

Package ‘puremoe’

May 9, 2026

Type Package

Title Pubmed Unified REtrieval for Multi-Output Exploration

Version 1.0.4

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Description Access a variety of 'PubMed' data through a single, user-friendly interface, including abstracts, bibliometrics from 'iCite', pubtations from 'PubTator3', and full-text records from 'PMC'.

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Encoding UTF-8

LazyData true

LazyDataCompression xz

Depends R (>= 3.5)

Imports rentrez, textshape, xml2, data.table, httr, pbapply, jsonlite, rappdirs

Suggests knitr, rmarkdown, DT, dplyr

VignetteBuilder knitr

RoxygenNote 7.3.3

URL <https://github.com/jaytimm/puremoe>,
<https://jaytimm.github.io/puremoe/>

BugReports <https://github.com/jaytimm/puremoe/issues>

NeedsCompilation no

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Repository CRAN

Date/Publication 2026-04-21 17:02:08 UTC

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data_mesh_frequencies *MeSH Descriptor Frequencies Across PubMed*

Description

Baseline frequencies for MeSH descriptors computed from a local PostgreSQL mirror of PubMed (April 2026). For each descriptor, counts reflect the number of distinct PMIDs indexed with that term; proportions use the full PubMed corpus of 39,703,112 PMIDs as denominator. Descriptor UI and canonical name are joined from the NLM MeSH thesaurus. Intended as a baseline for MeSH term enrichment analyses against arbitrary PubMed subsets.

Usage

data_mesh_frequencies

Format

A data.table with 30,521 rows and 4 columns:

DescriptorUI MeSH descriptor unique identifier (e.g., D000001)

DescriptorName Canonical MeSH descriptor name

n_pmid Number of distinct PubMed records indexed with this descriptor

prop_total Proportion of all 39,703,112 PubMed PMIDs indexed with this descriptor

Source

Computed from mesh_descriptor table in a local PubMed PostgreSQL mirror; descriptor meta-data from the NLM MeSH Thesaurus (April 2026).

data_mesh_thesaurus *Download and Combine 'MeSH' and Supplemental Thesauruses*

Description

This function downloads and combines the 'MeSH' (Medical Subject Headings) Thesaurus and a supplemental concept thesaurus. The data is sourced from specified URLs and stored locally for subsequent use. By default, the data is stored in a temporary directory. Users can opt into persistent storage by setting 'use_persistent_storage' to TRUE and optionally specifying a path.

Usage

```
data_mesh_thesaurus(  
  path = NULL,  
  use_persistent_storage = FALSE,  
  force_install = FALSE  
)
```

Arguments

path	A character string specifying the directory path where data should be stored. If not provided and persistent storage is requested, it defaults to a system-appropriate persistent location managed by 'rappdirs'.
use_persistent_storage	A logical value indicating whether to use persistent storage. If TRUE and no path is provided, data will be stored in a system-appropriate location. Defaults to FALSE, using a temporary directory.
force_install	A logical value indicating whether to force re-downloading of the data even if it already exists locally.

Value

A data.table containing the combined MeSH and supplemental thesaurus data.

Examples

```
if (interactive()) {  
  data <- data_mesh_thesaurus()  
}
```

Description

This function downloads and loads the 'MeSH' (Medical Subject Headings) Trees data.

Usage

```
data_mesh_trees(  
  path = NULL,  
  use_persistent_storage = FALSE,  
  force_install = FALSE  
)
```

Arguments

path	A character string specifying the directory path where data should be stored. If not provided and persistent storage is requested, it defaults to a system-appropriate persistent location managed by 'rappdirs'.
use_persistent_storage	A logical value indicating whether to use persistent storage. If TRUE and no path is provided, data will be stored in a system-appropriate location. Defaults to FALSE, using a temporary directory.
force_install	A logical value indicating whether to force re-downloading of the data even if it already exists locally.

Details

The data is sourced from specified URLs and stored locally for subsequent use. By default, the data is stored in a temporary directory. Users can opt into persistent storage by setting 'use_persistent_storage' to TRUE and optionally specifying a path.

Value

A data frame containing the MeSH Trees data.

Examples

```
if (interactive()) {  
  data <- data_mesh_trees()  
}
```

data_pmc_list	<i>Download and Load 'PMC' Open Access File List</i>
---------------	--

Description

This function downloads and loads the 'PMC' (PubMed Central) Open Access file list. The file list contains mappings between PMC IDs, PMIDs, and file paths for open access articles available for download.

Usage

```
data_pmc_list(  
  path = NULL,  
  use_persistent_storage = FALSE,  
  force_install = FALSE  
)
```

Arguments

path	A character string specifying the directory path where data should be stored. If not provided and persistent storage is requested, it defaults to a system-appropriate persistent location managed by 'rappdirs'.
use_persistent_storage	A logical value indicating whether to use persistent storage. If TRUE and no path is provided, data will be stored in a system-appropriate location. Defaults to FALSE, using a temporary directory.
force_install	A logical value indicating whether to force re-downloading of the data even if it already exists locally.

Details

The data is sourced from NCBI's FTP server and stored locally for subsequent use. By default, the data is stored in a temporary directory. Users can opt into persistent storage by setting 'use_persistent_storage' to TRUE and optionally specifying a path.

Value

A data.table containing the PMC file list with columns: file_path, citation, pmcid, pmid, and license_code.

Examples

```
if (interactive()) {  
  data <- data_pmc_list()  
}
```

endpoint_info	<i>Get Information About Available Endpoints</i>
---------------	--

Description

This function provides detailed information about the available endpoints in the package, including column descriptions, parameters, rate limits, and usage notes.

Usage

```
endpoint_info(endpoint = NULL, format = c("list", "json"))
```

Arguments

endpoint	Character string specifying which endpoint to get information about. If NULL (default), returns a list of all available endpoints.
format	Character string specifying the output format. Either "list" (default) or "json" for JSON-formatted output.

Value

If endpoint is NULL, returns a character vector of available endpoint names. If endpoint is specified, returns a list (or JSON string) with detailed information about that endpoint including description, columns, parameters, rate limits, and notes.

Examples

```
if (interactive()) {  
  # List all available endpoints  
  endpoint_info()  
  
  # Get information about a specific endpoint  
  endpoint_info("pubmed_abstracts")  
  
  # Get information in JSON format  
  endpoint_info("icites", format = "json")  
}
```

`get_records`*Retrieve Data from 'NLM'/'PubMed' databases Based on PMIDs*

Description

This function retrieves different types of data (like 'PubMed' records, affiliations, 'iCites' data, etc.) from 'PubMed' based on provided PMIDs. It supports parallel processing for efficiency.

Usage

```
get_records(  
  pmids,  
  endpoint = c("pubtations", "icites", "pubmed_affiliations", "pubmed_abstracts",  
              "pmc_fulltext"),  
  cores = 3,  
  sleep = 1,  
  ncbi_key = NULL  
)
```

Arguments

<code>pmids</code>	A vector of PMIDs for which data is to be retrieved. For 'pmc_fulltext' endpoint, provide full URLs instead (e.g., from <code>pmid_to_pmc()</code> \$url).
<code>endpoint</code>	A character vector specifying the type of data to retrieve ('pubtations', 'icites', 'pubmed_affiliations', 'pubmed_abstracts', 'pmc_fulltext').
<code>cores</code>	Number of cores to use for parallel processing (default is 3).
<code>sleep</code>	Duration (in seconds) to pause after each batch
<code>ncbi_key</code>	(Optional) NCBI API key for authenticated access.

Details

For the 'pmc_fulltext' endpoint, provide full URLs to PMC tar.gz files. Use [pmid_to_pmc](#) to convert PMIDs to PMC IDs and full URLs first.

Value

A data.table containing combined results from the specified endpoint.

Examples

```
pmids <- c("38136652")  
results <- get_records(pmids, endpoint = "pubmed_abstracts", cores = 1)
```

pmid_to_ftp

Convert PubMed IDs (PMIDs) to PMC IDs and Full-Text URLs

Description

This function converts PMIDs to PMC IDs, then fetches the full-text file URLs from the PMC Open Access service. It combines both steps into a single workflow.

Usage

```
pmid_to_ftp(
  pmids,
  batch_size = 200L,
  sleep = 0.5,
  verbose = FALSE,
  ncbi_key = NULL
)
```

Arguments

pmids	A character or numeric vector of PubMed IDs (PMIDs) to convert.
batch_size	An integer specifying the number of PMIDs to process per batch for ID conversion. Defaults to 200L. The NCBI API has limitations on batch sizes.
sleep	A numeric value specifying the number of seconds to pause between API requests for ID conversion (Step 1). Defaults to 0.5 seconds. For OA API calls (Step 2), sleep time is automatically adjusted based on rate limits: 0.11s with API key (10 req/sec), 0.34s without (3 req/sec).
verbose	Logical, whether to print progress messages. Defaults to FALSE.
ncbi_key	(Optional) NCBI API key for authenticated access.

Value

A data.table with columns:

- pmid: The input PubMed ID
- pmcid: The corresponding PMC ID
- doi: The corresponding DOI (NA if not available)
- url: The full HTTPS URL for downloading PMC full text

Results are filtered to only include rows with valid URLs (open access articles), ordered by PMID. Returns NULL with a message if the API is unavailable or returns invalid data.

Examples

```
if (interactive()) {  
  # Convert PMIDs to PMC IDs and get full-text URLs  
  result <- pmid_to_ftp(c("11250746", "11573492"))  
}
```

pmid_to_pmc

Convert PubMed IDs (PMIDs) to PMC IDs

Description

This function converts a vector of PubMed IDs (PMIDs) to their corresponding PubMed Central (PMC) IDs and DOIs using the NCBI ID Converter API.

Usage

```
pmid_to_pmc(pmids, batch_size = 200L, sleep = 0.5)
```

Arguments

pmids	A character or numeric vector of PubMed IDs (PMIDs) to convert.
batch_size	An integer specifying the number of PMIDs to process per batch. Defaults to 200L. The NCBI API has limitations on batch sizes.
sleep	A numeric value specifying the number of seconds to pause between API requests. Defaults to 0.5 seconds to respect API rate limits.

Value

A data.table with columns:

- pmid: The input PubMed ID
- pmcid: The corresponding PMC ID (NA if not available in PMC)
- doi: The corresponding DOI (NA if not available)

Results are ordered by PMID. Returns NULL with a message if the API is unavailable or returns invalid data.

Examples

```
if (interactive()) {  
  # Convert a single PMID to PMC ID  
  result <- pmid_to_pmc("12345678")  
  
  # Convert multiple PMIDs  
  pmids <- c("12345678", "23456789", "34567890")  
  result <- pmid_to_pmc(pmids, batch_size = 10, sleep = 1)
```

```
}
```

```
search_pubmed
```

```
Search 'PubMed' Records
```

Description

Performs a 'PubMed' search based on a query, optionally filtered by publication years. Returns a unique set of 'PubMed' IDs matching the query.

Usage

```
search_pubmed(  
  x,  
  start_year = NULL,  
  end_year = NULL,  
  retmax = 9999,  
  use_pub_years = FALSE  
)
```

Arguments

x	Character string, the search query.
start_year	Integer, the start year of publication date range (used if 'use_pub_years' is TRUE).
end_year	Integer, the end year of publication date range (used if 'use_pub_years' is TRUE).
retmax	Integer, maximum number of records to retrieve, defaults to 9999.
use_pub_years	Logical, whether to filter search by publication years, defaults to TRUE.

Value

Numeric vector of unique PubMed IDs.

Examples

```
ethnob1 <- search_pubmed("ethnobotany", 2010, 2012)
```

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