LowMACAAnnotation

February 26, 2025

getLowMACAAnnotationData

Retrieve Data from LowMACAAnnotation Package

Description

Functions to retrieve protein level, Pfam level and aliases data from LowMACAAnnotation Package.

Usage

Retrieve the dataset containing protein level information
getMyUni()

Retrieve the dataset containing Pfam level information
getMyPfam()

Retrieve the dataset containing Gene Symbols aliases
getMyAlias()

Value

All the functions return dataframes

See Also

myUni myPfam myAlias

Examples

```
myUni <- getMyUni()
myPfam <- getMyPfam()
myAlias <- getMyAlias()</pre>
```

myAlias

Description

A simple parsing of the HGNC database to map aliases and previous symbols to correct official Gene Symbols

Usage

data("myAlias")

Format

A data frame with the following 2 variables.

- Alias a character vector representing all the possible aliases and previous symbols for official Gene Symbols
- Official_Gene_Symbol a character vector representing the approved and official Gene Symbol for HGNC database
- Locus_Group a character vector representing all the possible locus groups in HGNC database, like protein coding, RNA, pseudogene etc.
- Locus_Type a character vector representing all the possible locus types in HGNC database. It is a specification of locus group
- MappedByLowMACA a character vector of yes and no if the gene is included in myUni.RData

Source

HGNC

Examples

```
#Load myAlias and show its structure
myAlias <- getMyAlias()
str(myAlias)</pre>
```

myPfam

Pfam-A for LowMACA package

Description

This dataset comprises all the Pfam-A entries mapped for LowMACA

Usage

data("myPfam")

myUni

Format

A data frame with the following 11 variables.

Entry a character vector of Uniprot entries

Envelope_Start a numeric vector of starts of the pfam domain relative to the reference protein **Envelope_End** a numeric vector of ends of the pfam domain relative to the reference protein

Pfam_ID a character vector of Pfam IDs in the form of PF###### supported by LowMACA

Pfam_Name a character vector of full Pfam domain names

Type a character vector. One of the following: "Domain" "Family" "Repeat" or "Motif"

Clan_ID a numeric vector of Clan IDs, a sort of families of Pfam domains

Entrez a numeric vector of Entrez IDs

UNIPROT a character vector of Uniprot entries in format "name_HUMAN"

Gene_Symbol a character vector of official Gene Symbols

Pfam_Fasta a character vector of amino acid sequences of corresponding Pfam

Details

This is the result of a merge between Pfam-A, Uniprot, HGNC databases

Source

Pfam website

Examples

#Load and show contents of myPfam
data(myPfam)
str(myPfam)

myUni

Uniprot database for LowMACA

Description

This dataset comprises all the Uniprot entries mapped for LowMACA

Usage

data("myUni")

Format

A data frame with 9 variables.

Gene_Symbol a character vector of official Gene Symbols

Entrez a numeric vector of Entrez IDs

UNIPROT a character vector of Uniprot entries in "name_HUMAN" format

Entry a character vector of Uniprot entries

HGNC a character vector of gene names as HGNC numbers
Approved_Name a character vector of approved extended gene names
Protein.name a character vector of approved extended protein names
Chromosome a character vector of chromosomic cytoband positions
AMINO_SEQ a character vector of amino acid sequences for Uniprot entries

Details

This dataset is a 1 to 1 mapping of genes to their canonical proteins or main proteins

Source

Uniprot

Examples

```
#Load dataset and show its structure
data(myUni)
str(myUni)
```

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