# Package 'restez'

October 25, 2023

```
Type Package
Title Create and Query a Local Copy of 'GenBank' in R
Version 2.1.4
Maintainer Joel H. Nitta < joelnitta@gmail.com>
Description Download large sections of
      'GenBank' <a href="https://www.ncbi.nlm.nih.gov/genbank/">https://www.ncbi.nlm.nih.gov/genbank/</a> and generate a local
      SQL-based database. A user can then query this database using 'restez'
      functions or through 'rentrez' <a href="https://CRAN.R-project.org/package=rentrez">https://CRAN.R-project.org/package=rentrez</a>
      wrappers.
URL https://github.com/ropensci/restez,
      https://docs.ropensci.org/restez/
BugReports https://github.com/ropensci/restez/issues
License MIT + file LICENSE
Encoding UTF-8
LazyData true
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      duckdb, fs, assertthat, ape
Suggests sessioninfo, testthat, knitr, R.utils, rmarkdown, mockery
RoxygenNote 7.2.3
NeedsCompilation no
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```

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add\_rcrd\_log

Log files added to the SQL database in the restez path

# Description

This function is called whenever sequence files have been successfully added to the nucleotide SQL database. Row entries are added to 'add\_lot.tsv' in the user's restez path containing the filename, GB release numbers and the time of successful adding. The log is to help users keep track of when sequences have been added.

# Usage

```
add_rcrd_log(fl)
```

#### **Arguments**

f1

filename, character

#### See Also

Other private: cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_seqrecpart(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

cat\_line

Cat lines

# Description

Helper function for printing lines to console. Automatically formats lines by adding newlines.

# Usage

```
cat_line(...)
```

char 5

# Arguments

... Text to print, character

#### See Also

Other private: add\_rcrd\_log(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

char

Print green

# Description

Print to console green text to indicate a name/filepath/text

### Usage

char(x)

#### **Arguments**

Χ

Text to print, character

# Value

coloured character encoding, character

```
Other private: add_rcrd_log(), cat_line(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(),
```

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gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(),
last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(),
message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(),
predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(),
restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(),
status\_class(), stat(), testdatadir\_get()

check\_connection

Helper function to test if a stable internet connection can be established.

# **Description**

All retrieval functions need a stable internet connection to work properly. This internal function pings the google homepage and throws an error if it cannot be reached.

### Usage

check\_connection()

# Author(s)

Hajk-Georg Drost

### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

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cleanup

Clean up test data

# **Description**

Removes all temporary test data created.

#### Usage

cleanup()

# See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

connected

Is restez connected?

# **Description**

Returns TRUE if a restez SQL database has been connected.

# Usage

connected()

#### Value

Logical

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### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

connection\_get

Retrieve restez connection

### Description

Safely acquire the restez connection. Raises error if no connection set.

#### Usage

connection\_get()

# Value

connection

### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

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count\_db\_ids

Return the number of ids

# Description

Return the number of ids in a user's restez database.

# Usage

```
count_db_ids(db = "nucleotide")
```

# Arguments

db

character, database name

# **Details**

Requires an open connection. If no connection or db 0 is returned.

# Value

integer

# See Also

```
Other database: db_create(), db_delete(), db_download(), demo_db_create(), is_in_db(), list_db_ids()
```

# **Examples**

```
library(restez)
restez_path_set(filepath = tempdir())
demo_db_create(n = 5)
(count_db_ids())

# delete demo after example
db_delete(everything = TRUE)
```

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db\_create

Create new NCBI database

### **Description**

Create a new local SQL database from downloaded files. Currently only GenBank/nucleotide/nuccore database is supported.

# Usage

```
db_create(
  db_type = "nucleotide",
  min_length = 0,
  max_length = NULL,
  acc_filter = NULL,
  invert = FALSE,
  alt_restez_path = NULL,
  scan = FALSE
)
```

# **Arguments**

db\_type character, database type

min\_length Minimum sequence length, default 0.

max\_length Maximum sequence length, default NULL.

acc\_filter Character vector; accessions to include or exclude from the database as specified

by invert.

invert Logical vector of length 1; if TRUE, accessions in acc\_filter will be excluded

from the database; if FALSE, only accessions in acc\_filter will be included

in the database. Default FALSE.

alt\_restez\_path

Alternative restez path if you would like to use the downloads from a different

restez path.

scan Logical vector of length 1; should the sequence file be scanned for accessions

in acc\_filter prior to processing? Requires zgrep to be installed (so does not work on Windows). Only used if acc\_filter is not NULL and invert is

FALSE. Default FALSE.

# **Details**

All .seq.gz files are added to the database by default. A user can specify minimum/maximum sequence lengths or accession numbers to limit the sequences to be added to the database – smaller databases are faster to search. The final selection of sequences is the result of applying all filters (acc\_filter, min\_length, max\_length) in combination.

The scan option can decrease the time needed to build a database if only a small number of sequences should be written to the database compared to the number of the sequences downloaded

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from GenBank; i.e., if many of the files downloaded from GenBank do not contain any sequences that should be written to the database. When set to TRUE, if a file does not contain any of the accessions in acc\_filter, further processing of that file will be skipped and none of the sequences it contains will be added to the database.

Alternatively, a user can use the alt\_restez\_path to add the files from an alternative restez file path. For example, you may wish to have a database of all environmental sequences but then an additional smaller one of just the sequences with lengths below 100 bp. Instead of having to download all environmental sequences twice, you can generate multiple restez databases using the same downloaded files from a single restez path.

This function will not overwrite a pre-existing database. Old databases must be deleted before a new one can be created. Use db\_delete() with everything=FALSE to delete an SQL database.

Connections/disconnections to the database are made automatically.

#### See Also

```
Other database: count_db_ids(), db_delete(), db_download(), demo_db_create(), is_in_db(), list_db_ids()
```

# **Examples**

```
## Not run:
# Example of general usage
library(restez)
restez_path_set(filepath = 'path/for/downloads/and/database')
db_download()
db_create()
# Example of using `acc_filter`
# Download files to temporary directory
temp_dir <- paste0(tempdir(), "/restez", collapse = "")</pre>
dir.create(temp_dir)
restez_path_set(filepath = temp_dir)
# Choose GenBank domain 20 ('unannotated'), the smallest
db_download(preselection = 20)
# Only include three accessions in database
 acc_filter = c("AF000122", "AF000123", "AF000124")
list_db_ids()
db_delete()
unlink(temp_dir)
## End(Not run)
```

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db\_delete

Delete database

# **Description**

Delete the local SQL database and/or restez folder.

# Usage

```
db_delete(everything = FALSE)
```

# **Arguments**

everything

T/F, delete the whole restez folder as well?

# **Details**

Any connected database will be automatically disconnected.

# See Also

```
Other database: count_db_ids(), db_create(), db_download(), demo_db_create(), is_in_db(), list_db_ids()
```

# **Examples**

```
library(restez)
fp <- tempdir()
restez_path_set(filepath = fp)
demo_db_create(n = 10)
db_delete(everything = FALSE)
# Will not run: gb_sequence_get(id = 'demo_1')
# only the SQL database is deleted
db_delete(everything = TRUE)
# Now returns NULL
(restez_path_get())</pre>
```

db\_download

Download database

# **Description**

Download .seq.tar files from the latest GenBank release.

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# Usage

```
db_download(
  db = "nucleotide",
  overwrite = FALSE,
  preselection = NULL,
  max_tries = 1
)
```

# **Arguments**

db Database type, only 'nucleotide' currently available.

overwrite T/F, overwrite pre-existing downloaded files?

preselection Character vector of length 1; GenBank domains to download. If not specified

(default), a menu will be provided for selection. To specify, provide either a single number or a single character string of numbers separated by spaces, e.g.

"19 20" for 'Phage' (19) and 'Unannotated' (20).

max\_tries Numeric vector of length 1; maximum number of times to attempt downloading

database (default 1).

#### **Details**

In default mode, the user interactively selects the parts (i.e., "domains") of GenBank to download (e.g. primates, plants, bacteria ...). Alternatively, the selected domains can be provided as a character string to preselection.

The max\_tries argument is useful for large databases that may otherwise fail due to periodic lapses in internet connectivity. This value can be set to Inf to continuously try until the database download succeeds (not recommended if you do not have an internet connection!).

#### Value

T/F, if all files download correctly, TRUE else FALSE.

# See Also

```
ncbi_acc_get()
Other database: count_db_ids(), db_create(), db_delete(), demo_db_create(), is_in_db(),
list_db_ids()
```

# **Examples**

```
## Not run:
library(restez)
restez_path_set(filepath = 'path/for/downloads')
db_download()
## End(Not run)
```

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db_download_intern	Download database (internal version)
o_download_intern	Download database (internal version

### **Description**

Download .seq.tar files from the latest GenBank release. The user interactively selects the parts of GenBank to download (e.g. primates, plants, bacteria ...). This is an internal function so the download can be wrapped in while() to enable persistent downloading.

### Usage

```
db_download_intern(db = "nucleotide", overwrite = FALSE, preselection = NULL)
```

### **Arguments**

db Database type, only 'nucleotide' currently available.

overwrite T/F, overwrite pre-existing downloaded files?

preselection Character vector of length 1; GenBank domains to download. If not specified

(default), a menu will be provided for selection. To specify, provide either a single number or a single character string of numbers separated by spaces, e.g.

"19 20" for 'Phage' (19) and 'Unannotated' (20).

### Details

The downloaded files will appear in the restez filepath under downloads.

### Value

T/F, if all files download correctly, TRUE else FALSE.

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

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db\_sqlngths\_get

Return the minimum and maximum sequence lengths in db

### **Description**

Returns the maximum and minimum sequence lengths as set by the user upon db creation.

# Usage

```
db_sqlngths_get()
```

#### **Details**

If no file found, returns empty character vector.

#### Value

vector of integers

# See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

db\_sqlngths\_log

Log the min and max sequence lengths

# **Description**

Log the min and maximum sequence length used in the created db.

# Usage

```
db_sqlngths_log(min_lngth, max_lngth)
```

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# **Arguments**

min\_lngth Minimum length
max\_lngth Maximum length

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

 $demo\_db\_create$ 

Create demo database

# **Description**

Creates a local mock SQL database from package test data for demonstration purposes. No internet connection required.

#### **Usage**

```
demo_db_create(db_type = "nucleotide", n = 100)
```

# **Arguments**

```
db_type character, database type

n integer, number of mock sequences
```

```
Other database: count_db_ids(), db_create(), db_delete(), db_download(), is_in_db(), list_db_ids()
```

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# **Examples**

```
library(restez)
# set the restez path to a temporary dir
restez_path_set(filepath = tempdir())
# create demo database
demo_db_create(n = 5)
# in the demo, IDs are 'demo_1', 'demo_2' ...
(gb_sequence_get(id = 'demo_1'))
# Delete a demo database after an example
db_delete(everything = TRUE)
```

dir\_size

Calculate the size of a directory

# **Description**

Returns the size of directory in GB

#### **Usage**

```
dir_size(fp)
```

#### **Arguments**

fp

File path, character

# Value

numeric

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

18 dwnld\_rcrd\_log

dwnld\_path\_get

Get dwnld path

# Description

Return path to folder where raw .seq files are stored.

# Usage

```
dwnld_path_get()
```

#### Value

character

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

dwnld\_rcrd\_log

Log a downloaded file in the restez path

# **Description**

This function is called whenever a file is successfully downloaded. A row entry is added to the 'download\_log.tsv' in the user's restez path containing the file name, the GB release number and the time of successfully download. The log is to help users keep track of when they downloaded files and to determine if the downloaded files are out of date.

# Usage

```
dwnld_rcrd_log(fl)
```

entrez\_fasta\_get 19

# Arguments

fl file name, character

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

entrez\_fasta\_get

Get Entrez fasta

# **Description**

Return fasta format as expected from an Entrez call. If not all IDs are returned, will run rentrez::entrez\_fetch.

#### Usage

```
entrez_fasta_get(id, ...)
```

#### **Arguments**

id vector, unique ID(s) for record(s)... arguments passed on to rentrez

# Value

character string containing the file created

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence
```

20 entrez\_fetch

```
extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(),
gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(),
gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(),
last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(),
mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(),
readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(),
search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(),
stat(), testdatadir_get()
```

entrez\_fetch

Entrez fetch

# **Description**

Wrapper for rentrez::entrez\_fetch.

# Usage

```
entrez_fetch(db, id = NULL, rettype, retmode = "", ...)
```

# **Arguments**

db character, name of the database id vector, unique ID(s) for record(s)

rettype character, data format retmode character, data mode

... Arguments to be passed on to rentrez

#### **Details**

Attempts to first search local database with user-specified parameters, if the record is missing in the database, the function then calls rentrez::entrez\_fetch to search GenBank remotely.

rettype='fasta' and rettype='gb' are respectively equivalent to gb\_fasta\_get() and gb\_record\_get().

#### Value

character string containing the file created

#### Supported return types and modes

XML retmode is not supported. Rettypes 'seqid', 'ft', 'acc' and 'uilist' are also not supported.

# Note

It is advisable to call restez and rentrez functions with '::' notation rather than library() calls to avoid namespace issues. e.g. restez::entrez\_fetch().

entrez\_gb\_get 21

# See Also

```
rentrez::entrez_fetch()
```

# **Examples**

```
library(restez)
restez_path_set(tempdir())
demo_db_create(n = 5)
# return fasta record
fasta_res <- entrez_fetch(db = 'nucleotide',</pre>
                           id = c('demo_1', 'demo_2'),
                            rettype = 'fasta')
cat(fasta_res)
# return whole GB record in text format
gb_res <- entrez_fetch(db = 'nucleotide',</pre>
                        id = c('demo_1', 'demo_2'),
                         rettype = 'gb')
cat(gb_res)
# NOT RUN
# whereas these request would go through rentrez
# fasta_res <- entrez_fetch(db = 'nucleotide',
# id = c('S71333', 'S71334'),
                              rettype = 'fasta')
# gb_res <- entrez_fetch(db = 'nucleotide',</pre>
                          id = c('S71333', 'S71334'),
#
                           rettype = 'gb')
# delete demo after example
db_delete(everything = TRUE)
```

entrez\_gb\_get

Get Entrez GenBank record

# **Description**

Return gb and gbwithparts format as expected from an Entrez call. If not all IDs are returned, will run rentrez::entrez\_fetch.

# Usage

```
entrez_gb_get(id, ...)
```

# Arguments

```
id vector, unique ID(s) for record(s)... arguments passed on to rentrez
```

# Value

character string containing the file created

22 extract\_accession

### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_seqrecpart(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

extract\_accession

Extract accession

# **Description**

Return accession ID from GenBank record

### Usage

extract\_accession(record)

#### **Arguments**

record

GenBank record in text format, character

# Details

If element is not found, "returned.

# Value

character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(),
```

extract\_by\_patterns 23

```
gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(),
last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(),
mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(),
readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(),
search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(),
stat(), testdatadir_get()
```

extract\_by\_patterns

Extract by keyword

### Description

Search through GenBank record for a keyword and return text up to the end\_pattern.

### Usage

```
extract_by_patterns(record, start_pattern, end_pattern = "\n")
```

# **Arguments**

record GenBank record in text format, character

start\_pattern REGEX pattern indicating the point to start extraction, character end\_pattern REGEX pattern indicating the point to stop extraction, character

### **Details**

The start\_pattern should be any of the capitalized elements in a GenBank record (e.g. LOCUS, DESCRIPTION, ACCESSION). The end\_pattern depends on how much of the selected element a user wants returned. By default, the extraction will stop at the next newline. If keyword or end pattern not found, returns NULL.

#### Value

character or NULL

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_seqrecpart(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(),
```

```
readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(),
search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(),
stat(), testdatadir_get()
```

extract\_clean\_sequence

Extract clean sequence from sequence part

# **Description**

Return clean sequence from seqrecpart of a GenBank record

#### Usage

```
extract_clean_sequence(seqrecpart, max_len = 1e+08)
```

### Arguments

seqrecpart Sequence part of a GenBank record, character

max\_len Number: maximum number of characters allowed in a single record before split-

ting the record into parts. Does not affect output, but only internal calculations,

so generally should not be changed. Default = 1e8.

# **Details**

If element is not found, "returned.

#### Value

character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_seqrecpart(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

extract\_definition 25

extract\_definition

Extract definition

# Description

Return definition from GenBank record.

# Usage

```
extract_definition(record)
```

# **Arguments**

record

GenBank record in text format, character

#### **Details**

If element is not found, "returned.

### Value

character

### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

26 extract\_features

extract\_features

Extract features

### **Description**

Return feature table as list from GenBank record

# Usage

```
extract_features(record)
```

# **Arguments**

record

GenBank record in text format, character

#### **Details**

If element is not found, empty list returned.

### Value

list of lists

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

extract\_inforecpart 27

# Description

Return information part from GenBank record

# Usage

```
extract_inforecpart(record)
```

# **Arguments**

record

GenBank record in text format, character

#### **Details**

If element is not found, "returned.

### Value

character

### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

28 extract\_keywords

extract\_keywords

Extract keywords

# **Description**

Return keywords as list from GenBank record

# Usage

```
extract_keywords(record)
```

# **Arguments**

record

GenBank record in text format, character

#### **Details**

If element is not found, "returned.

### Value

character vector

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_locus(), extract_organism(), extract_seqrecpart(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

extract\_locus 29

extract\_locus

Extract locus

### Description

Return locus information from GenBank record

# Usage

```
extract_locus(record)
```

# Arguments

record

GenBank record in text format, character

#### **Details**

If element is not found, "returned.

# Value

named character vector

### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_organism(), extract\_sequence(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

30 extract\_organism

extract\_organism

Extract organism

# Description

Return organism name from GenBank record

# Usage

```
extract_organism(record)
```

# **Arguments**

record

GenBank record in text format, character

#### **Details**

If element is not found, "returned.

# Value

character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_seqrecpart(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

extract\_seqrecpart 31

extract\_segrecpart

Extract the sequence record part

# Description

Return sequence part from GenBank record

# Usage

```
extract_seqrecpart(record)
```

# **Arguments**

record

GenBank record in text format, character

#### **Details**

If element is not found, "returned.

### Value

character

### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

32 extract\_sequence

extract\_sequence

Extract sequence

### **Description**

Return sequence from GenBank record

# Usage

```
extract_sequence(record)
```

# **Arguments**

record

GenBank record in text format, character

#### **Details**

If element is not found, "returned.

# Value

character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_seqrecpart(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

extract\_version 33

extract\_version

Extract version

### Description

Return accession + version ID from GenBank record

# Usage

```
extract_version(record)
```

# **Arguments**

record

GenBank record in text format, character

#### **Details**

If element is not found, "returned.

# Value

character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

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filename\_log

Write filenames to log files

# **Description**

Record a filename in a log file along with GB release and time.

# Usage

```
filename_log(fl, fp)
```

# **Arguments**

fl file name, character

fp filepath to log file, character

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

file\_download

Download a file

# **Description**

Download a GenBank .seq.tar file. Check the file has downloaded properly. If not, returns FALSE. If overwrite is true, any previous file will be overwritten.

# Usage

```
file_download(f1, overwrite = FALSE)
```

flatfile\_read 35

# **Arguments**

fl character, base filename (e.g. gbpri9.seq) to be downloaded

overwrite T/F

#### Value

T/F

### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

flatfile\_read

Read flatfile sequence records

#### **Description**

Read records from a .seq file.

# Usage

flatfile\_read(flpth)

# **Arguments**

flpth

Path to .seq file

#### Value

list of GenBank records in text format

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### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

gbrelease\_check

Check if the last GenBank release number is the latest

#### **Description**

Returns TRUE if the GenBank release number is the most recent GenBank release available.

#### Usage

gbrelease\_check()

# Value

logical

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

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gbrelease\_get

Get the GenBank release number in the restez path

#### **Description**

Returns the GenBank release number. Returns empty character if none found.

#### Usage

```
gbrelease_get()
```

#### **Details**

If no file found, returns empty character vector.

#### Value

character

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

gbrelease\_log

Log the GenBank release number in the restez path

#### **Description**

This function is called whenever db\_download is run. It logs the GB release number in the 'gb\_release.txt' in the user's restez path. The log is to help users keep track of whether their database if out of date.

## Usage

```
gbrelease_log(release)
```

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## Arguments

release GenBank release number, character

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

gb\_build

Read and add .seq files to database

## **Description**

Given a list of seq\_files, read and add the contents of the files to a SQL-like database. If any errors during the process, FALSE is returned.

## Usage

```
gb_build(
   dpth,
   seq_files,
   max_length,
   min_length,
   acc_filter = NULL,
   invert = FALSE,
   scan = FALSE
)
```

## **Arguments**

dpth Download path (where seq\_files are stored)
seq\_files .seq.tar seq file names
max\_length Maximum sequence length, default NULL.
min\_length Minimum sequence length, default 0.

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acc\_filter Character vector; accessions to include or exclude from the database as specified

by invert.

invert Logical vector of length 1; if TRUE, accessions in acc\_filter will be excluded

from the database; if FALSE, only accessions in acc\_filter will be included

in the database. Default FALSE.

scan Logical vector of length 1; should the sequence file be scanned for accessions

in acc\_filter prior to processing? Requires zgrep to be installed (so does not work on Windows). Only used if acc\_filter is not NULL and invert is

FALSE. Default FALSE.

#### **Details**

This function will automatically connect to the restez database.

#### Value

Logical

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

gb\_definition\_get

Get definition from GenBank

## **Description**

Return the definition line for an accession ID.

# Usage

```
gb_definition_get(id)
```

# **Arguments**

id

character, sequence accession ID(s)

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## Value

named vector of definitions, if no results found NULL

## See Also

```
ncbi_acc_get()
Other get: gb_fasta_get(), gb_organism_get(), gb_record_get(), gb_sequence_get(), gb_version_get()
```

# **Examples**

```
library(restez)
restez_path_set(filepath = tempdir())
demo_db_create(n = 5)
(def <- gb_definition_get(id = 'demo_1'))
(defs <- gb_definition_get(id = c('demo_1', 'demo_2')))

# delete demo after example
db_delete(everything = TRUE)</pre>
```

gb\_df\_create

Create GenBank data.frame

## **Description**

Make data.frame from columns vectors for nucleotide entries. As part of gb\_df\_generate().

# Usage

```
gb_df_create(accessions, versions, organisms, definitions, sequences, records)
```

# Arguments

```
accessions character, vector of accessions

versions character, vector of accessions + versions

organisms character, vector of organism names

definitions character, vector of sequence definitions

sequences character, vector of sequences

records character, vector of GenBank records in text format
```

#### Value

data.frame

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#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

gb\_df\_generate

Generate GenBank records data.frame

#### **Description**

For a list of records, construct a data.frame for insertion into SQL database.

## Usage

```
gb_df_generate(
  records,
  min_length = 0,
  max_length = NULL,
  acc_filter = NULL,
  invert = FALSE
)
```

#### **Arguments**

records	character, vector of GenBank records in text format
min_length	Minimum sequence length, default 0.
max_length	Maximum sequence length, default NULL.
acc_filter	Character vector; accessions to include or exclude from the database as specified by invert. $ \\$
invert	Logical vector of length 1; if TRUE, accessions in acc_filter will be excluded from the database; if FALSE, only accessions in acc_filter will be included in the database. Default FALSE.

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#### **Details**

The resulting data.frame has five columns: accession, organism, raw\_definition, raw\_sequence, raw\_record. The prefix 'raw\_' indicates the data has been converted to the raw format, see ?char-ToRaw, in order to save on RAM. The raw\_record contains the entire GenBank record in text format

Use acc\_filter and max and min sequence lengths to minimize the size of the database. All sequences have to be at least as long as min and less than or equal in length to max, unless max is NULL in which there is no maximum length. The final selection of sequences is the result of applying all filters (acc\_filter, min\_length, max\_length) in combination.

#### Value

data.frame, or NULL if no records pass filters

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

gb\_extract

Extract elements of a GenBank record

## Description

Return elements of GenBank record e.g. sequence, definition ...

#### Usage

```
gb_extract(
  record,
  what = c("accession", "version", "organism", "sequence", "definition", "locus",
        "features", "keywords")
)
```

gb\_fasta\_get 43

# **Arguments**

record GenBank record in text format, character

what Which element to extract

## **Details**

This function uses a REGEX to extract particular elements of a GenBank record. All of the what options return a single character with the exception of 'locus' or 'keywords' that return character vectors and 'features' that returns a list of lists for all features.

The accuracy of these functions cannot be guaranteed due to the enormity of the GenBank database. But the function is regularly tested on a range of GenBank records.

Note: all non-latin1 characters are converted to '-'.

## Value

character or list of lists (what='features') or named character vector (what='locus')

## **Examples**

```
library(restez)
data('record')
(gb_extract(record = record, what = 'locus'))
```

gb\_fasta\_get

Get fasta from GenBank

# Description

Get sequence and definition data in FASTA format. Equivalent to rettype='fasta' in rentrez::entrez\_fetch().

#### Usage

```
gb_fasta_get(id, width = 70)
```

# Arguments

id character, sequence accession ID(s)

width integer, maximum number of characters in a line

#### Value

named vector of fasta sequences, if no results found NULL

```
ncbi_acc_get()
Other get: gb_definition_get(), gb_organism_get(), gb_record_get(), gb_sequence_get(),
gb_version_get()
```

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## **Examples**

```
library(restez)
restez_path_set(filepath = tempdir())
demo_db_create(n = 5)
(fasta <- gb_fasta_get(id = 'demo_1'))
(fastas <- gb_fasta_get(id = c('demo_1', 'demo_2')))

# delete demo after example
db_delete(everything = TRUE)</pre>
```

gb\_organism\_get

Get organism from GenBank

## **Description**

Return the organism name for an accession ID.

# Usage

```
gb_organism_get(id)
```

# Arguments

id

character, sequence accession ID(s)

## Value

named vector of definitions, if no results found NULL

# See Also

```
ncbi_acc_get()
Other get: gb_definition_get(), gb_fasta_get(), gb_record_get(), gb_sequence_get(),
gb_version_get()
```

# **Examples**

```
library(restez)
restez_path_set(filepath = tempdir())
demo_db_create(n = 5)
(org <- gb_organism_get(id = 'demo_1'))
(orgs <- gb_organism_get(id = c('demo_1', 'demo_2')))
# delete demo after example
db_delete(everything = TRUE)</pre>
```

gb\_record\_get 45

gb\_record\_get

Get record from GenBank

# Description

Return the entire GenBank record for an accession ID. Equivalent to rettype='gb' in rentrez::entrez\_fetch().

# Usage

```
gb_record_get(id)
```

## **Arguments**

id

character, sequence accession ID(s)

## Value

named vector of records, if no results found NULL

#### See Also

```
ncbi_acc_get()
Other get: gb_definition_get(), gb_fasta_get(), gb_organism_get(), gb_sequence_get(),
gb_version_get()
```

# **Examples**

```
library(restez)
restez_path_set(filepath = tempdir())
demo_db_create(n = 5)
(rec <- gb_record_get(id = 'demo_1'))
(recs <- gb_record_get(id = c('demo_1', 'demo_2')))

# delete demo after example
db_delete(everything = TRUE)</pre>
```

gb\_sequence\_get

Get sequence from GenBank

# **Description**

Return the sequence(s) for a record(s) from the accession ID(s).

# Usage

```
gb_sequence_get(id, dnabin = FALSE)
```

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# **Arguments**

id character, sequence accession ID(s)

dnabin Logical vector of length 1; should the sequences be returned using the bit-level

coding scheme of the ape package? Default FALSE.

# **Details**

For more information about the dnabin format, see ape::DNAbin().

#### Value

named vector of sequences, if no results found NULL

# See Also

```
ncbi_acc_get()
Other get: gb_definition_get(), gb_fasta_get(), gb_organism_get(), gb_record_get(),
gb_version_get()
```

# **Examples**

```
library(restez)
restez_path_set(filepath = tempdir())
demo_db_create(n = 5)
(seq <- gb_sequence_get(id = 'demo_1'))
(seqs <- gb_sequence_get(id = c('demo_1', 'demo_2')))
(fasta_dnabin <- gb_sequence_get(id = 'demo_1', dnabin = TRUE))
# delete demo after example
db_delete(everything = TRUE)</pre>
```

gb\_sql\_add

Add to GenBank SQL database

## **Description**

Add records data.frame to SQL-like database.

# Usage

```
gb_sql_add(df)
```

# **Arguments**

df

Records data.frame

 $gb\_sql\_query$  47

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

gb\_sql\_query

Query the GenBank SQL

#### **Description**

Generic query function for retrieving data from the SQL database for the get functions.

#### Usage

```
gb_sql_query(nm, id)
```

#### **Arguments**

nm character, column name

id character, sequence accession ID(s)

#### Value

data.frame

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(),
```

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```
readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(),
search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(),
stat(), testdatadir_get()
```

gb\_version\_get

Get version from GenBank

# **Description**

Return the accession version for an accession ID.

# Usage

```
gb_version_get(id)
```

# **Arguments**

id

character, sequence accession ID(s)

## Value

named vector of versions, if no results found NULL

## See Also

```
ncbi_acc_get()
Other get: gb_definition_get(), gb_fasta_get(), gb_organism_get(), gb_record_get(),
gb_sequence_get()
```

# **Examples**

```
library(restez)
restez_path_set(filepath = tempdir())
demo_db_create(n = 5)
(ver <- gb_version_get(id = 'demo_1'))
(vers <- gb_version_get(id = c('demo_1', 'demo_2')))

# delete demo after example
db_delete(everything = TRUE)</pre>
```

has\_data 49

has\_data

Does the connected database have data?

#### Description

Returns TRUE if a restez SQL database has data.

## Usage

has\_data()

#### Value

Logical

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

identify\_downloadable\_files

Identify downloadable files

#### **Description**

Searches through the release notes for a GenBank release to find all listed .seq files. Returns a data.frame for all .seq files and their description.

## Usage

```
identify_downloadable_files()
```

## Value

data.frame

50 is\_in\_db

## See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

is\_in\_db

Is in db

## **Description**

Determine whether an id(s) is/are present in a database.

## Usage

```
is_in_db(id, db = "nucleotide")
```

#### Arguments

id character, sequence accession ID(s)
db character, database name

#### Value

named vector of booleans

# See Also

```
Other database: count_db_ids(), db_create(), db_delete(), db_download(), demo_db_create(),
list_db_ids()
```

## **Examples**

```
library(restez)
# set the restez path to a temporary dir
restez_path_set(filepath = tempdir())
# create demo database
demo_db_create(n = 5)
# in the demo, IDs are 'demo_1', 'demo_2' ...
```

last\_add\_get 51

```
ids <- c('thisisnotanid', 'demo_1', 'demo_2')
(is_in_db(id = ids))

# delete demo after example
db_delete(everything = TRUE)</pre>
```

last\_add\_get

Return date and time of the last added sequence

## **Description**

Return the date and time of the last added sequence as determined using the 'add\_log.tsv'.

#### Usage

```
last_add_get()
```

#### **Details**

If no file found, returns empty character vector.

#### Value

character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

52 last\_entry\_get

last\_dwnld\_get

Return date and time of the last download

#### **Description**

Return the date and time of the last download as determined using the 'download\_log.tsv'.

# Usage

```
last_dwnld_get()
```

#### **Details**

If no file found, returns empty character vector.

#### Value

character

## See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

last\_entry\_get

Return the last entry

## **Description**

Return the last entry from a tab-delimited log file.

## Usage

```
last_entry_get(fp)
```

latest\_genbank\_release 53

## Arguments

fp Filepath, character

#### Value

vector

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_add_get(), last_dwnld_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

latest\_genbank\_release

Retrieve latest GenBank release number

## **Description**

Downloads the latest GenBank release number and returns it.

## Usage

latest\_genbank\_release()

#### Value

character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(),
```

54 list\_db\_ids

```
gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(),
gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(),
last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), message_missing(),
mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(),
readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(),
search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(),
stat(), testdatadir_get()
```

latest\_genbank\_release\_notes

Download the latest GenBank Release Notes

## **Description**

Downloads the latest GenBank release notes to a user's restez download path.

#### Usage

latest\_genbank\_release\_notes()

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

list\_db\_ids

List database IDs

## **Description**

Return a vector of all IDs in a database.

## Usage

```
list_db_ids(db = "nucleotide", n = 100)
```

message\_missing 55

# **Arguments**

db character, database name

n Maximum number of IDs to return, if NULL returns all

## **Details**

Warning: can return very large vectors for large databases.

## Value

vector of characters

## See Also

```
Other database: count_db_ids(), db_create(), db_delete(), db_download(), demo_db_create(), is_in_db()
```

# **Examples**

```
library(restez)
restez_path_set(filepath = tempdir())
demo_db_create(n = 5)
# Warning: not recommended for real databases
# with potentially millions of IDs
all_ids <- list_db_ids()</pre>
# What shall we do with these IDs?
# ... how about make a mock fasta file
seqs <- gb_sequence_get(id = all_ids)</pre>
defs <- gb_definition_get(id = all_ids)</pre>
# paste together
fasta_seqs <- paste0('>', defs, '\n', seqs)
fasta_file <- paste0(fasta_seqs, collapse = '\n')</pre>
cat(fasta_file)
# delete after example
db_delete(everything = TRUE)
```

message\_missing

Produce message of missing IDs

# Description

Sends message to console stating number of missing IDs.

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#### Usage

```
message_missing(n)
```

# **Arguments**

n

Number of missing IDs

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

mock\_def

Mock def

# **Description**

Make a mock sequence definition. Designed to be part of a loop.

# Usage

```
mock_def(i)
```

## Arguments

i

integer, iterator

#### Value

character

mock\_gb\_df\_generate

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

mock\_gb\_df\_generate

Generate mock GenBank records data.frame

## **Description**

Make a mock nucleotide data.frame for entry into a demonstration SQL database.

#### **Usage**

```
mock_gb_df_generate(n)
```

#### Arguments

n

integer, number of entries

#### Value

data.frame

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(),
```

58 mock\_org

```
search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(),
stat(), testdatadir_get()
```

mock\_org

Mock org

# **Description**

Make a mock sequence organism. Designed to be part of a loop.

# Usage

mock\_org(i)

## Arguments

i

integer, iterator

## Value

character

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

mock\_rec 59

mock\_rec Mock rec

# Description

Create a mock GenBank record for demo-ing and testing purposes. Designed to be part of a loop. Accession, organism... etc. are optional arguments.

# Usage

```
mock_rec(
    i,
    definition = NULL,
    accession = NULL,
    version = NULL,
    organism = NULL,
    sequence = NULL
)
```

# Arguments

i integer, iterator
definition character
accession character
version character
organism character
sequence character

# Value

character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

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mock\_seq

Mock seq

## Description

Make a mock sequence. Designed to be part of a loop.

## Usage

```
mock_seq(i, sqlngth = 10)
```

## **Arguments**

i integer, iterator

sqlngth integer, sequence length

#### Value

character

## See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

ncbi\_acc\_get

Get accession numbers by querying NCBI GenBank

# Description

The query string can be formatted using GenBank advanced query terms to obtain accession numbers corresponding to a specific set of criteria.

predict\_datasizes 61

## Usage

```
ncbi_acc_get(query, strict = TRUE, drop_ver = TRUE)
```

#### **Arguments**

query Character vector of length 1; query string to search GenBank.

strict Logical vector of length 1; should an error be issued if the number of unique

accessions retrieved does not match the number of hits from GenBank? Default

TRUE.

drop\_ver Logical vector of length 1; should the version part of the accession number (e.g.,

'.1' in 'AB001538.1') be dropped? Default TRUE.

#### **Details**

Note this queries NCBI GenBank, not the local database generated with restez.

It can be used either to restrict the accessions used to construct the local database (acc\_filter argument of db\_create()) or to specify accessions to read from the local database (id argument of gb\_fasta\_get() and other gb\_\*\_get() functions).

## Value

Character vector; accession numbers resulting from query.

#### See Also

```
db_create(), gb_fasta_get()
```

## **Examples**

```
## Not run:
    # requires an internet connection
    cmin_accs <- ncbi_acc_get("Crepidomanes minutum")
    length(cmin_accs)
    head(cmin_accs)
## End(Not run)</pre>
```

predict\_datasizes

Print file size predictions to screen

## Description

Predicts the file sizes of the downloads and the database from the GenBank filesize information. Conversion factors are based on previous restez downloads.

## Usage

```
predict_datasizes(uncompressed_filesize)
```

62 print.status

#### **Arguments**

```
uncompressed_filesize

GBs of the stated filesize, numeric
```

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

print.status

Print method for status class

## **Description**

Prints to screen the three sections of the status class. Not meant to be used interactively.

# Usage

```
## S3 method for class 'status'
print(x, ...)
```

## **Arguments**

x Status object

... Other arguments (not used by this function)

readme\_log 63

readme\_log

Create README in restez\_path

#### **Description**

Write notes for the curious sorts who peruse the restez\_path.

# Usage

```
readme_log()
```

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

record

Example GenBank record

#### **Description**

Example GenBank record in text format for demonstration purposes.

## Usage

```
data("record")
```

## **Format**

A large character object containing record information and DNA sequence.

## Source

```
https://www.ncbi.nlm.nih.gov/nuccore/AY952423.1
```

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## References

GenBank

#### **Examples**

data(record)
cat(record)

restez\_connect

Connect to the restez database

## **Description**

Sets a connection to the local database.

#### Usage

```
restez_connect(read_only = FALSE)
```

# **Arguments**

read\_only

Logical; should the connection be made in read-only mode? Read-only mode is required for multiple R processes to access the database simultaneously. Default FALSE.

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

restez\_disconnect 65

restez\_disconnect

Disconnect from restez database

# **Description**

Safely disconnect from the restez connection

#### Usage

restez\_disconnect()

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

restez\_path\_check

Check restez filepath

# Description

Raises error if restez path does not exist.

#### Usage

restez\_path\_check()

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(),
```

restez\_path\_set

```
gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(),
gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(),
last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(),
message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(),
predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_rl(),
search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(),
stat(), testdatadir_get()
```

restez\_path\_get

Get restez path

## **Description**

Return filepath to where the restez database is stored.

#### Usage

```
restez_path_get()
```

## Value

character

#### See Also

```
Other setup: restez_path_set(), restez_path_unset(), restez_ready(), restez_status()
```

# **Examples**

```
library(restez)
# set a restez path with a tempdir
restez_path_set(filepath = tempdir())
# check what the set path is
(restez_path_get())
```

restez\_path\_set

Set restez path

# **Description**

Specify the filepath for the local GenBank database.

# Usage

```
restez_path_set(filepath)
```

restez\_path\_unset 67

# **Arguments**

filepath

character, valid filepath to the folder where the database should be stored.

#### **Details**

Adds 'restez\_path' to options(). In this path the folder 'restez' will be created and all downloaded and database files will be stored there.

#### See Also

```
Other setup: restez_path_get(), restez_path_unset(), restez_ready(), restez_status()
```

# **Examples**

```
## Not run:
library(restez)
restez_path_set(filepath = 'path/to/where/you/want/files/to/download')
## End(Not run)
```

restez\_path\_unset

Unset restez path

# **Description**

Set the restez path to NULL

# Usage

```
restez_path_unset()
```

```
Other setup: restez_path_get(), restez_path_set(), restez_ready(), restez_status()
```

68 restez\_rl

restez\_ready

Is restez ready?

# **Description**

Returns TRUE if a restez SQL database is available. Use restez\_status() for more information.

# Usage

```
restez_ready()
```

## Value

Logical

# See Also

```
Other setup: restez_path_get(), restez_path_set(), restez_path_unset(), restez_status()
```

# **Examples**

```
library(restez)
fp <- tempdir()
restez_path_set(filepath = fp)
demo_db_create(n = 5)
(restez_ready())
db_delete(everything = TRUE)
(restez_ready())</pre>
```

restez\_rl

Restez readline

# Description

Wrapper for base readline.

# Usage

```
restez_rl(prompt)
```

# Arguments

prompt

character, display text

# Value

character

restez\_status 69

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

restez\_status

Check restez status

## Description

Report to console current setup status of restez.

# Usage

```
restez_status(gb_check = FALSE)
```

## **Arguments**

gb\_check

Check whether last download was from latest GenBank release? Default FALSE.

#### **Details**

Set gb\_check=TRUE to see if your downloads are up-to-date.

#### Value

Status class

```
Other setup: restez_path_get(), restez_path_set(), restez_path_unset(), restez_ready()
```

70 search\_gz

## **Examples**

```
library(restez)
fp <- tempdir()
restez_path_set(filepath = fp)
demo_db_create(n = 5)
restez_status()
db_delete(everything = TRUE)
# Errors:
# restez_status()</pre>
```

search\_gz

Scan a gzipped file for text

# **Description**

Scans a zipped file for text strings and returns TRUE if any are present.

## Usage

```
search_gz(terms, path)
```

# **Arguments**

terms Character vector; search terms (most likely GenBank accession numbers)

path Path to the gzipped file to scan

#### Value

Logical

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

seshinfo\_log 71

seshinfo\_log

Log the system session information in restez path

# **Description**

Records the session and system information to file.

## Usage

seshinfo\_log()

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

setup

Set up test common test data

# Description

Creates temporary test folders.

#### Usage

setup()

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(),
```

72 slctn\_get

```
gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(),
gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(),
last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(),
message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(),
predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(),
restez_rl(), search_gz(), seshinfo_log(), slctn_get(), slctn_log(), sql_path_get(),
status_class(), stat(), testdatadir_get()
```

slctn\_get

Retrieve GenBank selections made by user

#### **Description**

Returns the selections made by the user.

## Usage

slctn\_get()

#### **Details**

If no file found, returns empty character vector.

#### Value

character vector

## See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_log(), sql\_path\_get(), status\_class(), stat(), testdatadir\_get()

slctn\_log 73

slctn\_log

Log the GenBank selection made by a user

## **Description**

This function is called whenever a user makes a selection with the db\_download(). It records GenBank numbers selections.

## Usage

```
slctn_log(selection)
```

## **Arguments**

selection

selected GenBank sequences, named vector

#### See Also

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(), restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), sql_path_get(), status_class(), stat(), testdatadir_get()
```

sql\_path\_get

Get SQL path

## **Description**

Return path to where SQL database is stored.

## Usage

```
sql_path_get()
```

## Value

character

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#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), status\_class(), stat(), testdatadir\_get()

stat

Print blue

#### **Description**

Print to console blue text to indicate a number/statistic.

## Usage

stat(...)

#### Arguments

Any number of text arguments to print, character

#### Value

coloured character encoding, character

```
Other private: add_rcrd_log(), cat_line(), char(), check_connection(), cleanup(), connected(), connection_get(), db_download_intern(), db_sqlngths_get(), db_sqlngths_log(), dir_size(), dwnld_path_get(), dwnld_rcrd_log(), entrez_fasta_get(), entrez_gb_get(), extract_accession(), extract_by_patterns(), extract_clean_sequence(), extract_definition(), extract_features(), extract_inforecpart(), extract_keywords(), extract_locus(), extract_organism(), extract_sequence(), extract_sequence(), extract_version(), file_download(), filename_log(), flatfile_read(), gb_build(), gb_df_create(), gb_df_generate(), gb_sql_add(), gb_sql_query(), gbrelease_check(), gbrelease_get(), gbrelease_log(), has_data(), identify_downloadable_files(), last_add_get(), last_dwnld_get(), last_entry_get(), latest_genbank_release_notes(), latest_genbank_release(), message_missing(), mock_def(), mock_gb_df_generate(), mock_org(), mock_rec(), mock_seq(), predict_datasizes(), readme_log(), restez_connect(), restez_disconnect(), restez_path_check(),
```

status\_class 75

```
restez_rl(), search_gz(), seshinfo_log(), setup(), slctn_get(), slctn_log(), sql_path_get(),
status_class(), testdatadir_get()
```

status\_class

Generate a list class for storing status information

## **Description**

Creates a three-part list for holding information on the status of the restez file path.

## Usage

```
status_class()
```

#### Value

Status class

#### See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), stat(), testdatadir\_get()

testdatadir\_get

Get test data directory

# Description

Get the folder containing test data.

## Usage

```
testdatadir_get()
```

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## See Also

Other private: add\_rcrd\_log(), cat\_line(), char(), check\_connection(), cleanup(), connected(), connection\_get(), db\_download\_intern(), db\_sqlngths\_get(), db\_sqlngths\_log(), dir\_size(), dwnld\_path\_get(), dwnld\_rcrd\_log(), entrez\_fasta\_get(), entrez\_gb\_get(), extract\_accession(), extract\_by\_patterns(), extract\_clean\_sequence(), extract\_definition(), extract\_features(), extract\_inforecpart(), extract\_keywords(), extract\_locus(), extract\_organism(), extract\_sequence(), extract\_version(), file\_download(), filename\_log(), flatfile\_read(), gb\_build(), gb\_df\_create(), gb\_df\_generate(), gb\_sql\_add(), gb\_sql\_query(), gbrelease\_check(), gbrelease\_get(), gbrelease\_log(), has\_data(), identify\_downloadable\_files(), last\_add\_get(), last\_add\_get(), last\_dwnld\_get(), last\_entry\_get(), latest\_genbank\_release\_notes(), latest\_genbank\_release(), message\_missing(), mock\_def(), mock\_gb\_df\_generate(), mock\_org(), mock\_rec(), mock\_seq(), predict\_datasizes(), readme\_log(), restez\_connect(), restez\_disconnect(), restez\_path\_check(), restez\_rl(), search\_gz(), seshinfo\_log(), setup(), slctn\_get(), slctn\_log(), sql\_path\_get(), status\_class(), stat()

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