

Package ‘ccdR’

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Type Package

Title Utilities for Interacting with the 'CTX' APIs

Version 1.1.0

Description Access chemical, hazard, bioactivity, and exposure data from the Computational Toxicology and Exposure ('CTX') APIs
<<https://api-ccte.epa.gov/docs/>>. 'ccdR' was developed to streamline the process of accessing the information available through the 'CTX' APIs without requiring prior knowledge of how to use APIs. Most data is also available on the CompTox Chemical Dashboard ('CCD')
<<https://comptox.epa.gov/dashboard/>> and other resources found at the EPA Computational Toxicology and Exposure Online Resources
<<https://www.epa.gov/comptox-tools>>.

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bioactivity_api_server	<i>Bioactivity API Server url</i>
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Description

A section of url used in Bioactivity API Endpoints

Usage

bioactivity_api_server

Format

An object of class character of length 1.

ccdr_options	<i>ccdr Options</i>
--------------	---------------------

Description

ccdr stores options as a named list in R’s global options, i.e. `getOption('ccdr')`. It currently stores two such options, one for CCTE credentialing and one to suppress private API information in the URLs printed to the screen when web queries are placed. For both of those, see [register_ccdr\(\)](#).

Usage

`set_ccdr_option(...)`

`has_ccdr_options()`

`has_ccdr_option(option)`

Arguments

- | | |
|--------|---|
| ... | a named listing of options to set |
| option | a specific option to query, e.g. <code>display_api_key</code> |

Value

- `set_ccdr_option()` does not have a return value but has the side effect of setting options used by other functions.
- `has_ccdr_option()` returns a Boolean.
- `has_ccdr_options()` returns a Boolean.

See Also

[register_ccdr\(\)](#)

Examples

```
# Set ccdr options
set_ccdr_option('display_api_key' = FALSE)

# Check if there are options registered to 'ccdr'
has_ccdr_options()

# Check if a specific option is registered for 'ccdr'
has_ccdr_option('display_api_key')
```

<code>chemical_api_server</code>	<i>Chemical API Server url</i>
----------------------------------	--------------------------------

Description

A section of url used in Chemical API Endpoints

Usage

```
chemical_api_server
```

Format

An object of class character of length 1.

chemical_contains	<i>Chemical contains</i>
-------------------	--------------------------

Description

Chemical contains

Usage

```
chemical_contains(  
  word = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE,  
  top = NULL  
)
```

Arguments

word	A character string of a chemical name or portion of a chemical name
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.
top	The number of results to return if there are multiple results available

Value

A data.frame of chemicals and related values matching the query parameters

Examples

```
# Pull chemicals that contain substring  
substring_chemicals <- chemical_contains(word = 'TXSID702018')
```

chemical_contains_batch	<i>Chemical contains batch search</i>
-------------------------	---------------------------------------

Description

Chemical contains batch search

Usage

```
chemical_contains_batch(  
  word_list = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = verbose,  
  top = NULL  
)
```

Arguments

word_list	A list of character strings of chemical names or portion of chemical names
API_key	User-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.
top	The number of results to return if there are multiple results available

Value

A named list of data.frames of chemicals and related values matching the query parameters

Examples

```
# Pull chemicals that contain substrings  
substring_chemicals <- chemical_contains_batch(word_list = c('TXDIS702018',  
                                                             'DTXSID70201'))
```

chemical_equal	<i>Chemical equal</i>
----------------	-----------------------

Description

Chemical equal

Usage

```
chemical_equal(  
  word = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

word	A character string of a chemical name or portion of a chemical name
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame of chemicals and related values matching the query parameters

Examples

```
# Pull chemicals with matching DTXSID
bpa_dtxsid <- chemical_equal(word = 'DTXSID7020182')
```

chemical_equal_batch *Chemical equal batch search*

Description

Chemical equal batch search

Usage

```
chemical_equal_batch(  
  word_list = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

word_list	A list of character strings of chemical names or portion of chemical names, DTXSIDs, CASRNs, InChIKeys.
API_key	User-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames of chemicals and related values matching the query parameters

Examples

```
# Pull chemicals that match input strings
bpa <- chemical_equal_batch(word_list = c('DTXSID7020182', 'DTXCID30182'))
```

chemical_starts_with	<i>Chemical starts with</i>
----------------------	-----------------------------

Description

Chemical starts with

Usage

```
chemical_starts_with(  
  word = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE,  
  top = NULL  
)
```

Arguments

word	A character string of a chemical name or portion of a chemical name
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.
top	The number of results to return if there are multiple results available

Value

A data.frame of chemicals and related values matching the query parameters

Examples

```
# Pull chemicals that start with a fragment DTXSID
dtxsid_fragment <- chemical_starts_with(word = 'DTXSID702018')
```

chemical_starts_with_batch

Chemical starts with batch search

Description

Chemical starts with batch search

Usage

```
chemical_starts_with_batch(
    word_list = NULL,
    API_key = NULL,
    rate_limit = 0L,
    verbose = FALSE,
    top = NULL
)
```

Arguments

word_list	A list of character strings of chemical names or portion of chemical names
API_key	User-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.
top	The number of results to return if there are multiple results available

Value

A named list of data.frames of chemicals and related values matching the query parameters

Examples

[illegible]

`create_data.table_chemical_details`*Create chemical details data.table helper function*

Description

Create chemical details data.table helper function

Usage

```
create_data.table_chemical_details(index = -1)
```

Arguments

`index` Determine which format should be used.

Value

An empty data.table with columns matching the expected format of the get_chemical_details API call.

`exposure_api_server` *Exposure API Server url*

Description

A section of url used in Exposure API Endpoints

Usage

```
exposure_api_server
```

Format

An object of class character of length 1.

get_all_assays	<i>Retrieve all assays</i>
----------------	----------------------------

Description

Retrieve all assays

Usage

```
get_all_assays(  
  API_key = NULL,  
  Server = bioactivity_api_server,  
  verbose = FALSE  
)
```

Arguments

API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing all the assays and associated information

Examples

```
# Retrieve all assays  
assays <- get_all_assays()
```

get_all_public_chemical_lists	<i>Get all public chemical lists</i>
-------------------------------	--------------------------------------

Description

Get all public chemical lists

Usage

```
get_all_public_chemical_lists(  
  Projection = "",  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

Projection	Optional parameter controlling return type. It takes values 'chemicallistall' and 'chemicallistname' with the former as the default value.
API_key	The user-specific api key
Server	The root address for the API endpoint
verbose	A logical indicating if some "progress report" should be given.

Value

A data.frame containing information on all public chemical lists available from the CTX chemical api.

Examples

```
# Pull all chemical lists
all_lists <- get_all_public_chemical_lists()
```

get_annotation_by_aeid

Retrieve annotations for AEID

Description

Retrieve annotations for AEID

Usage

```
get_annotation_by_aeid(
  AEID = NULL,
  API_key = NULL,
  Server = bioactivity_api_server,
  verbose = FALSE
)
```

Arguments

AEID	The assay endpoint identifier AEID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some "progress report" should be given.

Value

A data.frame containing the annotated assays corresponding to the input AEID parameter

Examples

```
# Retrieve annotation for an assay
annotation <- get_annotation_by_aeid(AEID = 159)
```

```
get_annotation_by_aeid_batch
      Retrieve annotations for AEID batch
```

Description

Retrieve annotations for AEID batch

Usage

```
get_annotation_by_aeid_batch(
  AEID = NULL,
  API_key = NULL,
  Server = NULL,
  rate_limit = 0L,
  verbose = FALSE
)
```

Arguments

AEID	A list of AEID identifiers
API_key	The user-specific API key
Server	The root address for the API endpoint
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing annotation information for the assays with AEID matching the input parameter.

Examples

```
# Get annotations for multiple auids
aeid_annotations <- get_annotation_by_aeid_batch(AEID = c(159, 160))
```

`get_bioactivity_details`*Retrieve bioactivity data from DTXSID, AEID, SPID, or m4id*

Description

Retrieve bioactivity data from DTXSID, AEID, SPID, or m4id

Usage

```
get_bioactivity_details(  
  DTXSID = NULL,  
  AEID = NULL,  
  SPID = NULL,  
  m4id = NULL,  
  API_key = NULL,  
  Server = bioactivity_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
AEID	The assay endpoint identifier AEID
SPID	The ChemSpider chemical input
m4id	The chemical identifier m4id
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing bioactivity information for the chemical or assay endpoint with identifier matching the input parameter.

Examples

```
# Pull BPA bioactivity details  
bpa <- get_bioactivity_details(DTXSID = 'DTXSID7020182')  
# Pull assay bioactivity details  
assay <- get_bioactivity_details(AEID = 159)
```

`get_bioactivity_details_batch`*Retrieve bioactivity data from DTXSID or AEID batch*

Description

Retrieve bioactivity data from DTXSID or AEID batch

Usage

```
get_bioactivity_details_batch(  
  DTXSID = NULL,  
  AEID = NULL,  
  SPID = NULL,  
  m4id = NULL,  
  API_key = NULL,  
  Server = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	A list of chemical identifier DTXSIDs.
AEID	A list of assay endpoint identifiers AEIDs.
SPID	A list of ChemSpider chemical inputs
m4id	A list of chemical identifier m4ids
API_key	The user-specific API key.
Server	The root address for the API endpoint
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing bioactivity information for the chemicals with DTXSID or assays with AEID matching the input parameter.

Examples

```
# Pull bioactivity details for multiple chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
batch_bioactivity <- get_bioactivity_details_batch(DTXSID = dtxsid)  
# Pull bioactivity details for multiple assays  
batch_bioactivity <- get_bioactivity_details_batch(AEID = c(159, 160))
```

get_bioactivity_endpoint_status
Bioactivity API Endpoint status

Description

Bioactivity API Endpoint status

Usage

```
get_bioactivity_endpoint_status()
```

Value

Status of Bioactivity API Endpoints

Examples

```
status <- get_bioactivity_endpoint_status()
print(status)
```

get_bioactivity_summary
Retrieve bioactivity summary for AEID

Description

Retrieve bioactivity summary for AEID

Usage

```
get_bioactivity_summary(  
  AEID = NULL,  
  API_key = NULL,  
  Server = bioactivity_api_server,  
  verbose = FALSE  
)
```

Arguments

AEID	The assay endpoint identifier AEID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing summary information corresponding to the input AEID

Examples

```
# Pull an assay bioactivity summary
aaid_1386 <- get_bioactivity_summary(AEID = 1386)
```

```
get_bioactivity_summary_batch
```

Retrieve bioactivity summary data from AEID batch

Description

Retrieve bioactivity summary data from AEID batch

Usage

```
get_bioactivity_summary_batch(  
  AEID = NULL,  
  API_key = NULL,  
  Server = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

AEID	A list of AEID identifiers
API_key	The user-specific API key.
Server	The root address for the API endpoint
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing bioactivity summary information for the assays with AEID matching the input parameter.

Examples

```
# Get bioactivity summary for multiple aoids
aoids <- get_bioactivity_summary_batch(AEID = c(159, 160))
```

get_cancer_hazard	<i>Get cancer hazard</i>
-------------------	--------------------------

Description

Get cancer hazard

Usage

```
get_cancer_hazard(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame of cancer hazard data related to the input DTXSID.

Examples

```
# Pull cancer hazard data for BPA  
bpa_cancer <- get_cancer_hazard(DTXSID = 'DTXSID7020182')
```

get_cancer_hazard_batch	<i>Get cancer hazard batch</i>
-------------------------	--------------------------------

Description

Get cancer hazard batch

Usage

```
get_cancer_hazard_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between requests
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table containing cancer hazard and related data for each input DTXSID.

Examples

```
# Pull cancer hazard data for multiples chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
dtxsid_cancer_hazard <- get_cancer_hazard_batch(DTXSID = dtxsid)
```

get_cancer_hazard_batch_old

Get cancer hazard batch

Description

Get cancer hazard batch

Usage

```
get_cancer_hazard_batch_old(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between requests
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames, each containing cancer hazard and related data for each input DTXSID.

get_chemicals_in_list *Get chemicals in a given chemical list*

Description

Get chemicals in a given chemical list

Usage

```
get_chemicals_in_list(  
  list_name = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

list_name	The name of the list of chemicals
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame of the chemical list

Examples

```
# Retrieve chemicals contained in chemical list 'CCL4'  
ccl4_chemicals <- get_chemicals_in_list(list_name = 'CCL4')
```

`get_chemicals_in_list_batch`*Get chemicals in a given chemical list batch*

Description

Get chemicals in a given chemical list batch

Usage

```
get_chemicals_in_list_batch(  
  list_names = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

<code>list_names</code>	A list of names of chemical lists.
<code>API_key</code>	The user-specific API key.
<code>rate_limit</code>	Number of seconds to wait between each request
<code>verbose</code>	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames each containing chemicals in the corresponding chemical lists.

Examples

```
# Pull chemicals in lists for multiple lists  
chemicals_in_lists <- get_chemicals_in_list_batch(list_names = c('CCL4', 'NATADB'))
```

`get_chemical_by_property_range`*Get chemicals by property and its value range*

Description

Get chemicals by property and its value range

Usage

```
get_chemical_by_property_range(  
  start = NULL,  
  end = NULL,  
  property = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

start	A numeric value, the beginning of the range
end	A numeric value, the end of the range
property	A string, the property in question
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing chemical information for chemicals matching the search criteria.

Examples

```
# Pull chemicals with a given property in a set range  
density <- get_chemical_by_property_range(start = 1.311, end = 1.313,  
                                          property = 'Density')
```

get_chemical_by_property_range_batch

Retrieve chemicals by property and value range in batch search

Description

Retrieve chemicals by property and value range in batch search

Usage

```
get_chemical_by_property_range_batch(  
  start_list = NULL,  
  end_list = NULL,  
  property_list = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

start_list	Numeric values, the beginning of the range
end_list	Numeric values, the end of the range
property_list	Strings, the properties being queried
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing chemical information for the chemicals matching the search criteria.

Examples

```
# Pull chemicals by property ranges
prop_ranges <- get_chemical_by_property_range_batch(start_list = c(1.311,
                                                                    211.99),
                                                    end_list = c(1.313,
                                                                    212.01),
                                                    property_list = c('Density',
                                                                    'Boiling Point'))
```

get_chemical_details	<i>Retrieve chemical details from DTXSID of DTXCID</i>
----------------------	--

Description

Retrieve chemical details from DTXSID of DTXCID

Usage

```
get_chemical_details(
  DTXSID = NULL,
  DTXCID = NULL,
  Projection = "chemicaldetailstandard",
  API_key = NULL,
  Server = chemical_api_server,
  verbose = FALSE
)
```


Arguments

DTXSID	The chemical identifier DTXSID
DTXCID	The chemical identifier DTXCID
Projection	The format and chemical detail data returned. Allowed values are 'chemicaldetailall', 'chemicaldetailstandard', 'chemicalidentifier', 'chemicalstructure', 'nta-toolkit', 'ccdchemicaldetails'. If left empty or there is a mismatch, the default format will be 'chemicaldetailstandard'.
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some "progress report" should be given.

Value

A data.table containing chemical information for the chemical with DTXSID matching the input parameter.

Examples

```
# Pull chemical details for BPA
bpa <- get_chemical_details(DTXSID = 'DTXSID7020182')
```

`get_chemical_details_batch`

Retrieve chemical details from DTXSID of DTXCID in batch search

Description

Retrieve chemical details from DTXSID of DTXCID in batch search

Usage

```
get_chemical_details_batch(  
  DTXSID = NULL,  
  DTXCID = NULL,  
  Projection = "chemicaldetailstandard",  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
DTXCID	The chemical identifier DTXCID
Projection	The format and chemical detail data returned. Allowed values are 'chemicaldetailall', 'chemicaldetailstandard', 'chemicalidentifier', 'chemicalstructure', 'nta-toolkit', 'ccdchemicaldetails'. If left empty or there is a mismatch, the default format will be 'chemicaldetailstandard'.
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some "progress report" should be given.

Value

A data.table (DTXSID) or a named list of data.tables (DTXCID) containing chemical information for the chemicals with DTXSID or DTXCID matching the input parameter.

Examples

```
# Pull chemical details for multiple chemicals by dtxsid
dtxsids <- c('DTXSID7020182', 'DTXSID2021315')
dtxsid_details <- get_chemical_details_batch(DTXSID = dtxsids)
# Pull chemical details for multiple chemicals by dtxcid
dtxcids <- c('DTXCID30182', 'DTXCID001315')
dtxcid_details <- get_chemical_details_batch(DTCID = dtxcids)
```

```
get_chemical_endpoint_status
```

Chemical API Endpoint status

Description

Chemical API Endpoint status

Usage

```
get_chemical_endpoint_status()
```

Value

Status of Chemical API Endpoints

Examples

```
status <- get_chemical_endpoint_status()
print(status)
```

get_chemical_image	Get image file by DTXSID or DTXCID
--------------------	------------------------------------

Description

Get image file by DTXSID or DTXCID

Usage

```
get_chemical_image(  
  DTXSID = NULL,  
  DTXCID = NULL,  
  SMILES = NULL,  
  format = "",  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	Chemical identifier DTXSID
DTXCID	Chemical identifier DTXCID
SMILES	Chemical identifier SMILES
format	The image type, either "png" or "svg". If left blank, will default to "png".
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some "progress report" should be given.

Value

A Large array of three dimensions representing an image. For displaying this, one may use `png::writePNG()` or `countcolors::plotArrayAsImage()` among many such functions.

Examples

```
# Pull chemical image for BPA by dtxsid  
bpa_image_matrix <- get_chemical_image(DTXSID = 'DTXSID7020182')  
if (requireNamespace("countcolors", quietly = TRUE)){  
  countcolors::plotArrayAsImage(bpa_image_matrix)  
}  
# Pull chemical image for BPA by dtxcid  
bpa_image_matrix <- get_chemical_image(DTXCID = 'DTXCID30182')  
if (requireNamespace("countcolors", quietly = TRUE)){  
  countcolors::plotArrayAsImage(bpa_image_matrix)  
}
```

`get_chemical_image_batch`*Get image file by DTXSID or DTXCID batch*

Description

Get image file by DTXSID or DTXCID batch

Usage

```
get_chemical_image_batch(  
  DTXSID = NULL,  
  DTXCID = NULL,  
  SMILES = NULL,  
  format = "",  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	A list of chemical identifier DTXSIDs.
DTXCID	A list of chemical identifier DTXCIDs.
SMILES	A list of chemical identifier SMILES.
format	The image type, either "png" or "svg". If left blank, will default to "png".
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some "progress report" should be given.

Value

A named list of Large arrays of three dimensions representing an image. For displaying an image, one may use `png::writePNG()` or `countcolors::plotArrayAsImage()` among many such functions.

Examples

```
# Pull images for multiple chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
images <- get_chemical_image_batch(DTXSID = dtxsid)  
if (requireNamespace("countcolors", quietly = TRUE)){  
  countcolors::plotArrayAsImage(images[[1]])  
  countcolors::plotArrayAsImage(images[[2]])  
}
```

```
get_chemical_lists_by_type
```

Get chemical lists by type

Description

Get chemical lists by type

Usage

```
get_chemical_lists_by_type(  
  type = NULL,  
  Projection = "",  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

type	The type of list. This is a case sensitive parameter and returns lists only for values of "federal", "international", "state", and "other".
Projection	Optional parameter controlling return type. It takes values 'chemicallistall' and 'chemicallistname' with the former as the default value.
API_key	The user-specified API key.
Server	The root address for the API endpoint
verbose	A logical indicating if some "progress report" should be given.

Value

A data.frame containing information about lists that meet the search criteria.

Examples

```
# Pull chemical lists by type  
federal <- get_chemical_lists_by_type(type = 'Federal')
```

get_chemical_lists_by_type_batch

Get chemical lists by type batch search

Description

Get chemical lists by type batch search

Usage

```
get_chemical_lists_by_type_batch(
    type_list = NULL,
    Projection = "",
    API_key = NULL,
    rate_limit = 0L,
    verbose = FALSE
)
```

Arguments

type_list	A list of list types. This is a case sensitive parameter and returns lists only for values of "federal", "international", "state", and "other".
Projection	Optional parameter controlling return type. It takes values 'chemicallistall' and 'chemicallistname' with the former as the default value.
API_key	The user-specified API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some "progress report" should be given.

Value

A named list of data.frames containing information about lists that meet the search criteria.

Examples

[illegible]

get_chemical_mol	<i>Get mol file by DTXSID or DTXCID</i>
------------------	---

Description

Get mol file by DTXSID or DTXCID

Usage

```
get_chemical_mol(  
  DTXSID = NULL,  
  DTXCID = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	Chemical identifier DTXSID
DTXCID	Chemical identifier DTXCID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A character string giving a mol file representation

Examples

```
# Pull mol file for BPA by dtxsid  
bpa_mol <- get_chemical_mol(DTXSID = 'DTXSID7020182')  
# Pull mol file for BPA by dtxcid  
bpa_mol <- get_chemical_mol(DTCID = 'DTXCID30182')
```

`get_chemical_mol_batch`*Get mol file by DTXSID or DTXCID batch*

Description

Get mol file by DTXSID or DTXCID batch

Usage

```
get_chemical_mol_batch(  
  DTXSID = NULL,  
  DTXCID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	A list of the chemical identifier DTXSIDs.
DTXCID	A list of the chemical identifier DTXCIDs.
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of character strings giving a mol file representations of the given input chemicals.

Examples

```
# Pull mol files for multiple chemicals by DTXSID  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
mol_files <- get_chemical_mol_batch(DTXSID = dtxsid)  
# Pull mol files for multiple chemicals by DTXCID  
dtxcid <- c('DTXCID30182', 'DTXCID001315')  
mol_files <- get_chemical_mol_batch(DTXCID = dtxcid)
```

get_chemical_mrv	<i>Get mrv file by DTXSID or DTXCID</i>
------------------	---

Description

Get mrv file by DTXSID or DTXCID

Usage

```
get_chemical_mrv(  
  DTXSID = NULL,  
  DTXCID = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
DTXCID	The chemical identifier DTXCID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

XML file format for representing a mrv file.

Examples

```
# Pull mrv file for BPA by dtxsid  
bpa_mrv <- get_chemical_mrv(DTxsID = 'DTXSID7020182')  
# Pull mrv file for BPA by dtxcid  
bpa_mrv <- getchemical_mrv(DTXCID = 'DTXCID30182')
```

`get_chemical_mrv_batch`*Ger mrv file by DTXSID or DTXCID batch*

Description

Ger mrv file by DTXSID or DTXCID batch

Usage

```
get_chemical_mrv_batch(  
  DTXSID = NULL,  
  DTXCID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	A list of the chemical identifier DTXSIDs.
DTXCID	A list of the chemical identifier DTXCIDs.
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of XML file format for representing a mrv file for each chemicals.

Examples

```
# Pull mrv files for multiple chemicals by DTXSID  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
mrv_files <- get_chemical_mrv_batch(DTxsid = dtxsid)  
# Pull mrv files for multiple chemicals by DTXCID  
dtxcid <- c('DTXCID30182', 'DTXCID001315')  
mrv_files <- get_chemical_mrv_batch(DTxcid = dtxcid)
```

get_chemical_synonym *Get chemical synonym*

Description

Get chemical synonym

Usage

```
get_chemical_synonym(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of synonym information for the input DTXSID

Examples

```
# Pull synonyms for BPA  
bpa_synonym <- get_chemical_synonym(DTXSID = 'DTXSID7020182')
```

get_chemical_synonym_batch
 Get chemical synonym batch

Description

Get chemical synonym batch

Usage

```
get_chemical_synonym_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	A list of chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	The number of seconds to wait between requests.
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of lists containing synonym information for each input DTXSID.

Examples

```
# Pull synonyms for multiple chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
batch_synonyms <- get_chemical_synonym_batch(DTXSID = dtxsid)
```

get_chem_info

Retrieve chemical information

Description

Retrieve chemical information

Usage

```
get_chem_info(  
  DTXSID = NULL,  
  type = "",  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
type	This specifies whether to only grab predicted or experimental results. If not specified, it will grab all details. The allowable input values are "predicted" or "experimental".
API_key	The user-specific API Key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing chemical information for the chemical with DTXSID matching the input parameter.

Examples

```
# Pull chemical information for BPA
bpa <- get_chem_info(DTXSID = 'DTXSID7020182')
```

get_chem_info_batch *Retrieve chemical information in batch search*

Description

Retrieve chemical information in batch search

Usage

```
get_chem_info_batch(
  DTXSID = NULL,
  type = "",
  API_key = NULL,
  rate_limit = 0L,
  Server = chemical_api_server,
  verbose = FALSE
)
```

Arguments

DTXSID	A vector of chemical identifier DTXSIDs
type	A vector of type used in get_chem_info(). This specifies whether to only grab predicted or experimental results. If not specified, it will grab all details. The allowable input values are "", predicted", or "experimental".
API_key	The user-specific API key.

rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table containing chemical information for the chemicals with DTXSID matching the input parameter.

Examples

```
# Pull chemical info for multiple chemicals
chem_info <- get_chem_info_batch(DTXSID = c('DTXSID7020182',
                                           'DTXSID2021315'))
```

```
get_chem_info_batch_old
```

Retrieve chemical information in batch search

Description

Retrieve chemical information in batch search

Usage

```
get_chem_info_batch_old(
  DTXSID = NULL,
  type = "",
  API_key = NULL,
  rate_limit = 0L,
  verbose = FALSE
)
```

Arguments

DTXSID	A vector of chemical identifier DTXSIDs
type	A vector of type used in get_chem_info(). This specifies whether to only grab predicted or experimental results. If not specified, it will grab all details. The allowable input values are "", predicted", or "experimental".
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing chemical information for the chemicals with DTXSID matching the input parameter.

`get_ecotox_hazard_by_dtxsid`*Get ecotox hazard data by DTXSID*

Description

Get ecotox hazard data by DTXSID

Usage

```
get_ecotox_hazard_by_dtxsid(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing chemical (ecotox) hazard data

Examples

```
# Pull ecotox hazard data for BPA  
bpa_ecotox <- get_ecotox_hazard_by_dtxsid(DTXSID = 'DTXSID7020182')
```

`get_ecotox_hazard_by_dtxsid_batch`*Get ecotox hazard data by DTXSID batch*

Description

Get ecotox hazard data by DTXSID batch

Usage

```
get_ecotox_hazard_by_dtxsid_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	A list of chemical identifier DTXSIDs.
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table containing chemical ecotox hazard data.

Examples

```
# Pull ecotox hazard data for multiples chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
dtxsid_ecotox_hazard <- get_ecotox_hazard_by_dtxsid_batch(DTXSID = dtxsid)
```

```
get_ecotox_hazard_by_dtxsid_batch_old
```

Get ecotox hazard data by DTXSID batch

Description

Get ecotox hazard data by DTXSID batch

Usage

```
get_ecotox_hazard_by_dtxsid_batch_old(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```


Arguments

DTXSID	A list of chemical identifier DTXSIDs.
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing chemical ecotox hazard data.

get_exposure_endpoint_status
Exposure API Endpoint status

Description

Exposure API Endpoint status

Usage

```
get_exposure_endpoint_status()
```

Value

Status of Exposure API Endpoints

Examples

```
status <- get_exposure_endpoint_status()
print(status)
```

get_exposure_functional_use
Retrieve exposure related functional use data

Description

Retrieve exposure related functional use data

Usage

```
get_exposure_functional_use(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = exposure_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	Chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame of functional use data.

Examples

```
# Pull functional use data for BPA  
bpa <- get_exposure_functional_use(DTXSID = 'DTXSID7020182')
```

get_exposure_functional_use_batch

Retrieve exposure related functional use data batch

Description

Retrieve exposure related functional use data batch

Usage

```
get_exposure_functional_use_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = exposure_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	Chemical identifier DTXSID
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames, each containing exposure functional use data for each input DTXSID.

Examples

```
has_ctx_key() & is.na(ctx_key() == 'FAKE_KEY')
# Pull exposure functional use data for multiple chemicals
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')
dtxsid_func_use <- get_exposure_functional_use_batch(DTXSID = dtxsid)
```

`get_exposure_functional_use_category`*Retrieve functional use categories*

Description

Retrieve functional use categories

Usage

```
get_exposure_functional_use_category(
  API_key = NULL,
  Server = exposure_api_server,
  verbose = FALSE
)
```

Arguments

API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame of functional use categories.

Examples

```
# Pull functional use category data for BPA
functional_use_categories <- get_exposure_functional_use_category()
```

```
get_exposure_functional_use_probability
```

Retrieve probability of exposure for functional use category

Description

Retrieve probability of exposure for functional use category

Usage

```
get_exposure_functional_use_probability(
  DTXSID = NULL,
  API_key = NULL,
  Server = exposure_api_server,
  verbose = FALSE
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame with probabilities corresponding to various routes of exposure related to functional use.

Examples

```
# Pull functional use probability data for BPA
bpa <- get_exposure_functional_use_probability(DTXSID = 'DTXSID7020182')
```

`get_exposure_functional_use_probability_batch`*Retrieve exposure functional use probability data batch*

Description

Retrieve exposure functional use probability data batch

Usage

```
get_exposure_functional_use_probability_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = exposure_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	Chemical identifier DTXSID
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table, with each row containing exposure functional use probability data for each input DTXSID. NA values are filled in for categories that have probability of 0

Examples

```
has_ctx_key() & is.na(ctx_key() == 'FAKE_KEY')  
# Pull exposure functional use probability data for multiple chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
dtxsid_func_use_prob <- get_exposure_functional_use_batch(DTXSID = dtxsid)
```

get_exposure_list_presence_tags
Retrieve list presence tags

Description

Retrieve list presence tags

Usage

```
get_exposure_list_presence_tags(  
  API_key = NULL,  
  Server = exposure_api_server,  
  verbose = FALSE  
)
```

Arguments

API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame with all the list presence tags and associated data.

Examples

```
# Pull list presence tags  
tags <- get_exposure_list_presence_tags()
```

get_exposure_list_presence_tags_by_dtxsid
Retrieve document data and list presence tags for a chemical

Description

Retrieve document data and list presence tags for a chemical

Usage

```
get_exposure_list_presence_tags_by_dtxsid(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = exposure_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame of document information and list presence tags

Examples

```
# Pull list presence tags for BPA
bpa <- get_exposure_list_presence_tags(DTXSID = 'DTXSID7020182')
```

```
get_exposure_list_presence_tags_by_dtxsid_batch
```

Retrieve document data and list presence tags for chemicals batch

Description

Retrieve document data and list presence tags for chemicals batch

Usage

```
get_exposure_list_presence_tags_by_dtxsid_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = exposure_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	Chemical identifier DTXSID
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames, each containing exposure list presence tags use data for each input DTXSID.

Examples

```
has_ctx_key() & is.na(ctx_key()) == 'FAKE_KEY')
# Pull exposure functional use data for multiple chemicals
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')
exp_list_tags <- get_exposure_list_presence_tags_by_dtxsid_batch(DTXSID = dtxsid)
```

get_exposure_product_data

Retrieve product data for exposure purposes

Description

Retrieve product data for exposure purposes

Usage

```
get_exposure_product_data(
  DTXSID = NULL,
  API_key = NULL,
  Server = exposure_api_server,
  verbose = FALSE
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame with product information relating to exposure to the given chemical

Examples

```
# Pull exposure product data for BPA
bpa <- get_exposure_product_data(DTXSID = 'DTXSID7020182')
```

`get_exposure_product_data_batch`*Retrieve product data for exposure purposes batch*

Description

Retrieve product data for exposure purposes batch

Usage

```
get_exposure_product_data_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = exposure_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	Chemical identifier DTXSID
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames, each containing exposure product data for each input DTXSID.

Examples

```
has_ctx_key() & is.na(ctx_key()) == 'FAKE_KEY')  
# Pull exposure functional use data for multiple chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
dtxsid_product_data <- get_exposure_product_data_batch(DTXSID = dtxsid)
```

`get_exposure_product_data_puc`*Retrieve product use categories related to exposure*

Description

Retrieve product use categories related to exposure

Usage

```
get_exposure_product_data_puc(  
  API_key = NULL,  
  Server = exposure_api_server,  
  verbose = FALSE  
)
```

Arguments

API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame consisting of all the product use categories

Examples

```
# Pull product data use categories for BPA  
puc_categories <- get_exposure_product_data_puc()
```

`get_fate_by_dtxsid`*Get fate by DTXSID*

Description

Get fate by DTXSID

Usage

```
get_fate_by_dtxsid(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

DTGSID	The chemical identifier DTGSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

@return A data.frame containing chemical information for the chemical with DTGSID matching the input parameter.

Examples

```
# Pull chemical fate data for BPA
bpa <- get_fate_by_dtgsid(DTGSID = 'DTGSID7020182')
```

`get_fate_by_dtgsid_batch`*Retrieve chemical fate data in batch search*

Description

Retrieve chemical fate data in batch search

Usage

```
get_fate_by_dtgsid_batch(  
  DTGSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

DTGSID	A vector of chemicals identifier DTGSIDs
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table containing chemical fate information for the chemicals with DTXSID matching the input parameter.

Examples

```
# Pull chemical fate by dtxsids
chemical_fates <- get_fate_by_dtxsid_batch(DTXSID = c('DTXSID7020182',
                                                    'DTXSID2021315'))
```

```
get_fate_by_dtxsid_batch_old
```

Retrieve chemical fate data in batch search

Description

Retrieve chemical fate data in batch search

Usage

```
get_fate_by_dtxsid_batch_old(
  DTXSID = NULL,
  API_key = NULL,
  rate_limit = 0L,
  verbose = FALSE
)
```

Arguments

DTXSID	A vector of chemicals identifier DTXSIDs
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing chemical fate information for the chemicals with DTXSID matching the input parameter.

get_genetox_details	<i>Get genetox details</i>
---------------------	----------------------------

Description

Get genetox details

Usage

```
get_genetox_details(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame of genetox detail data related to the input DTXSID.

Examples

```
# Pull genetox details for BPA  
bpa_genetox_details <- get_genetox_details(DTXSID = 'DTXSID7020182')
```

get_genetox_details_batch	<i>Get genetox details batch</i>
---------------------------	----------------------------------

Description

Get genetox details batch

Usage

```
get_genetox_details_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between requests
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table of genetox detail data for each input DTXSID.

Examples

```
# Pull genetox details data for multiples chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
dtxsid_genetox_details_hazard <- get_genetox_details_batch(DTXSID = dtxsid)
```

get_genetox_details_batch_old

Get genetox details batch

Description

Get genetox details batch

Usage

```
get_genetox_details_batch_old(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between requests
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames of genetox detail data for each input DTXSID.

get_genetox_summary	<i>Get genetox summary</i>
---------------------	----------------------------

Description

Get genetox summary

Usage

```
get_genetox_summary(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame of genetox summary data related to the input DTXSID.

Examples

```
# Pull genetox summary for BPA  
bpa_genetox_summary <- get_genetox_summary(DTXSID = 'DTXSID7020182')
```

```
get_genetox_summary_batch
```

Get genetox summary batch

Description

Get genetox summary batch

Usage

```
get_genetox_summary_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between requests
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table of genetox summary data for each input DTXSID.

Examples

```
# Pull genetox summary data for multiples chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
dtxsid_genetox_summary_hazard <- get_genetox_summary_batch(DTxsID = dtxsid)
```

get_genetox_summary_batch_old
Get genetox summary batch

Description

Get genetox summary batch

Usage

```
get_genetox_summary_batch_old(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between requests
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames of genetox summary data for each input DTXSID.

get_hazard_by_dtxsid *Get hazard data by DTXSID*

Description

Get hazard data by DTXSID

Usage

```
get_hazard_by_dtxsid(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing chemical (human and eco) hazard data

Examples

```
# Pull hazard data for BPA
bpa <- get_hazard_by_dtxsid(DTXSID = 'DTXSID7020182')
```

```
get_hazard_by_dtxsid_batch
```

Get hazard data by DTXSID batch

Description

Get hazard data by DTXSID batch

Usage

```
get_hazard_by_dtxsid_batch(
  DTXSID = NULL,
  API_key = NULL,
  rate_limit = 0L,
  Server = hazard_api_server,
  verbose = FALSE
)
```

Arguments

DTXSID	A list of chemical identifier DTXSIDs
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table containing chemical (human and eco) hazard data for each input chemical.

Examples

```
# Pull hazard data for multiple chemicals
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')
batch_hazard <- get_hazard_by_dtxsid_batch(DTXSID = dtxsid)
```

```
get_hazard_by_dtxsid_batch_old
```

Get hazard data by DTXSID batch

Description

Get hazard data by DTXSID batch

Usage

```
get_hazard_by_dtxsid_batch_old(
  DTXSID = NULL,
  API_key = NULL,
  rate_limit = 0L,
  verbose = FALSE
)
```

Arguments

DTXSID	A list of chemical identifier DTXSIDs
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing chemical (human and eco) hazard data for each input chemical.

get_hazard_endpoint_status
Hazard API Endpoint status

Description

Hazard API Endpoint status

Usage

```
get_hazard_endpoint_status()
```

Value

Status of Hazard API Endpoints

Examples

```
status <- get_hazard_endpoint_status()
print(status)
```

get_human_hazard_by_dtgsid
Get human hazard data by DTGSID

Description

Get human hazard data by DTGSID

Usage

```
get_human_hazard_by_dtgsid(
  DTGSID = NULL,
  API_key = NULL,
  Server = hazard_api_server,
  verbose = FALSE
)
```

Arguments

DTGSID	The chemical identifier DTGSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing chemical human hazard data

Examples

```
# Pull human hazard data for BPA
bpa_human <- get_human_hazard_by_dtxsid(DTXSID = 'DTXSID7020182')
```

```
get_human_hazard_by_dtxsid_batch
```

Get human hazard data by DTXSID batch

Description

Get human hazard data by DTXSID batch

Usage

```
get_human_hazard_by_dtxsid_batch(
  DTXSID = NULL,
  API_key = NULL,
  rate_limit = 0L,
  Server = hazard_api_server,
  verbose = FALSE
)
```

Arguments

DTXSID	A list of chemical identifier DTXSIDs.
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.table containing chemical human hazard data.

Examples

```
# Pull human hazard data for multiples chemicals
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')
dtxsid_human_hazard <- get_human_hazard_by_dtxsid_batch(DTXSID = dtxsid)
```

`get_human_hazard_by_dtxsid_batch_old`*Get human hazard data by DTXSID batch*

Description

Get human hazard data by DTXSID batch

Usage

```
get_human_hazard_by_dtxsid_batch_old(  
    DTXSID = NULL,  
    API_key = NULL,  
    rate_limit = 0L,  
    verbose = FALSE  
)
```

Arguments

DTXSID	A list of chemical identifier DTXSIDs.
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing chemical human hazard data.

`get_inchi`*Get InChI*

Description

Get InChI

Usage

```
get_inchi(  
    name = NULL,  
    API_key = NULL,  
    Server = chemical_api_server,  
    verbose = FALSE  
)
```

Arguments

name	Chemical name
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A string giving the associated inchi string.

Examples

```
bpa_inchi <- get_inchi(name = "Bisphenol A")
```

get_inchikey	<i>Get InChIKey</i>
--------------	---------------------

Description

Get InChIKey

Usage

```
get_inchikey(  
  name = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

name	Chemical name
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A string giving the associated InChIKey.

Examples

```
bpa_inchikey <- get_inchikey(name = "Bisphenol A")
```

`get_lists_containing_chemical`*Get chemical lists containing given chemical*

Description

Get chemical lists containing given chemical

Usage

```
get_lists_containing_chemical(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A list of names of chemical lists that contain the given chemical

Examples

```
# Pull chemical lists containing BPA  
bpa_lists <- get_lists_containing_chemical(DTXSID = 'DTXSID7020182')
```

`get_lists_containing_chemical_batch`*Get chemical lists containing given chemical batch*

Description

Get chemical lists containing given chemical batch

Usage

```
get_lists_containing_chemical_batch(
  chemical_list = NULL,
  API_key = NULL,
  rate_limit = 0L,
  verbose = FALSE
)
```

Arguments

`chemical_list` A list of the chemical identifier DTXSIDs.
`API_key` The user-specific API key.
`rate_limit` Number of seconds to wait between each request
`verbose` A logical indicating if some “progress report” should be given.

Value

A named list of chemical lists that contain the given chemicals.

Examples

```
# Pull lists containing chemicals for multiple chemicals
lists <- get_lists_containing_chemical_batch(chemical_list = c('DTXSID7020182',
                                                             'DTXSID2021315'))
```

`get_msready_by_dtxcid` *Get msready by DTXCID*

Description

Get msready by DTXCID

Usage

```
get_msready_by_dtxcid(
  DTXCID = NULL,
  API_key = NULL,
  Server = chemical_api_server,
  verbose = FALSE
)
```

Arguments

`DTXCID` The chemical identifier DTXCID
`API_key` The user-specific API key
`Server` The root address for the API endpoint
`verbose` A logical indicating if some “progress report” should be given.

A character list of DTXSIDs with DTXCIDs matching the search criteria

```
# Pull chemicals with matching DTXCID
dtxcid_msready <- get_msready_by_dtxcid(DTXSID = 'DTXCID30182')
```

get_msready_by_dtxcid_batch
Get msready by DTXCID batch search

Get msready by DTXCID batch search

```
get_msready_by_dtxcid_batch(
    DTXCID = NULL,
    API_key = NULL,
    rate_limit = 0L,
    verbose = FALSE
)
```

DTXCID	A list of chemical identifier DTXCIDs
API_key	A user-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

A named list of character lists of DTXSIDs with DTXCIDs matching the search criteria

[illegible]

`get_msready_by_formula`*Get msready by formula*

Description

Get msready by formula

Usage

```
get_msready_by_formula(  
  formula = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

<code>formula</code>	A string denoting the input chemical formula
<code>API_key</code>	The user-specific API key
<code>Server</code>	The root address for the API endpoint
<code>verbose</code>	A logical indicating if some “progress report” should be given.

Value

A character list of DTXSIDs with chemical formulas matching the search criteria

Examples

```
# Pull chemicals that match input formula  
mass_formula <- get_msready_by_formula(formula = 'C16H24N2O5S')
```

`get_msready_by_formula_batch`*Get msready by formula batch search*

Description

Get msready by formula batch search

Usage

```
get_msready_by_formula_batch(  
  formula_list = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

formula_list	A list of strings denoting the input chemicals formulas
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of character lists of DTXIDs with chemical formulas matching the search criteria

Examples

```
# Pull msready data for several chemical formulas  
msready_data <- get_msready_by_formula_batch(formula_list = c('C16H24N2O5S',  
                                                             'C15H16O2'))
```

get_msready_by_mass	<i>Get msready by mass</i>
---------------------	----------------------------

Description

Get msready by mass

Usage

```
get_msready_by_mass(  
  start = NULL,  
  end = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

start	The starting value for mass range
end	The ending value for mass range
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A list of DTXSIDs with msready mass falling within the given range.

Examples

```
# Pull chemicals with msready mass in given range
mass_range <- get_msready_by_mass(start = 200.9, end = 200.95)
```

```
get_msready_by_mass_batch
```

Get ms ready by mass batch search

Description

Get ms ready by mass batch search

Usage

```
get_msready_by_mass_batch(
  start_list = NULL,
  end_list = NULL,
  API_key = NULL,
  rate_limit = 0L,
  verbose = FALSE
)
```

Arguments

start_list	A numeric list of starting values for mass range
end_list	A numeric list of ending values for mass range
API_key	The user-specific API key
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of character lists with DTXSIDs with msready masses falling within the given ranges.

[illegible]

get_msready_by_mass_with_error_batch

Get msready by mass and error offset

Get msready by mass and error offset

```
get_msready_by_mass_with_error_batch(
    masses = NULL,
    error = NULL,
    API_key = NULL,
    rate_limit = 0,
    verbose = FALSE
)
```

masses	A numeric list of masses.
error	The mass offset value.
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some "progress report" should be given.

A list (of lists) of DTXSIDs, with a list returned for each input mass value.

[illegible]

```
get_public_chemical_list_by_name
```

Get chemical list by name

Description

Get chemical list by name

Usage

```
get_public_chemical_list_by_name(  
  list_name = NULL,  
  Projection = "",  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

list_name	The name of the list of chemicals
Projection	Optional parameter controlling return type. It takes values 'chemicallistall' and 'chemicallistname' with the former as the default value.
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some "progress report" should be given.

Value

A data.frame containing information about the chemical list. Note, this is not the chemical list itself. To access the chemicals in the list, use [get_chemicals_in_list](#).

See Also

[get_chemicals_in_list](#)

Examples

```
# Pull chemical list by list name  
ccl4 <- get_public_chemical_list_by_name(list_name = 'CCL4')
```

get_skin_eye_hazard	<i>Get skin and eye hazard</i>
---------------------	--------------------------------

Description

Get skin and eye hazard

Usage

```
get_skin_eye_hazard(  
  DTXSID = NULL,  
  API_key = NULL,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSID
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A data.frame containing skin and eye hazard data.

Examples

```
# Pull skin and eye hazard data for BPA  
bpa_skin_eye <- get_skin_eye_hazard_batch(DTXSID = 'DTXSID7020182')
```

get_skin_eye_hazard_batch	<i>Get skin and eye hazard batch</i>
---------------------------	--------------------------------------

Description

Get skin and eye hazard batch

Usage

```
get_skin_eye_hazard_batch(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  Server = hazard_api_server,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing skin and eye hazard data for each input DTXSID.

Examples

```
# Pull skin eye hazard data for multiples chemicals  
dtxsid <- c('DTXSID7020182', 'DTXSID2021315')  
dtxsid_skin_eye_hazard <- get_skin_eye_hazard_batch(DTxsid = dtxsid)
```

```
get_skin_eye_hazard_batch_old
```

Get skin and eye hazard batch

Description

Get skin and eye hazard batch

Usage

```
get_skin_eye_hazard_batch_old(  
  DTXSID = NULL,  
  API_key = NULL,  
  rate_limit = 0L,  
  verbose = FALSE  
)
```

Arguments

DTXSID	The chemical identifier DTXSIDs
API_key	The user-specific API key.
rate_limit	Number of seconds to wait between each request
verbose	A logical indicating if some “progress report” should be given.

Value

A named list of data.frames containing skin and eye hazard data for each input DTXSID.

get_smiles

Get Smiles

Description

Get Smiles

Usage

```
get_smiles(  
  name = NULL,  
  API_key = NULL,  
  Server = chemical_api_server,  
  verbose = FALSE  
)
```

Arguments

name	Chemical name
API_key	The user-specific API key
Server	The root address for the API endpoint
verbose	A logical indicating if some “progress report” should be given.

Value

A string giving a SMILES string for the input chemical.

Examples

```
bpa_smiles <- get_smiles(name = "Bisphenol A")
```

hazard_api_server	<i>Hazard API Server url</i>
-------------------	------------------------------

Description

A section of url used in Hazard API Endpoints

Usage

hazard_api_server

Format

An object of class character of length 1.

prepare_word	<i>Prepare url helper function</i>
--------------	------------------------------------

Description

Prepare url helper function

Usage

prepare_word(word)

Arguments

word	A character string
------	--------------------

Value

A character string that is ready for use in http request

register_ccdr	<i>Register a CTX API</i>
---------------	---------------------------

Description

This page contains documentation tools related to enabling CTX API services in R.

Usage

```
showing_key()

ccdr_show_api_key()

ccdr_hide_api_key()

register_ccdr(key, write = FALSE)

## S3 method for class 'ctx_credentials'
print(...)

ctx_key()

has_ctx_key()
```

Arguments

key	an API key
write	if TRUE, stores the secrets provided in the .Renviron file
...	a dumped formal argument to the generic print method

Details

To obtain an API key and enable services, go to <https://www.epa.gov/comptox-tools/computational-toxicology-and>. This documentation shows you how to obtain an API key to allow access to the CTX APIs.

To tell ccdR about your API key, use `register_ccdr()`, e.g. `register_ccdr(key = 'grbwigbwoginrowgbwibgdibdvirgi` (that's a fake key). This will set your API key for the current session, but if you restart R, you'll need to do it again. You can set it permanently by setting `write = TRUE` see the examples. If you set it permanently it will be stored in a local file, and that will be accessed by ccdR persistently across sessions.

Users should be aware that the API key, a string of garbled characters/numbers/symbols, is a PRIVATE key - it uniquely identifies and authenticates you to CTX's services. If anyone gets your API key, they can use it to masquerade as you to CTX. To mitigate against users inadvertently sharing their keys, by default ccdR never displays a user's key in messages displayed to the console.

Users should be aware that ccdR has no mechanism with which to safeguard the private key once registered with R. That is to say, once you register your API key, any R function will have access to it. As a consequence, ccdR will not know if another function, potentially from a compromised

package, accesses the key and uploads it to a third party. For this reason, when using ccdR we recommend a heightened sense of security and self-awareness: only use trusted packages, do not save the API keys in script files, etc.

Value

- `showing_key` returns a Boolean.
- `ccdr_show_api_key()` has no return value but has the side effect of changing the display settings of the API key.
- `ccdr_hide_api_key()` has no return value but has the side effect of changing the display settings of the API key.
- `register_ccdr()` has no return value but has the side effect of storing the API key.
- `print.ctx_credentials()` has no return value and is an S3 method for printing the `ctx_credentials` class.
- `ctx_key()` returns a string, either the stored API key or `NA_character_`.
- `has_ctx_key()` returns a Boolean.

Examples

```
# Check if API key is showing
showing_key()

# Toggle API key to display
ccdr_show_api_key()

# Toggle API key to be hidden
ccdr_hide_api_key()

# Register key for this session
register_ccdr(key = 'YOUR API KEY')
# Register key over sessions
register_ccdr(key = 'YOUR API KEY', write = TRUE)

# Print function for ctx_credentials class
print.ctx_credentials()

# Display ctx API key
ctx_key()

# Check whether API key is registered
has_ctx_key()
```

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