

# Package ‘HPiP’

August 20, 2025

**Type** Package

**Title** Host-Pathogen Interaction Prediction

**Version** 1.14.0

**Description** HPiP (Host-Pathogen Interaction Prediction) uses an ensemble learning algorithm for prediction of host-pathogen protein-protein interactions (HP-PPIs) using structural and physicochemical descriptors computed from amino acid-composition of host and pathogen proteins. The proposed package can effectively address data shortages and data unavailability for HP-PPI network reconstructions. Moreover, establishing computational frameworks in that regard will reveal mechanistic insights into infectious diseases and suggest potential HP-PPI targets, thus narrowing down the range of possible candidates for subsequent wet-lab experimental validations.

**Depends** R (>= 4.1)

**Imports** dplyr (>= 1.0.6), httr (>= 1.4.2), readr, tidyr, tibble, utils, stringr, magrittr, caret, corrplot, ggplot2, pROC, PRROC, igraph, graphics, stats, purrr, grDevices, protr, MCL

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**Encoding** UTF-8

**URL** <https://github.com/mrbakhsh/HPiP>

**BugReports** <https://github.com/mrbakhsh/HPiP/issues>

**VignetteBuilder** knitr

**Suggests** rmarkdown, colorspace, e1071, kernlab, ranger, SummarizedExperiment, Biostrings, randomForest, gprofiler2, gridExtra, ggthemes, BiocStyle, BiocGenerics, RUnit, tools, knitr

**biocViews** Proteomics, SystemsBiology, NetworkInference, StructuralPrediction, GenePrediction, Network

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## Contents

|                                 |    |
|---------------------------------|----|
| calculateAAC . . . . .          | 3  |
| calculateAutocor . . . . .      | 4  |
| calculateBE . . . . .           | 5  |
| calculateCTDC . . . . .         | 6  |
| calculateCTDD . . . . .         | 7  |
| calculateCTDT . . . . .         | 8  |
| calculateCTriad . . . . .       | 9  |
| calculateDC . . . . .           | 10 |
| calculateF . . . . .            | 11 |
| calculateKSAAP . . . . .        | 12 |
| calculateQD_Sm . . . . .        | 13 |
| calculateTC . . . . .           | 14 |
| calculateTC_Sm . . . . .        | 15 |
| corr_plot . . . . .             | 16 |
| enrich.df . . . . .             | 17 |
| enrichfindP . . . . .           | 17 |
| enrichfind_cpx . . . . .        | 18 |
| enrichfind_hp . . . . .         | 19 |
| enrichplot . . . . .            | 20 |
| example_data . . . . .          | 21 |
| filter_missing_values . . . . . | 22 |
| FreqInteractors . . . . .       | 22 |
| FSmethod . . . . .              | 23 |
| getFASTA . . . . .              | 25 |
| getHPI . . . . .                | 26 |
| get_negativePPI . . . . .       | 27 |
| get_positivePPI . . . . .       | 28 |
| Gold_ReferenceSet . . . . .     | 29 |
| host_se . . . . .               | 30 |
| impute_missing_data . . . . .   | 30 |
| plotPPI . . . . .               | 31 |
| predicted_PPIs . . . . .        | 32 |
| pred_ensembl . . . . .          | 32 |
| run_clustering . . . . .        | 35 |
| unlabel_data . . . . .          | 36 |
| UP000464024_df . . . . .        | 36 |

|                     |           |
|---------------------|-----------|
| <i>calculateAAC</i> | 3         |
| var_imp . . . . .   | 37        |
| viral_se . . . . .  | 38        |
| <b>Index</b>        | <b>39</b> |

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|              |                                                          |
|--------------|----------------------------------------------------------|
| calculateAAC | <i>Calculate Amino Acid Composition (AAC) Descriptor</i> |
|--------------|----------------------------------------------------------|

---

**Description**

This function calculates Amino Acid Composition (AAC) descriptor for the data input.

**Usage**

```
calculateAAC(x)
```

**Arguments**

x                      A data.frame containing gene/protein names and their fasta sequences.

**Details**

```
calculateAAC
```

**Value**

A length 20 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Dey, L., Chakraborty, S., and Mukhopadhyay, A. (2020). Machine learning techniques for sequence-based prediction of viral–host interactions between SARS-CoV-2 and human proteins. *Biomed. J.* 43, 438–450.

**See Also**

See [calculateDC](#) and [calculateTC](#) for Dipeptide Composition and Tripeptide Composition descriptors.

**Examples**

```
data(UP000464024_df)
x_df <- calculateAAC(UP000464024_df)
head(x_df, n = 2L)
```

calculateAutocor

*Calculate Autocorrelation Descriptors*

## Description

This function calculates autocorrelation descriptors:

- moran - moran autocorrelation, (Dim: length(target.props) \* nlag).
- geary - geary autocorrelation, (Dim: length(target.props) \* nlag).
- moreaubroto - moreau-broto autocorrelation, (Dim: length(target.props) \* nlag).

## Usage

```
calculateAutocor(
  x,
  target.props = c("CIDH920105", "BHAR880101", "CHAM820101", "CHAM820102",
    "CHOC760101", "BIGC670101", "CHAM810101", "DAYM780201"),
  nlag = 30L,
  type = c("moran", "geary", "moreaubroto")
)
```

## Arguments

|              |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |
|--------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| x            | A data.frame containing gene/protein names and their fasta sequences.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| target.props | A character vector, specifying the accession number of the target properties. 8 properties are used by default, as listed below:<br><b>AccNo. CIDH920105</b> Normalized average hydrophobicity scales (Cid et al., 1992)<br><b>AccNo. BHAR880101</b> Average flexibility indices (Bhaskaran-Ponnuswamy, 1988)<br><b>AccNo. CHAM820101</b> Polarizability parameter (Charton-Charton, 1982)<br><b>AccNo. CHAM820102</b> Free energy of solution in water, kcal/mole (Charton-Charton, 1982)<br><b>AccNo. CHOC760101</b> Residue accessible surface area in tripeptide (Chothia, 1976)<br><b>AccNo. BIGC670101</b> Residue volume (Bigelow, 1967)<br><b>AccNo. CHAM810101</b> Steric parameter (Charton, 1981)<br><b>AccNo. DAYM780201</b> Relative mutability (Dayhoff et al., 1978b) |
| nlag         | Maximum value of the lag parameter. Default is 30.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| type         | The autocorrelation type: moran, geary, or moreaubroto.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |

## Details

calculateAutocor

**Value**

A length nlag named vector for data input.

**Author(s)**

Matineh Rahmatbakhsh <<matinerb.94@gmail.com>>, Nan Xiao

**References**

- AAindex: Amino acid index database. <http://www.genome.ad.jp/dbget/aaindex.html>
- Feng, Z.P. and Zhang, C.T. (2000) Prediction of membrane protein types based on the hydrophobic index of amino acids. *Journal of Protein Chemistry*, 19, 269-275.
- Horne, D.S. (1988) Prediction of protein helix content from an autocorrelation analysis of sequence hydrophobicities. *Biopolymers*, 27, 451-477.
- Sokal, R.R. and Thomson, B.A. (2006) Population structure inferred by local spatial autocorrelation: an Usage from an Amerindian tribal population. *American Journal of Physical Anthropology*, 129, 121-131.

**Examples**

```
data(UP000464024_df)
x_df <- calculateAutocor(UP000464024_df,type = 'moran')
head(x_df, n = 2L)
```

---

calculateBE

*Transform a Sequence into Binary Encoding (BE)*

---

**Description**

This function transform each residue in a peptide into 20 coding values.

**Usage**

```
calculateBE(x)
```

**Arguments**

x                      A data.frame containing gene/protein names and their fasta sequences.

**Details**

```
calculateBE
```

**Value**

A length 400 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Al-Barakati, H. J., Saigo, H., and Newman, R. H. (2019). RF-GlutarySite: a random forest based predictor for glutarylation sites. *Mol. Omi.* 15, 189–204.

---

calculateCTDC

*Calculate CTD Descriptors - Composition (C)*

---

**Description**

This function calculates Composition (C) descriptor for data input.

**Usage**

```
calculateCTDC(x)
```

**Arguments**

`x` A data.frame containing gene/protein names and their fasta sequences.

**Details**

```
calculateCTDC
```

**Value**

A length 21 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Dubchak, I., Muchnik, I., Holbrook, S. R., and Kim, S.-H. (1995). Prediction of protein folding class using global description of amino acid sequence. *Proc. Natl. Acad. Sci.* 92, 8700–8704.

**See Also**

See [calculateCTDT](#) and [calculateCTDD](#) for Transition and Distribution descriptors.

**Examples**

```
data(UP000464024_df)
x_df <- calculateCTDC(UP000464024_df)
head(x_df, n = 2L)
```

---

calculateCTDD*Calculate CTD Descriptors - Distribution (D)*

---

**Description**

This function calculates Distribution (D) descriptor for data input.

**Usage**

```
calculateCTDD(x)
```

**Arguments**

**x** A data.frame containing gene/protein names and their fasta sequences.

**Details**

```
calculateCTDD
```

**Value**

A length 105 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Dubchak, I., Muchnik, I., Holbrook, S. R., and Kim, S.-H. (1995). Prediction of protein folding class using global description of amino acid sequence. *Proc. Natl. Acad. Sci.* 92, 8700–8704.

**See Also**

See [calculateCTDC](#) and [calculateCTDT](#) for Composition and Transition descriptors.

**Examples**

```
data(UP000464024_df)
x_df <- calculateCTDD(UP000464024_df)
head(x_df, n = 1L)
```

---

|               |                                                   |
|---------------|---------------------------------------------------|
| calculateCTDT | <i>Calculate CTD Descriptors - Transition (T)</i> |
|---------------|---------------------------------------------------|

---

**Description**

This function calculates Transition (T) descriptor for data input.

**Usage**

```
calculateCTDT(x)
```

**Arguments**

|   |                                                                       |
|---|-----------------------------------------------------------------------|
| x | A data.frame containing gene/protein names and their fasta sequences. |
|---|-----------------------------------------------------------------------|

**Details**

calculateCTDT

**Value**

A length 21 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Dubchak, I., Muchnik, I., Holbrook, S. R., and Kim, S.-H. (1995). Prediction of protein folding class using global description of amino acid sequence. *Proc. Natl. Acad. Sci.* 92, 8700–8704.

**See Also**

See [calculateCTDC](#) and [calculateCTDD](#) for Composition and Distribution descriptors.

**Examples**

```
data(UP000464024_df)
x_df <- calculateCTDT(UP000464024_df)
head(x_df, n = 2L)
```

---

|                 |                                            |
|-----------------|--------------------------------------------|
| calculateCTriad | <i>Calculate Conjoint Triad Descriptor</i> |
|-----------------|--------------------------------------------|

---

## Description

This function calculates Conjoint Triad descriptor for data input.

## Usage

```
calculateCTriad(x)
```

## Arguments

x                      A data.frame containing gene/protein names and their fasta sequences.

## Details

calculateCTriad

## Value

A length 343 named vector for the data input.

## Author(s)

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

## References

Shen, J., Zhang, J., Luo, X., Zhu, W., Yu, K., Chen, K., et al. (2007). Predicting protein–protein interactions based only on sequences information. *Proc. Natl. Acad. Sci.* 104, 4337–4341.

## Examples

```
data(UP000464024_df)
x_df <- calculateCTriad(UP000464024_df)
head(x_df, n = 2L)
```

---

`calculateDC`*Calculate Dipeptide Composition (DC) Descriptor*

---

**Description**

This function calculates Dipeptide Composition (DC) descriptor for data input.

**Usage**

```
calculateDC(x)
```

**Arguments**

`x` A data.frame containing gene/protein names and their fasta sequences.

**Details**

`calculateDC`

**Value**

A length 400 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Bhasin, M., and Raghava, G. P. S. (2004). Classification of nuclear receptors based on amino acid composition and dipeptide composition. *J. Biol. Chem.* 279, 23262–23266.

**See Also**

See [calculateAAC](#) and [calculateTC](#) for Amino Acid Composition and Tripeptide Composition descriptors.

**Examples**

```
data(UP000464024_df)
x_df <- calculateDC(UP000464024_df)
head(x_df, n = 2L)
```

---

|            |                                       |
|------------|---------------------------------------|
| calculateF | <i>Calculate F1 or F2 Descriptors</i> |
|------------|---------------------------------------|

---

**Description**

This function calculates F1 or F2 descriptors:

- F1 - sum of squared length of Single Amino Acid Repeats (SARs) in the entire protein sequence.
- F2 - maximum of the sum of Single Amino Acid Repeats (SARs) in a window of 6 residues.

**Usage**

```
calculateF(x, type = c("F1", "F2"))
```

**Arguments**

|      |                                                                       |
|------|-----------------------------------------------------------------------|
| x    | A data.frame containing gene/protein names and their fasta sequences. |
| type | The descriptor type: F1 or F2.                                        |

**Details**

calculateF

**Value**

A length 20 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Alguwaizani, S., Park, B., Zhou, X., Huang, D.-S., and Han, K. (2018). Predicting interactions between virus and host proteins using repeat patterns and composition of amino acids. *J. Healthc. Eng.* 2018.

**Examples**

```
data(UP000464024_df)
x_df <- calculateF(UP000464024_df, type = "F1")
head(x_df, n = 2L)
```

---

 calculateKSAAP

*Calculate k-spaced Amino Acid Pairs (KSAAP) Descriptor*


---

### Description

This function calculates k-spaced Amino Acid Pairs (KSAAP) Descriptor for data input. This function is adapted from the [CkSAAPair](#) function in the ftrCOOL package.

### Usage

```
calculateKSAAP(x, spc = 3)
```

### Arguments

|     |                                                                                                                                                                |
|-----|----------------------------------------------------------------------------------------------------------------------------------------------------------------|
| x   | A data.frame containing gene/protein names and their fasta sequences.                                                                                          |
| spc | A number of spaces separating two adjacent residues by a distance of spc, which can be any number up to two less than the length of the peptide; default to 3. |

### Details

calculateKSAAP

### Value

A length 400 named vector for the data input.

### Author(s)

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

### References

Kao, H.-J., Nguyen, V.-N., Huang, K.-Y., Chang, W.-C., and Lee, T.-Y. (2020). SuccSite: incorporating amino acid composition and informative k-spaced amino acid pairs to identify protein succinylation sites. *Genomics. Proteomics Bioinformatics* 18, 208–219.

### Examples

```
data(UP000464024_df)
x_df <- calculateKSAAP(UP000464024_df)
head(x_df, n = 2L)
```

---

|                |                                                                                             |
|----------------|---------------------------------------------------------------------------------------------|
| calculateQD_Sm | <i>Calculate Quadruples Composition (QC) Descriptor from Biochemical Similarity Classes</i> |
|----------------|---------------------------------------------------------------------------------------------|

---

## Description

This function calculates Quadruples Composition (QC) descriptor from biochemical similarity classes.

## Usage

```
calculateQD_Sm(x)
```

## Arguments

x                      A data.frame containing gene/protein names and their fasta sequences.

## Details

calculateQD\_Sm

## Value

A length 1296 named vector for the data input.

## Author(s)

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

## References

Ahmed, I., Witbooi, P., and Christoffels, A. (2018). Prediction of human-Bacillus anthracis protein-protein interactions using multi-layer neural network. *Bioinformatics* 34, 4159–4164.

## Examples

```
data(UP000464024_df)
x_df <- calculateQD_Sm(UP000464024_df)
head(x_df, n = 2L)
```

---

`calculateTC`*Calculate Tripeptide Composition (TC) Descriptor*

---

**Description**

This function calculates Tripeptide Composition (TC) descriptor for data input.

**Usage**

```
calculateTC(x)
```

**Arguments**

`x` A data.frame containing gene/protein names and their fasta sequences.

**Details**

`calculateTC`

**Value**

A length 8,000 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Liao, B., Jiang, J.-B., Zeng, Q.-G., and Zhu, W. (2011). Predicting apoptosis protein subcellular location with PseAAC by incorporating tripeptide composition. *Protein Pept. Lett.* 18, 1086–1092

**See Also**

See [calculateAAC](#), [calculateDC](#) and [calculateTC\\_Sm](#) for Amino Acid Composition, Dipeptide Composition and Tripeptide Composition (TC) Descriptor from Biochemical Similarity Classes.

**Examples**

```
data(UP000464024_df)
x_df <- calculateTC(UP000464024_df)
head(x_df, n = 2L)
```

---

|                |                                                                                             |
|----------------|---------------------------------------------------------------------------------------------|
| calculateTC_Sm | <i>Calculate Tripeptide Composition (TC) Descriptor from Biochemical Similarity Classes</i> |
|----------------|---------------------------------------------------------------------------------------------|

---

**Description**

This function calculates Tripeptide Composition (TC) descriptor from biochemical similarity classes.

**Usage**

```
calculateTC_Sm(x)
```

**Arguments**

x                      A data.frame containing gene/protein names and their fasta sequences.

**Details**

calculateTC\_Sm

**Value**

A length 216 named vector for the data input.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**References**

Ahmed, I., Witbooi, P., and Christoffels, A. (2018). Prediction of human-Bacillus anthracis protein-protein interactions using multi-layer neural network. *Bioinformatics* 34, 4159–4164.

Cui, G., Fang, C., and Han, K. (2012). Prediction of protein-protein interactions between viruses and human by an SVM model. *BMC bioinformatics*, 1–10.

**See Also**

See [calculateTC](#) for Tripeptide Composition descriptor.

**Examples**

```
data(UP000464024_df)
x_df <- calculateTC_Sm(UP000464024_df)
head(x_df, n = 2L)
```

---

`corr_plot`*Plot Correlation Matrix between Input Features*

---

**Description**

A graphical display of a correlation matrix.

**Usage**

```
corr_plot(cormat, method = "number", cex = 0.9)
```

**Arguments**

|                     |                                                                                                                    |
|---------------------|--------------------------------------------------------------------------------------------------------------------|
| <code>cormat</code> | A correlation matrix.                                                                                              |
| <code>method</code> | The visualization method of correlation matrix; defaults to number. See <a href="#">corrplot</a> for more details. |
| <code>cex</code>    | The size of x/y axis label.                                                                                        |

**Details**

`corr_plot`

**Value**

A correlation plot.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>.

**Examples**

```
data('example_data')
x <- na.omit(example_data)
#perform feature selection
s <- FSmethod(x, type = 'both',
cor.cutoff = 0.7, resampling.method = "repeatedcv",
iter = 5, repeats = 3, metric = "ROC", verbose = TRUE)
corr_plot(s$cor.result$corProfile, method = 'square' , cex = 0.5)
```

---

|           |                          |
|-----------|--------------------------|
| enrich.df | <i>Enrichment Result</i> |
|-----------|--------------------------|

---

### Description

Input data for [enrichplot](#)

### Usage

```
data(enrich.df)
```

### Details

To construct this dataset, predicted interactions, generated from [pred\\_ensembl](#) was used as data input for [enrichplot](#).

---

|             |                                                                                                |
|-------------|------------------------------------------------------------------------------------------------|
| enrichfindP | <i>Functional Enrichment Analysis for Pathogen Interactors in the High-Confidence Network.</i> |
|-------------|------------------------------------------------------------------------------------------------|

---

### Description

This function uses [gost](#) function in gprofiler2 package to perform functional enrichment analysis for pathogen interactors in the high-confidence network.

### Usage

```
enrichfindP(
  ppi,
  threshold = 0.05,
  sources = c("GO", "KEGG"),
  p.correction.method = "bonferroni",
  org = "hsapiens"
)
```

### Arguments

|                     |                                                                                                                         |
|---------------------|-------------------------------------------------------------------------------------------------------------------------|
| ppi                 | A data.frame containing pathogen proteins in the first column and host proteins in the second column.                   |
| threshold           | Custom p-value threshold for significance.                                                                              |
| sources             | A vector of data sources to use. See <a href="#">gost</a> for more details.                                             |
| p.correction.method | The algorithm used for multiple testing correction;defaults to 'bonferroni'. See <a href="#">gost</a> for more details. |
| org                 | An organism name;defaults to 'hsapiens'. See <a href="#">gost</a> for more details.                                     |

**Details**

enrichfindP

**Value**

A data.frame with the enrichment analysis results.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**See Also**

See [enrichplot](#) for plotting enrichment analysis.

**Examples**

```
data('predicted_PPIS')
#perform enrichment
enrich.df <- enrichfindP(predicted_PPIS,
  threshold = 0.05,
  sources = c("GO", "KEGG"),
  p.correction.method = "bonferroni",
  org = "hsapiens")
```

---

enrichfind\_cpx

*Functional Enrichment Analysis for Predicted Modules*

---

**Description**

This function uses [gost](#) function in gprofiler2 package to perform functional enrichment analysis for predicted modules.

**Usage**

```
enrichfind_cpx(
  predcpx,
  threshold = 0.05,
  sources = c("GO", "KEGG"),
  p.correction.method = "bonferroni",
  org = "hsapiens"
)
```

**Arguments**

|                     |                                                                                                                         |
|---------------------|-------------------------------------------------------------------------------------------------------------------------|
| predcpx             | Predicted modules resulted from <a href="#">run_clustering</a> .                                                        |
| threshold           | Custom p-value threshold for significance.                                                                              |
| sources             | A vector of data sources to use. See <a href="#">gost</a> for more details.                                             |
| p.correction.method | The algorithm used for multiple testing correction;defaults to 'bonferroni'. See <a href="#">gost</a> for more details. |
| org                 | An organism name;defaults to 'hsapiens'. See <a href="#">gost</a> for more details.                                     |

**Details**

enrichfind\_cpx

**Value**

A data.frame with the enrichment analysis results.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

---

enrichfind\_hp

*Functional Enrichment Analysis of all Host Proteins*

---

**Description**

This function uses [gost](#) function in gprofiler2 package to perform functional enrichment analysis for all predicted host proteins in the high-confidence network.

**Usage**

```
enrichfind_hp(
  ppi,
  threshold = 0.05,
  sources = c("GO", "KEGG"),
  p.correction.method = "bonferroni",
  org = "hsapiens"
)
```

**Arguments**

|           |                                                                                                       |
|-----------|-------------------------------------------------------------------------------------------------------|
| ppi       | A data.frame containing pathogen proteins in the first column and host proteins in the second column. |
| threshold | Custom p-value threshold for significance.                                                            |
| sources   | A vector of data sources to use. See <a href="#">gost</a> for more details.                           |

`p.correction.method`

The algorithm used for multiple testing correction;defaults to 'bonferroni'. See [gost](#) for more details.

`org`

An organism name;defaults to 'hsapiens'. See [gost](#) for more details.

## Details

`enrichfind_hp`

## Value

A data.frame with the enrichment analysis results.

## Author(s)

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

## See Also

See [enrichplot](#) for plotting enrichment analysis.

## Examples

```
data('predicted_PPIs')
#perform enrichment
enrich.df <- enrichfind_hp(predicted_PPIs,
  threshold = 0.05,
  sources = c("GO", "KEGG"),
  p.correction.method = "bonferroni",
  org = "hsapiens")
```

---

enrichplot

*Plot the Enrichment Result*

---

## Description

This function plots the enrichment result.

## Usage

```
enrichplot(x, low = "blue", high = "red", cex.size = 15)
```

## Arguments

|                       |                                                    |
|-----------------------|----------------------------------------------------|
| <code>x</code>        | A data.frame with the enrichment analysis results. |
| <code>low</code>      | Colours for low.                                   |
| <code>high</code>     | Colours for high.                                  |
| <code>cex.size</code> | Text size.                                         |

**Details**

enrichplot

**Value**

An enrichment plot.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**See Also**

See [enrichfindP](#) for functional enrichment analysis.

**Examples**

```
data('enrich.df')
#select enrichment for one of the example (e.g., E protein)
enrich.df <-
enrich.df[enrich.df$id == "E:P0DTC4", ]
enrichplot(enrich.df, low = "blue", high = "red", cex.size = 10)
```

---

example\_data

*Input Data for Prediction Algorithm*

---

**Description**

Input data for [pred\\_ensemble](#)

**Usage**

```
data(example_data)
```

**Format**

a data.frame containing unlabeled or labeled HP-PPIs and pre-computed numerical features.

---

`filter_missing_values` *Drop the Missing Values Above a Certain Threshold*

---

### Description

Given an input matrix, compute the missingness rate for each features and keep only features with missing rate more than user-defined percentage.

### Usage

```
filter_missing_values(x, max_miss_rate = 20)
```

### Arguments

`x` A numeric matrix as input.  
`max_miss_rate` Maximal missing rate allowed for a feature; default is 20.

### Details

`filter_missing_values`

### Value

A dataframe with features with missingness rate of more than user-defined threshold.

### Author(s)

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

### Examples

```
x <- matrix(1:10, ncol = 2)
x[, 2] <- NA
filter_missing_values(x, 30)
```

---

`FreqInteractors` *Plot the Pathogen Proteins' frequency of Interactions with Host Proteins*

---

### Description

This function plots the pathogen proteins' Frequency of interactions with host proteins

### Usage

```
FreqInteractors(ppi, cex.size = 12)
```

**Arguments**

`ppi` A data.frame containing pathogen proteins in the first column and host proteins in the second column.

`cex.size` Text size.

**Details**

FreqInteractors

**Value**

A frequency plot.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**Examples**

```
ppi <- data.frame(
  node1 = c("A", "A", "A", "B", "B", "B", "B"),
  node2 = c("C", "E", "D", "F", "G", "H", "I")
)
FreqInteractors(ppi)
```

---

FSmethod

---

*Feature Selection via Matrix Correlation and Recursive Feature Elimination (RFE)*


---

**Description**

This function performs feature selections via two approaches

- `filter.corr` - compute matrix correlation between features and filter using a threshold.
- `rfeFS` - perform recursive feature elimination (RFE) method wrapped with a Random Forest (RF) algorithm for feature importance evaluation.

**Usage**

```
FSmethod(
  x,
  type = c("cor", "rfe", "both"),
  cor.cutoff = 0.7,
  resampling.method = "cv",
  iter = 2,
  repeats = 3,
  metric = "Accuracy",
  verbose = TRUE
)
```

**Arguments**

|                                |                                                                                                                                                                                                                                                                                                                                             |
|--------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>x</code>                 | A data.frame containing protein-protein interactions, class labels and features.                                                                                                                                                                                                                                                            |
| <code>type</code>              | The feature selection type, one or two of <code>filter.corr</code> and <code>rfeFS</code> .                                                                                                                                                                                                                                                 |
| <code>cor.cutoff</code>        | Correlation coefficient cutoff used for filtering. See <code>filter.corr</code> for more details.                                                                                                                                                                                                                                           |
| <code>resampling.method</code> | The resampling method for RFE : <code>'boot'</code> , <code>'boot632'</code> , <code>'optimism_boot'</code> , <code>'boot_all'</code> , <code>'cv'</code> , <code>'repeatedcv'</code> , <code>'LOOCV'</code> , <code>'LGOCV'</code> ; defaults to <code>cv</code> . See <code>rfeFS</code> and <a href="#">rfeControl</a> for more details. |
| <code>iter</code>              | Number of partitions for cross-validation; defaults to 2. See <code>rfeFS</code> and <a href="#">rfeControl</a> for more details.                                                                                                                                                                                                           |
| <code>repeats</code>           | For repeated k-fold cross validation only; defaults to 3. See <code>rfeFS</code> and <a href="#">rfeControl</a> for more details.                                                                                                                                                                                                           |
| <code>metric</code>            | A string that specifies what summary metric will be used to select the optimal feature ; default to <code>ROC</code> . See <code>rfeFS</code> and <a href="#">rfe</a> for more details.                                                                                                                                                     |
| <code>verbose</code>           | Make the output verbose. See <code>rfeFS</code> and <a href="#">rfeControl</a> for more details.                                                                                                                                                                                                                                            |

**Details**

FSmethod

**Value**

If the type set to `filter.corr` , the output includes the following elements:

- `corProfile` - A correlation matrix.
- `corSelectedFeatures` - Name of features that retained after the correlation analysis.
- `cordf` - A data.frame filtered.

If the type set to `rfeFS` , the output includes the following elements:

- `rfProfile` - A list of elements. See [rfe](#) for more details.
- `rfSelectedFeatures` - Name of features that retained in the feature selection process.
- `rfdf` - A data.frame filtered.

If type set to both the output includes the following elements:

- `rfdf` - The final data.frame that includes the selected features retained after both `filter.corr` and `rfeFS` analysis.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>.

**Examples**

```
data('example_data')
x <- na.omit(example_data)
s <- FSmetho(x, type = 'both',
cor.cutoff = 0.7, resampling.method = "repeatedcv",
iter = 5, repeats = 3, metric = "ROC", verbose = TRUE)
```

getFASTA

*Fetch FASTA Sequence from the UniProt Database***Description**

This function retrieves protein sequences in FASTA format directly from the UniProt database via UniProt protein IDs. This function also checks if the amino-acid composition of protein sequences is in the 20 default types.

**Usage**

```
getFASTA(uniprot.id, filename = "FASTA.RData", path = "FASTASeq")
```

**Arguments**

|            |                                                                                                                                                                                                   |
|------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| uniprot.id | A character vector of UniProt identifiers.                                                                                                                                                        |
| filename   | A character string, indicating the output filename as an RData object to store the retrieved sequences.                                                                                           |
| path       | A character string indicating the path to the project directory that contains the interaction data. If the directory is missing, it will be stored in the current directory. Default is FASTASeq. |

**Details**

getFASTA

**Value**

A list containing protein FASTA sequences.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**Examples**

```
# get fasta sequences for three proteins of SARS-Cov-2
local = tempdir()
uniprot.id <- c("P0DTC4", "P0DTC5", "P0DTC9")
fasta_df <- getFASTA(uniprot.id, filename = 'FASTA.RData', path = local)
head(fasta_df)
```

getHPI

*Generating Host-Pathogen Protein-Protein Interaction (HP-PPI) Descriptors***Description**

This function calculates Host-Pathogen Protein-Protein Interaction (HP-PPI) descriptors via two approaches

- `combine` - combine the two descriptor matrix, result has  $(p1 + p2)$  columns
- `kron.prod` - if A has  $m \times n$  matrix and B is  $q \times p$  matrix, then the Kronecker product is the  $(pm \times qn)$  block matrix

**Usage**

```
getHPI(pathogenData, hostData, type = c("combine", "kron.prod"))
```

**Arguments**

|                           |                                                                |
|---------------------------|----------------------------------------------------------------|
| <code>pathogenData</code> | The pathogen descriptor matrix.                                |
| <code>hostData</code>     | The host descriptor matrix.                                    |
| <code>type</code>         | The interaction type, one or two of "combine" and "kron.prod". |

**Details**

getHPI

**Value**

A matrix containing the Host-Pathogen Protein-Protein Interaction (HP-PPI) descriptors.

**Author(s)**

Matineh Rahmatbakhsh <matinerb.94@gmail.com>

**Examples**

```
x <- matrix(c(1, 2, 3, 1), nrow = 2, ncol = 2, byrow = TRUE)
y <- matrix(c(0, 3, 2, 1), nrow = 2, ncol = 2, byrow = TRUE)
getHPI(x, y, "combine")
getHPI(x, y, "kron.prod")
```

---

|                 |                                                                                          |
|-----------------|------------------------------------------------------------------------------------------|
| get_negativePPI | <i>Construct Negative Reference Host-Pathogen Protein-Protein Interactions (HP-PPIs)</i> |
|-----------------|------------------------------------------------------------------------------------------|

---

## Description

Construct true negative protein-protein interactions from the positive interactions. In the context of PPI prediction, a negative interaction is a pair of proteins that unlikely to interact. Since there is no experimentally verified non-interacting pair, the negative sampling can be used to construct the negative reference set. The negative sampling can be constructed from a set of host proteins, a set of pathogen proteins, and a list of positive reference interactions between members of host and pathogen proteins (Eid et al., 2016).

## Usage

```
get_negativePPI(prot1, prot2, TPset)
```

## Arguments

|       |                                                                |
|-------|----------------------------------------------------------------|
| prot1 | A character vector containing pathogen proteins.               |
| prot2 | A character vector containing host proteins.                   |
| TPset | A character vector containing positive reference interactions. |

## Details

get\_negativePPI

## Value

A Data.frame containing true negative interactions.

## Author(s)

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

## References

Eid, F.-E., ElHefnawi, M., and Heath, L. S. (2016). DeNovo: virus-host sequence-based protein-protein interaction prediction. *Bioinformatics* 32, 1144–1150.

## See Also

See [get\\_positivePPI](#) for generating positive protein-protein interaction.

**Examples**

```
prot1 <- c("P0DTC4", "P0DTC5", "P0DTC9")
prot2 <- c("Q9Y679", "Q9NW15", "Q9NXF8")
TPset <- c("P0DTC4~P31948", "P0DTC8~Q13438")
TN_PPI <- get_negativePPI(prot1, prot2, TPset)
head(TN_PPI)
```

---

|                 |                                                                                                                |
|-----------------|----------------------------------------------------------------------------------------------------------------|
| get_positivePPI | <i>Fetch Positive Reference Host-Pathogen Protein-Protein Interactions (HP-PPIs) from the BioGRID Database</i> |
|-----------------|----------------------------------------------------------------------------------------------------------------|

---

**Description**

This function retrieves positive reference host-pathogen protein-protein interactions directly from BioGRID database.

**Usage**

```
get_positivePPI(
  organism.taxID,
  access.key,
  filename = "PositiveInt.RData",
  path = "PositiveInt"
)
```

**Arguments**

|                |                                                                                                                                                                                                                                            |
|----------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| organism.taxID | Taxonomy identifier for the pathogen.                                                                                                                                                                                                      |
| access.key     | Access key for using BioGRID webpage. To retrieve interactions from the BioGRID database, the users are first required to register for access key at <a href="https://webservice.thebiogrid.org/">https://webservice.thebiogrid.org/</a> . |
| filename       | A character string, indicating the output filename as an RData object to store the retrieved interactions.                                                                                                                                 |
| path           | A character string indicating the path to the project directory that contains the interaction data. If the directory is missing, it will be stored in the current directory. Default is PositiveInt.                                       |

**Details**

get\_positivePPI

**Value**

A Data.frame containing true positive protein-protein interactions for the selected pathogen.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**See Also**

See [get\\_negativePPI](#) for generating negative protein-protein interaction.

**Examples**

```
local = tempdir()
try(get_positivePPI(organism.taxID = 2697049,
  access.key = 'XXXX',
  filename = "PositiveInt.RData",
  path = local))
```

---

Gold\_ReferenceSet

*Gold-standard Reference Set of Inter-Species PPIs*

---

**Description**

This dataset consists of experimentally validated human-SARS-CoV-2 interactions (positive set) and non-interacting pairs (negative set). The following data consists of:

- PPI: SARS-CoV-2-human protein-protein interactions (PPIs)
- Official Symbol Interactor A: SARS-CoV-2 gene names
- official Symbol Interactor B: human host gene names
- Pathogen\_Protein: UniProt identifiers for SARS-CoV-2 virus
- Host\_Protein: UniProt identifiers for human proteins
- class: labeled examples (both positive and negative)

**Usage**

```
data(Gold_ReferenceSet)
```

**Format**

a data.frame containing 500 validated pairs (i.e., positive set) and 500 non-interacting pairs (i.e., negative set).

**Details**

To construct this dataset, validated interactions (positive set) were retrieved from BioGrid database and were further filtered to only include those interactions provided by (Samavarchi-Tehrani et al., 2020). In this study, the authors mapped interaction between 27 SARS-CoV-2 and human proteins via the proximity-dependent biotinylation (BioID) approach. 500 SARS-CoV-2-host interaction pairs then randomly selected from all pairs to serve as positive examples. To construct negative examples, negative sampling were used using [get\\_negativePPI](#).

**Source**

<https://www.biorxiv.org/content/10.1101/2020.09.03.282103v1>

**References**

Samavarchi-Tehrani, P. et al. (2020) A SARS-CoV-2-host proximity interactome. BioRxiv.

---

|         |                                         |
|---------|-----------------------------------------|
| host_se | <i>Host SummarizedExperiment object</i> |
|---------|-----------------------------------------|

---

**Description**

SummarizedExperiment object of numerical features for host proteins.

**Usage**

```
data(host_se)
```

**Details**

To construct this object, first protein sequences were converted to numerical features using (CTD) descriptors provided in the HPiP package, followed by converting each numerical features matrix to SummarizedExperiment object. Each object is then merged into one object using 'cbind()'.

---

|                     |                                                           |
|---------------------|-----------------------------------------------------------|
| impute_missing_data | <i>Impute missing Values per Features (i.e., Columns)</i> |
|---------------------|-----------------------------------------------------------|

---

**Description**

Given an input matrix, impute the missing values via three approaches including mean, median or zero.

**Usage**

```
impute_missing_data(x, method = c("mean", "median", "zero"))
```

**Arguments**

|        |                                                              |
|--------|--------------------------------------------------------------|
| x      | A numeric matrix as input.                                   |
| method | Imputation method for missing values (mean, median or zero). |

**Details**

```
impute_missing_data
```

**Value**

Imputed matrix.

**Author(s)**

Matineh Rahmatbakhsh <matinerb.94@gmail.com>

**Examples**

```
x <- matrix(1:10, ncol = 2)
x[1:3, 2] <- NA
row.names(x) <- c("A", "B", "C", "D", "E")
colnames(x) <- c("col1", "col2")
impute_missing_data(x, method = "mean")
impute_missing_data(x, method = "median")
impute_missing_data(x, method = "zero")
```

---

plotPPI

---

*Plot the Predicted PPI*


---

**Description**

Plot the predicted PPIs. This function uses the plot function of the igraph.

**Usage**

```
plotPPI(
  ppi,
  edge.name = "ensemble_score",
  node.color = "grey",
  edge.color = "orange",
  cex.node = 4,
  node.label.dist = 1.5
)
```

**Arguments**

|                 |                                                                       |
|-----------------|-----------------------------------------------------------------------|
| ppi             | A data.frame containing protein-protein interactions with edge score. |
| edge.name       | A character string giving an edge attribute name.                     |
| node.color      | The fill color of the node.                                           |
| edge.color      | The color of the edge.                                                |
| cex.node        | The size of the node.                                                 |
| node.label.dist | The distance of the label from the center of the node.                |

**Details**

plotPPI

**Value**

A PPI plot.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**Examples**

```
df <- data.frame(
  node1 = c("A", "B", "C", "D", "E"),
  node2 = c("C", "E", "E", "E", "A"),
  edge.scores = c(0.5, 0.4, 0.3, 0.2, 0.7)
)
plotPPI(df, edge.name = "edge.scores")
```

---

|                |                          |
|----------------|--------------------------|
| predicted_PPIs | <i>Predicted HP-PPIs</i> |
|----------------|--------------------------|

---

**Description**

Input data for [enrichfindP](#)

**Usage**

```
data(predicted_PPIs)
```

---

|              |                                                          |
|--------------|----------------------------------------------------------|
| pred_ensembl | <i>Predict Interactions via Ensemble Learning Method</i> |
|--------------|----------------------------------------------------------|

---

**Description**

This function uses an ensemble of classifiers to predict interactions from the sequence-based dataset. This ensemble algorithm combines different results generated from individual classifiers within the ensemble via average to enhance prediction.

**Usage**

```
pred_ensembl(
  features,
  gold_standard,
  classifier = c("avNNet", "svmRadial", "ranger"),
  resampling.method = "cv",
  ncross = 2,
  repeats = 2,
  verboseIter = TRUE,
  plots = TRUE,
  filename = "plots.pdf"
)
```

**Arguments**

|                   |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
|-------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| features          | A data frame with host-pathogen protein-protein interactions (HP-PPIs) in the first column, and features to be passed to the classifier in the remaining columns.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
| gold_standard     | A data frame with gold_standard HP-PPIs and class label indicating if such PPIs are positive or negative.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
| classifier        | The type of classifier to use. See caret for the available classifiers.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
| resampling.method | The resampling method: 'boot', 'boot632', 'optimism_boot', 'boot_all', 'cv', 'repeatedcv', 'LOOCV', 'LGOCV'; defaults to cv. See <a href="#">trainControl</a> for more details.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| ncross            | Number of partitions for cross-validation; defaults to 5. See <a href="#">trainControl</a> for more details.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
| repeats           | for repeated k-fold cross validation only; defaults to 3. See <a href="#">rfeControl</a> for more details.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| verboseIter       | Logical value, indicating whether to check the status of training process; defaults to FALSE.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| plots             | Logical value, indicating whether to plot the performance of ensemble learning algorithm as compared to individual classifiers; defaults to TRUE. If the argument set to TRUE, plots will be saved in the current working directory. These plots are : <ul style="list-style-type: none"> <li>• pr_plot - Precision-recall plot of ensemble classifier vs selected individual classifiers.</li> <li>• roc_plot - ROC plot of ensemble classifier vs selected individual classifiers.</li> <li>• points_plot - Plot accuracy, F1-score, positive predictive value (PPV), sensitivity (SE), and Matthews correlation coefficient (MCC) of ensemble classifier vs selected individual classifiers.</li> </ul> |
| filename          | A character string, indicating the output filename as a pdf object.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |

**Details**

pred\_ensembl

**Value**

Ensemble\_training\_output

- prediction score - Prediction scores for whole dataset from each individual classifier.
- Best - Selected hyper parameters.
- Parameter range - Tested hyper parameters.
- prediction\_score\_test - Scores probabilities for test data from each individual classifier.
- class\_label - Class probabilities for test data from each individual classifier.

classifier\_performance

- cm - A confusion matrix.
- ACC - Accuracy.
- SE - Sensitivity.
- SP - Specificity.
- PPV - Positive Predictive Value.
- F1 - F1-score.
- MCC - Matthews correlation coefficient.
- Roc\_Object - A list of elements. See [roc](#) for more details.
- PR\_Object - A list of elements. See [pr.curve](#) for more details.

predicted\_interactions - The input data frame of pairwise interactions, including classifier scores averaged across all models.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

**Examples**

```
data('example_data')
features <- example_data[, -2]
gd <- example_data[, c(1,2)]
gd <- na.omit(gd)
ppi <- pred_ensembel(features,gd,
  classifier = c("avNNet", "svmRadial", "ranger"),
  resampling.method = "cv",ncross = 2,verboseIter = FALSE,plots = FALSE,
  filename = "plots.pdf")
#extract predicted interactions
pred_interaction <- ppi[["predicted_interactions"]]
```

---

|                |                         |
|----------------|-------------------------|
| run_clustering | <i>Module Detection</i> |
|----------------|-------------------------|

---

**Description**

This function contains five module detection algorithms including fast-greedy algorithm (FC), walk-trap algorithm (RW), multi-level community algorithm (ML), label propagation algorithm (clp), and markov clustering (MCL).

**Usage**

```
run_clustering(  
  ppi,  
  method = c("FC", "RW", "ML", "clp", "MCL"),  
  expans = 2,  
  infla = 5,  
  iter = 50  
)
```

**Arguments**

|        |                                                                                                                                                                                                                                                                                         |
|--------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| ppi    | A data.frame containing pathogen proteins in the first column, host proteins in the second column, and edge weight in the third column.                                                                                                                                                 |
| method | Module detection algorithms including: <ul style="list-style-type: none"><li>• FC - fast-greedy algorithm.</li><li>• RW - walktrap algorithm.</li><li>• ML - multi-level community algorithm.</li><li>• clp - label propagation algorithm.</li><li>• MCL - markov clustering.</li></ul> |
| expans | Numeric value > 1 for the expansion parameter. See <a href="#">mcl</a> for more details.                                                                                                                                                                                                |
| infla  | Numeric value > 0 for the inflation power coefficient. See <a href="#">mcl</a> for more details.                                                                                                                                                                                        |
| iter   | An integer, the maximum number of iterations for the MCL. See <a href="#">mcl</a> for more details.                                                                                                                                                                                     |

**Details**

run\_clustering

**Value**

A data.frame with the enrichment analysis results.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>

---

unlabel\_data

*HP-PPIs with Unknown Class Labels*


---

### Description

This dataset consists of interactions between SARS-CoV-2 and human proteins, achieved by AP-MS (affinity purification mass spectrometry).

### Usage

```
data(unlabel_data)
```

### Format

A data.frame containing 700 SARS-CoV-2-Human protein-protein interactions (PPIs) with pre-computed numerical features using CTD (composition/transition/distribution) descriptors.

### Details

To construct this dataset, data (supplementary table 1) containing SARS-CoV-2-human PPIs was retrieved from (Gordon et al., 2020) and 700 pairs were randomly selected from total pairs, followed by converting protein sequences of host or viral proteins to numerical features and finally concatenating the computed features in order to construct host-pathogen PPIs.

### Source

<https://www.nature.com/articles/s41586-020-2286-9#Sec36>

### References

Gordon,D.E. et al. (2020) A SARS-CoV-2 protein interaction map reveals targets for drug repurposing. Nature, 583, 459–468.

---

UP000464024\_df

*Data.frame Containing SARS-CoV-2 FASTA Sequences*


---

### Description

This data includes one protein sequence per SARS-CoV-2 gene, retrieved directly from UniProt database using [getFASTA](#).

### Usage

```
data(UP000464024_df)
```

**Format**

A data.frame with two columns:(1) UniprotKBID, UniProt identifier.(2) FASTASEQ, sequences per SARS-CoV-2 gene.

**Source**

<https://www.uniprot.org/uniprot/?query=proteome:UP000464024>

---

|         |                                 |
|---------|---------------------------------|
| var_imp | <i>Variable Importance Plot</i> |
|---------|---------------------------------|

---

**Description**

A graphical display of variable importance of selected features.

**Usage**

```
var_imp(x, cex.x = 1, cex.y = 2)
```

**Arguments**

|       |                                                                                         |
|-------|-----------------------------------------------------------------------------------------|
| x     | A list of elements returned from RFE analysis. See <a href="#">rfe</a> for more details |
| cex.x | The size of x axis label.                                                               |
| cex.y | The size of y axis label.                                                               |

**Details**

var\_imp

**Value**

Variable Importance Plot.

**Author(s)**

Matineh Rahmatbakhsh, <matinerb.94@gmail.com>.

**Examples**

```
data('example_data')
x <- na.omit(example_data)
#perform feature selection
s <- FSmetho(x, type = 'both',
cor.cutoff = 0.7, resampling.method = "repeatedcv",
iter = 5, repeats = 3, metric = "ROC", verbose = TRUE)
var_imp(s$rf.result$rfProfile, cex.x = 10, cex.y = 10)
```

---

|          |                                          |
|----------|------------------------------------------|
| viral_se | <i>Viral SummarizedExperiment object</i> |
|----------|------------------------------------------|

---

**Description**

SummarizedExperiment object of numerical features for SARS-CoV-2 proteins.

**Usage**

```
data(viral_se)
```

**Details**

To construct this object, first protein sequences were converted to numerical features using (CTD) descriptors provided in the HPiP package, followed by converting each numerical features matrix to SummarizedExperiment object. Each object is then merged into one object using ‘cbind()’.

# Index

calculateAAC, [3](#), [10](#), [14](#)  
calculateAutocor, [4](#)  
calculateBE, [5](#)  
calculateCTDC, [6](#), [7](#), [8](#)  
calculateCTDD, [6](#), [7](#), [8](#)  
calculateCTDT, [6](#), [7](#), [8](#)  
calculateCTriad, [9](#)  
calculateDC, [3](#), [10](#), [14](#)  
calculateF, [11](#)  
calculateKSAAP, [12](#)  
calculateQD\_Sm, [13](#)  
calculateTC, [3](#), [10](#), [14](#), [15](#)  
calculateTC\_Sm, [14](#), [15](#)  
CkSAAPair, [12](#)  
corr\_plot, [16](#)  
corrplot, [16](#)  
  
enrich.df, [17](#)  
enrichfind\_cpx, [18](#)  
enrichfind\_hp, [19](#)  
enrichfindP, [17](#), [21](#), [32](#)  
enrichplot, [17](#), [18](#), [20](#), [20](#)  
example\_data, [21](#)  
  
filter\_missing\_values, [22](#)  
FreqInteractors, [22](#)  
FSmethod, [23](#)  
  
get\_negativePPI, [27](#), [29](#)  
get\_positivePPI, [27](#), [28](#)  
getFASTA, [25](#), [36](#)  
getHPI, [26](#)  
Gold\_ReferenceSet, [29](#)  
gost, [17–20](#)  
  
host\_se, [30](#)  
  
impute\_missing\_data, [30](#)  
  
mcl, [35](#)  
  
plotPPI, [31](#)  
pr.curve, [34](#)  
pred\_ensembl, [17](#), [21](#), [32](#)  
predicted\_PPIS, [32](#)  
  
rfe, [24](#), [37](#)  
rfeControl, [24](#), [33](#)  
roc, [34](#)  
run\_clustering, [19](#), [35](#)  
  
trainControl, [33](#)  
  
unlabel\_data, [36](#)  
UP000464024\_df, [36](#)  
  
var\_imp, [37](#)  
viral\_se, [38](#)