# Package 'easyPubMed'

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Title Search and Retrieve Scientific Publication Records from PubMed
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<b>Description</b> Query NCBI Entrez and retrieve PubMed records in XML or text format. Process PubMed records by extracting and aggregating data from selected fields. A large number of records can be easily downloaded via this simple-to-use interface to the NCBI PubMed API.
<pre>URL https://www.data-pulse.com/dev_site/easypubmed/</pre>
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R topics documented:
articles_to_list       2         article_to_df       3         batch_pubmed_download       5         custom_grep       6         EPMsamples       8         fetch_all_pubmed_ids       9         fetch_pubmed_data       10

2 articles\_to\_list

umi_address	18
get_pubmed_ids	<ul><li>. 13</li><li>. 14</li><li>. 15</li></ul>

articles\_to\_list

Cast PubMed Data into a List of Articles

# **Description**

Convert an XML object of PubMed records into a list of strings (character vector of length 1) corresponding to individual PubMed articles. PubMed records are identified by a "/PubmedArticle" XML tag. This automatically casts all the content of each PubMed record to a character-class object without removing XML tags.

# Usage

```
articles_to_list(pubmed_data, encoding = "UTF8", simplify = TRUE)
```

# **Arguments**

pubmed_data	String corresponding to the name of an XML file (typically, the result of a batch_pubmed_download() call). Alternatively, a string including PubMed records with XML tags, such as the object returned by a fetch_pubmed_data() call.
encoding	The encoding of an input/output connection can be specified by name (for example, "ASCII", or "UTF-8", in the same way as it would be given to the function base::iconv(). See iconv() help page for how to find out more about encodings that can be used on your platform. "UTF-8" is recommended.
simplify	Logical; should the result be simplified to a character vector. If FALSE, results are returned as a list.

# **Details**

The input is an XML object or a string including PubMed records (with XML tags). These are the output of easyPubMed functions: fetch\_pubmed\_data() or batch\_pubmed\_download(). The function returns a list or a character vector where each element is a different PubMed record.

## Value

List or character vector including all the records from the original XML object in text format. Elements in the list are not named and are only accessible via their numeric index.

# Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

article\_to\_df 3

#### References

https://www.data-pulse.com/dev\_site/easypubmed/

#### **Examples**

```
try({
    ## Retrieve PubMed data and return a list ot articles
    my_query <- "Damiano Fantini[AU]"
    my_query <- get_pubmed_ids(pubmed_query_string = my_query)
    my_data <- fetch_pubmed_data(my_query, encoding = "ASCII")
    listed_articles <- articles_to_list(my_data)
    custom_grep(listed_articles[[2]], "ArticleTitle", "char")
}, silent = TRUE)

## Not run:
## Download PubMed data and return a list ot articles
dami_query <- "Damiano Fantini[AU] AND 2018[PDAT]"
outfile <- batch_pubmed_download(dami_query, dest_file_prefix = "easyPM_ex001_")
listed_articles <- articles_to_list(pubmed_data = outfile)
custom_grep(listed_articles[[2]], "ArticleTitle", "char")

## End(Not run)</pre>
```

article\_to\_df

Extract Data from a PubMed Record

# Description

Extract publication-specific information from a PubMed record driven by XML tags. The input record is a string (character-class vector of length 1) and includes PubMed-specific XML tags. Data are returned as a data frame where each row corresponds to one of the authors of the PubMed article.

# Usage

#### **Arguments**

pubmedArticle String including one PubMed record.

autofill Logical. If TRUE, missing affiliations are automatically imputed based on other

non-NA addresses from the same record.

max\_chars Numeric (integer). Maximum number of characters to be extracted from the

Article Abstract field. Set max\_chars to -1 for extracting the full-length abstract.

Set max\_chars to 0 to extract no abstract.

4 article\_to\_df

getKeywords Logical. If TRUE, an attempt to extract article Keywords will be made.

Logical. If FALSE, author information won't be extracted. This will considerably speed up the operation.

#### **Details**

Given one Pubmed Article record, this function will automatically extract a set of features. Extracted information include: PMID, DOI, article title, article abstract, publication date (year, month, day), journal name (title, abbreviation), keywords, and a set of author-specific info (names, affiliation, email address). Each row of the output data frame corresponds to one of the authors of the PubMed record. Author-independent info (publication ID, title, journal, date) are identical across all rows. If information about authors are not required, set 'getAuthors' = TRUE.

#### Value

Data frame including the extracted features. Each row correspond a different author.

#### Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

### References

https://www.data-pulse.com/dev\_site/easypubmed/

```
try({
  ## Display some contents
 data("EPMsamples")
 #display Query String used for collecting the data
 print(EPMsamples$NUBL_1618$qry_st)
 #Get records
 BL_list <- EPMsamples$NUBL_1618$rec_lst
 cat(BL_list[[1]])
 # cast PM recort to data.frame
 BL_df <- article_to_df(BL_list[[1]], max_chars = 0)
 print(BL_df)
}, silent = TRUE)
## Not run:
## Query PubMed, retrieve a selected citation and format it as a data frame
dami_query <- "Damiano Fantini[AU] AND 2017[PDAT]"</pre>
dami_on_pubmed <- get_pubmed_ids(dami_query)</pre>
dami_abstracts_xml <- fetch_pubmed_data(dami_on_pubmed)</pre>
dami_abstracts_list <- articles_to_list(dami_abstracts_xml)</pre>
article_to_df(pubmedArticle = dami_abstracts_list[[1]], autofill = FALSE)
article_to_df(pubmedArticle = dami_abstracts_list[[2]], autofill = TRUE, max_chars = 300)[1:2,]
## End(Not run)
```

## **Description**

Performs a PubMed Query (via the get\_pubmed\_ids() function), downloads the resulting data (via multiple fetch\_pubmed\_data() calls) and then saves data in a series of xml or txt files on the local drive. The function is suitable for downloading a very large number of records.

# Usage

#### **Arguments**

pubmed\_query\_string

String (character-vector of length 1): this is the string used for querying PubMed (the standard PubMed Query synthey applies)

(the standard PubMed Query synthax applies).

dest\_dir String (character-vector of length 1): this string corresponds to the name of the

existing folder where files will be saved. Existing files will be overwritten. If

NULL, the current working directory will be used.

dest\_file\_prefix

String (character-vector of length 1): this string is used as prefix for the files that

are written locally.

format String (character-vector of length 1): data will be requested from Entrez in this

format. Acceptable values are: c("medline","uilist","abstract","asn.1", "xml").

When format != "xml", data will be saved as text files (txt).

api\_key String (character vector of length 1): user-specific API key to increase the limit

of queries per second. You can obtain your key from NCBI.

batch\_size Integer (1 < batch\_size < 5000): maximum number of records to be saved in a

single xml or txt file.

res\_cn Integer (> 0): numeric index of the data batch to start downloading from. This

parameter is useful to resume an incomplete download job after a system crash.

encoding The encoding of an input/output connection can be specified by name (for exam-

ple, "ASCII", or "UTF-8", in the same way as it would be given to the function base::iconv(). See iconv() help page for how to find out more about encodings

that can be used on your platform. Here, we recommend using "UTF-8".

6 custom\_grep

#### **Details**

Download large number of PubMed records as a set of xml or txt files that are saved in the folder specified by the user. This function enforces data integrity. If a batch of downloaded data is corrupted, it is discarded and downloaded again. Each download cycle is monitored until the download job is successfully completed. This function should allow to download a whole copy of PubMed, if desired. The function informs the user about the current progress by constantly printing to console the number of batches still in queue for download. pubmed\_query\_string accepts standard PubMed synthax. The function will query PubMed multiple times using the same query string. Therefore, it is recommended to use a [EDAT] or a [PDAT] filter in the query if you want to ensure reproducible results.

#### Value

Character vector including the names of files downloaded to the local system

### Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

#### References

```
https://www.data-pulse.com/dev_site/easypubmed/
```

# **Examples**

```
## Not run:
## Example 01: retrieve data from PubMed and save as XML file
ml_query <- "Machine Learning[TI] AND 2016[PD]"
out1 <- batch_pubmed_download(pubmed_query_string = ml_query, batch_size = 180)
readLines(out1[1])[1:30]
##
## Example 02: retrieve data from PubMed and save as TXT file
ml_query <- "Machine Learning[TI] AND 2016[PD]"
out2 <- batch_pubmed_download(pubmed_query_string = ml_query, batch_size = 180, format = "medline")
readLines(out2[1])[1:30]
## End(Not run)</pre>
```

custom\_grep

Retrieve Text Between XML Tags

# **Description**

Extract text form a string containing XML or HTML tags. Text included between tags of interest will be returned. If multiple tagged substrings are found, they will be returned as different elements of a list or character vector.

custom\_grep 7

## Usage

```
custom_grep(xml_data, tag, format = "list")
```

# **Arguments**

xml\_data String (of class character and length 1): corresponds to the PubMed record or

any string including XML/HTML tags.

tag String (of class character and length 1): the tag of interest (does NOT include <

> chars).

format c("list", "char"): specifies the format for the output.

#### **Details**

The input string has to be a character string (length 1) containing tags (HTML or XML format). If an XML Document is provided as input, the function will rise an error.

#### Value

List or vector where each element corresponds to an in-tag substring.

# Author(s)

Damiano Fantini <damiano .fantini@gmail.com>

# References

```
https://www.data-pulse.com/dev_site/easypubmed/
```

```
try({
    ## extract substrings based on regular expressions
    string_01 <- "I can't wait to watch the <strong>Late Night Show with"
    string_01 <- paste(string_01, "Seth Meyers</strong> tonight at <strong>11:30</strong>pm CT!")
    print(string_01)
    custom_grep(xml_data = string_01, tag = "strong", format = "char")
    custom_grep(xml_data = string_01, tag = "strong", format = "list")
}, silent = TRUE)
```

8 EPMsamples

**EPMsamples** 

PubMed Records downloaded and analyzed via easyPubMed

# **Description**

This dataset includes a collection of 4 examples showing how to download and analyze records from PubMed by using easyPubMed. Each element in the EPMsamples list corresponds to a different query and/or analysis. Also, each element of EPMsamples is a list including intermediates and notes about the analysis.

# Usage

```
data("EPMsamples")
```

#### **Format**

The dataset is formatted as a list including 4 elements:

```
* 'DF_papers_abs': List of 4

* 'DF_papers_std': List of 4

* 'NUBL_dw18': List of 3

* 'NUBL_1618': List of 5
```

## **Details**

The dataset was built as described in this vignette: https://www.data-pulse.com/projects/Rlibs/vignettes/building\_the\_easyPubMed\_EPMsamples\_dataset.html

```
## Display some contents
data("EPMsamples")
# The following examples are focused on example query #4 (i.e., NUBL_1618)
# Display Query String used for collecting the data
print(EPMsamples$NUBL_1618$qry_st)
# show one PubMed record element from the IL vector
NU_records <- EPMsamples$NUBL_1618$rec_lst
cat(NU_records[[1]])
# cast PM recort to data.frame
BL_df <- article_to_df(NU_records[[6]], max_chars = 0)
print(BL_df)</pre>
```

fetch\_all\_pubmed\_ids 9

# **Description**

Retrieve PubMed record identifiers from Entrez following a search performed via the get\_pubmed\_ids() function. Identifiers are returned as a character vector.

# Usage

```
fetch_all_pubmed_ids(pubmed_id_list)
```

# **Arguments**

```
pubmed_id_list List: the result of a get_pubmed_ids() call.
```

#### **Details**

Retrieve PubMed identifiers, without any other information (such as article title, authors, publication date, and so on). The PubMed IDs can be stored or used with other software.

#### Value

Character vector including all PMID (PubMed Identifiers) returned by the current query.

#### Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

# References

```
https://www.data-pulse.com/dev_site/easypubmed/
```

```
## Not run:
## Fetch only PubMed Record IDs (PMIDs)
dami_query_string <- "Damiano Fantini[AU]"
dami_on_pubmed <- get_pubmed_ids(dami_query_string)
dami_pmids <- fetch_all_pubmed_ids(dami_on_pubmed)
print(dami_pmids)
## End(Not run)</pre>
```

10 fetch\_pubmed\_data

fetch\_pubmed\_data

Retrieve PubMed Data in XML or TXT Format

## Description

Retrieve PubMed records from Entrez following a search performed via the get\_pubmed\_ids() function. Data are downloaded in the XML or TXT format and are retrieved in batches of up to 5000 records.

#### **Usage**

#### **Arguments**

pubmed\_id\_list List: the result of a get\_pubmed\_ids() call.

retstart Integer (>=0): index of the first UID in the retrieved PubMed Search Result set

to be included in the output (default=0, corresponding to the first record of the

entire set).

retmax Integer (>=1): size of the batch of PubMed records to be retrieved at one time.

format Character: element specifying the output format. The following values are al-

lowed: c("asn.1", "xml", "medline", "uilist", "abstract").

encoding The encoding of an input/output connection can be specified by name (for exam-

ple, "ASCII", or "UTF-8", in the same way as it would be given to the function base::iconv(). See iconv() help page for how to find out more about encodings

that can be used on your platform. Here, we recommend using "UTF-8".

#### **Details**

Retrieve PubMed records based on the results of a get\_pubmed\_ids() query. Records are retrieved from Entrez via the PubMed API efetch function. The first entry to be retrieved may be adjusted via the retastart parameter (this allows the user to download large batches of PubMed data in multiple runs). The maximum number of entries to be retrieved can also be set adjusting the retmax parameter (1 < retmax < 5000). Data will be downloaded on the fly (no files are saved locally).

# Value

An object (vector) of class "character". If format is set to "xml" (default), a single String including all PubMed records (with XML tags embedded) is returned. If a different format is selected, a vector of strings is returned, where each row corresponds to a line of the output document.

get\_pubmed\_ids 11

## Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

#### References

https://www.data-pulse.com/dev\_site/easypubmed/https://www.ncbi.nlm.nih.gov/books/NBK25499/table/chapter4.T.\_valid\_values\_of\_\_retmode\_and/

# **Examples**

```
try({
 ## Example 01: retrieve data in TXT format
 library("easyPubMed")
 dami_query_string <- "Damiano Fantini[AU] AND 2018[PDAT]"</pre>
 dami_on_pubmed <- get_pubmed_ids(dami_query_string)</pre>
 Sys.sleep(1) # avoid server timeout
 dami_papers <- fetch_pubmed_data(dami_on_pubmed, format = "abstract")</pre>
 dami_papers[dami_papers == ""] <- "\n"
 cat(paste(dami_papers[1:65], collapse = ""))
}, silent = TRUE)
## Not run:
## Example 02: retrieve data in XML format
library("easyPubMed")
dami_query_string <- "Damiano Fantini[AU]"</pre>
dami_on_pubmed <- get_pubmed_ids(dami_query_string)</pre>
dami_papers <- fetch_pubmed_data(dami_on_pubmed)</pre>
titles <- custom_grep(dami_papers, "ArticleTitle", "char")
print(titles)
## End(Not run)
```

get\_pubmed\_ids

Simple PubMed Record Search

# **Description**

Query PubMed (Entrez) in a simple way via the PubMed API eSearch function. Calling this function results in posting the query results on the PubMed History Server. This allows later access to the resulting data via the fetch\_pubmed\_data() function, or other easyPubMed functions.

#### Usage

```
get_pubmed_ids(pubmed_query_string, api_key = NULL)
```

12 get\_pubmed\_ids

#### **Arguments**

```
pubmed_query_string
is a string (character vector of length 1) that is used for querying PubMed (standard PubMed synthax, see reference for details).

api_key

String (character vector of length 1): user-specific API key to increase the limit
```

of queries per second. You can obtain your key from NCBI.

#### **Details**

This function will use the String provided as argument for querying PubMed via the eSearch function of the PubMed API. The Query Term can include one or multiple words, as well as the standard PubMed operators (AND, OR, NOT) and tags (i.e., [AU], [PDAT], [Affiliation], and so on). ESearch will post the UIDs resulting from the search operation onto the History server so that they can be used directly in a subsequent fetchPubmedData() call.

## Value

The function returns a list. The list includes the number of records found on PubMed and the first 20 PubMed IDs (UID) retrieved by the query. The list also includes QueryKey and WebEnv that are required for a subsequent fetch\_pubmed\_data() call.

## Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

## References

https://www.data-pulse.com/dev\_site/easypubmed/https://www.ncbi.nlm.nih.gov/books/NBK3827/#\_pubmedhelp\_Search\_Field\_Descriptions\_and\_

```
try({
    ## Search for scientific articles written by Damiano Fantini
    ## and print the number of retrieved records to screen.
    ## Also print the retrieved UIDs to screen.
    ##
    dami_on_pubmed <- get_pubmed_ids("Damiano Fantini[AU]")
    print(dami_on_pubmed$Count)
    print(unlist(dami_on_pubmed$IdList))
}, silent = TRUE)</pre>
```

```
get_pubmed_ids_by_fulltitle
```

Simple PubMed Record Search by Full-length Title

## **Description**

Query PubMed (Entrez) in a simple way via the PubMed API eSearch function. This function is designed to query PubMed using a full-length publication title as query string. It performs stopword removal from the query string before querying the PubMed server. Calling this function results in posting the results on the PubMed History Server. This allows later access to the resulting data via the fetch\_pubmed\_data() function, or other easyPubMed functions.

#### Usage

```
get_pubmed_ids_by_fulltitle(fulltitle, field = "[Title]", api_key = NULL)
```

# **Arguments**

fulltitle	String (character vector of length 1) that corresponds to the full-length publication title used for querying PubMed (titles should be used as is, without adding extra filters/tags).
field	String (character vector of length 1) with a tag indicating the PubMed record field where the full-length string (fulltitle) should be searched in. By default, this points to the 'Title' field. This field can be changed (use fields supported by PubMed) as required by the user (for example, to attempt an exact-match query using a specific sentence included in the abstract of a record).
api_key	String (character vector of length 1): user-specific API key to increase the limit of queries per second. You can obtain your key from NCBI.

#### **Details**

This function will use the String provided as argument for querying PubMed via the eSearch function of the PubMed API. The Query Term should include a full-length publication title, without other PubMed operators (AND, OR, NOT) nor tags (i.e., [AU], [PDAT], [Affiliation], and so on). ESearch will post the UIDs resulting from the search operation onto the History server so that they can be used directly in a subsequent fetchPubmedData() call.

# Value

The function returns a list. The list includes the number of records found on PubMed and the first 20 PubMed IDs (UID) retrieved by the query. The list also includes QueryKey and WebEnv that are required for a subsequent fetch\_pubmed\_data() call.

# Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

PubMed\_stopwords

## References

https://www.data-pulse.com/dev\_site/easypubmed/

# **Examples**

```
## Not run:
## Search for a scientific article matching a full-length title
my_query <- "Body mass index and cancer risk among Chinese patients with type 2 diabetes mellitus"
my_field <- "[Title]"
# Full-length title query (designed to query titles)
res0 <- get_pubmed_ids(my_query)
print(as.numeric(res0$Count))
# Weird count!
res <- get_pubmed_ids_by_fulltitle(my_query, field = my_field)
# Num results = 1 as expected
print(as.numeric(res$Count))
## End(Not run)</pre>
## End(Not run)
```

PubMed\_stopwords

PubMed Records about Bladder Research from Northwestern University

### **Description**

This dataset includes a collection of 87 PubMed Records of scientific publications about Bladder biology and pathology, published by clinical and research groups from Northwestern University (Chicago, IL), between 2016 and 2018.

# Usage

```
data("PubMed_stopwords")
```

#### **Format**

A character vector including all PubMed stopwords tat are typically filtered out from queries.

# **Details**

Number of stopwords included, n=133.

```
## Display some contents
data("PubMed_stopwords")
head(PubMed_stopwords)
```

table\_articles\_byAuth 15

table\_articles\_byAuth Extract Publication and Affiliation Data from PubMed Records

# Description

Extract Publication Info from PubMed records and cast data into a data.frame where each row corresponds to a different author. It is possible to limit data extraction to first authors or last authors only, or get information about all authors of each PubMed record.

# Usage

```
table_articles_byAuth(pubmed_data,
                              included_authors = "all",
                              max_chars = 500,
                              autofill = TRUE,
                              dest_file = NULL,
                              getKeywords = TRUE,
                              encoding = "UTF8")
```

# Arg

guments		
pubmed_data	PubMed Data in XML format: typically, an XML file resulting from a batch_pubmed_download() call or an XML object, result of a fetch_pubmed_data() call.	
included_authors		
	Character: c("first", "last", "all"). Only includes information from the first, the last or all authors of a PubMed record.	
max_chars	Numeric: maximum number of chars to extract from the AbstractText field.	
autofill	Logical. If TRUE, missing affiliations are imputed according to the available values (from the same article).	
dest_file	String (character of length 1). Name of the file that will be written for storing the output. If NULL, no file will be saved.	
getKeywords	Logical. If TRUE, the operation will attempt to extract PubMed record keywords (MESH topics, keywords).	
encoding	The encoding of an input/output connection can be specified by name (for example, "ASCII", or "UTF-8", in the same way as it would be given to the function base::iconv(). See iconv() help page for how to find out more about encodings that can be used on your platform. Here, we recommend using "UTF-8".	

## **Details**

Retrieve publication and author information from PubMed data, and cast them as a data.frame.

### Value

```
Data frame including the following fields: c("article.title", "article.abstract", "date.year", "date.month",
"date.day", "journal.abbrv", "journal.title", "keywords", "auth.last", "auth.fore", "auth.address", "auth.email").
```

16 trim\_address

## Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

## References

```
https://www.data-pulse.com/dev_site/easypubmed/
```

# **Examples**

```
## Not run:
## Cast PubMed record info into a data.frame
dami_query <- "Damiano Fantini[AU]"</pre>
dami_on_pubmed <- get_pubmed_ids(dami_query)</pre>
dami_abstracts_xml <- fetch_pubmed_data(dami_on_pubmed, encoding = "ASCII")</pre>
xx <- table_articles_byAuth(pubmed_data = dami_abstracts_xml,</pre>
                              included_authors = "first",
                              max_chars = 100,
                              autofill = TRUE)
print(xx[1:5, c("pmid", "lastname", "jabbrv")])
## Download records first
## Also, auto-fill disabled
dami_query <- "Damiano Fantini[AU]"</pre>
curr.file <- batch_pubmed_download(dami_query, dest_file_prefix = "test_bpd_", encoding = "ASCII")</pre>
xx <- table_articles_byAuth(pubmed_data = curr.file[1],</pre>
                              included_authors = "all",
                              max_chars = 20,
                              autofill = FALSE)
print(xx[1:5, c("pmid", "lastname", "jabbrv")])
## End(Not run)
```

trim\_address

Trim and Format Address Information

## **Description**

Set of rules for trimming and standardizing the format of address information retrieved from PubMed records. Affiliations including more than one address will be trimmend and only the first address will be returned.

# Usage

```
trim_address(addr)
```

trim\_address 17

# **Arguments**

addr

Character string including an address as extracted from PubMed records.

## Value

Character string including a formatted and trimmed address (if available).

# Author(s)

Damiano Fantini <damiano.fantini@gmail.com>

#### References

```
https://www.data-pulse.com/dev_site/easypubmed/
```

```
addr_string <- " 2 Dept of Urology, Feinberg School of Medicine,"
addr_string <- paste(addr_string, "Chicago, US; Dept of Mol Bio as well...")
print(addr_string)
print(trim_address(addr = addr_string))</pre>
```

# **Index**

```
* datasets
    EPMsamples, 8
    PubMed_stopwords, 14

article_to_df, 3
articles_to_list, 2

batch_pubmed_download, 5

custom_grep, 6

EPMsamples, 8

fetch_all_pubmed_ids, 9
fetch_pubmed_ids, 11
get_pubmed_ids, 11
get_pubmed_ids_by_fulltitle, 13

PubMed_stopwords, 14

table_articles_byAuth, 15
trim_address, 16
```