Package 'aliases2entrez'

October 12, 2022
Title Converts Human gene symbols to entrez IDs
Version 0.1.2
Description Queries multiple resources authors HGNC (2019) https://www.genenames.org , authors limma (2015) doi:10.1093/nar/gkv007 to find the correspondence between evolving nomenclature of human gene symbols, aliases, previous symbols or synonyms with stable, curated gene entrezID from NCBI database. This allows fast, accurate and up-to-date correspondence between human gene expression datasets from various date and platform (e.g. gene symbol: BRCA1 - ID: 672).
<pre>BugReports https://github.com/peyronlab/aliases2entrez/issues</pre>
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Author Raphael Bonnet [aut, cre] (Université Côte d'Azur), Lee Mariault [ctb] (Université Côte d'Azur), Jean-François Peyron [aut] (Inserm)
Maintainer Raphael Bonnet <raphael.bonnet@univ-cotedazur.fr></raphael.bonnet@univ-cotedazur.fr>
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convert_symbols

Multi resources gene symbols conversion to entrez ID (Human)

Description

This function is used to convert gene symbols, previous symbols or aliases to gene entrez ID It performs :

- -a gene query to limma::alias2Symbol to map gene alias to official symbols
- -looks for LOC* symbols
- -tries to find correspondence within HGNC database
- -queries org.Hs.eg.db
- -checks again with adaptive symbol parsing (e.g. transforms BRCA-1 to BRCA1)

Usage

```
convert_symbols(symbols, HGNC, c=1)
```

Arguments

symbols gene matrix from which rownames (gene symbols) are extracted

HGNC correspondence file.

c number of cores to use for parallel processes

Value

returns a vector containing IDs if match were found or NA if unknown or withdrawn symbol

Examples

```
# import the correspondence file
file <- system.file("extdata", "HGNC.txt", package = "aliases2entrez")
HGNC <- read.delim(file)
# alternatively update a new one with update_symbols()
symbols <- c("BRCA1", "TP53")
# run the main function
ids <- convert_symbols(symbols, HGNC)</pre>
```

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update_symbols

Update last HGNC correspondence database

Description

This function is used to update gene symbol correspondence from HGNC database

Usage

```
update_symbols(url=NULL)
```

Arguments

url

user can provide url (default is NULL)

Value

returns a data.frame containing gene symbols with status, previous symbols and synonyms as well as their corresponding entrezIDs

Examples

```
HGNC <- update_symbols()</pre>
```

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