

Package ‘PEIMAN2’

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Title Post-Translational Modification Enrichment, Integration, and Matching Analysis

Version 0.1.0

Description Functions and mined database from 'UniProt' focusing on post-translational modifications to do single enrichment analysis (SEA) and protein set enrichment analysis (PSEA). Payman Nickchi, Mehdi Mirzaie, Marc Baumann, Amir Ata Saei, Mohieddin Jafari (2022) <bioRxiv:10.1101/2022.11.09.515610>.

License GPL (>= 3)

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Suggests knitr, rmarkdown, testthat (>= 3.0.0)

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exmplData1	<i>Example dataset1</i>
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Description

A dataset with randomly selected proteins from UniProt.

Usage

exmplData1

Format

A list with 2 elements:

p11 97 randomly selected Homo sapiens (Human) proteins randomly selected from UniProt.

p12 45 randomly selected Homo sapiens (Human) proteins randomly selected from UniProt. ...

Source

<https://www.uniprot.org/>

exmplData2	<i>Example dataset 2</i>
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Description

Proteins of rat hippocampus proteome.

Usage

exmplData2

Format

A dataframe with 209 rows and 2 columns.

UniProtAC UniProt accession code of proteins

Score Check with MJ ...

Source

[doi:molpharm.120.000210](https://doi.org/10.26434/chemrxiv-2019-07-01-molpharm.120.000210)

getTaxonomyName *Return the exact taxonomy name for list of protein*

Description

getTaxonomyName get a character vector of proteins with their UniProt accession code and returns the exact taxonomy code.

Usage

```
getTaxonomyName(x)
```

Arguments

x A character vector with each entry presenting a protein UniProt accession code.

Value

The exact taxonomy name

Examples

```
getTaxonomyName(x = exmplData1$p11)
```

mod_ont *Database of protein modifications*

Description

Ontology database for post-translational modification terms. For more details, see the reference.

Usage

```
data(mod_ont)
```

Format

A data frame with 2102 rows and 3 variables

Details

- id
- name
- def

Source

<https://raw.githubusercontent.com/HUPO-PSI/psi-mod-CV/master/PSI-MOD.obo>

plotEnrichment	<i>Plot and match singular enrichment results</i>
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Description

This function can be used to plot results of singular enrichment analysis for one set of protein. It can also be used to integrate and match the results of two separate singular enrichment analysis and plot the common PTMs. For more details please see examples.

Usage

```
plotEnrichment(x, y = NULL, sig.level = 0.05, number.rep = NULL)
```

Arguments

x	A data frame that contains singular enrichment results generated by runEnrichment
y	Default value is NULL. If provided by a singular enrichment results, the matching results of x and y are plotted.
sig.level	The significance level to select post-translational modification (based on their corrected p-value). Note that sig.level applies to both x and y simultaneously.
number.rep	Only plot PTM terms that occurred more than a specific number of times in UniProt database. This number is set by number.rep parameter. The default value is NULL.

Value

Plot.

Examples

```
## Enrichment analysis for the first protein list
enrich1 <- runEnrichment(protein = exmplData1$pl1, os.name = 'Homo sapiens (Human)')
## Plot results for first protein list
plotEnrichment(x = enrich1)

## Enrichment analysis for the second protein list
enrich2 <- runEnrichment(protein = exmplData1$pl2, os.name = 'Homo sapiens (Human)')
## Plot results for second protein list
plotEnrichment(x = enrich2)

## Integrate and match the results of two separate singular enrichment analysis
plotEnrichment(x = enrich1, y = enrich2)
plotEnrichment(x = enrich1, y = enrich2, number.rep = 100)
```

plotPSEA	<i>Plot the results of protein set enrichment analysis (PSEA)</i>
----------	---

Description

plotPSEA can be used to plot the results of protein set enrichment analysis (psea) for a set of proteins obtained from an experiment.

Usage

```
plotPSEA(x, y = NULL, sig.level = 0.05, number.rep = NULL)
```

Arguments

x	A data frame returned by runPSEA function.
y	Default value is NULL. If provided by a protein set enrichment results, the matching results of x and y are plotted.
sig.level	The significance level applied on adjusted p-value by permutation to filter pathways for plotting. The default value is 0.05
number.rep	Only plot PTM terms that occurred more than a specific number of times in UniProt. This number is set by number.rep parameter. The default value is NULL.

Value

Plot

Examples

```
# We recommend at least nperm = 1000.  
# The number of permutations was reduced to 10  
# to accommodate CRAN policy on examples (run time <= 5 seconds).  
psea_res <- runPSEA(protein = exmplData2, os.name = 'Rattus norvegicus (Rat)', nperm = 10)  
plotPSEA(psea_res, sig.level = 0.05)
```

plotRunningScore	<i>Plot running score plot for the results of psea</i>
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Description

This function takes results generated by [runPSEA](#). It plots running enrichment score of ranked protein for each PTM.

Usage

```
plotRunningScore(  
  x,  
  nplot = length(x$psea.result),  
  type = "l",  
  lty = 1,  
  lwd = 3,  
  cex = 1.2,  
  cex.axis = 1.2,  
  cex.lab = 1.1,  
  col = "blue"  
)
```

Arguments

x	A list of 6 generated by runPSEA function.
nplot	An integer that defines the number of running score plots to show. Default value is the number of enriched PTMs in x.
type	Type of line used in the plot.
lty	A list of 6 generated by runPSEA function.
lwd	line width
cex	Specify the size of the title text
cex.axis	Specify the size of the tick label
cex.lab	Specify the size of the axis label text
col	Color of running enrichment score line

Value

Plot

Examples

```
# We recommend at least nperm = 1000.  
# The number of permutations was reduced to 10  
# to accommodate CRAN policy on examples (run time <= 5 seconds).  
psea_res <- runPSEA(protein = exmplData2, os.name = 'Rattus norvegicus (Rat)', nperm = 10)  
plotRunningScore(x = psea_res)
```

psea2mass

Translate PSEA results for Mass Spectrometry searching tools

Description

This function translates protein set enrichment analysis results and extracts the required information for mass spectrometry searching tools. The subset of protein modifications is from <https://raw.githubusercontent.com/HUPO-PSI/psi-mod-CV/master/PSI-MOD.obo>.

Usage

```
psea2mass(x, sig.level = 0.05, number.rep = NULL)
```

Arguments

x	A list of psea results generated by runPSEA function.
sig.level	The significance level to filter PTMs (applies on adjusted p-value). Default value is 0.05
number.rep	Only consider PTM terms that occurred more than a specific number of times in UniProt. This number is set by number.rep parameter. The default value is NULL.

Value

A database of subset of protein modifications:

- id: a unique identification for each subset of protein modifications, PSI-MOD.
- name: the name of modification.
- def: definition of PSI-MOD definition

Examples

```
# We recommend at least nperm = 1000.  
# The number of permutations was reduced to 10  
# to accommodate CRAN policy on examples (run time <= 5 seconds).  
psea_res <- runPSEA(protein = exmplData2, os.name = 'Rattus norvegicus (Rat)', nperm = 10)  
MS <- psea2mass(x = psea_res, sig.level = 0.05)
```

ptmlist	<i>Controlled vocabulary for post-translational modifications (PTM) terms</i>
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Description

This dataframe lists the posttranslational modifications used in the UniProt knowledgebase (Swiss-Prot and TrEMBL). The columns in this dataframe are as follows:

Usage

```
data(ptmlist)
```

Format

A data frame with 686 rows and 5 variables

Details

- ID Identifier (FT description)
- AC Accession (PTM-xxxx)
- KW Keyword
- FT Feature key
- DR Cross-reference to external databases

Source

<https://ftp.uniprot.org/pub/databases/uniprot/knowledgebase/complete/docs/ptmlist.txt>

runEnrichment	<i>Run singular enrichment analysis (SEA) for a given list of protein</i>
---------------	---

Description

This function takes proteins with their UniProt accession code, runs singular enrichment (SEA) analysis, and returns enrichment results.

Usage

```
runEnrichment(protein, os.name, p.adj.method = "BH")
```


Arguments

protein	A character vector with protein UniProt accession codes.
os.name	A character vector of length one with exact taxonomy name of species. If you do not know the the exact taxonomy name of species you are working with, please read getTaxonomyName .
p.adj.method	The adjustment method to correct for multiple testing. The default value is 'BH'. Run/see p.adjust.methods to get a list of possible methods.

Value

The result is a dataframe with the following columns:

- PTM: Post-translational modification (PTM) keyword
- FreqinUniprot: The total number of proteins in UniProt with this PTM.
- FreqinList: The total number of proteins in the given list with this PTM.
- Sample: Number of proteins in the given list.
- Population: Total number of proteins in the current version of PEIMAN database with this PTM.
- pvalue: The p-value obtained from hypergeometric test (enrichment analysis).
- corrected pvalue: Adjusted p-value to correct for multiple testing.
- AC: Uniprot accession code (AC) of proteins with each PTM.

Examples

```
enrich1 <- runEnrichment(protein = exmplData1$p11, os.name = 'Homo sapiens (Human)')
```

runPSEA

Run Protein Set Enrichment Analysis (PSEA)

Description

This is the main function to run protein set enrichment analysis for a list of proteins and their score.

Usage

```
runPSEA(
  protein,
  os.name,
  pexponent = 1,
  nperm = 1000,
  p.adj.method = "fdr",
  sig.level = 0.05,
  minSize = 1
)
```

Arguments

protein	A dataframe with two columns. First column should be protein accession code, second column is the score.
os.name	A character vector of length one with exact taxonomy name of species. If you do not know the exact taxonomy name of species you are working with, please read getTaxonomyName .
pexponent	Enrichment weighting exponent, p. For values of $p < 1$, one can detect incoherent patterns in a set of protein. If one expects a small number of proteins to be coherent in a large set, then $p > 1$ is a good choice.
nperm	Number of permutation to estimate false discovery rate (FDR). Default value is 1000.
p.adj.method	The adjustment method to correct p-values for multiple testing in enrichment. Run <code>p.adjust.methods()</code> to get a list of possible methods.
sig.level	The significance level to filter PTM (applies on adjusted p-value)
minSize	PTMs with the number of proteins below this threshold are excluded.

Value

Returns a list of 6: 1: A dataframe with protein set enrichment analysis (PSEA) results. Every row corresponds to a post-translational modification (PTM) pathway.

- PTM: PTM keyword
- pval: p-value for singular enrichment analysis
- pvaladj: adjusted p-value
- FreqinUniProt: The frequency of PTM in UniProt
- FreqinList: The frequency of PTM in the given list
- ES: enrichment score
- NES: enrichment score normalized to mean enrichment of random samples of the same size
- nMoreExtreme: number of times the permuted sample resulted in a profile with a larger ES value than $\text{abs}(\text{ES})$
- size: Number of proteins with the PTM
- Enrichment: Whether the proteins in the pathway have been enriched in the list.
- AC: Uniprot accession code (AC) of proteins with each PTM.
- leadingEdge:

Examples

```
# We recommend at least nperm = 1000.
# The number of permutations was reduced to 10
# to accommodate CRAN policy on examples (run time <= 5 seconds).
psea_res <- runPSEA(protein = exmplData2, os.name = 'Rattus norvegicus (Rat)', nperm = 10)
```

`sea2mass`*Translate SEA results for Mass Spectrometry searching tools*

Description

This function translates singular enrichment analysis results and extracts the required information for mass spectrometry searching tools. The subset of protein modifications is from <https://raw.githubusercontent.com/HUPO-PSI/psi-mod-CV/master/PSI-MOD.obo>.

Usage

```
sea2mass(x, sig.level = 0.05, number.rep = NULL)
```

Arguments

<code>x</code>	A dataframe of single enrichment analysis results generated by <code>runEnrichment</code> function.
<code>sig.level</code>	The significance level to filter pathways (applies on adjusted p-value). Default value is 0.05.
<code>number.rep</code>	Only consider PTM terms that occurred more than a specific number of times in UniProt. This number is set by <code>number.rep</code> parameter. The default value is NULL.

Value

A database of subset of protein modifications:

- `id`: a unique identification for each subset of protein modifications, PSI-MOD.
- `name`: the name of modification.
- `def`: definition of PSI-MOD definition

Examples

```
enrich1 <- runEnrichment(protein = exmplData1$p11, os.name = 'Homo sapiens (Human)')
MS      <- sea2mass(x = enrich1, sig.level = 0.05)
```

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