Package: protr (via r-universe)

October 27, 2024

```
Version 1.7-4
Title Generating Various Numerical Representation Schemes for Protein
      Sequences
Description Comprehensive toolkit for generating various numerical
      features of protein sequences described in Xiao et al. (2015)
      <DOI:10.1093/bioinformatics/btv042>. For full functionality,
      the software 'ncbi-blast+' is needed, see
      <https://blast.ncbi.nlm.nih.gov/doc/blast-help/downloadblastdata.html>
      for more information.
License BSD 3 clause + file LICENSE
URL https://nanx.me/protr/, https://github.com/nanxstats/protr,
      http://protr.org
BugReports https://github.com/nanxstats/protr/issues
Encoding UTF-8
LazyData true
SystemRequirements ncbi-blast+ (see
      <a href="https://blast.ncbi.nlm.nih.gov/doc/blast-help/downloadblastdata.html">https://blast.ncbi.nlm.nih.gov/doc/blast-help/downloadblastdata.html</a>)
VignetteBuilder knitr
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Suggests knitr, rmarkdown, Biostrings, GOSemSim, foreach, doParallel,
      org.Hs.eg.db
RoxygenNote 7.3.2
NeedsCompilation no
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```

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Description

This dataset includes the 2D autocorrelations descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AA2DACOR)

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3D-MoRSE Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the 3D-MoRSE descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AA3DMoRSE)

AAACF

Atom-Centred Fragments Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the atom-centred fragments descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAACF)

AABLOSUM100

BLOSUM100 Matrix for 20 Amino Acids

Description

 $BLOSUM100\ Matrix$ for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AABLOSUM100)

AABLOSUM45 5

AABLOSUM45

BLOSUM45 Matrix for 20 Amino Acids

Description

BLOSUM45 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AABLOSUM45)

AABLOSUM50

BLOSUM50 Matrix for 20 Amino Acids

Description

BLOSUM50 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AABLOSUM50)

AABLOSUM62

BLOSUM62 Matrix for 20 Amino Acids

Description

BLOSUM62 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AABLOSUM62)

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AABLOSUM80 BLOSUM80 Matrix for 20 Amino Acids

Description

BLOSUM80 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AABLOSUM80)

	AABurden	Burden Eigenvalues Descriptors for 20 Amino Acids calculated by Dragon
--	----------	--

Description

This dataset includes the Burden eigenvalues descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AABurden)

-	
AAConn	Connectivity Indices Descriptors for 20 Amino Acids calculated by
	Dragon

Description

This dataset includes the connectivity indices descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAConn)

AAConst 7

AAConst	Constitutional Descriptors for 20 Amino Acids calculated by Dragon
AACONST	Constitutional Descriptors for 20 Amino Acias calculated by Dragon

Description

This dataset includes the constitutional descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAConst)

AACPSA

CPSA Descriptors for 20 Amino Acids calculated by Discovery Studio

Description

This dataset includes the CPSA descriptors of the 20 amino acids calculated by Discovery Studio (version 2.5) used for scales extraction in this package.

Details

All amino acid molecules had also been optimized with MOE 2011.10 (semiempirical AM1) before calculating these CPSA descriptors. The SDF file containing the information of the optimized amino acid molecules is included in this package. See OptAA3d for more information.

Examples

data(AACPSA)

AADescAll

All 2D Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes all the 2D descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AADescAll)

8 AAFGC

AAEdgeAdj	Edge Adjacency Indices Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the edge adjacency indices descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAEdgeAdj)

AAEigIdx	Eigenvalue-Based Indices Descriptors for 20 Amino Acids calculated
	by Dragon

Description

This dataset includes the eigenvalue-based indices descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAEigIdx)

AAFGC	Functional Group Counts Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the functional group counts descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAFGC)

AAGeom 9

AAGeom	Geometrical Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the geometrical descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAGeom)

AAGETAWAY

GETAWAY Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the GETAWAY descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAGETAWAY)

AAindex Data of 544 Physicochemical and Biological Properties for 20 Amino Acids

Description

The data was extracted from the AAindex1 database ver 9.1 (https://www.genome.jp/ftp/db/community/aaindex/old/ver9.1/aaindex1) as of November, 2012 (data last modified on 2006-08-14).

Details

With this dataset, users can investigate each property's accession number and other details. See https://www.genome.jp/aaindex/ for more information.

Examples

data(AAindex)

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AAInfo	Information Indices Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the information indices descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAInfo)

AAMetaInfo

Meta Information for the 20 Amino Acids

Description

This dataset includes the meta information of the 20 amino acids used for the 2D and 3D descriptor calculation in this package. Each column represents:

AAName Amino acid name
Short One-letter representation
Abbreviation Three-letter representation
mol SMILES representation
PUBCHEM_COMPOUND_CID PubChem CID for the amino acid
PUBCHEM_LINK PubChem link for the amino acid

Examples

data(AAMetaInfo)

AAMOE2D

2D Descriptors for 20 Amino Acids calculated by MOE 2011.10

Description

This dataset includes the 2D descriptors of the 20 amino acids calculated by MOE 2011.10 used for scales extraction in this package.

Examples

data(AAMOE2D)

AAMOE3D 11

AAMOE3D

3D Descriptors for 20 Amino Acids calculated by MOE 2011.10

Description

This dataset includes the 3D descriptors of the 20 amino acids calculated by MOE 2011.10 used for scales extraction in this package. All amino acid molecules had also been optimized with MOE (semiempirical AM1) before calculating these 3D descriptors. The SDF file containing the information of the optimized amino acid molecules is included in this package. See OptAA3d for more information.

Examples

data(AAMOE3D)

AAMolProp

Molecular Properties Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the molecular properties descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAMolProp)

AAPAM120

PAM120 Matrix for 20 Amino Acids

Description

PAM120 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AAPAM120)

12 AAPAM40

AAPAM250

PAM250 Matrix for 20 Amino Acids

Description

PAM250 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AAPAM250)

AAPAM30

PAM30 Matrix for 20 Amino Acids

Description

PAM30 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AAPAM30)

AAPAM40

PAM40 Matrix for 20 Amino Acids

Description

PAM40 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AAPAM40)

AAPAM70 13

AAPAM70

PAM70 Matrix for 20 Amino Acids

Description

PAM70 Matrix for the 20 amino acids. The matrix was extracted from the Biostrings package of Bioconductor.

Examples

data(AAPAM70)

AARandic

Randic Molecular Profiles Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the Randic molecular profiles descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AARandic)

AARDF

RDF Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the RDF descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AARDF)

14 AAWalk

ААТоро	Topological Descriptors for 20 Amino Acids calculated by Dragon
ААТоро	Topological Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the topological descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AATopo)

AATopoChg	Topological Charge Indices Descriptors for 20 Amino Acids calculated by Dragon
	and by Bragon

Description

This dataset includes the topological charge indices descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AATopoChg)

AAWalk	Walk and Path Counts Descriptors for 20 Amino Acids calculated by Dragon

Description

This dataset includes the walk and path counts descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAWalk)

AAWHIM 15

AAWHIM	WHIM Descriptors for 20 Amino Acids calculated by Dragon	
--------	--	--

Description

This dataset includes the WHIM descriptors of the 20 amino acids calculated by Dragon (version 5.4) used for scales extraction in this package.

Examples

data(AAWHIM)

acc

Auto Cross Covariance (ACC) for Generating Scales-Based Descriptors of the Same Length

Description

This function calculates the auto covariance and auto cross covariance for generating scale-based descriptors of the same length.

Usage

```
acc(mat, lag)
```

Arguments

mat A p * n matrix. Each row represents one scale (total p scales), each column

represents one amino acid position (total n amino acids).

lag The lag parameter. Must be less than the amino acids.

Value

A length $lag * p^2$ named vector, the element names are constructed by: the scales index (crossed scales index) and lag index.

Note

Please see the references for details about auto cross covariance.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

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References

Wold, S., Jonsson, J., Sjorstrom, M., Sandberg, M., & Rannar, S. (1993). DNA and peptide sequences and chemical processes multivariately modelled by principal component analysis and partial least-squares projections to latent structures. *Analytica chimica acta*, 277(2), 239–253.

Sjostrom, M., Rannar, S., & Wieslander, A. (1995). Polypeptide sequence property relationships in *Escherichia coli* based on auto cross covariances. *Chemometrics and intelligent laboratory systems*, 29(2), 295–305.

See Also

See extractScales for scales-based descriptors. For more details, see extractDescScales and extractProtFP.

Examples

```
p <- 8 # p is the scales number
n <- 200 # n is the amino acid number
lag <- 7 # the lag paramter
mat <- matrix(rnorm(p * n), nrow = p, ncol = n)
acc(mat, lag)</pre>
```

crossSetSim

Parallel Protein Sequence Similarity Calculation Between Two Sets Based on Sequence Alignment (In-Memory Version)

Description

Parallel calculation of protein sequence similarity based on sequence alignment between two sets of protein sequences.

Usage

```
crossSetSim(
  protlist1,
  protlist2,
  type = "local",
  cores = 2,
  batches = 1,
  verbose = FALSE,
  submat = "BLOSUM62",
  gap.opening = 10,
  gap.extension = 4
)
```

crossSetSim 17

Arguments

protlist1	A length n list containing n protein sequences, each component of the list is a character string, storing one protein sequence. Unknown sequences should be represented as "".
protlist2	A length n list containing m protein sequences, each component of the list is a character string, storing one protein sequence. Unknown sequences should be represented as "".
type	Type of alignment, default is "local", can be "global" or "local", where "global" represents Needleman-Wunsch global alignment; "local" represents Smith-Waterman local alignment.
cores	Integer. The number of CPU cores to use for parallel execution, default is 2. Users can use the availableCores() function in the parallelly package to see how many cores they could use.
batches	Integer. How many batches should we split the similarity computations into. This is useful when you have a large number of protein sequences, enough number of CPU cores, but not enough RAM to compute and fit all the similarities into a single batch. Defaults to 1.
verbose	Print the computation progress? Useful when batches > 1.
submat	Substitution matrix, default is "BLOSUM62", can be one of "BLOSUM45", "BLOSUM50", "BLOSUM62", "BLOSUM80", "BLOSUM100", "PAM30", "PAM40", "PAM70", "PAM120", or "PAM250".
gap.opening	The cost required to open a gap of any length in the alignment. Defaults to 10.

Value

A n x m similarity matrix.

Author(s)

Sebastian Mueller https://alva-genomics.com

Examples

```
## Not run:

# Be careful when testing this since it involves parallelization
# and might produce unpredictable results in some environments

library("Biostrings")
library("foreach")
library("doParallel")

s1 <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
s2 <- readFASTA(system.file("protseq/P08218.fasta", package = "protr"))[[1]]
s3 <- readFASTA(system.file("protseq/P10323.fasta", package = "protr"))[[1]]
s4 <- readFASTA(system.file("protseq/P20160.fasta", package = "protr"))[[1]]</pre>
```

gap.extension The cost to extend the length of an existing gap by 1. Defaults to 4.

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crossSetSimDisk

Parallel Protein Sequence Similarity Calculation Between Two Sets Based on Sequence Alignment (Disk-Based Version)

Description

Parallel calculation of protein sequence similarity based on sequence alignment between two sets of protein sequences. This version offloads the partial results in each batch to the hard drive and merges the results together in the end, which reduces the memory usage.

Usage

```
crossSetSimDisk(
  protlist1,
  protlist2,
  cores = 2,
  batches = 1,
  path = tempdir(),
  verbose = FALSE,
  type = "local",
  submat = "BLOSUM62",
  gap.opening = 10,
  gap.extension = 4
)
```

Arguments

protlist1

A length n list containing n protein sequences, each component of the list is a character string, storing one protein sequence. Unknown sequences should be represented as "".

protlist2

A length n list containing m protein sequences, each component of the list is a character string, storing one protein sequence. Unknown sequences should be represented as "".

crossSetSimDisk 19

cores Integer. The number of CPU cores to use for parallel execution, default is 2.

Users can use the availableCores() function in the parallelly package to see

how many cores they could use.

batches Integer. How many batches should we split the pairwise similarity computations

into. This is useful when you have a large number of protein sequences, enough number of CPU cores, but not enough RAM to compute and fit all the pairwise

similarities into a single batch. Defaults to 1.

path Directory for caching the results in each batch. Defaults to the temporary direc-

tory.

verbose Print the computation progress? Useful when batches > 1.

type Type of alignment, default is "local", can be "global" or "local", where

"global" represents Needleman-Wunsch global alignment; "local" represents

Smith-Waterman local alignment.

submat Substitution matrix, default is "BLOSUM62", can be one of "BLOSUM45", "BLOSUM50",

"BLOSUM62", "BLOSUM80", "BLOSUM100", "PAM30", "PAM40", "PAM70", "PAM120",

or "PAM250".

gap.opening The cost required to open a gap of any length in the alignment. Defaults to 10.

gap.extension The cost to extend the length of an existing gap by 1. Defaults to 4.

Value

An x m similarity matrix.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

See Also

See crossSetSim for the in-memory version.

```
## Not run:

# Be careful when testing this since it involves parallelization
# and might produce unpredictable results in some environments

library("Biostrings")
library("foreach")
library("doParallel")

s1 <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
s2 <- readFASTA(system.file("protseq/P08218.fasta", package = "protr"))[[1]]
s3 <- readFASTA(system.file("protseq/P10323.fasta", package = "protr"))[[1]]
s4 <- readFASTA(system.file("protseq/P20160.fasta", package = "protr"))[[1]]
s5 <- readFASTA(system.file("protseq/Q9NZP8.fasta", package = "protr"))[[1]]
set.seed(1010)</pre>
```

20 extractAAC

```
plist1 <- as.list(c(s1, s2, s3, s4, s5)[sample(1:5, 100, replace = TRUE)])
plist2 <- as.list(c(s1, s2, s3, s4, s5)[sample(1:5, 100, replace = TRUE)])
psimmat <- crossSetSimDisk(
   plist1, plist2,
   cores = 2, batches = 10, verbose = TRUE,
   type = "local", submat = "BLOSUM62"
)
## End(Not run)</pre>
```

extractAAC

Amino Acid Composition Descriptor

Description

This function calculates the Amino Acid Composition descriptor (dim: 20).

Usage

```
extractAAC(x)
```

Arguments

Х

A character vector, as the input protein sequence.

Value

A length 20 named vector

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

M. Bhasin, G. P. S. Raghava. Classification of Nuclear Receptors Based on Amino Acid Composition and Dipeptide Composition. *Journal of Biological Chemistry*, 2004, 279, 23262.

See Also

See extractDC and extractTC for Dipeptide Composition and Tripeptide Composition descriptors.

```
x \leftarrow readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]] extractAAC(x)
```

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extractAPAAC

Amphiphilic Pseudo Amino Acid Composition (APseAAC) Descriptor

Description

This function calculates the Amphiphilic Pseudo Amino Acid Composition (APseAAC, or APAAC) descriptor (dim: 20 + (n * lambda), n is the number of properties selected, default is 80).

Usage

```
extractAPAAC(
    x,
    props = c("Hydrophobicity", "Hydrophilicity"),
    lambda = 30,
    w = 0.05,
    customprops = NULL
)
```

Arguments

lambda

x A character vector, as the input protein sequence.

props A character vector, specifying the properties used. 2 properties are used by

default, as listed below:

'Hydrophobicity' Hydrophobicity value of the 20 amino acids 'Hydrophilicity' Hydrophilicity value of the 20 amino acids

The lambda parameter for the APAAC descriptors, default is 30.

w The weighting factor, default is 0.05.

customprops A n x 21 named data frame contains n customized property. Each row contains

one property. The column order for different amino acid types is 'AccNo', 'A', 'R', 'N', 'D', 'C', 'E', 'Q', 'G', 'H', 'I', 'L', 'K', 'M', 'F', 'P', 'S', 'T', 'W', 'Y', 'V', and the columns should also be *exactly* named like this. The AccNo column contains the properties' names. Then users should explicitly specify these properties with these names in the argument props. See the examples below for a demonstration. The default value for customprops is NULL.

Value

A length 20 + n * lambda named vector, n is the number of properties selected.

Note

Note the default 20 * 2 prop values have already been independently given in the function. Users can also specify other (up to 544) properties with the Accession Number in the AAindex data, with or without the default three properties, which means users should explicitly specify the properties to use. For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use

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some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Kuo-Chen Chou. Prediction of Protein Cellular Attributes Using Pseudo-Amino Acid Composition. *PROTEINS: Structure, Function, and Genetics*, 2001, 43: 246-255.

Kuo-Chen Chou. Using Amphiphilic Pseudo Amino Acid Composition to Predict Enzyme Subfamily Classes. *Bioinformatics*, 2005, 21, 10-19.

```
JACS, 1962, 84: 4240-4246. (C. Tanford). (The hydrophobicity data) PNAS, 1981, 78:3824-3828 (T.P.Hopp & K.R.Woods). (The hydrophilicity data)
```

See Also

See extractPAAC for the pseudo amino acid composition (PseAAC) descriptor.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]</pre>
extractAPAAC(x)
myprops <- data.frame(</pre>
 AccNo = c("MyProp1", "MyProp2", "MyProp3"),
 A = c(0.62, -0.5, 15), R = c(-2.53, 3, 101),
 N = c(-0.78, 0.2, 58), D = c(-0.9, 3, 59),
 C = c(0.29, -1, 47), E = c(-0.74, 3, 73),
 Q = c(-0.85, 0.2, 72), G = c(0.48, 0, 1),
 H = c(-0.4, -0.5, 82), I = c(1.38, -1.8, 57),
 L = c(1.06, -1.8, 57), K = c(-1.5, 3, 73),
 M = c(0.64, -1.3, 75), F = c(1.19, -2.5, 91),
 P = c(0.12, 0, 42), S = c(-0.18, 0.3, 31),
 T = c(-0.05, -0.4, 45), W = c(0.81, -3.4, 130),
 Y = c(0.26, -2.3, 107), V = c(1.08, -1.5, 43)
)
# use 2 default properties, 4 properties from the
# AAindex database, and 3 cutomized properties
extractAPAAC(
 customprops = myprops,
 props = c(
    "Hydrophobicity", "Hydrophilicity",
    "CIDH920105", "BHAR880101",
    "CHAM820101", "CHAM820102",
    "MyProp1", "MyProp2", "MyProp3"
```

extractBLOSUM 23

extractBLOSUM	BLOSUM and PAM Matrix-Derived Descriptors	

Description

This function calculates BLOSUM matrix-derived descriptors. For users' convenience, protr provides the BLOSUM45, BLOSUM50, BLOSUM62, BLOSUM80, BLOSUM100, PAM30, PAM40, PAM70, PAM120, and PAM250 matrices for the 20 amino acids to select from.

Usage

```
extractBLOSUM(x, submat = "AABLOSUM62", k, lag, scale = TRUE, silent = TRUE)
```

Arguments

x	A character vector, as the input protein sequence.
submat	Substitution matrix for the 20 amino acids. Should be one of AABLOSUM45, AABLOSUM50, AABLOSUM62, AABLOSUM80, AABLOSUM100, AAPAM30, AAPAM40, AAPAM70, AAPAM120, or AAPAM250. Default is "AABLOSUM62".
k	Integer. The number of selected scales (i.e. the first k scales) derived by the substitution matrix. This can be selected according to the printed relative importance values.
lag	The lag parameter. Must be less than the amino acids.
scale	Logical. Should we auto-scale the substitution matrix (submat) before doing eigen decomposition? Default is TRUE.
silent	Logical. Whether we print the relative importance of each scales (diagnal value of the eigen decomposition result matrix B) or not. Default is TRUE.

Value

A length lag * p^2 named vector, p is the number of scales selected.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Georgiev, A. G. (2009). Interpretable numerical descriptors of amino acid space. Journal of Computational Biology, 16(5), 703–723.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
blosum <- extractBLOSUM(x, submat = "AABLOSUM62", k = 5, lag = 7, scale = TRUE, silent = FALSE)</pre>
```

24 extractCTDC

extractCTDC

CTD Descriptors - Composition

Description

This function calculates the Composition descriptor of the CTD descriptors (dim: 21).

Usage

```
extractCTDC(x)
```

Arguments

Х

A character vector, as the input protein sequence.

Value

A length 21 named vector

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Inna Dubchak, Ilya Muchink, Stephen R. Holbrook and Sung-Hou Kim. Prediction of protein folding class using global description of amino acid sequence. *Proceedings of the National Academy of Sciences*. USA, 1995, 92, 8700-8704.

Inna Dubchak, Ilya Muchink, Christopher Mayor, Igor Dralyuk and Sung-Hou Kim. Recognition of a Protein Fold in the Context of the SCOP classification. *Proteins: Structure, Function and Genetics*, 1999, 35, 401-407.

See Also

See extractCTDT and extractCTDD for Transition and Distribution of the CTD descriptors.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
extractCTDC(x)</pre>
```

extractCTDCClass 25

extractCTDCClass	CTD Descriptors - Composition (with customized amino acid classification support)

Description

This function calculates the Composition descriptor of the CTD descriptors, with customized amino acid classification support.

Usage

```
extractCTDCClass(x, aagroup1, aagroup2, aagroup3)
```

Arguments

x	A character vector, as the input protein sequence.
aagroup1	A named list which contains the first group of customized amino acid classification. See example below.
aagroup2	A named list which contains the second group of customized amino acid classification. See example below.
aagroup3	A named list which contains the third group of customized amino acid classification. See example below.

Value

A length k * 3 named vector, k is the number of amino acid properties used.

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Inna Dubchak, Ilya Muchink, Stephen R. Holbrook and Sung-Hou Kim. Prediction of protein folding class using global description of amino acid sequence. *Proceedings of the National Academy of Sciences*. USA, 1995, 92, 8700-8704.

Inna Dubchak, Ilya Muchink, Christopher Mayor, Igor Dralyuk and Sung-Hou Kim. Recognition of a Protein Fold in the Context of the SCOP classification. *Proteins: Structure, Function and Genetics*, 1999, 35, 401-407.

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See Also

See extractCTDTClass and extractCTDDClass for Transition and Distribution of the CTD descriptors with customized amino acid classification support.

Examples

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]</pre>
# using five customized amino acid property classification
group1 <- list(</pre>
  "hydrophobicity" = c("R", "K", "E", "D", "Q", "N"),
  "normwaalsvolume" = c("G", "A", "S", "T", "P", "D", "C"),
  "polarizability" = c("G", "A", "S", "D", "T"),
  "secondarystruct" = c("E", "A", "L", "M", "Q", "K", "R", "H"),
  "solventaccess" = c("A", "L", "F", "C", "G", "I", "V", "W")
)
group2 <- list(</pre>
  "hydrophobicity" = c("G", "A", "S", "T", "P", "H", "Y"),
  "normwaalsvolume" = c("N", "V", "E", "Q", "I", "L"),
"polarizability" = c("C", "P", "N", "V", "E", "Q", "I", "L"),
  "secondarystruct" = c("V", "I", "Y", "C", "W", "F", "T"),
  "solventaccess" = c("R", "K", "Q", "E", "N", "D")
)
group3 <- list(</pre>
  "hydrophobicity" = c("C", "L", "V", "I", "M", "F", "W"),
  "normwaalsvolume" = c("M", "H", "K", "F", "R", "Y", "W"),
  "polarizability" = c("K", "M", "H", "F", "R", "Y", "W"),
"secondarystruct" = c("G", "N", "P", "S", "D"),
  "solventaccess" = c("M", "S", "P", "T", "H", "Y")
)
extractCTDCClass(x, aagroup1 = group1, aagroup2 = group2, aagroup3 = group3)
```

extractCTDD

CTD Descriptors - Distribution

Description

This function calculates the Distribution descriptor of the CTD descriptors (dim: 105).

Usage

```
extractCTDD(x)
```

Arguments

x A character vector, as the input protein sequence.

extractCTDDClass 27

Value

A length 105 named vector

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Inna Dubchak, Ilya Muchink, Stephen R. Holbrook and Sung-Hou Kim. Prediction of protein folding class using global description of amino acid sequence. *Proceedings of the National Academy of Sciences*. USA, 1995, 92, 8700-8704.

Inna Dubchak, Ilya Muchink, Christopher Mayor, Igor Dralyuk and Sung-Hou Kim. Recognition of a Protein Fold in the Context of the SCOP classification. *Proteins: Structure, Function and Genetics*, 1999, 35, 401-407.

See Also

See extractCTDC and extractCTDT for Composition and Transition of the CTD descriptors.

Examples

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
extractCTDD(x)</pre>
```

extractCTDDClass *CTD Descriptors - Distribution (with customized amino acid classification support)*

Description

This function calculates the Distribution descriptor of the CTD descriptors, with customized amino acid classification support.

Usage

```
extractCTDDClass(x, aagroup1, aagroup2, aagroup3)
```

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Arguments

X	A character vector, as the input protein sequence.
aagroup1	A named list which contains the first group of customized amino acid classification. See example below.
aagroup2	A named list which contains the second group of customized amino acid classification. See example below.
aagroup3	A named list which contains the third group of customized amino acid classification. See example below.

Value

A length k * 15 named vector, k is the number of amino acid properties used.

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <https://nanx.me>
```

References

Inna Dubchak, Ilya Muchink, Stephen R. Holbrook and Sung-Hou Kim. Prediction of protein folding class using global description of amino acid sequence. *Proceedings of the National Academy of Sciences*. USA, 1995, 92, 8700-8704.

Inna Dubchak, Ilya Muchink, Christopher Mayor, Igor Dralyuk and Sung-Hou Kim. Recognition of a Protein Fold in the Context of the SCOP classification. *Proteins: Structure, Function and Genetics*, 1999, 35, 401-407.

See Also

See extractCTDCClass and extractCTDTClass for Composition and Transition of the CTD descriptors with customized amino acid classification support.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
# using five customized amino acid property classification
group1 <- list(
   "hydrophobicity" = c("R", "K", "E", "D", "Q", "N"),
   "normwaalsvolume" = c("G", "A", "S", "T", "P", "D", "C"),
   "polarizability" = c("G", "A", "S", "D", "T"),
   "secondarystruct" = c("E", "A", "L", "M", "Q", "K", "R", "H"),
   "solventaccess" = c("A", "L", "F", "C", "G", "I", "V", "W")
)</pre>
```

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```
group2 <- list(
    "hydrophobicity" = c("G", "A", "S", "T", "P", "H", "Y"),
    "normwaalsvolume" = c("N", "V", "E", "Q", "I", "L"),
    "polarizability" = c("C", "P", "N", "V", "E", "Q", "I", "L"),
    "secondarystruct" = c("V", "I", "Y", "C", "W", "F", "T"),
    "solventaccess" = c("R", "K", "Q", "E", "N", "D")
)

group3 <- list(
    "hydrophobicity" = c("C", "L", "V", "I", "M", "F", "W"),
    "normwaalsvolume" = c("M", "H", "K", "F", "R", "Y", "W"),
    "polarizability" = c("K", "M", "H", "F", "R", "Y", "W"),
    "secondarystruct" = c("G", "N", "P", "S", "D"),
    "solventaccess" = c("M", "S", "P", "T", "H", "Y")
)

extractCTDDClass(x, aagroup1 = group1, aagroup2 = group2, aagroup3 = group3)</pre>
```

extractCTDT

CTD Descriptors - Transition

Description

This function calculates the Transition descriptor of the CTD descriptors (dim: 21).

Usage

```
extractCTDT(x)
```

Arguments

Х

A character vector, as the input protein sequence.

Value

A length 21 named vector

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

30 extractCTDTClass

References

Inna Dubchak, Ilya Muchink, Stephen R. Holbrook and Sung-Hou Kim. Prediction of protein folding class using global description of amino acid sequence. *Proceedings of the National Academy of Sciences*. USA, 1995, 92, 8700-8704.

Inna Dubchak, Ilya Muchink, Christopher Mayor, Igor Dralyuk and Sung-Hou Kim. Recognition of a Protein Fold in the Context of the SCOP classification. *Proteins: Structure, Function and Genetics*, 1999, 35, 401-407.

See Also

See extractCTDC and extractCTDD for Composition and Distribution of the CTD descriptors.

Examples

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
extractCTDT(x)</pre>
```

	TD Descriptors - Transition (with customized amino acid classifica- on support)
--	--

Description

This function calculates the Transition descriptor of the CTD descriptors, with customized amino acid classification support.

Usage

```
extractCTDTClass(x, aagroup1, aagroup2, aagroup3)
```

Arguments

X	A character vector, as the input protein sequence.
aagroup1	A named list which contains the first group of customized amino acid classification. See example below.
aagroup2	A named list which contains the second group of customized amino acid classification. See example below.
aagroup3	A named list which contains the third group of customized amino acid classification. See example below.

Value

A length k * 3 named vector, k is the number of amino acid properties used.

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Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Inna Dubchak, Ilya Muchink, Stephen R. Holbrook and Sung-Hou Kim. Prediction of protein folding class using global description of amino acid sequence. *Proceedings of the National Academy of Sciences*. USA, 1995, 92, 8700-8704.

Inna Dubchak, Ilya Muchink, Christopher Mayor, Igor Dralyuk and Sung-Hou Kim. Recognition of a Protein Fold in the Context of the SCOP classification. *Proteins: Structure, Function and Genetics*, 1999, 35, 401-407.

See Also

See extractCTDCClass and extractCTDDClass for Composition and Distribution of the CTD descriptors with customized amino acid classification support.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]</pre>
# using five customized amino acid property classification
group1 <- list(</pre>
  "hydrophobicity" = c("R", "K", "E", "D", "Q", "N"),
  "normwaalsvolume" = c("G", "A", "S", "T", "P", "D", "C"),
  "polarizability" = c("G", "A", "S", "D", "T"),
  "secondarystruct" = c("E", "A", "L", "M", "Q", "K", "R", "H"),
  "solventaccess" = c("A", "L", "F", "C", "G", "I", "V", "W")
)
group2 <- list(</pre>
  "hydrophobicity" = c("G", "A", "S", "T", "P", "H", "Y"),
  "normwaalsvolume" = c("N", "V", "E", "Q", "I", "L"),
  "polarizability" = c("C", "P", "N", "V", "E", "Q", "I", "L"),
  "secondarystruct" = c("V", "I", "Y", "C", "W", "F", "T"),
  "solventaccess" = c("R", "K", "Q", "E", "N", "D")
)
group3 <- list(
  "hydrophobicity" = c("C", "L", "V", "I", "M", "F", "W"),
  "normwaalsvolume" = c("M", "H", "K", "F", "R", "Y", "W"),
  "polarizability" = c("K", "M", "H", "F", "R", "Y", "W"),
  "secondarystruct" = c("G", "N", "P", "S", "D"),
  "solventaccess" = c("M", "S", "P", "T", "H", "Y")
)
```

32 extractCTriad

```
extractCTDTClass(x, aagroup1 = group1, aagroup2 = group2, aagroup3 = group3)
```

extractCTriad

Conjoint Triad Descriptor

Description

This function calculates the Conjoint Triad descriptor (dim: 343).

Usage

```
extractCTriad(x)
```

Arguments

Χ

A character vector, as the input protein sequence.

Value

A length 343 named vector

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

J.W. Shen, J. Zhang, X.M. Luo, W.L. Zhu, K.Q. Yu, K.X. Chen, Y.X. Li, H.L. Jiang. Predicting Protein-protein Interactions Based Only on Sequences Information. *Proceedings of the National Academy of Sciences*. 007, 104, 4337–4341.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
extractCTriad(x)</pre>
```

extractCTriadClass 33

extractCTriadClass	Conjoint Triad Descriptor (with customized amino acid classification
	support)

Description

This function calculates the Conjoint Triad descriptor, with customized amino acid classification support.

Usage

```
extractCTriadClass(x, aaclass)
```

Arguments

x A character vector, as the input protein sequence.

aaclass A list containing the customized amino acid classification. See example below.

Value

A length k^3 named vector, where k is the number of customized classes of the amino acids.

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

J.W. Shen, J. Zhang, X.M. Luo, W.L. Zhu, K.Q. Yu, K.X. Chen, Y.X. Li, H.L. Jiang. Predicting Protein-protein Interactions Based Only on Sequences Information. *Proceedings of the National Academy of Sciences*. 007, 104, 4337–4341.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
# use customized amino acid classification (normalized van der Waals volume)
newclass <- list(
    c("G", "A", "S", "T", "P", "D", "C"),
    c("N", "V", "E", "Q", "I", "L"),
    c("M", "H", "K", "F", "R", "Y", "W")
)</pre>
```

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```
extractCTriadClass(x, aaclass = newclass)
```

extractDC

Dipeptide Composition Descriptor

Description

This function calculates the Dipeptide Composition descriptor (dim: 400).

Usage

```
extractDC(x)
```

Arguments

Х

A character vector, as the input protein sequence.

Value

A length 400 named vector

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

M. Bhasin, G. P. S. Raghava. Classification of Nuclear Receptors Based on Amino Acid Composition and Dipeptide Composition. *Journal of Biological Chemistry*, 2004, 279, 23262.

See Also

See extractAAC and extractTC for Amino Acid Composition and Tripeptide Composition descriptors.

```
x \leftarrow readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]] extractDC(x)
```

extractDescScales 35

extractDescScales

Scales-Based Descriptors with 20+ classes of Molecular Descriptors

Description

This function calculates the scales-based descriptors with molecular descriptors sets calculated by Dragon, Discovery Studio and MOE. Users can specify which molecular descriptors to select from one of these descriptor sets by specify the numerical or character index of the molecular descriptors in the descriptor set.

Usage

```
extractDescScales(
    x,
    propmat,
    index = NULL,
    pc,
    lag,
    scale = TRUE,
    silent = TRUE
)
```

Arguments

_		
	х	A character vector, as the input protein sequence.
	propmat	The matrix containing the descriptor set for the amino acids, which can be chosen from AAMOE2D, AAMOE3D, AACPSA, AADescall, AA2DACOR, AA3DMORSE, AAACF, AABurden, AAConn, AAConst, AAEdgeAdj, AAEigIdx, AAFGC, AAGeom, AAGETAWAY, AAInfo, AAMolProp, AARandic, AARDF, AATopo, AATopoChg, AAWalk and AAWHIM.
	index	Integer vector or character vector. Specify which molecular descriptors to select from one of these descriptor sets by specify the numerical or character index of the molecular descriptors in the descriptor set. Default is NULL, which means selecting all the molecular descriptors in this descriptor set.
	pc	Integer. The maximum dimension of the space which the data are to be represented in. Must be no greater than the number of amino acid properties provided.
	lag	The lag parameter. Must be less than the amino acids.
	scale	Logical. Should we auto-scale the property matrix (propmat) before doing MDS? Default is TRUE.
	silent	Logical. Whether we print the standard deviation, proportion of variance and the cumulative proportion of the selected principal components or not. Default is TRUE.

Value

A length lag * p^2 named vector, p is the number of scales selected.

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Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

Examples

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
descscales <- extractDescScales(
    x,
    propmat = "AATopo", index = c(37:41, 43:47),
    pc = 5, lag = 7, silent = FALSE
)</pre>
```

extractFAScales

Scales-Based Descriptors derived by Factor Analysis

Description

This function calculates scales-based descriptors derived by Factor Analysis (FA). Users can provide customized amino acid property matrices.

Usage

```
extractFAScales(
    x,
    propmat,
    factors,
    scores = "regression",
    lag,
    scale = TRUE,
    silent = TRUE
)
```

Arguments

X	A character vector, as the input protein sequence.
propmat	A matrix containing the properties for the amino acids. Each row represent one amino acid type, each column represents one property. Note that the one-letter row names must be provided for we need them to seek the properties for each AA type.
factors	Integer. The number of factors to be fitted. Must be no greater than the number of AA properties provided.
scores	Type of scores to produce. The default is "regression", which gives Thompson's scores, "Bartlett" given Bartlett's weighted least-squares scores.
lag	The lag parameter. Must be less than the amino acids number in the protein sequence.

extractGeary 37

scale	Logical. Should we auto-scale the property matrix (propmat) before doing Factor Analysis? Default is TRUE.
silent	Logical. Whether we print the SS loadings, proportion of variance and the cumulative proportion of the selected factors or not. Default is TRUE.

Value

A length lag * p^2 named vector, p is the number of scales (factors) selected.

Author(s)

```
Nan Xiao <https://nanx.me>
```

References

Atchley, W. R., Zhao, J., Fernandes, A. D., & Druke, T. (2005). Solving the protein sequence metric problem. Proceedings of the National Academy of Sciences of the United States of America, 102(18), 6395-6400.

Examples

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
data(AATopo)
tprops <- AATopo[, c(37:41, 43:47)] # select a set of topological descriptors
fa <- extractFAScales(x, propmat = tprops, factors = 5, lag = 7, silent = FALSE)</pre>
```

extractGeary

Geary Autocorrelation Descriptor

Description

This function calculates the Geary autocorrelation descriptor (dim: length(props) * nlag).

Usage

```
extractGeary(
    x,
    props = c("CIDH920105", "BHAR880101", "CHAM820101", "CHAM820102", "CHOC760101",
        "BIGC670101", "CHAM810101", "DAYM780201"),
        nlag = 30L,
        customprops = NULL
)
```

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Arguments

X

A character vector, as the input protein sequence.

props

A character vector, specifying the Accession Number of the target properties. 8 properties are used by default, as listed below:

AccNo. CIDH920105 Normalized average hydrophobicity scales (Cid et al., 1992)

AccNo. BHAR880101 Average flexibility indices (Bhaskaran-Ponnuswamy, 1988)

AccNo. CHAM820101 Polarizability parameter (Charton-Charton, 1982)

AccNo. CHAM820102 Free energy of solution in water, kcal/mole (Charton-Charton, 1982)

AccNo. CHOC760101 Residue accessible surface area in tripeptide (Chothia, 1976)

AccNo. BIGC670101 Residue volume (Bigelow, 1967) **AccNo. CHAM810101** Steric parameter (Charton, 1981)

AccNo. DAYM780201 Relative mutability (Dayhoff et al., 1978b)

nlag

Maximum value of the lag parameter. Default is 30.

customprops

A n x 21 named data frame contains n customized property. Each row contains one property. The column order for different amino acid types is 'AccNo', 'A', 'R', 'N', 'D', 'C', 'E', 'Q', 'G', 'H', 'I', 'L', 'K', 'M', 'F', 'P', 'S', 'T', 'W', 'Y', 'V', and the columns should also be *exactly* named like this. The AccNo column contains the properties' names. Then users should explicitly specify these properties with these names in the argument props. See the examples below for a demonstration. The default value for customprops is NULL.

Value

A length length(props) * nlag named vector.

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

Nan Xiao <https://nanx.me>

References

AAindex: Amino acid index database. https://www.genome.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.

extractMDSScales 39

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.

See Also

See extractMoreauBroto and extractMoran for Moreau-Broto autocorrelation descriptors and Moran autocorrelation descriptors.

Examples

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]</pre>
extractGeary(x)
myprops <- data.frame(</pre>
 AccNo = c("MyProp1", "MyProp2", "MyProp3"),
 A = c(0.62, -0.5, 15), R = c(-2.53, 3, 101),
 N = c(-0.78, 0.2, 58), D = c(-0.9, 3, 59),
 C = c(0.29, -1, 47), E = c(-0.74, 3, 73),
 Q = c(-0.85, 0.2, 72), G = c(0.48, 0, 1),
 H = c(-0.4, -0.5, 82), I = c(1.38, -1.8, 57),
 L = c(1.06, -1.8, 57), K = c(-1.5, 3, 73),
 M = c(0.64, -1.3, 75), F = c(1.19, -2.5, 91),
 P = c(0.12, 0, 42), S = c(-0.18, 0.3, 31),
 T = c(-0.05, -0.4, 45), W = c(0.81, -3.4, 130),
 Y = c(0.26, -2.3, 107), V = c(1.08, -1.5, 43)
# Use 4 properties in the AAindex database, and 3 cutomized properties
extractGeary(
 х,
 customprops = myprops,
 props = c(
    "CIDH920105", "BHAR880101",
    "CHAM820101", "CHAM820102",
    "MyProp1", "MyProp2", "MyProp3"
 )
)
```

extractMDSScales

Scales-Based Descriptors derived by Multidimensional Scaling

Description

This function calculates scales-based descriptors derived by Multidimensional Scaling (MDS). Users can provide customized amino acid property matrices.

Usage

```
extractMDSScales(x, propmat, k, lag, scale = TRUE, silent = TRUE)
```

40 extractMDSScales

Arguments

Х	A character vector, as the input protein sequence.
propmat	A matrix containing the properties for the amino acids. Each row represent one amino acid type, each column represents one property. Note that the one-letter row names must be provided for we need them to seek the properties for each AA type.
k	Integer. The maximum dimension of the space which the data are to be represented in. Must be no greater than the number of AA properties provided.
lag	The lag parameter. Must be less than the amino acids.
scale	Logical. Should we auto-scale the property matrix (propmat) before doing MDS? Default is TRUE.
silent	Logical. Whether to print the k eigenvalues computed during the scaling process or not. Default is TRUE.

Value

A length $lag * p^2$ named vector, p is the number of scales (dimensionality) selected.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Venkatarajan, M. S., & Braun, W. (2001). New quantitative descriptors of amino acids based on multidimensional scaling of a large number of physical-chemical properties. Molecular modeling annual, 7(12), 445–453.

See Also

See extractScales for scales-based descriptors derived by Principal Components Analysis.

```
x \leftarrow \text{readFASTA}(\text{system.file}(\text{"protseq/P00750.fasta"}, \text{package = "protr"}))[[1]]  data(AATopo) tprops <- AATopo[, c(37:41, 43:47)] # select a set of topological descriptors mds <- extractMDSScales(x, propmat = tprops, k = 5, lag = 7, silent = FALSE)
```

extractMoran 41

extractMoran

Moran Autocorrelation Descriptor

Description

This function calculates the Moran autocorrelation descriptor (dim: length(props) * nlag).

Usage

```
extractMoran(
    x,
    props = c("CIDH920105", "BHAR880101", "CHAM820101", "CHAM820102", "CHOC760101",
        "BIGC670101", "CHAM810101", "DAYM780201"),
    nlag = 30L,
    customprops = NULL
)
```

Arguments

v

A character vector, as the input protein sequence.

props

A character vector, specifying the Accession Number of the target properties. 8 properties are used by default, as listed below:

AccNo. CIDH920105 Normalized average hydrophobicity scales (Cid et al., 1992)

AccNo. BHAR880101 Average flexibility indices (Bhaskaran-Ponnuswamy, 1988)

AccNo. CHAM820101 Polarizability parameter (Charton-Charton, 1982)

AccNo. CHAM820102 Free energy of solution in water, kcal/mole (Charton-Charton, 1982)

AccNo. CHOC760101 Residue accessible surface area in tripeptide (Chothia, 1976)

AccNo. BIGC670101 Residue volume (Bigelow, 1967)

AccNo. CHAM810101 Steric parameter (Charton, 1981)

AccNo. DAYM780201 Relative mutability (Dayhoff et al., 1978b)

nlag

Maximum value of the lag parameter. Default is 30.

customprops

A n x 21 named data frame contains n customized property. Each row contains one property. The column order for different amino acid types is 'AccNo', 'A', 'R', 'N', 'D', 'C', 'E', 'Q', 'G', 'H', 'I', 'L', 'K', 'M', 'F', 'P', 'S', 'T', 'W', 'Y', 'V', and the columns should also be *exactly* named like this. The AccNo column contains the properties' names. Then users should explicitly specify these properties with these names in the argument props. See the examples below for a demonstration. The default value for customprops is NULL.

Value

A length length(props) * nlag named vector.

42 extractMoran

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

AAindex: Amino acid index database. https://www.genome.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.

See Also

See extractMoreauBroto and extractGeary for Moreau-Broto autocorrelation descriptors and Geary autocorrelation descriptors.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]</pre>
extractMoran(x)
myprops <- data.frame(</pre>
 AccNo = c("MyProp1", "MyProp2", "MyProp3"),
 A = c(0.62, -0.5, 15), R = c(-2.53, 3, 101),
 N = c(-0.78, 0.2, 58), D = c(-0.9, 3, 59),
 C = c(0.29, -1, 47), E = c(-0.74, 3, 73),
 Q = c(-0.85, 0.2, 72), G = c(0.48, 0, 1),
 H = c(-0.4, -0.5, 82), I = c(1.38, -1.8, 57),
 L = c(1.06, -1.8, 57), K = c(-1.5, 3, 73),
 M = c(0.64, -1.3, 75), F = c(1.19, -2.5, 91),
 P = c(0.12, 0, 42), S = c(-0.18, 0.3, 31),
 T = c(-0.05, -0.4, 45), W = c(0.81, -3.4, 130),
 Y = c(0.26, -2.3, 107), V = c(1.08, -1.5, 43)
# Use 4 properties in the AAindex database, and 3 cutomized properties
extractMoran(
 Х,
 customprops = myprops,
 props = c(
    "CIDH920105", "BHAR880101",
```

extractMoreauBroto 43

```
"CHAM820101", "CHAM820102",
"MyProp1", "MyProp2", "MyProp3"
)
```

extractMoreauBroto

Normalized Moreau-Broto Autocorrelation Descriptor

Description

This function calculates the normalized Moreau-Broto autocorrelation descriptor (dim: length(props) * nlag).

Usage

```
extractMoreauBroto(
    x,
    props = c("CIDH920105", "BHAR880101", "CHAM820101", "CHAM820102", "CHOC760101",
        "BIGC670101", "CHAM810101", "DAYM780201"),
    nlag = 30L,
    customprops = NULL
)
```

Arguments

Χ

A character vector, as the input protein sequence.

props

A character vector, specifying the Accession Number of the target properties. 8 properties are used by default, as listed below:

AccNo. CIDH920105 Normalized average hydrophobicity scales (Cid et al., 1992)

AccNo. BHAR880101 Average flexibility indices (Bhaskaran-Ponnuswamy, 1988)

AccNo. CHAM820101 Polarizability parameter (Charton-Charton, 1982)

AccNo. CHAM820102 Free energy of solution in water, kcal/mole (Charton-Charton, 1982)

AccNo. CHOC760101 Residue accessible surface area in tripeptide (Chothia, 1976)

AccNo. BIGC670101 Residue volume (Bigelow, 1967)

AccNo. CHAM810101 Steric parameter (Charton, 1981)

AccNo. DAYM780201 Relative mutability (Dayhoff et al., 1978b)

nlag

Maximum value of the lag parameter. Default is 30.

customprops

A n x 21 named data frame contains n customized property. Each row contains one property. The column order for different amino acid types is 'AccNo', 'A', 'R', 'N', 'D', 'C', 'E', 'Q', 'G', 'H', 'I', 'L', 'K', 'M', 'F', 'P', 'S', 'T', 'W', 'Y', 'V', and the columns should also be *exactly* named like this. The AccNo column contains the properties' names. Then users should explicitly specify these properties with these names in the argument props. See the examples below for a demonstration. The default value for customprops is NULL.

44 extractMoreauBroto

Value

A length length(props) * nlag named vector.

Note

For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <https://nanx.me>
```

References

AAindex: Amino acid index database. https://www.genome.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.

See Also

See extractMoran and extractGeary for Moran autocorrelation descriptors and Geary autocorrelation descriptors.

Examples

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
extractMoreauBroto(x)
myprops <- data.frame(</pre>
 AccNo = c("MyProp1", "MyProp2", "MyProp3"),
 A = c(0.62, -0.5, 15), R = c(-2.53, 3, 101),
 N = c(-0.78, 0.2, 58), D = c(-0.9, 3, 59),
 C = c(0.29, -1, 47), E = c(-0.74, 3, 73),
 Q = c(-0.85, 0.2, 72), G = c(0.48, 0, 1),
 H = c(-0.4, -0.5, 82), I = c(1.38, -1.8, 57),
 L = c(1.06, -1.8, 57), K = c(-1.5, 3, 73),
 M = c(0.64, -1.3, 75), F = c(1.19, -2.5, 91),
 P = c(0.12, 0, 42), S = c(-0.18, 0.3, 31),
 T = c(-0.05, -0.4, 45), W = c(0.81, -3.4, 130),
 Y = c(0.26, -2.3, 107), V = c(1.08, -1.5, 43)
# Use 4 properties in the AAindex database, and 3 cutomized properties
```

extractMoreauBroto(

extractPAAC 45

```
x,
customprops = myprops,
props = c(
   "CIDH920105", "BHAR880101",
   "CHAM820101", "CHAM820102",
   "MyProp1", "MyProp2", "MyProp3"
)
)
```

extractPAAC

Pseudo Amino Acid Composition (PseAAC) Descriptor

Description

This function calculates the Pseudo Amino Acid Composition (PseAAC) descriptor (dim: 20 + lambda, default is 50).

Usage

```
extractPAAC(
    x,
    props = c("Hydrophobicity", "Hydrophilicity", "SideChainMass"),
    lambda = 30,
    w = 0.05,
    customprops = NULL
)
```

Arguments

x A character vector, as the input protein sequence.

props A character vector, specifying the properties used. 3 properties are used by

default, as listed below:

'Hydrophobicity' Hydrophobicity value of the 20 amino acids

'Hydrophilicity' Hydrophilicity value of the 20 amino acids

'SideChainMass' Side-chain mass of the 20 amino acids

1ambda The lambda parameter for the PseAAC descriptors, default is 30.

w The weighting factor, default is 0.05.

customprops A n x 21 named data frame contains n customized property. Each row contains

one property. The column order for different amino acid types is 'AccNo', 'A', 'R', 'N', 'D', 'C', 'E', 'Q', 'G', 'H', 'I', 'L', 'K', 'M', 'F', 'P', 'S', 'T', 'W', 'Y', 'V', and the columns should also be *exactly* named like this. The AccNo column contains the properties' names. Then users should explicitly specify these properties with these names in the argument props. See the examples below for a demonstration. The default value for customprops is NULL.

46 extractPAAC

Value

A length 20 + lambda named vector

Note

Note the default 20 * 3 prop values have already been independently given in the function. Users can also specify other (up to 544) properties with the Accession Number in the AAindex data, with or without the default three properties, which means users should explicitly specify the properties to use. For this descriptor type, users need to intelligently evaluate the underlying details of the descriptors provided, instead of using this function with their data blindly. It would be wise to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Kuo-Chen Chou. Prediction of Protein Cellular Attributes Using Pseudo-Amino Acid Composition. *PROTEINS: Structure, Function, and Genetics*, 2001, 43: 246-255.

Kuo-Chen Chou. Using Amphiphilic Pseudo Amino Acid Composition to Predict Enzyme Subfamily Classes. *Bioinformatics*, 2005, 21, 10-19.

JACS, 1962, 84: 4240-4246. (C. Tanford). (The hydrophobicity data)

PNAS, 1981, 78:3824-3828 (T.P.Hopp & K.R.Woods). (The hydrophilicity data)

CRC Handbook of Chemistry and Physics, 66th ed., CRC Press, Boca Raton, Florida (1985). (The side-chain mass data)

R.M.C. Dawson, D.C. Elliott, W.H. Elliott, K.M. Jones, Data for Biochemical Research 3rd ed., Clarendon Press Oxford (1986). (The side-chain mass data)

See Also

See extractAPAAC for amphiphilic pseudo amino acid composition descriptor.

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
extractPAAC(x)

myprops <- data.frame(
    AccNo = c("MyProp1", "MyProp2", "MyProp3"),
    A = c(0.62, -0.5, 15), R = c(-2.53, 3, 101),
    N = c(-0.78, 0.2, 58), D = c(-0.9, 3, 59),
    C = c(0.29, -1, 47), E = c(-0.74, 3, 73),
    Q = c(-0.85, 0.2, 72), G = c(0.48, 0, 1),
    H = c(-0.4, -0.5, 82), I = c(1.38, -1.8, 57),
    L = c(1.06, -1.8, 57), K = c(-1.5, 3, 73),
    M = c(0.64, -1.3, 75), F = c(1.19, -2.5, 91),</pre>
```

extractProtFP 47

extractProtFP

Amino Acid Properties Based Scales Descriptors (Protein Fingerprint)

Description

This function calculates amino acid properties based scales descriptors (protein fingerprint). Users can specify which AAindex properties to select from the AAindex database by specify the numerical or character index of the properties in the AAindex database.

Usage

```
extractProtFP(x, index = NULL, pc, lag, scale = TRUE, silent = TRUE)
```

Arguments

X	A character vector, as the input protein sequence.
index	Integer vector or character vector. Specify which AAindex properties to select from the AAindex database by specify the numerical or character index of the properties in the AAindex database. Default is NULL, means selecting all the AA properties in the AAindex database.
рс	Integer. Use the first pc principal components as the scales. Must be no greater than the number of AA properties provided.
lag	The lag parameter. Must be less than the amino acids.
scale	Logical. Should we auto-scale the property matrix before PCA? Default is TRUE.
silent	Logical. Whether we print the standard deviation, proportion of variance and the cumulative proportion of the selected principal components or not. Default is TRUE.

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Value

A length lag * p^2 named vector, p is the number of scales (principal components) selected.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

Examples

```
x \leftarrow readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]] fp <- extractProtFP(x, index = c(160:165, 258:296), pc = 5, lag = 7, silent = FALSE)
```

extractProtFPGap Amino Acid Properties Based Scales Descriptors (Protein Fingerprint) with Gap Support

Description

This function calculates amino acid properties based scales descriptors (protein fingerprint) with gap support. Users can specify which AAindex properties to select from the AAindex database by specify the numerical or character index of the properties in the AAindex database.

Usage

```
extractProtFPGap(x, index = NULL, pc, lag, scale = TRUE, silent = TRUE)
```

Arguments

A character vector, as the input protein sequence. Use '-' to represent gaps in the sequence.
Integer vector or character vector. Specify which AAindex properties to select from the AAindex database by specify the numerical or character index of the properties in the AAindex database. Default is NULL, means selecting all the AA properties in the AAindex database.
Integer. Use the first pc principal components as the scales. Must be no greater than the number of AA properties provided.
The lag parameter. Must be less than the amino acids.
$Logical. \ Should we auto-scale the property matrix before PCA? \ Default is \ TRUE.$
Logical. Whether we print the standard deviation, proportion of variance and the cumulative proportion of the selected principal components or not. Default is TRUE.

Value

A length lag * p^2 named vector, p is the number of scales (principal components) selected.

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Author(s)

```
Nan Xiao <https://nanx.me>
```

Examples

```
# amino acid sequence with gaps
x <- readFASTA(system.file("protseq/align.fasta", package = "protr"))$`IXI_235`
fp <- extractProtFPGap(x, index = c(160:165, 258:296), pc = 5, lag = 7, silent = FALSE)</pre>
```

extractPSSM

Compute PSSM (Position-Specific Scoring Matrix) for given protein sequence

Description

This function calculates the PSSM (Position-Specific Scoring Matrix) derived by PSI-Blast for given protein sequence or peptides.

Usage

```
extractPSSM(
  seq,
  start.pos = 1L,
  end.pos = nchar(seq),
  psiblast.path = NULL,
 makeblastdb.path = NULL,
  database.path = NULL,
  iter = 5,
  silent = TRUE,
  evalue = 10L,
  word.size = NULL,
  gapopen = NULL,
  gapextend = NULL,
 matrix = "BLOSUM62",
  threshold = NULL,
  seg = "no",
  soft.masking = FALSE,
  culling.limit = NULL,
  best.hit.overhang = NULL,
  best.hit.score.edge = NULL,
  xdrop.ungap = NULL,
  xdrop.gap = NULL,
  xdrop.gap.final = NULL,
  window.size = NULL,
  gap.trigger = 22L,
  num.threads = 1L,
  pseudocount = 0L,
```

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```
inclusion.ethresh = 0.002
)
```

Arguments

seq Character vector, as the input protein sequence.

start.pos Optional integer denoting the start position of the fragment window. Default is

1, i.e. the first amino acid of the given sequence.

end.pos Optional integer denoting the end position of the fragment window. Default is

nchar(seq), i.e. the last amino acid of the given sequence.

psiblast.path Character string indicating the path of the psiblast program. If NCBI Blast+

was previously installed in the operation system, the path will be automatically

detected.

makeblastdb.path

Character string indicating the path of the makeblastdb program. If NCBI Blast+ was previously installed in the system, the path will be automatically

detected.

database.path Character string indicating the path of a reference database (a FASTA file).

iter Number of iterations to perform for PSI-Blast.

silent Logical. Whether the PSI-Blast running output should be shown or not (May

not work on some Windows versions and PSI-Blast versions), default is TRUE.

evalue Expectation value (E) threshold for saving hits. Default is 10.

word.size Word size for wordfinder algorithm. An integer >= 2.

gapopen Integer. Cost to open a gap.

gapextend Integer. Cost to extend a gap.

matrix Character string. The scoring matrix name (default is "BLOSUM62").

threshold Minimum word score such that the word is added to the BLAST lookup table.

A real value >= 0.

seg Character string. Filter query sequence with SEG ("yes", "window locut hicut",

or "no" to disable). Default is "no".

soft.masking Logical. Apply filtering locations as soft masks? Default is FALSE.

culling.limit An integer >= 0. If the query range of a hit is enveloped by that of at least this

many higher-scoring hits, delete the hit. Incompatible with best.hit.overhang

and best_hit_score_edge.

best.hit.overhang

Best Hit algorithm overhang value (A real value ≥ 0 and =< 0.5, recommended

value: 0.1). Incompatible with culling_limit.

best.hit.score.edge

Best Hit algorithm score edge value (A real value >=0 and =< 0.5, recommended

value: 0.1). Incompatible with culling_limit.

xdrop.ungap X-dropoff value (in bits) for ungapped extensions.

xdrop.gap X-dropoff value (in bits) for preliminary gapped extensions.

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xdrop.gap.final

X-dropoff value (in bits) for final gapped alignment.

window.size An integer >= 0. Multiple hits window size, To specify 1-hit algorithm, use 0.

gap. trigger Number of bits to trigger gapping. Default is 22.

num. threads Integer. Number of threads (CPUs) to use in the BLAST search. Default is 1.

pseudocount Integer. Pseudo-count value used when constructing PSSM. Default is 0.

inclusion.ethresh

E-value inclusion threshold for pairwise alignments. Default is 0.002.

Details

For given protein sequences or peptides, PSSM represents the log-likelihood of the substitution of the 20 types of amino acids at that position in the sequence. Note that the output value is not normalized.

Value

The original PSSM, a numeric matrix which has end.pos - start.pos + 1 columns and 20 named rows.

Note

The function requires the makeblastdb and psiblast programs to be properly installed in the operation system or their paths provided.

The two command-line programs are included in the NCBI-BLAST+ software package. To install NCBI Blast+ for your operating system, see https://blast.ncbi.nlm.nih.gov/doc/blast-help/downloadblastdata.html for detailed instructions.

Ubuntu/Debian users can directly use the command sudo apt-get install ncbi-blast+ to install NCBI Blast+. For OS X users, download ncbi-blast-....dmg then install. For Windows users, download ncbi-blast-....exe then install.

Author(s)

Nan Xiao https://nanx.me

References

Altschul, Stephen F., et al. "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs." *Nucleic acids research* 25.17 (1997): 3389–3402.

Ye, Xugang, Guoli Wang, and Stephen F. Altschul. "An assessment of substitution scores for protein profile-profile comparison." *Bioinformatics* 27.24 (2011): 3356–3363.

Rangwala, Huzefa, and George Karypis. "Profile-based direct kernels for remote homology detection and fold recognition." *Bioinformatics* 21.23 (2005): 4239–4247.

See Also

extractPSSMFeature extractPSSMAcc

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Examples

```
if (Sys.which("makeblastdb") == "" | Sys.which("psiblast") == "") {
  cat("Cannot find makeblastdb or psiblast. Please install NCBI Blast+ first")
} else {
  x <- readFASTA(system.file(
    "protseq/P00750.fasta",
    package = "protr"
))[[1]]
  dbpath <- tempfile("tempdb", fileext = ".fasta")
  invisible(file.copy(from = system.file(
    "protseq/Plasminogen.fasta",
    package = "protr"
), to = dbpath))

pssmmat <- extractPSSM(seq = x, database.path = dbpath)

dim(pssmmat) # 20 x 562 (P00750: length 562, 20 Amino Acids)
}</pre>
```

extractPSSMAcc

Profile-based protein representation derived by PSSM (Position-Specific Scoring Matrix) and auto cross covariance

Description

This function calculates the feature vector based on the PSSM by running PSI-Blast and auto cross covariance tranformation.

Usage

```
extractPSSMAcc(pssmmat, lag)
```

Arguments

pssmmat The PSSM computed by extractPSSM.

The lag parameter. Must be less than the number of amino acids in the sequence

(i.e. the number of columns in the PSSM matrix).

Value

A length lag * 20^2 named numeric vector, the element names are derived by the amino acid name abbreviation (crossed amino acid name abbreviation) and lag index.

Author(s)

Nan Xiao https://nanx.me

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References

Wold, S., Jonsson, J., Sjorstrom, M., Sandberg, M., & Rannar, S. (1993). DNA and peptide sequences and chemical processes multivariately modelled by principal component analysis and partial least-squares projections to latent structures. *Analytica chimica acta*, 277(2), 239–253.

See Also

extractPSSM extractPSSMFeature

Examples

```
if (Sys.which("makeblastdb") == "" | Sys.which("psiblast") == "") {
  cat("Cannot find makeblastdb or psiblast. Please install NCBI Blast+")
} else {
  x <- readFASTA(system.file(
    "protseq/P00750.fasta",
    package = "protr"
))[[1]]
  dbpath <- tempfile("tempdb", fileext = ".fasta")
  invisible(file.copy(from = system.file(
    "protseq/Plasminogen.fasta",
    package = "protr"
), to = dbpath))

  pssmmat <- extractPSSM(seq = x, database.path = dbpath)
  pssmacc <- extractPSSMAcc(pssmmat, lag = 3)
  tail(pssmacc)
}</pre>
```

extractPSSMFeature

Profile-based protein representation derived by PSSM (Position-Specific Scoring Matrix)

Description

This function calculates the profile-based protein representation derived by PSSM. The feature vector is based on the PSSM computed by extractPSSM.

Usage

```
extractPSSMFeature(pssmmat)
```

Arguments

pssmmat

The PSSM computed by extractPSSM.

54 extractPSSMFeature

Details

For a given sequence, the PSSM feature represents the log-likelihood of the substitution of the 20 types of amino acids at that position in the sequence.

Each PSSM feature value in the vector represents the degree of conservation of a given amino acid type. The value is normalized to interval (0, 1) by the transformation $1/(1+e^{-(-x)})$.

Value

A numeric vector which has 20 x N named elements, where N is the size of the window (number of rows of the PSSM).

Author(s)

```
Nan Xiao <https://nanx.me>
```

References

Ye, Xugang, Guoli Wang, and Stephen F. Altschul. "An assessment of substitution scores for protein profile-profile comparison." *Bioinformatics* 27.24 (2011): 3356–3363.

Rangwala, Huzefa, and George Karypis. "Profile-based direct kernels for remote homology detection and fold recognition." *Bioinformatics* 21.23 (2005): 4239–4247.

See Also

extractPSSM extractPSSMAcc

```
if (Sys.which("makeblastdb") == "" | Sys.which("psiblast") == "") {
  cat("Cannot find makeblastdb or psiblast. Please install NCBI Blast+")
} else {
  x <- readFASTA(system.file(
    "protseq/P00750.fasta",
    package = "protr"
))[[1]]
  dbpath <- tempfile("tempdb", fileext = ".fasta")
  invisible(file.copy(from = system.file(
    "protseq/Plasminogen.fasta",
    package = "protr"
), to = dbpath))

pssmmat <- extractPSSM(seq = x, database.path = dbpath)
  pssmfeature <- extractPSSMFeature(pssmmat)
  head(pssmfeature)
}</pre>
```

extractQSO 55

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Quasi-Sequence-Order (QSO) Descriptor

Description

This function calculates the Quasi-Sequence-Order (QSO) descriptor (dim: 20 + 20 + (2 * nlag), default is 100).

Usage

```
extractQSO(x, nlag = 30, w = 0.1)
```

Arguments

x A character vector, as the input protein sequence.

nlag The maximum lag, defualt is 30.

w The weighting factor, default is 0.1.

Value

```
A length 20 + 20 + (2 * nlag) named vector
```

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Kuo-Chen Chou. Prediction of Protein Subcellar Locations by Incorporating Quasi-Sequence-Order Effect. *Biochemical and Biophysical Research Communications*, 2000, 278, 477-483.

Kuo-Chen Chou and Yu-Dong Cai. Prediction of Protein Sucellular Locations by GO-FunD-PseAA Predictor. *Biochemical and Biophysical Research Communications*, 2004, 320, 1236-1239.

Gisbert Schneider and Paul Wrede. The Rational Design of Amino Acid Sequences by Artifical Neural Networks and Simulated Molecular Evolution: Do Novo Design of an Idealized Leader Cleavge Site. *Biophys Journal*, 1994, 66, 335-344.

See Also

See extractSOCN for sequence-order-coupling numbers.

```
x \leftarrow readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]] extractQSO(x)
```

56 extractScales

extractScales	Scales-Based Descriptors derived by Principal Components Analysis
CACI GCCCGTCC	Scares Basea Bescriptors derived by Frincipal Components Finallysis

Description

This function calculates scales-based descriptors derived by Principal Components Analysis (PCA). Users can provide customized amino acid property matrices. This function implements the core computation procedure needed for the scales-based descriptors derived by AA-Properties (AAindex) and scales-based descriptors derived by 20+ classes of 2D and 3D molecular descriptors (Topological, WHIM, VHSE, etc.) in the protr package.

Usage

```
extractScales(x, propmat, pc, lag, scale = TRUE, silent = TRUE)
```

Arguments

Х	A character vector, as the input protein sequence.
propmat	A matrix containing the properties for the amino acids. Each row represent one amino acid type, each column represents one property. Note that the one-letter row names must be provided for we need them to seek the properties for each AA type.
рс	Integer. Use the first pc principal components as the scales. Must be no greater than the number of AA properties provided.
lag	The lag parameter. Must be less than the amino acids.
scale	Logical. Should we auto-scale the property matrix (propmat) before PCA? Default is TRUE.
silent	Logical. Whether we print the standard deviation, proportion of variance and the cumulative proportion of the selected principal components or not. Default is TRUE.

Value

A length lag * p^2 named vector, p is the number of scales (principal components) selected.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

See Also

See extractDescScales scales descriptors based on 20+ classes of molecular descriptors, and extractProtFP for amino acid property based scales descriptors (protein fingerprint).

extractScalesGap 57

Examples

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
data(AAindex)
AAidxmat <- t(na.omit(as.matrix(AAindex[, 7:26])))
scales <- extractScales(x, propmat = AAidxmat, pc = 5, lag = 7, silent = FALSE)</pre>
```

extractScalesGap Scales-Based Descriptors derived by Principal Components Analysis (with Gap Support)

Description

This function calculates scales-based descriptors derived by Principal Components Analysis (PCA), with gap support. Users can provide customized amino acid property matrices. This function implements the core computation procedure needed for the scales-based descriptors derived by AA-Properties (AAindex) and scales-based descriptors derived by 20+ classes of 2D and 3D molecular descriptors (Topological, WHIM, VHSE, etc.) in the protr package.

Usage

```
extractScalesGap(x, propmat, pc, lag, scale = TRUE, silent = TRUE)
```

Arguments

x	A character vector, as the input protein sequence. Use '-' to represent gaps in the sequence.
propmat	A matrix containing the properties for the amino acids. Each row represent one amino acid type, each column represents one property. Note that the one-letter row names must be provided for we need them to seek the properties for each AA type.
рс	Integer. Use the first pc principal components as the scales. Must be no greater than the number of AA properties provided.
lag	The lag parameter. Must be less than the amino acids.
scale	Logical. Should we auto-scale the property matrix (propmat) before PCA? Default is TRUE.
silent	Logical. Whether to print the standard deviation, proportion of variance and the cumulative proportion of the selected principal components or not. Default is TRUE.

Value

A length lag * p^2 named vector, p is the number of scales (principal components) selected.

Author(s)

Nan Xiao https://nanx.me

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See Also

See extractProtFPGap for amino acid property based scales descriptors (protein fingerprint) with gap support.

Examples

```
# amino acid sequence with gaps
x <- readFASTA(system.file("protseq/align.fasta", package = "protr"))$`IXI_235`
data(AAindex)
AAidxmat <- t(na.omit(as.matrix(AAindex[, 7:26])))
scales <- extractScalesGap(x, propmat = AAidxmat, pc = 5, lag = 7, silent = FALSE)</pre>
```

extractSOCN

Sequence-Order-Coupling Numbers

Description

This function calculates the Sequence-Order-Coupling Numbers (dim: nlag * 2, default is 60).

Usage

```
extractSOCN(x, nlag = 30)
```

Arguments

x A character vector, as the input protein sequence.

nlag The maximum lag, defualt is 30.

Value

A length nlag * 2 named vector

Author(s)

```
Nan Xiao <https://nanx.me>
```

References

Kuo-Chen Chou. Prediction of Protein Subcellar Locations by Incorporating Quasi-Sequence-Order Effect. *Biochemical and Biophysical Research Communications*, 2000, 278, 477-483.

Kuo-Chen Chou and Yu-Dong Cai. Prediction of Protein Sucellular Locations by GO-FunD-PseAA Predictor. *Biochemical and Biophysical Research Communications*, 2004, 320, 1236-1239.

Gisbert Schneider and Paul Wrede. The Rational Design of Amino Acid Sequences by Artifical Neural Networks and Simulated Molecular Evolution: Do Novo Design of an Idealized Leader Cleavge Site. *Biophys Journal*, 1994, 66, 335-344.

extractTC 59

See Also

See extractQSO for quasi-sequence-order descriptors.

Examples

```
x \leftarrow readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]] extractSOCN(x)
```

extractTC

Tripeptide Composition Descriptor

Description

This function calculates the Tripeptide Composition descriptor (dim: 8,000).

Usage

```
extractTC(x)
```

Arguments

Χ

A character vector, as the input protein sequence.

Value

A length 8,000 named vector

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

M. Bhasin, G. P. S. Raghava. Classification of Nuclear Receptors Based on Amino Acid Composition and Dipeptide Composition. *Journal of Biological Chemistry*, 2004, 279, 23262.

See Also

See extractAAC and extractDC for Amino Acid Composition and Dipeptide Composition descriptors.

```
x \leftarrow readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]] extractTC(x)
```

getUniProt

getUniProt

Retrieve Protein Sequences from UniProt by Protein ID

Description

This function retrieves protein sequences from uniprot.org by protein ID(s).

Usage

```
getUniProt(id)
```

Arguments

id

A character vector, as the protein ID(s).

Value

A list, each component contains one protein sequence.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

See Also

See readFASTA for reading FASTA format files.

```
## Not run:
# Network latency may slow down this example
# Only test this when your connection is fast enough
ids <- c("P00750", "P00751", "P00752")
getUniProt(ids)
## End(Not run)</pre>
```

OptAA3d 61

OptAA3d.sdf - 20 Amino Acids Optimized with MOE 2011.10 (Semiempirical AM1)

Description

OptAA3d.sdf - 20 Amino Acids Optimized with MOE 2011.10 (Semiempirical AM1)

Examples

```
# this operation requires the rcdk package
# require(rcdk)
# optaa3d = load.molecules(system.file("sysdata/OptAA3d.sdf", package = "protr"))
# view.molecule.2d(optaa3d[[1]]) # view the first AA
```

parGOSim

Protein Similarity Calculation based on Gene Ontology (GO) Similarity

Description

This function calculates protein similarity based on Gene Ontology (GO) similarity.

Usage

```
parGOSim(
  golist,
  type = c("go", "gene"),
  ont = c("MF", "BP", "CC"),
  organism = "human",
  measure = "Resnik",
  combine = "BMA"
)
```

Arguments

A list, each component contains a character vector of GO terms or one Entrez Gene ID.

type Input type for golist, "go" for GO Terms, "gene" for gene ID.

ont Default is "MF", can be one of "MF", "BP", or "CC" subontologies.

organism organism name. Default is "human", can be one of "anopheles", "arabidopsis", "bovine", "canine", "chicken", "chimp", "coelicolor", "ecolik12", "ecsakai", "fly", "human", "malaria", "mouse", "pig", "rat", "rhesus", "worm", "xenopus", "yeast" or "zebrafish". Before specifying the organism, please install the corresponding genome wide annotation data package for the selected organism.

62 parGOSim

```
measure Default is "Resnik", can be one of "Resnik", "Lin", "Rel", "Jiang" or "Wang".

Combine Default is "BMA", can be one of "max", "average", "rcmax" or "BMA" for combining semantic similarity scores of multiple GO terms associated with proteins.
```

Value

A n x n similarity matrix.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

See Also

See twoGOSim for calculating the GO semantic similarity between two groups of GO terms or two Entrez gene IDs. See parSeqSim for paralleled protein similarity calculation based on Smith-Waterman local alignment.

```
## Not run:
# Be careful when testing this since it involves GO similarity computation
# and might produce unpredictable results in some environments
library("GOSemSim")
library("org.Hs.eg.db")
# By GO Terms
# AP4B1
go1 <- c(
  "GO:0005215", "GO:0005488", "GO:0005515",
  "GO:0005625", "GO:0005802", "GO:0005905"
# BCAS2
go2 <- c(
  "GO:0005515", "GO:0005634", "GO:0005681",
  "GO:0008380", "GO:0031202"
)
# PDE4DIP
go3 <- c(
  "GO:0003735", "GO:0005622", "GO:0005840",
  "GO:0006412"
)
golist <- list(go1, go2, go3)</pre>
parGOSim(golist, type = "go", ont = "CC", measure = "Wang")
# By Entrez gene id
genelist <- list(c("150", "151", "152", "1814", "1815", "1816"))</pre>
parGOSim(genelist, type = "gene", ont = "BP", measure = "Wang")
## End(Not run)
```

parSeqSim 63

parSeqSim	Parallel Protein Sequence Similarity Calculation Based on Sequence Alignment (In-Memory Version)

Description

Parallel calculation of protein sequence similarity based on sequence alignment.

Usage

```
parSeqSim(
  protlist,
  cores = 2,
  batches = 1,
  verbose = FALSE,
  type = "local",
  submat = "BLOSUM62",
  gap.opening = 10,
  gap.extension = 4
)
```

Arguments

protlist	A length n list containing n protein sequences, each component of the list is a character string, storing one protein sequence. Unknown sequences should be represented as "".
cores	Integer. The number of CPU cores to use for parallel execution, default is 2. Users can use the availableCores() function in the parallelly package to see how many cores they could use.
batches	Integer. How many batches should we split the pairwise similarity computations into. This is useful when you have a large number of protein sequences, enough number of CPU cores, but not enough RAM to compute and fit all the pairwise similarities into a single batch. Defaults to 1.
verbose	Print the computation progress? Useful when batches > 1.
type	Type of alignment, default is "local", can be "global" or "local", where "global" represents Needleman-Wunsch global alignment; "local" represents Smith-Waterman local alignment.
submat	Substitution matrix, default is "BLOSUM62", can be one of "BLOSUM45", "BLOSUM50", "BLOSUM62", "BLOSUM80", "BLOSUM100", "PAM30", "PAM40", "PAM70", "PAM120", or "PAM250".
gap.opening	The cost required to open a gap of any length in the alignment. Defaults to 10.
gap.extension	The cost to extend the length of an existing gap by 1. Defaults to 4.

Value

A n x n similarity matrix.

64 parSeqSimDisk

Author(s)

```
Nan Xiao <https://nanx.me>
```

See Also

See parSeqSimDisk for the disk-based version.

Examples

```
## Not run:

# Be careful when testing this since it involves parallelization
# and might produce unpredictable results in some environments

library("Biostrings")
library("foreach")
library("doParallel")

s1 <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
s2 <- readFASTA(system.file("protseq/P08218.fasta", package = "protr"))[[1]]
s3 <- readFASTA(system.file("protseq/P10323.fasta", package = "protr"))[[1]]
s4 <- readFASTA(system.file("protseq/P20160.fasta", package = "protr"))[[1]]
s5 <- readFASTA(system.file("protseq/Q9NZP8.fasta", package = "protr"))[[1]]
plist <- list(s1, s2, s3, s4, s5)
(psimmat <- parSeqSim(plist, cores = 2, type = "local", submat = "BLOSUM62"))

## End(Not run)</pre>
```

parSeqSimDisk

Parallel Protein Sequence Similarity Calculation Based on Sequence Alignment (Disk-Based Version)

Description

Parallel calculation of protein sequence similarity based on sequence alignment. This version offloads the partial results in each batch to the hard drive and merges the results together in the end, which reduces the memory usage.

Usage

```
parSeqSimDisk(
  protlist,
  cores = 2,
  batches = 1,
  path = tempdir(),
  verbose = FALSE,
  type = "local",
  submat = "BLOSUM62",
```

parSeqSimDisk 65

```
gap.opening = 10,
gap.extension = 4
)
```

Arguments

protlist A length n list containing n protein sequences, each component of the list is a

character string, storing one protein sequence. Unknown sequences should be

represented as "".

cores Integer. The number of CPU cores to use for parallel execution, default is 2.

Users can use the availableCores() function in the parallelly package to see

how many cores they could use.

batches Integer. How many batches should we split the pairwise similarity computations

into. This is useful when you have a large number of protein sequences, enough number of CPU cores, but not enough RAM to compute and fit all the pairwise

similarities into a single batch. Defaults to 1.

path Directory for caching the results in each batch. Defaults to the temporary direc-

tory.

verbose Print the computation progress? Useful when batches > 1.

type Type of alignment, default is "local", can be "global" or "local", where

"global" represents Needleman-Wunsch global alignment; "local" represents

Smith-Waterman local alignment.

submat Substitution matrix, default is "BLOSUM62", can be one of "BLOSUM45", "BLOSUM50",

"BLOSUM62", "BLOSUM80", "BLOSUM100", "PAM30", "PAM40", "PAM70", "PAM120",

or "PAM250".

gap.opening The cost required to open a gap of any length in the alignment. Defaults to 10.

gap. extension The cost to extend the length of an existing gap by 1. Defaults to 4.

Value

A n x n similarity matrix.

Author(s)

```
Nan Xiao <https://nanx.me>
```

See Also

See parSeqSim for the in-memory version.

```
## Not run:

# Be careful when testing this since it involves parallelization
# and might produce unpredictable results in some environments
library("Biostrings")
```

66 protcheck

```
library("foreach")
library("doParallel")

s1 <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
s2 <- readFASTA(system.file("protseq/P08218.fasta", package = "protr"))[[1]]
s3 <- readFASTA(system.file("protseq/P10323.fasta", package = "protr"))[[1]]
s4 <- readFASTA(system.file("protseq/P20160.fasta", package = "protr"))[[1]]
s5 <- readFASTA(system.file("protseq/Q9NZP8.fasta", package = "protr"))[[1]]
set.seed(1010)
plist <- as.list(c(s1, s2, s3, s4, s5)[sample(1:5, 100, replace = TRUE)])
psimmat <- parSeqSimDisk(
   plist,
   cores = 2, batches = 10, verbose = TRUE,
   type = "local", submat = "BLOSUM62"
)

## End(Not run)</pre>
```

protcheck

Protein sequence amino acid type sanity check

Description

This function checks if the protein sequence's amino acid types are in the 20 default types.

Usage

```
protcheck(x)
```

Arguments

Х

A character vector, as the input protein sequence.

Value

Logical. TRUE if all of the amino acid types of the sequence are within the 20 default types.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

```
x <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
protcheck(x) # TRUE
protcheck(paste(x, "Z", sep = "")) # FALSE</pre>
```

protseg 67

protseg

Protein Sequence Segmentation/Partition

Description

This function extracts the segmentations from the protein sequence.

Usage

```
protseg(
    x,
    aa = c("A", "R", "N", "D", "C", "E", "Q", "G", "H", "I", "L", "K", "M", "F", "P", "S",
        "T", "W", "Y", "V"),
    k = 7
)
```

Arguments

x A character vector, as the input protein sequence.

```
aa A character, the amino acid type. One of 'A', 'R', 'N', 'D', 'C', 'E', 'Q', 'G', 'H', 'I', 'L', 'K', 'M', 'F', 'P', 'S', 'T', 'W', 'Y', 'V'.
```

k A positive integer, specifys the window size (half of the window), default is 7.

Value

A named list, each component contains one of the segmentations (a character string), names of the list components are the positions of the specified amino acid in the sequence.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

```
x \leftarrow \text{readFASTA}(\text{system.file}(\text{"protseq/P00750.fasta"}, \text{package = "protr"}))[[1]] \text{protseg}(x, \text{aa = "R"}, \text{k = 5})
```

68 readFASTA

readFASTA

Read Protein Sequences in FASTA Format

Description

This function reads protein sequences in FASTA format.

Usage

```
readFASTA(
   file = system.file("protseq/P00750.fasta", package = "protr"),
   legacy.mode = TRUE,
   seqonly = FALSE
)
```

Arguments

file Path to the file containing the protein sequences in FASTA format. If it does

not contain an absolute or relative path, the file name is relative to the current working directory, getwd. The default here is to read the P00750.fasta file

which is present in the protseq directory of the protr package.

legacy.mode If set to TRUE, lines starting with a semicolon (;) are ignored. Default value is

TRUE.

segonly If set to TRUE, only sequences as returned without attempt to modify them or to

get their names and annotations (execution time is divided approximately by a

factor 3). Default value is FALSE.

Value

Character vector of the protein sequences.

The three returned argument are just different forms of the same output. If one is interested in a Mahalanobis metric over the original data space, the first argument is all she/he needs. If a transformation into another space (where one can use the Euclidean metric) is preferred, the second returned argument is sufficient. Using A and B is equivalent in the following sense.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Pearson, W.R. and Lipman, D.J. (1988) Improved tools for biological sequence comparison. *Proceedings of the National Academy of Sciences of the United States of America*, **85**: 2444–2448.

See Also

See getUniProt for retrieving protein sequences from uniprot.org.

readPDB 69

Examples

```
P00750 <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))
```

readPDB

Read Protein Sequences in PDB Format

Description

This function reads protein sequences in PDB (Protein Data Bank) format, and return the amino acid sequences represented by single-letter code.

Usage

```
readPDB(file = system.file("protseq/4HHB.pdb", package = "protr"))
```

Arguments

file

Path to the file containing the protein sequences in PDB format. If it does not contain an absolute or relative path, the file name is relative to the current working directory, getwd. The default here is to read the 4HHB.PDB file which is present in the protseq directory of the protr package.

Value

Character vector of the protein sequence.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

References

Protein Data Bank Contents Guide: Atomic Coordinate Entry Format Description, Version 3.30. Accessed 2013-06-26. https://files.wwpdb.org/pub/pdb/doc/format_descriptions/Format_v33_Letter.pdf

See Also

See readFASTA for reading protein sequences in FASTA format.

```
Seq4HHB <- readPDB(system.file("protseq/4HHB.pdb", package = "protr"))</pre>
```

70 removeGaps

Remove or replace gaps from protein sequences.

Description

Remove/replace gaps or any irregular characters from protein sequences, to make them suitable for feature extraction or sequence alignment based similarity computation.

Usage

```
removeGaps(x, pattern = "-", replacement = "", ...)
```

Arguments

character vector, containing the input protein sequence(s).
 character string contains the gap (or other irregular) character to be removed or replaced. Default is "-". For advanced usage, see gsub.
 a replacement a replacement for matched characters. Default is "" (remove the matched character).
 additional parameters for gsub.

Value

a vector of protein sequence(s) with gaps or irregular characters removed/replaced.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

```
# amino acid sequences that contain gaps ("-")
aaseq <- list(
    "MHGDTPTLHEYMLDLQPETTDLYCYEQLSDSSE-EEDEIDGPAGQAEPDRAHYNIVTFCCKCDSTLRLCVQS",
    "MHGDTPTLHEYMLDLQPETTDLYCYEQLNDSSE-EEDEIDGPAGQAEPDRAHYNIVTFCCKCDSTLRLCVQS"
)
## Not run:
#' # gaps create issues for alignment
parSeqSim(aaseq)

# remove the gaps
nogapseq <- removeGaps(aaseq)
parSeqSim(nogapseq)

## End(Not run)</pre>
```

twoGOSim 71

twoGOSim	Protein Similarity Calculation based on Gene Ontology (GO) Similarity
	•

Description

This function calculates the Gene Ontology (GO) similarity between two groups of GO terms or two Entrez gene IDs.

Usage

```
twoGOSim(
  id1,
  id2,
  type = c("go", "gene"),
  ont = c("MF", "BP", "CC"),
  organism = "human",
  measure = "Resnik",
  combine = "BMA"
)
```

Arguments

id1	Character vector. When length > 1: each element is a GO term; when length = 1: the Entrez Gene ID.
id2	Character vector. When length > 1: each element is a GO term; when length = 1: the Entrez Gene ID.
type	Input type of id1 and id2, 'go' for GO Terms, "gene" for gene ID.
ont	Default is "MF", can be one of "MF", "BP", or "CC" subontologies.
organism	Organism name. Default is "human", can be one of "anopheles", "arabidopsis", "bovine", "canine", "chicken", "chimp", "coelicolor", "ecolik12", "ecsakai", "fly", "human", "malaria", "mouse", "pig", "rat", "rhesus", "worm", "xenopus", "yeast" or "zebrafish". Before specifying the organism, please install the corresponding genome wide annotation data package for the selected organism.
measure	Default is "Resnik", can be one of "Resnik", "Lin", "Rel", "Jiang" or "Wang".
combine	Default is "BMA", can be one of "max", "average", "rcmax" or "BMA" for combining semantic similarity scores of multiple GO terms associated with proteins.

Value

Similarity value.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

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See Also

See parGOSim for protein similarity calculation based on Gene Ontology (GO) semantic similarity. See parSeqSim for paralleled protein similarity calculation based on Smith-Waterman local alignment.

Examples

```
## Not run:

# Be careful when testing this since it involves GO similarity computation
# and might produce unpredictable results in some environments

library("GOSemSim")
library("org.Hs.eg.db")

# By GO terms
go1 <- c("GO:0004022", "GO:0004024", "GO:0004023")
go2 <- c("GO:0009055", "GO:0020037")
twoGOSim(go1, go2, type = "go", ont = "MF", measure = "Wang")

# By Entrez gene id
gene1 <- "1956" # EGFR
gene2 <- "2261" # FGFR3
twoGOSim(gene1, gene2, type = "gene", ont = "BP", measure = "Lin")

## End(Not run)</pre>
```

twoSeqSim

Protein Sequence Alignment for Two Protein Sequences

Description

Sequence alignment between two protein sequences.

Usage

```
twoSeqSim(
  seq1,
  seq2,
  type = "local",
  submat = "BLOSUM62",
  gap.opening = 10,
  gap.extension = 4
)
```

twoSeqSim 73

Arguments

seq1	Character string, containing one protein sequence.		
seq2	Character string, containing another protein sequence.		
type	Type of alignment, default is "local", could be "global" or "local", where "global" represents Needleman-Wunsch global alignment; "local" represents Smith-Waterman local alignment.		
submat	Substitution matrix, default is "BLOSUM62", can be one of "BLOSUM45", "BLOSUM50", "BLOSUM62", "BLOSUM80", "BLOSUM100", "PAM30", "PAM40", "PAM70", "PAM120", or "PAM250".		
gap.opening	The cost required to open a gap of any length in the alignment. Defaults to 10.		
gap.extension	The cost to extend the length of an existing gap by 1. Defaults to 4.		

Value

A Biostrings object containing the alignment scores and other alignment information.

Author(s)

```
Nan Xiao <a href="https://nanx.me">https://nanx.me</a>
```

See Also

See parSeqSim for paralleled pairwise protein similarity calculation based on sequence alignment. See twoGOSim for calculating the GO semantic similarity between two groups of GO terms or two Entrez gene IDs.

```
## Not run:

# Be careful when testing this since it involves sequence alignment
# and might produce unpredictable results in some environments

library("Biostrings")

s1 <- readFASTA(system.file("protseq/P00750.fasta", package = "protr"))[[1]]
s2 <- readFASTA(system.file("protseq/P10323.fasta", package = "protr"))[[1]]
seqalign <- twoSeqSim(s1, s2)
summary(seqalign)
score(seqalign)

## End(Not run)</pre>
```

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