_bank, sla_geo

Value

A data frame with trait data.

Examples

```
if (interactive()) {
  # Age of first flowering
  leda(trait = "age_first_flowering")

  # Seed number
  leda("seed_number")

  # Releasing height
  leda(trait = "releasing_height")

  # Clonal growth organs
  leda(trait = "clonal_growth_organs")

  all <- c("age_first_flowering", "branching", "buds_seasonality",
    "buds_vertical_dist", "canopy_height",
    "dispersal_type", "leaf_distribution", "ldmc_geo", "leaf_mass",
    "leaf_size", "morphology_disperal", "growth_form", "life_span",
    "releasing_height", "seed_longevity", "seed_mass",
```

```

      "seed_number", "seed_shape", "shoot_growth_form",
      "snp", "ssd", "tv", "clonal_growth_organs")
out <- list()
for (i in seq_along(all)) {
  cat(all[i], sep = "\n")
  out[[i]] <- leda(all[i])
}
sapply(out, NROW)
}

```

ncbi_byid

*Retrieve gene sequences from NCBI by accession number.***Description**

Retrieve gene sequences from NCBI by accession number.

Usage

```
ncbi_byid(ids, format = NULL, verbose = TRUE)
```

Arguments

ids	(character) GenBank ids to search for. One or more. Required.
format	(character) Return type, e.g., "fasta". NOW IGNORED.
verbose	(logical) If TRUE (default), informative messages printed.

Details

If bad ids are included with good ones, the bad ones are silently dropped. If all ids are bad you'll get a stop with error message.

Value

data.frame of the form:

- taxon - taxonomic name (may include some junk, but hard to parse off)
- taxonomy - organism lineage
- gene_desc - gene description
- organelle - if mitochondrial or chloroplast
- gi_no - GI number
- acc_no - accession number
- keyword - if official DNA barcode
- specimen_voucher - museum/lab accession number of vouchered material
- lat_lon - longitude/latitude of specimen collection event

- country - country/location of specimen collection event
- paper_title - title of study
- journal - journal study published in (if published)
- first_author - first author of study
- uploaded_date - date sequence was uploaded to GenBank
- length - sequence length
- sequence - sequence character string

Author(s)

Scott Chamberlain, Rupert Collins

See Also

[ncbi_searcher\(\)](#), [ncbi_byname\(\)](#)

Examples

```
if (interactive()) {
  # A single gene
  ncbi_byid(ids = "360040093")

  # Many genes (with different accession numbers)
  ncbi_byid(ids = c("360040093", "347448433"))
}
```

ncbi_byname

Retrieve gene sequences from NCBI by taxon name and gene names.

Description

Retrieve gene sequences from NCBI by taxon name and gene names.

Usage

```
ncbi_byname(
  taxa,
  gene = "COI",
  seqrange = "1:3000",
  getrelated = FALSE,
  verbose = TRUE,
  batch_size = 100,
  ...
)
```

Arguments

taxa	(character) Scientific name to search for.
gene	(character) Gene or genes (in a vector) to search for. See examples.
seqrage	(character) Sequence range, as e.g., "1:1000". This is the range of sequence lengths to search for. So "1:1000" means search for sequences from 1 to 1000 characters in length.
getrelated	(logical) If TRUE, gets the longest sequences of a species in the same genus as the one searched for. If FALSE, returns nothing if no match found.
verbose	(logical) If TRUE (default), informative messages printed.
batch_size	An integer specifying the number of names to query per batch.
...	Curl options passed on to curl::verb-GET

Details

Removes predicted sequences so you don't have to remove them. Predicted sequences are those with accession numbers that have "XM_" or "XR_" prefixes. This function retrieves one sequences for each species, picking the longest available for the given gene.

Value

data.frame

Author(s)

Scott Chamberlain

See Also

[ncbi_searcher\(\)](#), [ncbi_byid\(\)](#)

Examples

```
if (interactive()) {
  # A single species
  ncbi_byname(taxa = "Acipenser brevirostrum")

  # Many species
  species <- c("Colletes similis", "Halictus ligatus", "Perdita californica")
  ncbi_byname(taxa = species, gene = c("coi", "co1"), seqrange = "1:2000")
}
```

ncbi_searcher

*Search for gene sequences available for taxa from NCBI.***Description**

Search for gene sequences available for taxa from NCBI.

Usage

```
ncbi_searcher(
  taxa = NULL,
  id = NULL,
  seqrange = "1:3000",
  getrelated = FALSE,
  fuzzy = FALSE,
  limit = 500,
  entrez_query = NULL,
  hypothetical = FALSE,
  verbose = TRUE,
  sleep = 0L
)
```

Arguments

taxa	(character) Scientific name to search for.
id	(character) Taxonomic id to search for. Not compatible with argument taxa.
seqrange	(character) Sequence range, as e.g., "1:1000". This is the range of sequence lengths to search for. So "1:1000" means search for sequences from 1 to 1000 characters in length.
getrelated	(logical) If TRUE, gets the longest sequences of a species in the same genus as the one searched for. If FALSE, returns nothing if no match found.
fuzzy	(logical) Whether to do fuzzy taxonomic ID search or exact search. If TRUE, we use <code>xXarbitraryXx[porgn:__txid<ID>]</code> , but if FALSE, we use <code>txid<ID></code> . Default: FALSE
limit	(numeric) Number of sequences to search for and return. Max of 10,000. If you search for 6000 records, and only 5000 are found, you will of course only get 5000 back.
entrez_query	(character; length 1) An Entrez-format query to filter results with. This is useful to search for sequences with specific characteristics. The format is the same as the one used to seach genbank. (https://www.ncbi.nlm.nih.gov/books/NBK3837/#EntrezHelp.Entrez_Searching_Options)
hypothetical	(logical; length 1) If FALSE, an attempt will be made to not return hypothetical or predicted sequences judging from accession number prefixes (XM and XR). This can result in less than the <code>limit</code> being returned even if there are more sequences available, since this filtering is done after searching NCBI.

verbose (logical) If TRUE (default), informative messages printed.

sleep (integer) number of seconds to sleep before each HTTP request. use if running to 429 Too Many Requests errors from NCBI. default: 0 (no sleep)

Value

data.frame of results if a single input is given. A list of data.frames if multiple inputs are given.

Authentication

NCBI rate limits requests. If you set an API key you have a higher rate limit. Set your API key like `Sys.setenv(ENTREZ_KEY="yourkey")` or you can use `?rentrez::set_entrez_key`. set verbose curl output (`curl::set_verbose()`) to make sure your api key is being sent in the requests

Author(s)

Scott Chamberlain, Zachary Foster <zacharyfoster1989@gmail.com>

See Also

[ncbi_byid](#), [ncbi_byname](#)

Examples

```
if (interactive()) {
  # A single species
  out <- ncbi_searcher(taxa = "Umbra limi", seqrage = "1:2000")
  # Get the same species information using a taxonomy id
  out <- ncbi_searcher(id = "75935", seqrage = "1:2000")
  # If the taxon name is unique, using the taxon name and id are equivalent
  all(ncbi_searcher(id = "75935") == ncbi_searcher(taxa = "Umbra limi"))
  # If the taxon name is not unique, use taxon id
  # "266948" is the uid for the butterfly genus, but there is also a genus
  # of orchids with the
  # same name
  nrow(ncbi_searcher(id = "266948")) == nrow(ncbi_searcher(taxa = "Satyrium"))
  # get list of genes available, removing non-unique
  unique(out$gene_desc)
  # does the string 'RAG1' exist in any of the gene names
  out[grep("RAG1", out$gene_desc, ignore.case = TRUE), ]

  # A single species without records in NCBI
  out <- ncbi_searcher(taxa = "Sequoia wellingtonia", seqrage = "1:2000",
    getrelated = TRUE)

  # Many species, can run in parallel or not using plyr
  species <- c("Salvelinus alpinus", "Ictalurus nebulosus", "Carassius auratus")
  out2 <- ncbi_searcher(taxa = species, seqrage = "1:2000")
  lapply(out2, head)
  library("plyr")
  out2df <- ldply(out2) # make data.frame of all
  unique(out2df$gene_desc) # get list of genes available, removing non-unique
}
```

```
out2df[grep("12S", out2df$gene_desc, ignore.case = TRUE), ]

# Using the getrelated and entrez_query options
ncbi_searcher(taxa = "Olpidiopsidales", limit = 5, getrelated = TRUE,
              entrez_query = "18S[title] AND 28S[title]")

# get refseqs
one <- ncbi_searcher(taxa = "Salmonella enterica",
                    entrez_query = "srcdb_refseq[PROP]")
two <- ncbi_searcher(taxa = "Salmonella enterica")
}
```

plantatt	<i>PLANTATT plant traits dataset</i>
----------	--------------------------------------

Description

PLANTATT plant traits dataset

taxa_search	<i>Search for traits by taxa names</i>
-------------	--

Description

Search for traits by taxa names

Usage

```
taxa_search(x, db, ...)
```

Arguments

- x (character) Taxonomic name(s) to search for
- db (character) only 'ncbi' for now - other options maybe in the future
- ... Curl options passed on to [GET](#)

Value

A data.frame

Author(s)

Scott Chamberlain

Examples

```
if (interactive()) {
  taxa_search("Poa annua", db = "ncbi")
}
```

tr_ernest	<i>Amniote life history dataset</i>
-----------	-------------------------------------

Description

Amniote life history dataset

Usage

```
tr_ernest(read = TRUE, ...)
```

Arguments

read (logical) read in csv files. Default: TRUE

... Curl options passed on to `curl::HttpClient()`

Details

When using this data, cite the paper:

Myhrvold, N. P., Baldrige, E., Chan, B., Sivam, D., Freeman, D. L. and Ernest, S. K. M. (2015), An amniote life-history database to perform comparative analyses with birds, mammals, and reptiles. *Ecology*, 96: 3109. <https://doi.org/10.1890/15-0846R.1>

As well as the Dryad data package:

L. Freeman, Daniel; P. Myhrvold, Nathan; Chan, Benjamin; Sivam, Dhileep; Ernest, S. K. Morgan; Baldrige, Elita (2016): Full Archive. figshare. <https://doi.org/10.6084/m9.figshare.3563457.v1>

Value

paths to the files (character) if read=FALSE or a list of data.frame's if read=TRUE

References

<https://doi.org/10.1890/15-0846R.1> <https://doi.org/10.6084/m9.figshare.3563457.v1>

Examples

```
if (interactive()) {
  res <- tr_ernest()
  res$data
  res$references
  res$sparse
  res$range_count
}
```

tr_zanne

*Zanne et al. plant dataset***Description**

Zanne et al. plant dataset

Usage

```
tr_zanne(read = TRUE, ...)
```

Arguments

read (logical) read in csv files. Default: TRUE

... Curl options passed on to `curl::HttpClient()`

Details

This data is a dataset stored on Dryad (doi: 10.5061/dryad.63q27). When using this data, cite the paper:

Zanne AE, Tank DC, Cornwell WK, Eastman JM, Smith SA, FitzJohn RG, McGlinn DJ, O'Meara BC, Moles AT, Reich PB, Royer DL, Soltis DE, Stevens PF, Westoby M, Wright IJ, Aarssen L, Bertin RI, Calaminus A, Govaerts R, Hemmings F, Leishman MR, Oleksyn J, Soltis PS, Swenson NG, Warman L, Beaulieu JM, Ordonez A (2014) Three keys to the radiation of angiosperms into freezing environments. *Nature* 506(7486): 89-92. <http://dx.doi.org/10.1038/nature12872>

As well as the Dryad data package:

Zanne AE, Tank DC, Cornwell WK, Eastman JM, Smith SA, FitzJohn RG, McGlinn DJ, O'Meara BC, Moles AT, Reich PB, Royer DL, Soltis DE, Stevens PF, Westoby M, Wright IJ, Aarssen L, Bertin RI, Calaminus A, Govaerts R, Hemmings F, Leishman MR, Oleksyn J, Soltis PS, Swenson NG, Warman L, Beaulieu JM, Ordonez A (2013) Data from: Three keys to the radiation of angiosperms into freezing environments. Dryad Digital Repository. <http://dx.doi.org/10.5061/dryad.63q27.2>

Value

paths to the files (character) if read=FALSE or a list of data.frame's if read=TRUE

References

<http://datadryad.org/resource/doi:10.5061/dryad.63q27>

Examples

```
if (interactive()) {  
  res <- tr_zanne()  
  res$tax_lookup  
  res$woodiness  
  res$freezing  
  res$leaf_phenology  
}
```

traitbank

Search for traits from EOL's Traitbank

Description

Search for traits from EOL's Traitbank

Usage

```
traitbank(query, key = NULL, ...)
```

Arguments

query	(character) a query to the EOL Cypher service that holds Traitbank data. required. no default query given. see examples
key	(character) EOL Cypher query API key. required, either passed in or as an environment variable
...	Curl options passed on to verb-GET

Details

traitbank is an interface to the EOL Cypher query. Note that the previous interface EOL had for Traits has been completely replaced - thus, this function is completely different. You no longer query by EOL page id, but using the query language for a database called Neo4J. See the docs for help. Later we plan to make a more user friendly interface to get Traitbank data that doesn't require knowing the Neo4J query syntax

Value

a list

Authentication

You'll need an EOL cypher key to use this function. Get one by signing in to your EOL account https://eol.org/users/sign_in then head to <https://eol.org/services/authenticate> to get a key. Store your key in your .Renviro file or similar under the name "EOL_CYPHER_KEY", and we will use that key in this function. Alternatively, you can pass in your key to the key parameter, but we do not recommend doing that as you risk accidentally committing your key to the public web.

References

https://github.com/EOL/eol_website/blob/master/doc/api.md https://github.com/EOL/eol_website/blob/master/doc/query-examples.md

Examples

```
if (interactive()) {
  # traitbank_query function
  traitbank(query = "MATCH (n:Trait) RETURN n LIMIT 1;")

  # traitbank function
  res <- traitbank(query = "MATCH (n:Trait) RETURN n LIMIT 2;")
  res
}
```

traits-defunct

Defunct functions in traits

Description

These functions have been removed.

Details

- `eol_invasive_`: This function has moved to a new package. See `originr::eol`
- `fe_native`: This function has moved to a new package. See `originr::flora_europaea`
- `g_invasive`: This function has moved to a new package. See `originr::gisd`
- `is_native`: This function has moved to a new package. See `originr::is_native`
- `tr_usda`: the API behind this function is down for good
- `coral_locations`: API down for good, as far as I can tell
- `coral_methodologies`: API down for good, as far as I can tell
- `coral_resources`: API down for good, as far as I can tell
- `coral_species`: API down for good, as far as I can tell
- `coral_taxa`: API down for good, as far as I can tell
- `coral_traits`: API down for good, as far as I can tell

traits_cache	<i>Caching</i>
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Description

Manage cached traits package files with **hoardr**

Details

The default cache directory is `paste0(rappdirs::user_cache_dir(), "/R/traits")`, but you can set your own path using `cache_path_set()`

`cache_delete` only accepts 1 file name, while `cache_delete_all` doesn't accept any names, but deletes all files. For deleting many specific files, use `cache_delete` in a `lapply()` type call

Value

A `hoardr` object with methods for managing cached files

Useful user functions

- `traits_cache$cache_path_get()` get cache path
- `traits_cache$cache_path_set()` set cache path
- `traits_cache$list()` returns a character vector of full path file names
- `traits_cache$files()` returns file objects with metadata
- `traits_cache$details()` returns files with details
- `traits_cache$delete()` delete specific files
- `traits_cache$delete_all()` delete all files, returns nothing

Examples

```
traits_cache

# list files in cache
traits_cache$list()

# delete certain database files
# traits_cache$delete("file path")
# traits_cache$list()

# delete all files in cache
# traits_cache$delete_all()
# traits_cache$list()

# set a different cache path from the default
```

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