Package 'mgi.report.reader'

July 7, 2024

```
Title Read Mouse Genome Informatics Reports
Version 0.1.3
Description Provides readers for easy and consistent importing of
       Mouse Genome Informatics (MGI) report files:
       <a href="https://www.informatics.jax.org/downloads/reports/index.html">https://www.informatics.jax.org/downloads/reports/index.html</a>. These data
       are provided by Baldarelli RM, Smith CL, Ringwald M, Richardson JE, Bult CJ,
       Mouse Genome Informatics Group (2024) <doi:10.1093/genetics/iyae031>.
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Description

chromosomes() returns mouse chromosome names.

Usage

```
chromosomes(autosomal = TRUE, sexual = TRUE, mitochondrial = TRUE)
```

Arguments

autosomal	Whether to include the autosomal chromosomes (1 thru 19).
sexual	Whether to include the sexual chromosomes (X and Y).
mitochondrial	Whether to include the mitochondrial chromosome (MT).

Value

A character vector of mouse chromosome names, or a subset thereof, or an empty character vector.

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Examples

```
# All chromosomes.
chromosomes()

# Autosomal chromosomes.
chromosomes(autosomal = TRUE, sexual = FALSE, mitochondrial = FALSE)
```

feature_types

Genome Feature types

Description

feature_types() returns different types of gene and genome features. For feature type definitions, see ?feature_type_definitions.

Usage

```
feature_types()
```

Value

A character vector of feature types' names.

Examples

```
feature_types()
```

feature_type_definitions

Genome Feature Type Definitions

Description

A dataset containing different types of gene and genome features along with their Sequence Ontology (SO) identifiers and definitions.

```
feature_type_definitions
```

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Format

A tibble with 71 rows and 3 variables:

feature_type Character. The type of gene or genome feature.

so_id Character. The Sequence Ontology identifier associated with the feature type.

definition Character. The definition of the feature type.

Source

The table in https://www.informatics.jax.org/userhelp/GENE_feature_types_help.shtml and a few other terms found in MGI reports.

Examples

```
print(feature_type_definitions, n = Inf)
```

marker_types

Genetic marker types

Description

marker_types() returns MGI marker types. See marker_type_definitions for the meaning of each type.

Usage

```
marker_types()
```

Value

A character vector.

```
marker_types()
```

marker_type_definitions

Genetic Marker Type Definitions

Description

A dataset of marker types definitions.

Use instead marker_types() for the marker type names as a single character vector.

Usage

```
marker_type_definitions
```

Format

A tibble with 10 rows and 2 variables:

```
marker_type Character. The type of genetic marker. definition Character. The definition of the marker type.
```

Source

The cross-references in the entry definition for marker at MGI glossary: https://www.informatics.jax.org/glossary/marker/.

Examples

```
print(marker_type_definitions, n = Inf)
```

```
open_marker_id_in_mgi Browse MGI markers identifiers online
```

Description

open_marker_id_in_mgi() launches the web browser and opens a tab for each MGI accession identifier on the Mouse Genome Informatics web interface: https://www.informatics.jax.org.

Usage

```
open_marker_id_in_mgi(marker_id)
```

Arguments

marker_id A character vector. MGI accession identifiers.

Value

Returns TRUE if successful, or FALSE otherwise. But note that this function is run for its side effect of launching the browser.

Examples

```
# Read about Acta1 (actin alpha 1, skeletal muscle) online.
open_marker_id_in_mgi("MGI:87902")

# `open_marker_id_in_mgi()` is vectorized, so you can open multiple pages.
# NB: think twice if you really need to open many tabs at once.
open_marker_id_in_mgi(c("MGI:87902", "MGI:87909"))
```

```
open_marker_symbol_in_mgi
```

Browse MGI markers symbols online

Description

open_marker_symbol_in_mgi() launches the web browser and opens a tab for each MGI symbol on the Mouse Genome Informatics web interface: https://www.informatics.jax.org.

Usage

```
open_marker_symbol_in_mgi(marker_symbol)
```

Arguments

marker_symbol A character vector. MGI marker symbols.

Value

Returns TRUE if successful, or FALSE otherwise. But note that this function is run for its side effect of launching the browser.

```
# Read about Acta1 (actin alpha 1, skeletal muscle) online.
open_marker_symbol_in_mgi("Acta1")

# `open_marker_symbol_in_mgi()` is vectorized, so you can open multiple pages.
# NB: think twice if you really need to open many tabs at once.
open_marker_symbol_in_mgi(c("Acta1", "Hes1"))
```

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read_report Read an MGI report

Description

read_report() imports data from an MGI report into R as a tidy data set.

You may call this function in two alternative ways:

- Using report_key: this is the easiest approach. A report key maps to a report currently hosted at MGI, e.g. read_report("marker_list2") reads MRK_List2.rpt directly from MGI server into R. See Supported Reports below for options.
- Using report_file and report_type: this approach is more flexible as you can read directly from a file or URL.

Supported Reports:

The set of currently supported reports:

```
reports
#> # A tibble: 13 x 4
     report_key
                              report_file
#>
                                                      report_type
                                                                    report_name
#>
      <chr>
                                 <chr>
                                                           <chr>
                                                                         <chr>
#> 1 marker list1
                              MRK_List1.rpt
                                                      MRK List1
                                                                    Mouse Gene~
#> 2 marker_list2
                              MRK_List2.rpt
                                                      MRK_List2
                                                                    Mouse Gene~
#> 3 marker_coordinates
                              MGI_MRK_Coord.rpt
                                                      MGI_MRK_Coord MGI Marker~
#> 4 gene_model_coordinates
                              MGI_Gene_Model_Coord.rpt MGI_Gene_Mod~ MGI Gene M~
#> 5 sequence_coordinates
                               MGI_GTGUP.gff
                                                      MGI_GTGUP
                                                                    MGI Sequen~
#> 6 genbank_refseq_ensembl_ids MRK_Sequence.rpt
                                                      MRK_Sequence MGI Marker~
#> 7 swiss_trembl_ids
                             MRK_SwissProt_TrEMBL.rpt MRK_SwissPro~ MGI Marker~
#> 8 swiss_prot_ids
                              MRK_SwissProt.rpt
                                                      MRK_SwissProt MGI Marker~
#> 9 gene_trap_ids
                              MRK_GeneTrap.rpt
                                                      MRK_GeneTrap MGI Marker~
#> 10 ensembl_ids
                              MRK_ENSEMBL.rpt
                                                      MRK_ENSEMBL MGI Marker~
#> 11 biotype_conflicts
                             MGI_BioTypeConflict.rpt MGI_BioTypeC~ MGI Marker~
#> 12 primers
                              PRB_PrimerSeq.rpt
                                                      PRB_PrimerSeg MGI Marker~
#> 13 interpro_domains
                             MGI_InterProDomains.rpt MGI_InterPro~ InterPro d~
```

```
read_report(
  report_key = NULL,
  report_file = NULL,
  report_type = NULL,
  n_max = Inf
)
```

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Arguments

report_key A character vector. A key used to uniquely refer to an MGI report.

report_file A character vector. The file path or URL to an MGI report file.

report_type A character vector. The type of an MGI report.

Maximum number of lines to read.

Value

A tibble with report data in tidy format. The set of variables is dependent on the specific report requested:

```
• For "marker_list1", see vignette("marker_list1").
```

- For "marker_list2", see vignette("marker_list2").
- For "marker_coordinates", see vignette("marker_coordinates").
- For "gene_model_coordinates", see vignette("gene_model_coordinates").
- For "sequence_coordinates", see vignette("sequence_coordinates").
- For "genbank_refseq_ensembl_ids", see vignette("genbank_refseq_ensembl_ids").
- For "swiss_trembl_ids", see vignette("swiss_trembl_ids").
- For "swiss_prot_ids", see vignette("swiss_prot_ids").
- For "gene_trap_ids", see vignette("gene_trap_ids").
- For "ensembl_ids", see vignette("ensembl_ids").
- For "biotype_conflicts", see vignette("biotype_conflicts").
- For "primers", see vignette("primers").
- For "interpro_domains", see vignette("interpro_domains").

report-attributes

Get MGI report specs by report key

Description

Set of functions to retrieve metadata details of a MGI report.

```
report_file(report_key)
report_name(report_key)
report_type(report_key)
report_url(report_key)
```

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Arguments

report_key A character vector. A key used to uniquely refer to an MGI report.

Value

A character vector:

```
    report_file(): report file name as hosted in https://www.informatics.jax.org/downloads/reports/.
```

```
• report_name(): report title.
```

- report_type(): report type.
- report_url(): report remote location.

Examples

```
report_file("marker_list1")
report_name("marker_list1")
report_type("marker_list1")
report_url("marker_list1")
```

reports

Supported MGI reports

Description

reports is a data set of supported MGI reports, meaning reports that {mgi.report.reader} can currently read into R. To browse all reports made available by MGI visit https://www.informatics.jax.org/downloads/reports/.

Usage

reports

Format

A tibble of 4 variables:

report_key A string key used to uniquely refer to an MGI report, which is only meaningful within the context of the {mgi.report.reader}.

report_file MGI report file name as hosted at https://www.informatics.jax.org/downloads/reports/.

report_type MGI report type. The type is used internally to find the appropriate reader for parsing, and is only meaningful within the context of {mgi.report.reader}.

report_name MGI report name. Report names are taken from https://www.informatics.jax.org/downloads/reports/index.html.

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Examples

reports

report_example

Report example

Description

report_example() returns the local path of an example report file. These files are typically very small and are useful for demonstrations. These are mostly used in the Examples section of functions and in unit tests.

Usage

```
report_example(report_file)
```

Arguments

```
report_file File basename.
```

Examples

```
report_example("MRK_List1-EX01.rpt")
report_example("MRK_List1-EX02.rpt")
report_example("MRK_List1-EX03.rpt")
```

report_last_modified Report last modification date

Description

report_last_modified() returns the last modified date and time of the report source: local file or remote file. If a local file, the modification date will be that indicated by the file system; if a remote file, the date of last update is that provided by HTTP header "last-modified".

MGI updates its reports weekly, every Thursday. However, not all reports are updated each week. The return value of this function is the closest you will get to a versioning of MGI report files.

```
report_last_modified(tbl)
```

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Arguments

tbl

Report data as a tibble.

Value

A last modified date-time as a POSIXct object.

Examples

```
if (FALSE) {
  markers <- read_report("marker_list1", n_max = 10L)

# When was the report file last updated?
  report_last_modified(markers)
}</pre>
```

report_source

Report source

Description

report_source() returns the source used to obtain the report data: a file path or an URL.

Usage

```
report_source(tbl)
```

Arguments

tbl

Report data as a tibble.

Value

A single string with an absolute path to a file on disk or an URL.

```
if (FALSE) {
  markers <- read_report("marker_list1", n_max = 10L)

# Where did the data come from?
  report_source(markers)
}</pre>
```

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```
symbol_to_identifier Convert marker symbols to updated marker identifiers
```

Description

symbol_to_identifier() remaps old marker symbols to, in-use, most up to date marker identifiers.

Usage

```
symbol_to_identifier(x, report_file = NULL, n_max = Inf)
```

Arguments

x A character vector of marker symbols to be remapped.

tomatically download the report from https://www.informatics.jax.org/

downloads/reports/MRK_List1.rpt.

n_max Maximum number of lines to read from the report_file.

Examples

```
rpt_ex01 <- report_example("MRK_List1-EX01.rpt")
read_report(report_file = rpt_ex01, report_type = "MRK_List1") |>
    dplyr::select("marker_status", "marker_symbol", "marker_id_now")

# NB:
# - "1700024N20Rik" has two conflicting mappings, so maps to `NA`.
# - "Hes1" is not present in MRK_List1-EX01.rpt, so maps to `NA`.
# - "Plpbp" (official) and "Prosc" (withdrawn) both map to "MGI:1891207"

marker_symbols <- c("2200002F22Rik", "Plpbp", "Prosc", "1700024N20Rik", "Hes1")
symbol_to_identifier(x = marker_symbols, report_file = rpt_ex01)</pre>
```

symbol_to_symbol

Update marker symbols

Description

```
symbol_to_symbol() remaps old marker symbols to, in-use, most up to date symbols.
```

```
symbol_to_symbol(x, report_file = NULL, n_max = Inf)
```

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Arguments

x A character vector of marker symbols to be remapped.

report_file The path to a MRK_List1.rpt file. Leave this as NULL and the function will automatically download the report from https://www.informatics.jax.org/downloads/reports/MRK_List1.rpt.

n_max Maximum number of lines to read from the report_file.

Value

A character vector of most up to date symbols.

```
rpt_ex01 <- report_example("MRK_List1-EX01.rpt")
read_report(report_file = rpt_ex01, report_type = "MRK_List1") |>
    dplyr::select("marker_status", "marker_symbol", "marker_symbol_now")

# NB:
# - "1700024N20Rik" has two conflicting mappings, so maps to `NA`.
# - "Hes1" is not present in MRK_List1-EX01.rpt, so maps to `NA`.
# - "Plpbp" (official) and "Prosc" (withdrawn) both map to "Plpbp"

marker_symbols <- c("2200002F22Rik", "Plpbp", "Prosc", "1700024N20Rik", "Hes1")
symbol_to_symbol(x = marker_symbols, report_file = rpt_ex01)</pre>
```

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