Package 'AlteredPQR'

July 21, 2025

Type Package Title Detection of Altered Protein Quantitative Relationships Version 0.1.0 Author Marija Buljan [aut, cre] Maintainer Marija Buljan <marija.buljan@empa.ch> Description Inference of protein complex states from quantitative proteomics data. The package takes information on known stable protein interactions (i.e. protein components of the same complex) and assesses how protein quantitative ratios change between different conditions. It reports protein pairs for which relative protein quantities to each other have been significantly altered in the tested condition. License GPL (>= 3) **Encoding** UTF-8 LazyData true NeedsCompilation no **Depends** R (>= 3.5.0) Suggests tools **Repository** CRAN Date/Publication 2023-06-10 14:00:02 UTC

Contents

Index																														5
	quant_data	_all	 	•	• •	•	•	 •	•	•	•	•		•	•	•	•	 •	•	•	•	•	•	•	•	• •	•	•	•	4
	int_pairs .		 		• •	•		 •		•	•	•	• •			•	•			•	•	•	•			• •	•	•	•	4
	CorShift .		 			•		 •		•	•	• •		•		•	•			•	•	•					•	•	•	3
	AlteredPQI	_																												

1

AlteredPQR_RB

Description

The function identifies outliers in protein pair log ratios compared to the reference set of measurements.

Usage

```
AlteredPQR_RB (modif_z_score_threshold = 3.5, fraction_of_samples_threshold = 0.10,
modif = 1, filter_variable_in_ref_set = "NO", write_table = "NO", print_recomm = "NO",
quant_data_all_local = quant_data_all,
cols_with_reference_data_local = cols_with_reference_data)
```

Arguments

<pre>modif_z_score_threshold</pre>										
	Numeric value defining a threshold to consider log ratio in the tested sample as a outlier.									
fraction_of_sa	ples_threshold									
	Numeric value defining a fraction of samples that need to be classified as outliers for the protein pair to be included in the results table.									
modif	Numeric value defining a modifier value that is used to idenify proteins in the pair that contributed to the outlier signal. The higher the variable 'modif', the higher the modified z score value the single protein needs to have (compared to its own values in the reference samples) to be considered as an outlier in at least half of the samples classified as outliers for the protein pair. Proteins classified as outliers in their own measuremnts are listed as 'driving the signal' in the reults table.									
filter_variable_in_ref_set										
	Option (T or F) to exclude from the results table protein pairs that strongly varied in the reference samples.									
write_table	Option (T or F) to save results table as a text file.									
print_recomm	Option (T or F) to get information on the distribution of all modified z-scores in the test samples and recommendation on the thresholds for the user defined qunatitative proteomics dataset.									
quant_data_all	_local									
cols with refe	A data matrix with quantiative proteomics measurements in which rows repre- sent uniprot protein identifiers, and columns samples. rence_data_local									
	Numeric vector with information on columns that contain reference data.									

Value

representative_pairs table

CorShift

Author(s)

Marija Buljan <marija.buljan@empa.ch>

Examples

```
data("int_pairs", package = "AlteredPQR")
data("quant_data_all", package = "AlteredPQR")
cols_with_reference_data = 1:23
RepresentativePairs = AlteredPQR_RB()
```

```
CorShift
```

Changes in correlation trends

Description

The function identifies instances in which two proteins correlate strongly only in one of the two studied groups.

Usage

```
CorShift(samplesA = samplesGroupA, samplesB = samplesGroupB, shift_threshold = 0.6,
writeTable = FALSE, min_cor_in_samples = 0.6, cor_signif = 0.01,
quant_data_all_local = quant_data_all, int_pairs_local = int_pairs)
```

Arguments

samplesA	Numeric vector with information on column numbers for the samples in the first group for the comparison.								
samplesB	Numeric vector with information on column numbers for the samples in second group for the comparison.								
shift_threshold									
	Numeric value defining a minimum thresold of the Pearson correlation value between the two sample groups in order for them to be included in the results table.								
writeTable	Option (T or F) to save results table as a text file.								
min_cor_in_samples									
	Numeric value defining a minimum Pearson correlation value of protein quanti- ties, which is taken as a threshold to consider that two proteins correlate in either of the two compared groups.								
cor_signif	Numeric value defining a maximum allowed p-value for the Pearson correlation, which is taken as a threshold to consider that quantitative measurements for the two proteins correlate significantly in either of the two compared groups.								
quant_data_all_local									
	A data matrix with quantiative proteomics measurements in which rows repre- sent uniprot protein identifiers, and columns samples.								
int_pairs_loca	1								
	A data matrix with two columns. Rows contain information on interacting pro- tein pairs.								

Value

cor_table table

Author(s)

Marija Buljan <marija.buljan@empa.ch>

Examples

```
data("int_pairs", package = "AlteredPQR")
data("quant_data_all", package = "AlteredPQR")
samplesGroupA = 1:23
samplesGroupB = (1+23):(23+18)
cor_results = CorShift()
```

int_pairs

Description

Protein pairs that can form stable interactions.

Author(s)

Marija Buljan <marija.buljan@empa.ch>

quant_data_all Proteomic measurements data matrix

Protein pairs

Description

Quantitative proteomics measurements; columns are samples, rows are proteins (Uniprot IDs).

Author(s)

Marija Buljan <marija.buljan@empa.ch>

Index

 $\texttt{AlteredPQR_RB, 2}$

 ${\tt CorShift}, {\tt 3}$

int_pairs,4

quant_data_all, 4