Package ‘ctrdata’

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URL https://cran.r-project.org/package=ctrdata
BugReports https://github.com/rfhb/ctrdata/issues
Description Provides functions for querying, retrieving and analyzing protocol- and results-related information on clinical trials from two public registers, the 'European Union Clinical Trials Register' ('EUCTR', <https://www.clinicaltrialsregister.eu/>), 'ClinicalTrials.gov' ('CTGOV', <https://clinicaltrials.gov/>) and the 'ISRCTN' (<http://www.isrctn.com/>).
Trial information is downloaded, converted and stored in a database ('SQLite' or 'MongoDB', via 'nodbi'). Functions are provided to identify de-duplicated records, to easily find and extract variables (fields) of interest even from complex nesting as used by the registers, and to update previous queries that users retrieved in a database. The package can be used for meta analysis and trend-analysis of the design and conduct as well as results of clinical trials.
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**Description**

An active substance can be identified by a recommended international nonproprietary name, a trade or product name, or a company code(s).

**Usage**

```r
ctrFindActiveSubstanceSynonyms(activesubstance = "")
```

**Arguments**

- `activesubstance`

  An active substance, in an atomic character vector

**Details**

At this time, this function uses the register ClinicalTrials.Gov to detect which substances were also searched for.

**Value**

A character vector of the active substance (input parameter) and synonyms, if any were found
ctrGetQueryUrl

Examples

```r
## Not run:
ctrFindActiveSubstanceSynonyms(
  activesubstance = "imatinib"
)
## End(Not run)
```

---

### Description

Extract query parameters and register name from input or from clipboard into which the URL of a register search was copied

### Usage

```r
ctrGetQueryUrl(url = "", register = "")
```

### Arguments

- **url**: URL such as from the browser address bar. If not specified, clipboard contents will be checked for a suitable URL. Can also contain a query term such as from `dbQueryHistory()$"query-term"`
- **register**: Optional name of register (i.e., "EUCTR" or "CTGOV") in case url is a query term

### Value

- A string of query parameters that can be used to retrieve data from the register.
- A data frame with column names query term and register name that can directly be used in `ctrLoadQueryIntoDb` and in `ctrOpenSearchPagesInBrowser`

### Examples

```r
## Not run:
db <- nodbi::src_sqlite(
  collection = "my_collection"
)

# user now copies into the clipboard the URL from
# the address bar of the browser that shows results
# from a query in one of the trial registers
```
ctrGetQueryUrlFromBrowser

Import from clipboard the URL of a search in one of the registers

Description

Import from clipboard the URL of a search in one of the registers

Usage

ctrGetQueryUrlFromBrowser(url = "", register = "")

Arguments

url URL such as from the browser address bar. If not specified, clipboard contents will be checked for a suitable URL. Can also contain a query term such as from dbQueryHistory()"query-term"

register Optional name of register (i.e., "EUCTR" or "CTGOV") in case url is a query term

Value

A string of query parameters that can be used to retrieve data from the register. A data frame with column names query term and register name that can directly be used in ctrLoadQueryIntoDb and in ctrOpenSearchPagesInBrowser

Examples

## Not run:
db <- nдобi::src_sqlite(
   collection = "my_collection"
)

# user now copies into the clipboard the URL from # the address bar of the browser that shows results # from a query in one of the trial registers ctrLoadQueryIntoDb( 
   ctrGetQueryUrlFromBrowser(), 
   con = db
)
ctrLoadQueryIntoDb

Retrieve or update information on clinical trials from register and store in database

Description

This is the main function of package ctrdata for accessing registers. Note that re-rerunning this function adds or updates trial records in a database, even if from different queries or different registers. Updating means that the previously stored record is overwritten; see annotation.text for persisting user comments added to a record.

Usage

ctrLoadQueryIntoDb(
  queryterm = "", 
  register = "", 
  querytoupdate = 0L, 
  forctoupdate = FALSE, 
  euctrresults = FALSE, 
  euctrresultshistory = FALSE, 
  euctrresultspdfpath = NULL, 
  annotation.text = "", 
  annotation.mode = "append", 
  parallelretrievals = 10L, 
  only.count = FALSE, 
  con = NULL, 
  verbose = FALSE
)

Arguments

queryterm Either a string with the full URL of a search in a register, or the data frame returned by the ctrGetQueryUrl or the dbQueryHistory functions, or, together with parameter register, a string with query elements of a search URL. The queryterm is recorded in the collection for later use to update records.

register String with abbreviation of register to query, either "EUCTR", "CTGOV" or "ISRCTN". Not needed if queryterm provide the information which register to query (see queryterm).

querytoupdate Either the word "last" or the number of the query (based on dbQueryHistory) that should be run to retrieve any trial records that are new or have been updated since this query was run the last time. This parameter takes precedence over queryterm. For EUCTR, updates are available only for the last seven days; the query is run again if more time has passed since it was run last.
**ctrLoadQueryIntoDb**

- **forcetoupdate**: If TRUE, run again the query given in `querytoupdate`, irrespective of when it was run last (default is FALSE).

- **euctrresults**: If TRUE, also download available results when retrieving and loading trials from EUCTR. This slows down this function. (For CTGOV, all available results are always retrieved and loaded.)

- **euctrresultshistory**: If TRUE, also download available history of results publication in EUCTR. This is quite time-consuming (default is FALSE).

- **euctrresultspdfpath**: If a valid directory is specified, save PDF files of result publications from EUCTR into this directory (default is NULL).

- **annotation.text**: Text to be including in the records retrieved with the current query, in the field "annotation".

- **annotation.mode**: One of "append" (default), "prepend" or "replace" for new `annotation.text` with respect to any existing annotation for the records retrieved with the current query.

- **parallelretrievals**: Number of parallel downloads of information from the register, defaults to 10.

- **only.count**: Set to TRUE to return only the number of trial records found in the register for the query. Does not load trial information into the database. Default is FALSE.

- **con**: A src connection object, as obtained with `nodbi::src_mongo()` or `nodbi::src_sqlite()`

- **verbose**: Printing additional information if set to TRUE; default is FALSE.

**Value**

A list with elements "n" (the number of trials that were newly imported or updated with this function call), "ids" (a vector of the _id[s] of these trials) and the "queryterm" used, with several attributes set (database connection details and a data frame of the query history in this database).

**Examples**

```r
## Not run:

db <- nodbi::src_sqlite(
  collection = "test"
)

# Retrieve protocol-related information on a
# single trial identified by EudraCT number
ctrLoadQueryIntoDb(
  queryterm = "2013-001291-38", con = db
)

# Retrieve protocol-related information on
# ongoing interventional cancer trials in children
ctrLoadQueryIntoDb(
  queryterm = "cancer&recr=Open&type=Intr&age=0",
  register = "CTGOV",
  con = db
)
```
ctrOpenSearchPagesInBrowser

Open advanced search pages of register(s) or execute search in browser

Description
Open advanced search pages of register(s) or execute search in browser

Usage
ctrOpenSearchPagesInBrowser(url = "", register = "", copyright = FALSE, ...)

Arguments

url of search results page to show in the browser. May be the output of ctrGetQueryUrl or from dbQueryHistory.

register Register(s) to open. Either "EUCTR" or "CTGOV" or a vector of both. Default is to open both registers’ advanced search pages. To open the browser with a previous search, the output of ctrGetQueryUrl or one row from dbQueryHistory can be used.

copyright (Optional) If set to TRUE, opens copyright pages of register(s).

... May include the deprecated input parameter.

Value
Is always true, invisibly.

Examples
## Not run:
ctrOpenSearchPagesInBrowser(
  "https://www.clinicaltrialsregister.eu/ctr-search/search?query=cancer"
)

# for this example, the clipboard has to
# contain the URL from a search in a register
ctrOpenSearchPagesInBrowser(
  ctrGetQueryUrl()
)

# open the last query that was
# loaded into the database
db <- nodbi::src_sqlite(
  collection = "previously_created"
dbFindFields

Find names of fields in the database collection

Description

Given part of the name of a field of interest to the user, this function returns the full field names as found in the database.

Usage

dbFindFields(namepart = "", con, verbose = FALSE)

Arguments

namepart A plain string (can include a regular expression, including Perl-style) to be searched for among all field names (keys) in the database.
con A src connection object, as obtained with nodbi::src_mongo() or nodbi::src_sqlite()
verbose If TRUE, prints additional information (default FALSE).

Details

For fields in EUCTR (protocol- and results-related information), see also the register’s documentation at https://eudract.ema.europa.eu/.

For fields in CTGOV (protocol-related information), see also the register’s definitions at https://prsinfo.clinicaltrials.gov/definitions.html.

Note: Generating a list of fields with this function may take some time, and may involve running a mapreduce function if using a MongoDB server. If the user is not not authorized to run such a function, random documents are sampled to generate a list of fields.

Value

Vector of names of found field(s) in alphabetical order (that is, not by register or field frequency)
Examples

```r
## Not run:
db <- nodbi::src_sqlite(
  collection = "my_collection"
)
dbFindFields(
  namepart = "date",
  con = db
)
## End(Not run)
```

dbFindIdsUniqueTrials  Deduplicate records to provide unique clinical trial identifiers

Description

If records for a clinical trial are found from more than one register, the record from EUCTR is returned. The function currently relies on CTGOV recording other identifiers such as the EudraCT number in the field "Other IDs".

Usage

```r
dbFindIdsUniqueTrials(
  preferregister = c("EUCTR", "CTGOV", "ISRCTN"),
  prefermemberstate = "GB",
  include3rdcountrytrials = TRUE,
  con,
  verbose = TRUE
)
```

Arguments

- `preferregister`  A vector of the sequence of preference for registers from which to generate unique _id’s, default c("EUCTR", "CTGOV", "ISRCTN")
- `prefermemberstate`  Code of single EU Member State for which records should returned. If not available, a record for GB or lacking this, any other record for the trial will be returned. For a list of codes of EU Member States, please see vector countriesEUCTR. Alternatively, "3RD" will lead to return a Third Country record of a trial, if available.
- `include3rdcountrytrials`  A logical value if trials should be retained that are conducted exclusively in third countries, that is, outside the European Union.
- `con`  A src connection object, as obtained with nodbi::src_mongo() or nodbi::src_sqlite()
- `verbose`  If set to TRUE, prints out information about numbers of records found at subsequent steps when searching for duplicates
dbGetFieldsIntoDf

Create data frame by extracting specified fields from database collection

Description

With this convenience function, fields in the database are retrieved into an R data frame. Note that fields within the record of a trial can be hierarchical and structured, that is, nested.

Usage

dbGetFieldsIntoDf(fields = "", con, verbose = FALSE, stopifnodata = TRUE)

Arguments

fields Vector of one or more strings, with names of the sought fields. See function dbFindFields for how to find names of fields. Regular expressions are possible. "item.subitem" notation is supported.
con A src connection object, as obtained with nodbi::src_mongo() or nodbi::src_sqlite()
verbose Printing additional information if set to TRUE; default is FALSE.
stopifnodata Stops with an error (TRUE, default) or with a warning (FALSE) if the sought field is empty in all, or not available in any of the records in the database collection.

Details

With both src_sqlite and src_mongo, the function returns a list of data for a field that includes nested content; use function dfTrials2Long followed by dfName2Value to extract desired nested variables. For more sophisticated data retrieval from the database, see vignette examples and other packages to query mongodb such as mongolite.
**Value**

A data frame with columns corresponding to the sought fields. Note: a column for the record _id will always be included. Each column can be either a simple data type (numeric, character, date) or a list (see example below): For complicated lists, use function `dfTrials2Long` followed by function `dfName2Value` to extract values for nested variables. The maximum number of rows of the returned data frame is equal to, or less than the number of records of trials in the database.

**Examples**

```r
## Not run:
db <- nodbi::src_sqlite(
  collection = "my_collection"
)

# access fields that are nested within another field
# and can have multiple values with the other field
getFieldsIntoDf(
  "b1_sponsor.b31_and_b32_status_of_the_sponsor",
  con = db
)[1,]
# _id b1_sponsor.b31_and_b32_status_of_the_sponsor
# 1 2004-000015-25-GB Non-commercial / Commercial

# access fields that include a list of values
# which are printed as comma separated values
getFieldsIntoDf(
  "keyword",
  con = db
)[1,]
# _id keyword
# 1 NCT00129259 T1D, type 1 diabetes, juvenile diabetes

str(.Last.value)
```

## End(Not run)

---

**dbQueryHistory**

Show the history of queries that were loaded into a database

**Description**

Show the history of queries that were loaded into a database
dfListExtractKey

**Usage**

\[\text{dbQueryHistory}(\text{con, \text{verbose = FALSE}})\]

**Arguments**

- **con**: A src connection object, as obtained with `nodbi::src_mongo()` or `nodbi::src_sqlite()`
- **verbose**: If TRUE, prints additional information (default FALSE).

**Value**

A data frame with columns: query-timestamp, query-register, query-records (note: this is the number of records loaded when last executing `ctrLoadQueryIntoDb`, not the total record number) and query-term, and with one row for each `ctrLoadQueryIntoDb` loading trial records in this collection.

**Examples**

```r
## Not run:
da <- nodbi::src_sqlite(
  collection = "my_collection"
)
dbQueryHistory(
  con = da
)
## End(Not run)
```

dfListExtractKey

**Extract named element(s) from list(s) into long-format data frame**

**Description**

The function uses a name (key) to extract an element from a list in a data.frame such as obtained with `dbGetFieldsIntoDf`. This helps to simplify working with nested lists and with complex structures.

**Usage**

\[\text{dfListExtractKey}(\text{df, \text{list.key = list(c("endPoints.endPoint", ",^title"))})\]

**Arguments**

- **df**: A data frame
- **list.key**: A list of pairs of list names and key names, where the list name corresponds to the name of a column in df that holds a list and the name of the key identifies the element to be extracted. See example.
dfMergeTwoVariablesRelevel

Value

A data frame in long format with columns name (identifying the full path in the data frame, "<list>.<key>", _id (of the trial record), value (of name per _id), item (number of value of name per _id).

Examples

```r
## Not run:
db <- nodbi::src_sqlite(
  collection = "my_collection"
)
df <- dbGetFieldsIntoDf(
  fields = c(
    "endPoints.endPoint",
    "subjectDisposition.postAssignmentPeriods"),
  con = db
)
dfListExtractKey(
  df = df,
  list.key = list(
    c("endPoints.endPoint", "title"),
    c("subjectDisposition.postAssignmentPeriods", "arms.arm.type.value")
  )
)
## End(Not run)
```

---

**dfMergeTwoVariablesRelevel**

*Merge two variables into one, optionally map values to new levels*

Description

Merge two variables into one, optionally map values to new levels

Usage

```r
dfMergeTwoVariablesRelevel(df = NULL, colnames = "", levellist = NULL, ...)
```

Arguments

- **df** A data frame in which there are two variables (columns) to be merged into one.
- **colnames** A vector of length two with names of the two columns that hold the variables to be merged. See colnames for how to obtain the names of columns of a data frame.
levelslist: A list with one slice each for a new value to be used for a vector of old values (optional).

... for deprecated varnames parameter (will be removed)

Value

A vector of strings

Examples

```r
## Not run:
statusvalues <- list(
  "ongoing" = c("Recruiting", "Active", "Ongoing",
                 "Active, not recruiting", "Enrolling by invitation"),
  "completed" = c("Completed", "Prematurely Ended", "Terminated"),
  "other" = c("Withdrawn", "Suspended",
               "No longer available", "Not yet recruiting"))

df$Recruitment
```

```r
dfMergeTwoVariablesRelevel(
  df = result,
  colnames = c("Recruitment", "x5_trial_status"),
  levelslist = statusvalues)
## End(Not run)
```

---

**dfName2Value**

*Extract information of interest (e.g., endpoint) from long data frame of protocol- or result-related trial information as returned by dfTrials2Long*

**Description**

Extract information of interest (e.g., endpoint) from long data frame of protocol- or result-related trial information as returned by dfTrials2Long

**Usage**

dfName2Value(df, valuename = "", wherename = "", wherevalue = "")

**Arguments**

- **df**: A data frame with four columns (`_id`, `identifier`, `name`, `value`) as returned by `dfTrials2Long`
- **valuename**: A character string for the name of the variable from which to extract information for the variable of interest
- **wherename**: A character string to identify the variable of interest
- **wherevalue**: A character string with the value of interest for the variable of interest
Value

A data frame with columns _id, identifier, name, value that only includes the values of interest, where value are strings unless all value elements are numbers.

Examples

```r
## Not run:
db <- nodbi::src_sqlite(
  collection = "my_collection"
)
df <- ctrdata::dbGetFieldsIntoDf(
  fields = c(
    # ctgov - typical results fields
    "clinical_results.baseline.analyzed_list.analyzed.count_list.count",
    "clinical_results.baseline.group_list.group",
    "clinical_results.baseline.analyzed_list.analyzed.units",
    "clinical_results.outcome_list.outcome",
    "study_design_info.allocation",
    # euctr - typical results fields
    "trialInformation.fullTitle",
    "subjectDispension.recruitmentDetails",
    "baselineCharacteristics.baselineReportingGroups.baselineReportingGroup",
    "endPoints.endPoint",
    "trialChanges.hasGlobalInterruptions",
    "subjectAnalysisSets",
    "adverseEvents.seriousAdverseEvents.seriousAdverseEvent"
  ), con = dbc
)
# convert to long
reslong <- ctrdata::dfTrials2Long(
  df = df
)
# get values for endpoint of interest, duration of response
ctrdata::dfValue2Name(
  df = df,
  valuename = paste0(
    "endPoints.endPoint.*armReportingGroup.tendencyValues.tendencyValue.value|",
    "clinical_results.*category.measurement_list.measurement.value|",
    "clinical_results.outcome.measure.units|endPoints.endPoint.unit"
  ),
  wherename = "clinical_results.*outcome.measure.title|endPoints.endPoint.title",
  wherevalue = "duration of response"
)
## End(Not run)
```

---

dfTrials2Long  
Extract trial information into long format
installCygwinWindowsDoInstall

Convenience function to install a minimal cygwin environment under MS Windows, including perl, sed and php

Description

Alternatively and in case of difficulties, download and run the cygwin setup yourself as follows:


installCygwinWindowsDoInstall

Description

The function works with protocol- and results- related information. It converts lists and other values into individual rows of a long data frame. From the resulting data frame, values of interest can then be selected (e.g. select an outcome and its analysis by the identifier of the measure which has "Hazard Ratio" in its name, see dfName2Value).

Usage

dfTrials2Long(df)

Arguments

df       Data frame with columns including the trial identifier (_id) and one or more variables as obtained from dbGetFieldsIntoDf

Value

A data frame with the four columns: _id, identifier, name, value

Examples

```r
## Not run:
db <- nodbi::src_sqlite(
  collection = "my_collection"
)
df <- dbGetFieldsIntoDf(
  fields = c(  
    "clinical_results.outcome_list.outcome"),
  con = db
)
dfTrials2Long(
  df = df
)
## End(Not run)
```
installCygwinWindowsDoInstall

Usage

installCygwinWindowsDoInstall(force = FALSE, proxy = "")

Arguments

force Set to TRUE to force updating and overwriting an existing installation in c:\cygwin

proxy Specify any proxy to be used for downloading via http, e.g. "host_or_ip:port". installCygwinWindowsDoInstall may detect and use the proxy configuration used in MS Windows to use an automatic proxy configuration script. Authenticated proxies are not supported at this time.
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