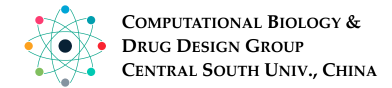
protr: R package for generating various numerical representation schemes of protein sequences

Nan Xiao, Qing-Song Xu, Dong-Sheng Cao

Package Version: 1.3-0 2017-05-07



Contents

1. The Computational Workflow	1
2. Package Overview	3
3. Commonly Used Descriptors	6
4. Descriptors for Proteochemometric Modeling	28
5. Similarity Calculation by Sequence Alignment	30
6. Similarity Calculation by GO Semantic Similarity Measures	31
7. ProtrWeb	32
8. Miscellaneous Tools	32
9. Summary	35
10. How to Cite	36

Abstract

The **protr** package offers a unique and comprehensive toolkit for generating various numerical representation schemes of protein sequence. The descriptors included are extensively utilized in Bioinformatics and Chemogenomics research. The commonly used descriptors listed in **protr** include amino acid composition, autocorrelation, CTD, conjoint traid, quasi-sequence order, pseudo amino acid composition, and profile-based descriptors derived by Position-Specific Scoring Matrix (PSSM). The descriptors for proteochemometric (PCM) modeling, includes the scales-based descriptors derived by principal components analysis, factor analysis, multidimensional scaling, amino acid properties (AAindex), 20+ classes of 2D and 3D molecular descriptors (Topological, WHIM, VHSE, etc.), and BLOSUM/PAM matrix-derived descriptors. The **protr** package also integrates the function of parallelized similarity computation derived by pairwise protein sequence alignment and Gene Ontology (GO) semantic similarity measures. **ProtrWeb**, the web server built on **protr**, can be accessed from: http://protr.org.

Keywords: protein sequence, amino acid, descriptor, structural similarity, functional similarity, sequence alignment, Gene Ontology

1. The Computational Workflow

Here we use the subcellular localization dataset of human proteins presented in Chou and Shen (2008) to demonstrate the workflow of using protr.

The complete dataset includes 3,134 protein sequences (2,750 different proteins), classified into 14 human subcellular locations. We selected two classes of proteins as our benchmark dataset. Class 1 contains 325 extracell proteins, and class 2 includes 307 mitochondrion proteins.

First, we load the **protr** package, then read the protein sequences stored in two separated FASTA files with readFASTA():

```
library("protr")

# load FASTA files
extracel1 = readFASTA(system.file(
    "protseq/extracel1.fasta", package = "protr"))
mitonchon = readFASTA(system.file(
    "protseq/mitochondrion.fasta", package = "protr"))
```

To read protein sequences stored in PDB format files, use readPDB() instead. The loaded sequences will be stored as two lists in R, and each component in the list is a character string representing one protein sequence. In this case, there are 325 extracell protein sequences and 306 mitonchon protein sequences:

```
length(extracell)
## [1] 325
```

```
length(mitonchon)
```

```
## [1] 306
```

To assure that the protein sequences only have the twenty standard amino acid types which is required for the descriptor computation, we use the protcheck() function in protr to do the amino acid type sanity checking and remove the *non-standard* sequences:

```
extracell = extracell[(sapply(extracell, protcheck))]
mitonchon = mitonchon[(sapply(mitonchon, protcheck))]
length(extracell)
## [1] 323
length(mitonchon)
## [1] 304
```

Two protein sequences were removed from each class. For the remaining sequences, we calculate the Type II PseAAC descriptor, i.e., the amphiphilic pseudo amino acid composition (APAAC) descriptor (Chou 2005) and make class labels for classification modeling.

```
# calculate APAAC descriptors
x1 = t(sapply(extracell, extractAPAAC))
x2 = t(sapply(mitonchon, extractAPAAC))
x = rbind(x1, x2)

# make class labels
labels = as.factor(c(rep(0, length(extracell)), rep(1, length(mitonchon))))
```

In **protr**, the functions of commonly used descriptors for protein sequences and proteochemometric (PCM) modeling descriptors are named after extract...().

Next, we will split the data into a 75% training set and a 25% test set.

```
set.seed(1001)
# split training and test set
tr.idx = c(
    sample(1:nrow(x1), round(nrow(x1) * 0.75)),
    sample(nrow(x1) + 1:nrow(x2), round(nrow(x2) * 0.75)))
te.idx = setdiff(1:nrow(x), tr.idx)

x.tr = x[tr.idx,]
x.te = x[te.idx,]
y.tr = labels[tr.idx]
y.te = labels[te.idx]
```

We will train a random forest classification model on the training set with 5-fold cross-validation, using the **randomForest** package.

```
library("randomForest")
rf.fit = randomForest(x.tr, y.tr, cv.fold = 5)
print(rf.fit)
The training result is:
## Call:
##
    randomForest(x = x.tr, y = y.tr, cv.fold = 5)
##
                  Type of random forest: classification
##
                         Number of trees: 500
## No. of variables tried at each split: 8
##
##
           OOB estimate of error rate: 25.11%
## Confusion matrix:
       0
           1 class.error
         46
               0.1900826
## 0 196
## 1 72 156
               0.3157895
```

With the model trained on the training set, we predict on the test set and plot the ROC curve with the **pROC** package, as is shown in figure 1.

```
# predict on test set
rf.pred = predict(rf.fit, newdata = x.te, type = "prob")[, 1]

# plot ROC curve
require(pROC)
plot.roc(y.te, rf.pred, col = "#0080ff", grid = TRUE, print.auc = TRUE)

The area under the ROC curve (AUC) is:

## Call:
## plot.roc.default(x = y.te, predictor = rf.pred, col = "#0080ff",
## grid = TRUE, print.auc = TRUE)

## Data: rf.pred in 81 controls (y.te 0) > 76 cases (y.te 1).
## Area under the curve: 0.8697
```

2. Package Overview

The **protr** package (Xiao *et al.* 2015) implemented most of the state-of-the-art protein sequence descriptors with R. The **protr** package is freely available from CRAN (https://cran.

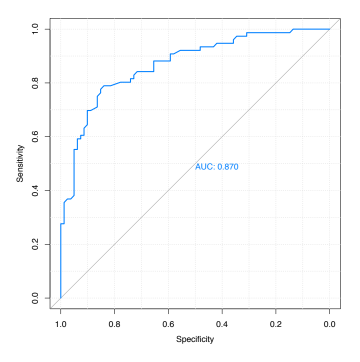


Figure 1: ROC curve for the test set of protein subcellular localization data

r-project.org/package=protr). This vignette corresponds to **protr** version 1.3-0 and was typeset on 2017-05-07.

Generally, each type of the descriptors (features) can be calculated with a function named extractX() in the protr package, where X stands for the abbrevation of the descriptor name. The descriptors and the function names implemented are listed below:

- Amino acid composition
 - extractAAC() Amino acid composition
 - extractDC() Dipeptide composition
 - extractTC() Tripeptide composition
- Autocorrelation
 - extractMoreauBroto() Normalized Moreau-Broto autocorrelation
 - extractMoran() Moran autocorrelation
 - extractGeary() Geary autocorrelation
- CTD
 - extractCTDC() Composition
 - extractCTDT() Transition
 - extractCTDD() Distribution

- Conjoint triad descriptors
 - extractCTriad() Conjoint triad descriptors
- Quasi-sequence-order descriptors
 - extractSOCN() Sequence-order-coupling number
 - extractQSO() Quasi-sequence-order descriptors
- Pseudo-amino acid composition
 - extractPAAC() Pseudo-amino acid composition
 - extractAPAAC() Amphiphilic pseudo-amino acid composition
- Profile-based descriptors
 - extractPSSM()
 - extractPSSMAcc()
 - extractPSSMFeature()

The descriptors commonly used in Proteochemometric Modeling (PCM) implemented in **protr** include:

- extractScales() and extractScalesGap() Scales-based descriptors derived by Principal Components Analysis
 - extractProtFP() and extractProtFPGap() Scales-based descriptors derived by amino acid properties from AAindex (a.k.a. Protein Fingerprint)
 - extractDescScales() Scales-based descriptors derived by 20+ classes of 2D and
 3D molecular descriptors (Topological, WHIM, VHSE, etc.)
- extractFAScales() Scales-based descriptors derived by Factor Analysis
- extractMDSScales() Scales-based descriptors derived by Multidimensional Scaling
- extractBLOSUM() BLOSUM and PAM matrix-derived descriptors

The **protr** package integrates the function of parallelized similarity score computation derived by local or global protein sequence alignment between a list of protein sequences, the sequence alignment computation is provided by **Biostrings**, the corresponding functions listed in the **protr** package include:

- twoSeqSim() Similarity calculation derived by sequence alignment between two protein sequences
- parSeqSim() Parallelized pairwise similarity calculation with a list of protein sequences

The **protr** package also integrates the function of parallelized similarity score computation derived by Gene Ontology (GO) semantic similarity measures between a list of GO terms / Entrez Gene IDs, the GO similarity computation is provided by **GOSemSim**, the corresponding functions listed in the **protr** package include:

- twoGOSim() Similarity calculation derived by GO-terms semantic similarity measures between two GO terms / Entrez Gene IDs;
- parGOSim() Pairwise similarity calculation with a list of GO terms / Entrez Gene IDs.

To use the parSeqSim() function, we suggest the users to install the packages foreach and doParallel first, in order to make the parallelized pairwise similarity computation available. In the next sections, we will introduce the descriptors and function usage in this order.

3. Commonly Used Descriptors

Note: Users of the **protr** package need to intelligently evaluate the underlying details of the descriptors provided, instead of using protr with their data blindly, especially for the descriptor types with more flexibility. It would be wise for the users to use some negative and positive control comparisons where relevant to help guide interpretation of the results.

A protein or peptide sequence with N amino acid residues can be generally represented as $\{R_1, R_2, \ldots, R_n\}$, where R_i represents the residue at the i-th position in the sequence. The labels i and j are used to index amino acid position in a sequence, and r, s, t are used to represent the amino acid type. The computed descriptors are roughly divided into 4 groups according to their known applications described in the literature.

A protein sequence can be divided equally into segments and the methods, described as follows for the global sequence, could be applied to each segment.

3.1. Amino Acid Composition (AAC)

The Amino Acid Composition (AAC) is the fraction of each amino acid type within a protein. The fractions of all 20 natural amino acids are calculated as:

$$f(r) = \frac{N_r}{N}$$
 $r = 1, 2, \dots, 20.$

where N_r is the number of the amino acid type r and N is the length of the sequence.

As was described above, we can use the function extractAAC() to extract the descriptors (features) from protein sequences:

- > library("protr")
 > x = readFASTA(system.file(
 + "protseq/P00750.fasta", package = "protr"))[[1]]
 > extractAAC(x)
- A R N D C E Q
 0.06405694 0.07117438 0.03914591 0.05160142 0.06761566 0.04804270 0.04804270
 G H I L K M F
 0.08185053 0.03024911 0.03558719 0.07651246 0.03914591 0.01245552 0.03202847
 P S T W Y V
 0.05338078 0.08896797 0.04448399 0.02313167 0.04270463 0.04982206

Here with the function readFASTA() we loaded a single protein sequence (P00750, Tissue-type plasminogen activator) from a FASTA format file. Then extracted the AAC descriptors with extractAAC(). The result returned is a named vector, whose elements are tagged with the name of each amino acid.

3.2. Dipeptide Composition (DC)

The Dipeptide Composition (DC) gives 400 descriptors, defined as:

$$f(r,s) = \frac{N_{rs}}{N-1}$$
 $r, s = 1, 2, \dots, 20.$

where N_{rs} is the number of dipeptide represented by amino acid type r and type s. Similar to extractAAC(), here we use extractDC() to compute the descriptors:

$$> dc = extractDC(x)$$

 $> head(dc, n = 30L)$

AA	RA	NA	DA	CA	EA
0.003565062	0.003565062	0.000000000	0.007130125	0.003565062	0.003565062
QA	GA	HA	IA	LA	KA
0.007130125	0.007130125	0.001782531	0.003565062	0.001782531	0.001782531
MA	FA	PA	SA	TA	WA
0.000000000	0.005347594	0.003565062	0.007130125	0.003565062	0.000000000
ΑY	VA	AR	RR	NR	DR
0.000000000	0.00000000	0.003565062	0.007130125	0.005347594	0.001782531
CR	ER	QR	GR	HR	IR
0.005347594	0.005347594	0.000000000	0.007130125	0.001782531	0.003565062

Here we only showed the first 30 elements of the result vector and omitted the rest of the result. The element names of the returned vector are self-explanatory as before.

3.3. Tripeptide Composition (TC)

The Tripeptide Composition (TC) gives 8000 descriptors, defined as:

$$f(r, s, t) = \frac{N_{rst}}{N - 2}$$
 $r, s, t = 1, 2, \dots, 20$

where N_{rst} is the number of tripeptides represented by amino acid type r, s and t. With function extractTC(), we can easily obtain the length-8000 descriptor, to save some space, here we also omitted the long outputs:

>
$$tc = extractTC(x)$$

> $head(tc, n = 36L)$

KAA	LAA	IAA	HAA	GAA	QAA
0.000000000	0.000000000	0.000000000	0.000000000	0.000000000	0.001785714
WAA	TAA	SAA	PAA	FAA	MAA
0.000000000	0.000000000	0.001785714	0.000000000	0.000000000	0.000000000
DRA	NRA	RRA	ARA	VAA	YAA
0.000000000	0.000000000	0.000000000	0.000000000	0.000000000	0.000000000
IRA	HRA	GRA	QRA	ERA	CRA
0.000000000	0.000000000	0.001785714	0.000000000	0.000000000	0.000000000
SRA	PRA	FRA	MRA	KRA	LRA
0.000000000	0.000000000	0.000000000	0.000000000	0.000000000	0.000000000

3.4. Autocorrelation Descriptors

Autocorrelation descriptors are defined based on the distribution of amino acid properties along the sequence. The amino acid properties used here are various types of amino acids index (Retrieved from AAindex Database: http://www.genome.jp/dbget/aaindex.html, see Kawashima et al. (1999), Kawashima and Kanehisa (2000), and Kawashima et al. (2008), see Figure 2 for an illustrated example). Three types of autocorrelation descriptors are defined here and described below.

All the amino acid indices are centralized and standardized before the calculation, i.e.

$$P_r = \frac{P_r - \bar{P}}{\sigma}$$

where \bar{P} is the average of the property of the 20 amino acids:

$$\bar{P} = \frac{\sum_{r=1}^{20} P_r}{20}$$
 and $\sigma = \sqrt{\frac{1}{2} \sum_{r=1}^{20} (P_r - \bar{P})^2}$

```
Database: AAindex
Entry: ANDN920101
LinkDB: ANDN920101
H ANDN920101
D alpha-CH chemical shifts (Andersen et al., 1992)
R LIT:1810048b PMID:1575719
A Andersen, N.H., Cao, B. and Chen, C.
T Peptide/protein structure analysis using the chemical shift index method:
  upfield alpha-CH values reveal dynamic helices and aL sites
J Biochem. and Biophys. Res. Comm. 184, 1008-1014 (1992)
C BUNA790102
                 0.949
     A/L
                                                                                       I/V
    4.35
                                4.76
             4.38
                                                                             4.63
                                                                                      3.95
             4.36
```

Figure 2: An illustrated example in the AAIndex database

Normalized Moreau-Broto Autocorrelation Descriptors

DBGET integrated database retrieval system

Moreau-Broto autocorrelation descriptors application to protein sequences can be defined as:

$$AC(d) = \sum_{i=1}^{N-d} P_i P_{i+d}$$
 $d = 1, 2, ..., \text{nlag}$

where d is called the lag of the autocorrelation and P_i and P_{i+d} are the properties of the amino acids at position i and i+d, respectively. nlag is the maximum value of the lag.

The normalized Moreau-Broto autocorrelation descriptors are defined as:

$$ATS(d) = \frac{AC(d)}{N-d}$$
 $d = 1, 2, \dots, \text{nlag}$

The corresponding function for this descriptor is extractMoreauBroto(). A typical call would be:

- > moreau = extractMoreauBroto(x)
- > head(moreau, n = 36L)

```
CIDH920105.lag1
                  CIDH920105.lag2
                                    CIDH920105.lag3
                                                      CIDH920105.lag4
     0.081573213
                     -0.016064817
                                       -0.015982990
                                                         -0.025739038
CIDH920105.lag5
                  CIDH920105.lag6
                                    CIDH920105.lag7
                                                     CIDH920105.lag8
     0.079058632
                     -0.042771564
                                       -0.036320847
                                                          0.024087298
CIDH920105.lag9 CIDH920105.lag10 CIDH920105.lag11 CIDH920105.lag12
    -0.005273958
                      0.052274763
                                        0.082170073
                                                          0.005419919
CIDH920105.lag13 CIDH920105.lag14 CIDH920105.lag15 CIDH920105.lag16
     0.083292042
                      0.004810584
                                        0.001872446
                                                         -0.001531495
CIDH920105.lag17 CIDH920105.lag18 CIDH920105.lag19
                                                    CIDH920105.lag20
    -0.011917230
                                                          0.026882737
                      0.071161551
                                        0.033473197
CIDH920105.lag21 CIDH920105.lag22 CIDH920105.lag23 CIDH920105.lag24
     0.073075402
                      0.115272790
                                        0.041517897
                                                         -0.027025993
CIDH920105.lag25 CIDH920105.lag26 CIDH920105.lag27 CIDH920105.lag28
     0.033477388
                     -0.003245255
                                        0.078117010
                                                         -0.028177304
CIDH920105.lag29 CIDH920105.lag30
                                    BHAR880101.lag1
                                                     BHAR880101.lag2
     0.046695832
                      0.020584423
                                        0.052740185
                                                          0.030804784
BHAR880101.lag3
                                                     BHAR880101.lag6
                  BHAR880101.lag4
                                    BHAR880101.lag5
     0.037170476
                     -0.058993771
                                        0.070641780
                                                         -0.089192490
```

The 8 default properties used here are:

- AccNo. CIDH920105 Normalized Average Hydrophobicity Scales
- AccNo. BHAR880101 Average Flexibility Indices
- AccNo. CHAM820101 Polarizability Parameter
- AccNo. CHAM820102 Free Energy of Solution in Water, kcal/mole
- AccNo. CHOC760101 Residue Accessible Surface Area in Tripeptide

- AccNo. BIGC670101 Residue Volume
- AccNo. CHAM810101 Steric Parameter
- AccNo. DAYM780201 Relative Mutability

Users can change the property names of AAindex database with the argument props. The AAindex data shipped with **protr** can be loaded by data(AAindex), which has the detailed information of each property. With the argument customprops and nlag, users can specify their own properties and lag value to calculate with. For illustration, we could use:

```
> # Define 3 custom properties
> 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),
                  -1, 47), E = c(-0.74,
                                            3, 73),
   C = c(0.29,
   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,
   M = c(0.64, -1.3, 75), F = c(1.19, -2.5, 91),
                    0, 42), S = c(-0.18, 0.3, 31),
   P = c(0.12,
   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
> moreau2 = extractMoreauBroto(
   x, customprops = myprops,
   props = c(
      "CIDH920105", "BHAR880101",
      "CHAM820101", "CHAM820102",
+
      "MyProp1", "MyProp2", "MyProp3"))
> head(moreau2, n = 36L)
CIDH920105.lag1 CIDH920105.lag2 CIDH920105.lag3
                                                    CIDH920105.lag4
     0.081573213
                     -0.016064817
                                      -0.015982990
                                                       -0.025739038
CIDH920105.lag5 CIDH920105.lag6 CIDH920105.lag7
                                                    CIDH920105.lag8
     0.079058632
                     -0.042771564
                                      -0.036320847
                                                        0.024087298
CIDH920105.lag9 CIDH920105.lag10 CIDH920105.lag11 CIDH920105.lag12
                      0.052274763
                                       0.082170073
    -0.005273958
                                                        0.005419919
CIDH920105.lag13 CIDH920105.lag14 CIDH920105.lag15 CIDH920105.lag16
     0.083292042
                      0.004810584
                                       0.001872446
                                                       -0.001531495
CIDH920105.lag17 CIDH920105.lag18 CIDH920105.lag19 CIDH920105.lag20
    -0.011917230
                     0.071161551
                                       0.033473197
                                                        0.026882737
CIDH920105.lag21 CIDH920105.lag22 CIDH920105.lag23 CIDH920105.lag24
                      0.115272790
                                       0.041517897
                                                       -0.027025993
     0.073075402
```

CIDH920105.lag25 CIDH920105.lag26 CIDH920105.lag27 CIDH920105.lag28

-0.028177304	0.078117010	-0.003245255	0.033477388
BHAR880101.lag2	BHAR880101.lag1	CIDH920105.lag30	CIDH920105.lag29
0.030804784	0.052740185	0.020584423	0.046695832
BHAR880101.lag6	BHAR880101.lag5	BHAR880101.lag4	BHAR880101.lag3
-0.089192490	0.070641780	-0.058993771	0.037170476

About the standard input format of props and customprops, see ?extractMoreauBroto for details.

Moran Autocorrelation Descriptors

Moran autocorrelation descriptors application to protein sequence may be defined as:

$$I(d) = \frac{\frac{1}{N-d} \sum_{i=1}^{N-d} (P_i - \bar{P}')(P_{i+d} - \bar{P}')}{\frac{1}{N} \sum_{i=1}^{N} (P_i - \bar{P}')^2} \quad d = 1, 2, \dots, 30$$

where d and P_i and P_{i+d} are defined in the same way as in the first place, and \bar{P}' is the considered property P along the sequence, i.e.,

$$\bar{P}' = \frac{\sum_{i=1}^{N} P_i}{N}$$

 d, P, P_i and P_{i+d} , nlag have the same meaning as above.

> # Use the 3 custom properties defined before

With extractMoran(), which has exactly the same arguments with extractMoreauBroto(), we can compute the Moran autocorrelation descriptors (only output the first 36 elements of the result):

```
> # and 4 properties in the AAindex database
> moran = extractMoran(
    x, customprops = myprops,
    props = c(
      "CIDH920105", "BHAR880101",
      "CHAM820101", "CHAM820102",
      "MyProp1", "MyProp2", "MyProp3"))
> head(moran, n = 36L)
CIDH920105.lag1
                  CIDH920105.lag2
                                   CIDH920105.lag3
                                                     CIDH920105.lag4
     0.062895724
                     -0.044827681
                                      -0.045065117
                                                        -0.055955678
CIDH920105.lag5
                 CIDH920105.lag6 CIDH920105.lag7
                                                     CIDH920105.lag8
     0.060586377
                     -0.074128412
                                      -0.067308852
                                                        -0.001293384
CIDH920105.lag9 CIDH920105.lag10 CIDH920105.lag11 CIDH920105.lag12
    -0.033747588
                      0.029392193
                                       0.061789800
                                                        -0.023368437
CIDH920105.lag13 CIDH920105.lag14 CIDH920105.lag15 CIDH920105.lag16
                     -0.024912264
                                      -0.028298043
     0.062769417
                                                        -0.031584063
CIDH920105.lag17 CIDH920105.lag18 CIDH920105.lag19 CIDH920105.lag20
```

```
-0.043466730
                     0.047830694
                                      0.005883901
                                                      -0.001769769
CIDH920105.lag21 CIDH920105.lag22 CIDH920105.lag23 CIDH920105.lag24
                                      0.015147594
    0.049334048
                     0.096427969
                                                      -0.060092509
CIDH920105.lag25 CIDH920105.lag26 CIDH920105.lag27 CIDH920105.lag28
                    -0.033987885
    0.007549152
                                      0.056307675
                                                      -0.061844453
CIDH920105.lag29 CIDH920105.lag30 BHAR880101.lag1 BHAR880101.lag2
    0.021484780
                    -0.008461776
                                      0.014229951
                                                      -0.009142419
BHAR880101.lag3 BHAR880101.lag4 BHAR880101.lag5 BHAR880101.lag6
    -0.003272262
                    -0.109613332
                                       0.033346233
                                                      -0.141538598
```

Geary Autocorrelation Descriptors

Geary autocorrelation descriptors for protein sequence can be defined as:

$$C(d) = \frac{\frac{1}{2(N-d)} \sum_{i=1}^{N-d} (P_i - P_{i+d})^2}{\frac{1}{N-1} \sum_{i=1}^{N} (P_i - \bar{P}')^2} \quad d = 1, 2, \dots, 30$$

where d, P, P_i and P_{i+d} , nlag have the same meaning as above.

> # Use the 3 custom properties defined before

For each amino acid index, there will be $3 \times \text{nlag}$ autocorrelation descriptors. The usage of extractGeary() is exactly the same with extractMoreauBroto() and extractMoran():

```
> # and 4 properties in the AAindex database
> geary = extractGeary(
    x, customprops = myprops,
    props = c(
      "CIDH920105", "BHAR880101",
      "CHAM820101", "CHAM820102",
      "MyProp1", "MyProp2", "MyProp3"))
> head(geary, n = 36L)
CIDH920105.lag1 CIDH920105.lag2 CIDH920105.lag3 CIDH920105.lag4
       0.9361830
                        1.0442920
                                         1.0452843
                                                           1.0563467
CIDH920105.lag5 CIDH920105.lag6 CIDH920105.lag7 CIDH920105.lag8
       0.9406031
                        1.0765517
                                         1.0675786
                                                          0.9991363
CIDH920105.lag9 CIDH920105.lag10 CIDH920105.lag11 CIDH920105.lag12
       1.0316555
                        0.9684585
                                         0.9353130
                                                           1.0201990
CIDH920105.lag13 CIDH920105.lag14 CIDH920105.lag15 CIDH920105.lag16
       0.9340933
                        1.0207373
                                         1.0251486
                                                           1.0290464
CIDH920105.lag17 CIDH920105.lag18 CIDH920105.lag19 CIDH920105.lag20
       1.0414375
                        0.9494403
                                         0.9905987
                                                          0.9987183
CIDH920105.lag21 CIDH920105.lag22 CIDH920105.lag23 CIDH920105.lag24
       0.9472542
                        0.9010009
                                         0.9828848
                                                           1.0574098
CIDH920105.lag25 CIDH920105.lag26 CIDH920105.lag27 CIDH920105.lag28
                        1.0290018
       0.9897955
                                         0.9400066
                                                          1.0584150
```

BHAR880101.lag2	BHAR880101.lag1	CIDH920105.lag30	CIDH920105.lag29
1.0051730	0.9818711	1.0029734	0.9762904
BHAR880101.lag6	BHAR880101.lag5	BHAR880101.lag4	BHAR880101.lag3
1.1337056	0.9595859	1.1012905	0.9967069

3.5. Composition / Transition / Distribution

These descriptors are developed and described by Dubchak *et al.* (1995) and Dubchak *et al.* (1999).

Sequence	M	\mathbf{T}	\mathbf{E}	Ι	\mathbf{T}	\mathbf{A}	\mathbf{S}	\mathbf{M}	\mathbf{V}	K	\mathbf{E}	\mathbf{L}	\mathbf{R}	\mathbf{E}	A	${f T}$	\mathbf{G}	${f T}$	\mathbf{G}	A
Sequence Index	1				5					10					15					20
Transformation	3	2	1	3	2	2	2	3	3	1	1	3	1	1	2	2	2	2	2	2
Index for 1			1							2	3		4	5						
Index for 2		1			2	3	4								5	6	7	8	9	10
Index for 3	1			2				3	4			5								
1/2 Transitions																				
1/3 Transitions																				
2/3 Transitions																				

Figure 3: The sequence of a hypothetic protein indicating the construction of composition, transition and distribution descriptors of a protein. Sequence index indicates the position of an amino acid in the sequence. The index for each type of amino acids in the sequence ('1', '2' or '3') indicates the position of the first, second, third, ... of that type of amino acid. 1/2 transition indicates the position of '12' or '21' pairs in the sequence (1/3 and 2/3 are defined) in the same way.).

Step 1: Sequence Encoding

The amino acids are divided in three classes according to its attribute and each amino acid is encoded by one of the indices 1, 2, 3 according to which class it belonged. The attributes used here include hydrophobicity, normalized van der Waals volume polarity, and polarizability, as in the references. The corresponding division is in Table 1.

For example, for a given sequence "MTEITAAMVKELRESTGAGA", it will be encoded as "32132223311311222222" according to its hydrophobicity division.

Step 2: Compute Composition, Transition and Distribution Descriptors

Three descriptors, Composition (C), Transition (T), and Distribution (D) were calculated for a given attribute as follows.

Composition

It is the global percent for each encoded class in the sequence. In the above example using hydrophobicity division, the numbers for encoded classes "1", "2", "3" are 5, 10, 5 respectively, so that the compositions for them are 5/20 = 25%, 10/20 = 10%, and 5/20 = 25% respectively, where 20 is the length of the protein sequence. Composition can be defined as

Table 1: Amino acid attributes and the division of the amino acids into three groups for each attribute

	Group 1	Group 2	Group 3
Hydrophobicity	Polar	Neutral	Hydrophobicity
	R, K, E, D, Q, N	G, A, S, T, P, H, Y	C, L, V, I, M, F, W
Normalized van der	0-2.78	2.95-4.0	4.03-8.08
Waals Volume	G, A, S, T, P, D, C	N, V, E, Q, I, L	M, H, K, F, R, Y, W
Polarity	4.9-6.2	8.0-9.2	10.4-13.0
	L, I, F, W, C, M, V, Y	P, A, T, G, S	H, Q, R, K, N, E, D
Polarizability	0-1.08	0.128-0.186	0.219-0.409
	G, A, S, D, T	C, P, N, V, E, Q, I, L	K, M, H, F, R, Y, W
Charge	Positive	Neutral	Negative
	K, R	A, N, C, Q, G, H, I, L, M, F, P, S, T, W, Y, V	D, E
Secondary	Helix	Strand	Coil
Structure	E, A, L, M, Q, K, R, H	V, I, Y, C, W, F, T	G, N, P, S, D
Solvent	Buried	Exposed	Intermediate
Accessibility	A, L, F, C, G, I, V, W	R, K, Q, E, N, D	M, S, P, T, H, Y

$$C_r = \frac{n_r}{n} \quad r = 1, 2, 3$$

where n_r is the number of amino acid type r in the encoded sequence and N is the length of the sequence. An example for extractCTDC():

> extractCTDC(x)

hydrophobicity.Group1	hydrophobicity.Group2	hydrophobicity.Group3
0.29715302	0.40569395	0.29715302
${\tt normwaalsvolume.Group1}$	${\tt normwaalsvolume.Group2}$	${\tt normwaalsvolume.Group3}$
0.45195730	0.29715302	0.25088968
polarity.Group1	polarity.Group2	polarity.Group3
0.33985765	0.33274021	0.32740214
polarizability.Group1	polarizability.Group2	polarizability.Group3
0.33096085	0.41814947	0.25088968
charge.Group1	charge.Group2	charge.Group3
0.11032028	0.79003559	0.09964413
${\tt secondarystruct.Group1}$	${\tt secondary struct.Group 2}$	${\tt secondary struct.Group 3}$
0.38967972	0.29537367	0.31494662
${\tt solventaccess.Group1}$	solventaccess.Group2	solventaccess.Group3
0.43060498	0.29715302	0.27224199

The result shows the elements whose names are PropertyNumber. GroupNumber in the returned vector.

Transition

A transition from class 1 to 2 is the percent frequency with which 1 is followed by 2 or 2 is

followed by 1 in the encoded sequence. Transition descriptor can be calculated as

$$T_{rs} = \frac{n_{rs} + n_{sr}}{N - 1}$$
 $rs = `12', `13', `23'$

where n_{rs} , n_{sr} is the numbers of dipeptide encoded as "rs" and "sr" respectively in the sequence and N is the length of the sequence. An example for extractCTDT():

> extractCTDT(x)

```
prop1.Tr1221 prop1.Tr1331 prop1.Tr2332 prop2.Tr1221 prop2.Tr1331 prop2.Tr2332
               0.16042781
  0.27094474
                            0.23351159
                                          0.26737968
                                                       0.22638146
                                                                    0.17112299
prop3.Tr1221 prop3.Tr1331 prop3.Tr2332 prop4.Tr1221 prop4.Tr1331 prop4.Tr2332
  0.21033868
               0.20499109
                            0.23707665
                                          0.27272727
                                                       0.15151515
                                                                    0.24598930
prop5.Tr1221 prop5.Tr1331 prop5.Tr2332 prop6.Tr1221 prop6.Tr1331 prop6.Tr2332
                                                       0.22816399
  0.18181818
               0.02139037
                            0.15686275
                                          0.21925134
                                                                    0.15864528
prop7.Tr1221 prop7.Tr1331 prop7.Tr2332
               0.21568627
  0.25133690
                            0.18003565
```

Distribution

The "distribution" descriptor describes the distribution of each attribute in the sequence.

There are five "distribution" descriptors for each attribute and they are the position percents in the whole sequence for the first residue, 25% residues, 50% residues, 75% residues and 100% residues, respectively, for a specified encoded class. For example, there are 10 residues encoded as "2" in the above example, the positions for the first residue "2", the 2nd residue "2" (25%*10=2), the 5th "2" residue (50%*10=5), the 7th "2" (75%*10=7) and the 10th residue "2" (100%*10) in the encoded sequence are 2, 5, 15, 17, 20, respectively, so the distribution descriptors for "2" are: $10.0 \ (2/20*100)$, $25.0 \ (5/20*100)$, $75.0 \ (15/20*100)$, $85.0 \ (17/20*100)$, $100.0 \ (20/20*100)$, respectively.

Finally, an example for extractCTDD():

> extractCTDD(x)

```
prop1.G1.residue0
                     prop1.G1.residue25
                                          prop1.G1.residue50
                                                               prop1.G1.residue75
          0.3558719
                               0.3558719
                                                    0.3558719
                                                                        0.3558719
                                          prop1.G2.residue25
prop1.G1.residue100
                      prop1.G2.residue0
                                                               prop1.G2.residue50
          0.3558719
                               0.5338078
                                                    0.5338078
                                                                        0.5338078
 prop1.G2.residue75 prop1.G2.residue100
                                           prop1.G3.residue0
                                                               prop1.G3.residue25
          0.5338078
                               0.5338078
                                                    0.1779359
                                                                        0.1779359
 prop1.G3.residue50
                     prop1.G3.residue75 prop1.G3.residue100
                                                                prop2.G1.residue0
          0.1779359
                               0.1779359
                                                    0.1779359
                                                                        0.3558719
                                          prop2.G1.residue75 prop2.G1.residue100
 prop2.G1.residue25
                     prop2.G1.residue50
          0.3558719
                               0.3558719
                                                    0.3558719
                                                                        0.3558719
                                                              prop2.G2.residue75
  prop2.G2.residue0
                     prop2.G2.residue25
                                          prop2.G2.residue50
```

1.4234875	1.4234875	1.4234875	1.4234875
prop2.G2.residue100	prop2.G3.residue0	prop2.G3.residue25	prop2.G3.residue50
1.4234875	0.1779359	0.1779359	0.1779359
	prop2.G3.residue100	prop3.G1.residue0	prop3.G1.residue25
0.1779359	0.1779359	0.1779359	0.1779359
		prop3.G1.residue100	
prop3.G1.residue50			prop3.G2.residue0
0.1779359	0.1779359	0.1779359	0.5338078
prop3.G2.residue25	prop3.G2.residue50		prop3.G2.residue100
0.5338078	0.5338078	0.5338078	0.5338078
prop3.G3.residue0	prop3.G3.residue25	prop3.G3.residue50	prop3.G3.residue75
0.3558719	0.3558719	0.3558719	0.3558719
<pre>prop3.G3.residue100</pre>	prop4.G1.residue0	prop4.G1.residue25	prop4.G1.residue50
0.3558719	0.3558719	0.3558719	0.3558719
prop4.G1.residue75	${\tt prop4.G1.residue100}$	prop4.G2.residue0	prop4.G2.residue25
0.3558719	0.3558719	1.4234875	1.4234875
prop4.G2.residue50	prop4.G2.residue75	<pre>prop4.G2.residue100</pre>	<pre>prop4.G3.residue0</pre>
1.4234875	1.4234875	1.4234875	0.1779359
prop4.G3.residue25	prop4.G3.residue50	prop4.G3.residue75	prop4.G3.residue100
0.1779359	0.1779359	0.1779359	0.1779359
prop5.G1.residue0	prop5.G1.residue25	prop5.G1.residue50	prop5.G1.residue75
0.8896797	0.8896797	0.8896797	0.8896797
prop5.G1.residue100	prop5.G2.residue0	prop5.G2.residue25	prop5.G2.residue50
0.8896797	0.1779359	0.1779359	0.1779359
prop5.G2.residue75	prop5.G2.residue100	prop5.G3.residue0	prop5.G3.residue25
0.1779359	0.1779359	0.3558719	0.3558719
prop5.G3.residue50		prop5.G3.residue100	prop6.G1.residue0
0.3558719	0.3558719	0.3558719	0.1779359
prop6.G1.residue25	prop6.G1.residue50		prop6.G1.residue100
0.1779359	0.1779359	0.1779359	0.1779359
prop6.G2.residue0	prop6.G2.residue25	prop6.G2.residue50	prop6.G2.residue75
1.6014235	1.6014235	1.6014235	1.6014235
prop6.G2.residue100	prop6.G3.residue0	prop6.G3.residue25	prop6.G3.residue50
1.6014235	0.3558719	0.3558719	0.3558719
	prop6.G3.residue100	prop7.G1.residue0	prop7.G1.residue25
0.3558719	0.3558719	0.5338078	0.5338078
prop7.G1.residue50		prop7.G1.residue100	prop7.G2.residue0
0.5338078	0.5338078	0.5338078	0.3558719
prop7.G2.residue25	prop7.G2.residue50		prop7.G2.residue100
0.3558719	0.3558719	0.3558719	0.3558719
prop7.G3.residue0	prop7.G3.residue25	prop7.G3.residue50 0.1779359	prop7.G3.residue75
0.1779359	0.1779359	0.1779359	0.1779359
prop7.G3.residue100			
0.1779359			

3.6. Conjoint Triad Descriptors

Conjoint triad descriptors are proposed by Shen *et al.* (2007). These conjoint triad descriptors abstracts the features of protein pairs based on the classification of amino acids. In this approach, each protein sequence is represented by a vector space consisting of descriptors of amino acids. To reduce the dimensions of vector space, the 20 amino acids were clustered into several classes according to their dipoles and volumes of the side chains. The conjoint triad descriptors are calculated as follows:

Step 1: Classification of Amino Acids

Electrostatic and hydrophobic interactions dominate protein-protein interactions. These two kinds of interactions may be reflected by the dipoles and volumes of the side chains of amino acids, respectively. Accordingly, these two parameters were calculated, respectively, by using the density-functional theory method B3LYP/6-31G and molecular modeling approach. Based on the dipoles and volumes of the side chains, the 20 amino acids can be clustered into seven classes (See Table 2). Amino acids within the same class likely involve synonymous mutations because of their similar characteristics.

No.	Dipole Scale ¹	Volume Scale ²	Class
1	_	_	Ala, Gly, Val
2	_	+	Ile, Leu, Phe, Pro
3	+	+	Tyr, Met, Thr, Ser
4	++	+	His, Asn, Gln, Tpr
5	+++	+	Arg, Lys
6	+' +' +'	+	Asp, Glu
7	$+^{3}$	+	Cys

Table 2: Classification of amino acids based on dipoles and volumes of the side chains

Step 2: Conjoint Triad Calculation

The conjoint triad descriptors considered the properties of one amino acid and its vicinal amino acids and regarded any three continuous amino acids as a unit. Thus, the triads can be differentiated according to the classes of amino acids, i.e., triads composed by three amino acids belonging to the same classes, such as ART and VKS, can be treated identically. To conveniently represent a protein, we first use a binary space (\mathbf{V}, \mathbf{F}) to represent a protein sequence. Here, \mathbf{V} is the vector space of the sequence features, and each feature v_i represents a sort of triad type; \mathbf{F} is the frequency vector corresponding to \mathbf{V} , and the value of the *i*-th dimension of $\mathbf{F}(f_i)$ is the frequency of type v_i appearing in the protein sequence. For the amino acids that have been catogorized into seven classes, the size of \mathbf{V} should be $7 \times 7 \times 7$; thus $i = 1, 2, \ldots, 343$. The detailed description for (\mathbf{V}, \mathbf{F}) is illustrated in Figure 4.

Clearly, each protein correlates to the length (number of amino acids) of protein. In general, a long protein would have a large value of f_i , which complicates the comparison between two heterogeneous proteins. Thus, we defined a new parameter, d_i , by normalizing f_i with the following equation:

 $^{^1}$ Dipole Scale (Debye): –, Dipole < 1.0; +, 1.0 < Dipole < 2.0; ++, 2.0 < Dipole < 3.0; ++ +, Dipole > 3.0; +' +' +', Dipole > 3.0 with opposite orientation.

²Volume Scale (Å³): -, Volume < 50; +, Volume > 50.

³Cys is separated from class 3 because of its ability to form disulfide bonds.

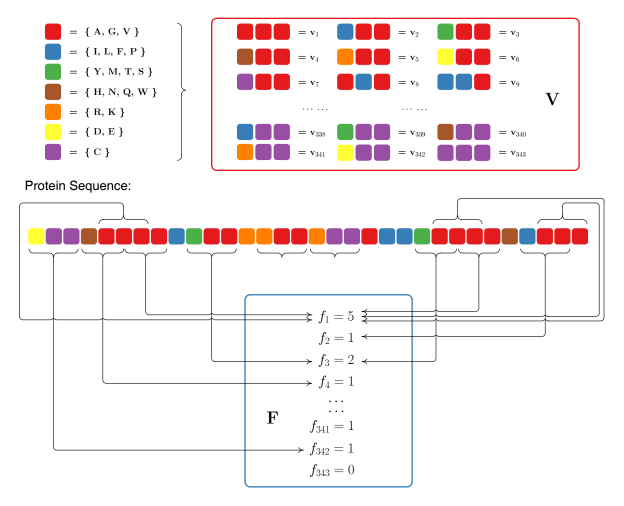


Figure 4: Schematic diagram for constructing the vector space (\mathbf{V}, \mathbf{F}) of protein sequence. \mathbf{V} is the vector space of the sequence features; each feature (v_i) represents a triad composed of three consecutive amino acids; \mathbf{F} is the frequency vector corresponding to \mathbf{V} , and the value of the *i*-th dimension of $\mathbf{F}(f_i)$ is the frequency that v_i triad appeared in the protein sequence.

$$d_i = \frac{f_i - \min\{f_1, f_2, \dots, f_{343}\}}{\max\{f_1, f_2, \dots, f_{343}\}}$$

The numerical value of d_i of each protein ranges from 0 to 1, which thereby enables the comparison between proteins. Accordingly, we obtain another vector space (designated **D**) consisting of d_i to represent protein.

To compute conjoint triads of protein sequences, we can simply use:

```
> ctriad = extractCTriad(x)
> head(ctriad, n = 65L)
```

VS111 VS211 VS311 VS411 VS511 VS611 VS711 VS121 VS221 VS321 VS421 VS521 VS621 0.6 0.0 0.1 0.3 0.6 0.2 0.4 0.0 0.3 1.0 0.5 0.2 VS721 VS131 VS231 VS331 VS431 VS531 VS631 VS731 VS141 VS241 VS341 VS441 VS541 0.0 0.2 0.4 0.5 0.2 0.3 0.3 0.1 0.3 0.3 0.2 0.2 0.0 VS641 VS741 VS151 VS251 VS351 VS451 VS551 VS651 VS751 VS161 VS261 VS361 VS461 0.1 0.2 0.2 0.2 0.5 0.1 0.2 0.0 0.0 0.1 0.4 0.2 0.3 VS561 VS661 VS761 VS171 VS271 VS371 VS471 VS571 VS671 VS771 VS112 VS212 VS312 0.2 0.0 0.1 0.1 0.3 0.1 0.0 0.1 0.0 0.1 0.8 0.4 0.4 VS412 VS512 VS612 VS712 VS122 VS222 VS322 VS422 VS522 VS622 VS722 VS132 VS232 0.6 0.1 0.5 0.2 0.8 0.5 0.2 0.3 0.2 0.0 0.2 0.1 0.3

by which we only outputted the first 65 of total 343 dimension to save space.

3.7. Quasi-sequence-order Descriptors

The quasi-sequence-order descriptors are proposed by Chou (2000). They are derived from the distance matrix between the 20 amino acids.

Sequence-order-coupling Number

The d-th rank sequence-order-coupling number is defined as:

$$\tau_d = \sum_{i=1}^{N-d} (d_{i,i+d})^2 \quad d = 1, 2, \dots, \text{maxlag}$$

where $d_{i,i+d}$ is the distance between the two amino acids at position i and i+d.

Note: maxlag is the maximum lag and the length of the protein must be not less than maxlag. The function extractSOCN(x) is used for computing the sequence-order-coupling numbers:

> extractSOCN(x)

Schneider.lag1 Schneider.lag2 Schneider.lag3 Schneider.lag4 Schneider.lag5 204.2036 199.8708 206.8102 197.4828 193.3366 Schneider.lag6 Schneider.lag7 Schneider.lag8 Schneider.lag9 Schneider.lag10

208.1936	195.5476	200.9789	196.7110	193.9931
Schneider.lag11	Schneider.lag12	Schneider.lag13	Schneider.lag14	Schneider.lag15
199.7031	204.9389	187.0140	198.4702	205.4526
Schneider.lag16	Schneider.lag17	${\tt Schneider.lag18}$	Schneider.lag19	Schneider.lag20
193.1274	187.3529	190.4949	202.8853	198.5299
Schneider.lag21	${\tt Schneider.lag22}$	${\tt Schneider.lag23}$	${\tt Schneider.lag24}$	Schneider.lag25
191.1013	185.0074	189.9857	202.7113	201.6267
Schneider.lag26	${\tt Schneider.lag27}$	${\tt Schneider.lag28}$	Schneider.lag29	Schneider.lag30
194.5770	185.9939	204.1297	191.1629	183.9073
${\tt Grantham.lag1}$	${\tt Grantham.lag2}$	${\tt Grantham.lag3}$	${\tt Grantham.lag4}$	Grantham.lag5
6674686.0000	6761609.0000	7138892.0000	6748261.0000	6291229.0000
${\tt Grantham.lag6}$	${\tt Grantham.lag7}$	${\tt Grantham.lag8}$	${\tt Grantham.lag9}$	Grantham.lag10
6839853.0000	6594164.0000	6556148.0000	6620183.0000	6770614.0000
Grantham.lag11	Grantham.lag12	Grantham.lag13	Grantham.lag14	Grantham.lag15
6495689.0000	6865537.0000	6297267.0000	6498247.0000	6615566.0000
Grantham.lag16	Grantham.lag17	Grantham.lag18	Grantham.lag19	Grantham.lag20
6572680.0000	6569081.0000	6173947.0000	6570829.0000	6471308.0000
Grantham.lag21	Grantham.lag22	Grantham.lag23	Grantham.lag24	Grantham.lag25
6461649.0000	5939432.0000	6532121.0000	6652472.0000	6480660.0000
Grantham.lag26	Grantham.lag27	Grantham.lag28	Grantham.lag29	Grantham.lag30
6382281.0000	6276521.0000	6537634.0000	6442991.0000	6350157.0000

Users can also specify the maximum lag value with the nlag argument.

Note: In addition to Schneider-Wrede physicochemical distance matrix (Schneider and Wrede 1994) used by Kuo-Chen Chou, another chemical distance matrix by Grantham (1974) is also used here. So the descriptors dimension will be nlag * 2. The quasi-sequence-order descriptors described next also utilized the two matrices.

Quasi-sequence-order Descriptors

For each amino acid type, a quasi-sequence-order descriptor can be defined as:

$$X_r = \frac{f_r}{\sum_{r=1}^{20} f_r + w \sum_{d=1}^{\text{maxlag}} \tau_d}$$
 $r = 1, 2, \dots, 20$

where f_r is the normalized occurrence for amino acid type i and w is a weighting factor (w = 0.1). These are the first 20 quasi-sequence-order descriptors. The other 30 quasi-sequence-order are defined as:

$$X_d = \frac{w\tau_{d-20}}{\sum_{r=1}^{20} f_r + w \sum_{d=1}^{\text{maxlag}} \tau_d}$$
 $d = 21, 22, \dots, 20 + \text{maxlag}$

An minimal example for extractQSO():

> extractQSO(x)

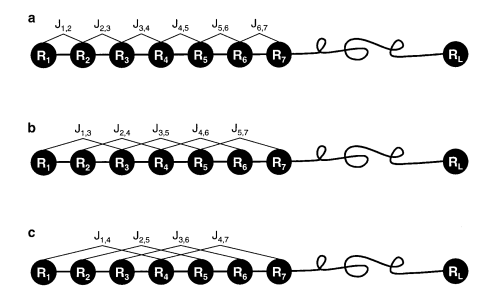


Figure 5: A schematic drawing to show (a) the 1st-rank, (b) the 2nd-rank, and (3) the 3rd-rank sequence-order-coupling mode along a protein sequence. (a) Reflects the coupling mode between all the most contiguous residues, (b) that between all the 2nd most contiguous residues, and (c) that between all the 3rd most contiguous residues. This figure is from Chou (2000).

Xr.C Ze-02 Xr.I
Xr.I
8e-02
Xr.P
2e-02
Xr.V
3e-02
Xr.C
e-06
Xr.I
Be-06
Xr.P
le-06
Xr.V
8e-06
Xd.5
le-02
d.10
8e-02
(d.15
le-02
d.20
8e-02

Schneider.Xd.21	Schneider.Xd.22	Schneider.Xd.23	Schneider.Xd.24	Schneider.Xd.25
3.236099e-02	3.132904e-02	3.217206e-02	3.432701e-02	3.414334e-02
Schneider.Xd.26	Schneider.Xd.27	Schneider.Xd.28	Schneider.Xd.29	Schneider.Xd.30
3.294954e-02	3.149609e-02	3.456720e-02	3.237140e-02	3.114275e-02
Grantham.Xd.1	Grantham.Xd.2	Grantham.Xd.3	Grantham.Xd.4	Grantham.Xd.5
3.402298e-02	3.446605e-02	3.638918e-02	3.439801e-02	3.206838e-02
Grantham.Xd.6	Grantham.Xd.7	Grantham.Xd.8	Grantham.Xd.9	Grantham.Xd.10
3.486488e-02	3.361253e-02	3.341875e-02	3.374516e-02	3.451195e-02
Grantham.Xd.11	Grantham.Xd.12	Grantham.Xd.13	Grantham.Xd.14	Grantham.Xd.15
3.311057e-02	3.499580e-02	3.209915e-02	3.312361e-02	3.372162e-02
Grantham.Xd.16	Grantham.Xd.17	Grantham.Xd.18	Grantham.Xd.19	Grantham.Xd.20
3.350302e-02	3.348467e-02	3.147055e-02	3.349358e-02	3.298629e-02
Grantham.Xd.21	Grantham.Xd.22	Grantham.Xd.23	Grantham.Xd.24	Grantham.Xd.25
3.293706e-02	3.027516e-02	3.329628e-02	3.390974e-02	3.303396e-02
Grantham.Xd.26	Grantham.Xd.27	Grantham.Xd.28	Grantham.Xd.29	Grantham.Xd.30
3.253250e-02	3.199340e-02	3.332438e-02	3.284195e-02	3.236875e-02

where users can also specify the maximum lag with argument nlag and the weighting factor with argument w.

3.8. Pseudo-Amino Acid Composition (PAAC)

This groups of descriptors are proposed in Chou (2001). PAAC descriptors are also called the type 1 pseudo-amino acid composition. Let $H_1^o(i)$, $H_2^o(i)$, $M^o(i)$ (i = 1, 2, 3, ..., 20) be the original hydrophobicity values, the original hydrophilicity values and the original side chain masses of the 20 natural amino acids, respectively. They are converted to following qualities by a standard conversion:

$$H_1(i) = \frac{H_1^o(i) - \frac{1}{20} \sum_{i=1}^{20} H_1^o(i)}{\sqrt{\frac{\sum_{i=1}^{20} [H_1^o(i) - \frac{1}{20} \sum_{i=1}^{20} H_1^o(i)]^2}{20}}}$$

 $H_0^o(i)$ and $M^o(i)$ are normalized as $H_2(i)$ and M(i) in the same way.

Then, a correlation function can be defined as

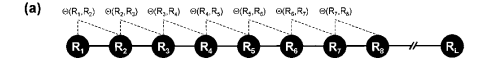
$$\Theta(R_i, R_j) = \frac{1}{3} \left\{ [H_1(R_i) - H_1(R_j)]^2 + [H_2(R_i) - H_2(R_j)]^2 + [M(R_i) - M(R_j)]^2 \right\}$$

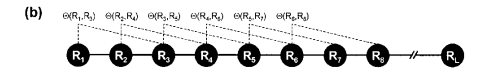
This correlation function is actually an averaged value for the three amino acid properties: hydrophobicity value, hydrophilicity value and side chain mass. Therefore we can extend this definition of correlation function for one amino acid property or for a set of n amino acid properties.

For one amino acid property, the correlation can be defined as:

$$\Theta(R_i, R_i) = [H_1(R_i) - H_1(R_i)]^2$$

where $H(R_i)$ is the amino acid property of amino acid R_i after standardization.





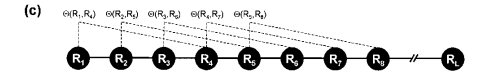


Figure 6: A schematic drawing to show (a) the first-tier, (b) the second-tier, and (3) the third-tiersequence order correlation mode along a protein sequence. Panel (a) reflects the correlation mode between all the most contiguous residues, panel (b) that between all the second-most contiguous residues, and panel (c) that between all the third-most contiguous residues. This figure is from Chou (2001).

For a set of n amino acid properties, it can be defined as: where $H_k(R_i)$ is the k-th property in the amino acid property set for amino acid R_i .

$$\Theta(R_i, R_j) = \frac{1}{n} \sum_{k=1}^{n} [H_k(R_i) - H_k(R_j)]^2$$

where $H_k(R_i)$ is the k-th property in the amino acid property set for amino acid R_i . A set of descriptors called sequence order-correlated factors are defined as:

$$\theta_{1} = \frac{1}{N-1} \sum_{i=1}^{N-1} \Theta(R_{i}, R_{i+1})$$

$$\theta_{2} = \frac{1}{N-2} \sum_{i=1}^{N-2} \Theta(R_{i}, R_{i+2})$$

$$\theta_{3} = \frac{1}{N-3} \sum_{i=1}^{N-3} \Theta(R_{i}, R_{i+3})$$
...
$$\theta_{\lambda} = \frac{1}{N-\lambda} \sum_{i=1}^{N-\lambda} \Theta(R_{i}, R_{i+\lambda})$$

 λ ($\lambda < L$) is a parameter to be chosen. Let f_i be the normalized occurrence frequency of the

20 amino acids in the protein sequence, a set of $20 + \lambda$ descriptors called the pseudo-amino acid composition for a protein sequence can be defines as:

$$X_c = \frac{f_c}{\sum_{r=1}^{20} f_r + w \sum_{j=1}^{\lambda} \theta_j} \quad (1 < c < 20)$$

$$X_c = \frac{w\theta_{c-20}}{\sum_{r=1}^{20} f_r + w \sum_{j=1}^{\lambda} \theta_j} \quad (21 < c < 20 + \lambda)$$

where w is the weighting factor for the sequence-order effect and is set to w = 0.05 in **protr** as suggested by Kuo-Chen Chou.

With extractPAAC(), we can compute the PAAC descriptors:

> extractPAAC(x)

The extractPAAC() function also provides the props and customprops arguments, which is similar to the functions for Moreau-Broto/Moran/Geary autocorrelation descriptors. For minor differences, see ?extracPAAC. Users can specify the lambda parameter and the weighting factor with arguments lambda and w.

Note: In the work of Kuo-Chen Chou, the definition for "normalized occurrence frequency" was not given. In this work, we define it as the occurrence frequency of amino acid in the sequence normalized to 100% and hence our calculated values are not the same as values by them.

3.9. Amphiphilic Pseudo-Amino Acid Composition (APAAC)

Amphiphilic Pseudo-Amino Acid Composition (APAAC) was proposed in Chou (2001). APAAC is also recognized as the *type 2 pseudo-amino acid composition*. The definitions of these qualities are similar to the PAAC descriptors. From $H_1(i)$ and $H_2(j)$ defined before, the hydrophobicity and hydrophilicity correlation functions are defined respectively as:

$$H_{i,j}^1 = H_1(i)H_1(j)$$

 $H_{i,j}^2 = H_2(i)H_2(j)$

From these qualities, sequence order factors can be defines as:

$$\tau_{1} = \frac{1}{N-1} \sum_{i=1}^{N-1} H_{i,i+1}^{1}$$

$$\tau_{2} = \frac{1}{N-1} \sum_{i=1}^{N-1} H_{i,i+1}^{2}$$

$$\tau_{3} = \frac{1}{N-2} \sum_{i=1}^{N-2} H_{i,i+2}^{1}$$

$$\tau_{4} = \frac{1}{N-2} \sum_{i=1}^{N-2} H_{i,i+2}^{2}$$

$$\dots$$

$$\tau_{2\lambda-1} = \frac{1}{N-\lambda} \sum_{i=1}^{N-\lambda} H_{i,i+\lambda}^{1}$$

$$\tau_{2\lambda} = \frac{1}{N-\lambda} \sum_{i=1}^{N-\lambda} H_{i,i+\lambda}^{2}$$

Then a set of descriptors called Amphiphilic Pseudo-Amino Acid Composition (APAAC) are defined as:

$$P_c = \frac{f_c}{\sum_{r=1}^{20} f_r + w \sum_{j=1}^{2\lambda} \tau_j} \quad (1 < c < 20)$$

$$P_c = \frac{w\tau_u}{\sum_{r=1}^{20} f_r + w \sum_{j=1}^{2\lambda} \tau_j} \quad (21 < u < 20 + 2\lambda)$$

where w is the weighting factor and is taken as w = 0.5 in **protr** as in the work of Chou KC. A minimal example for extracAPAAC() is:

> extractAPAAC(x)

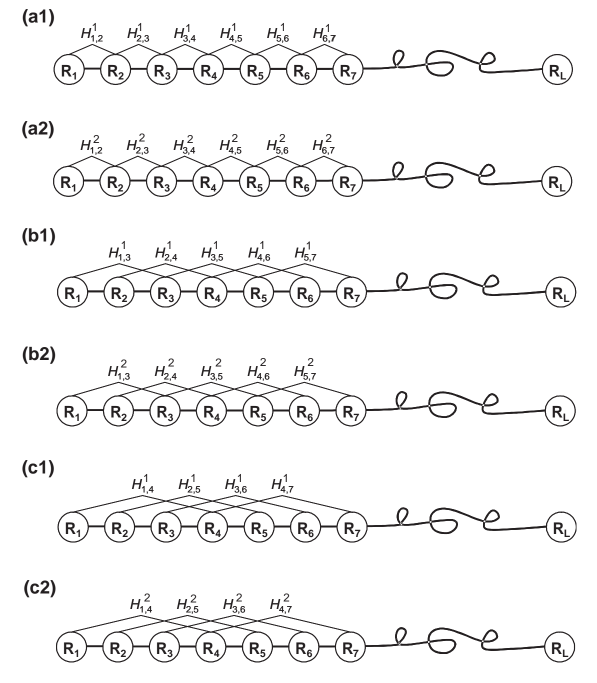


Figure 7: A schematic diagram to show $(\mathbf{a1/a2})$ the first-rank, $(\mathbf{b1/b2})$ the second-rank and $(\mathbf{c1/c2})$ the third-rank sequence-order-coupling mode along a protein sequence through a hydrophobicity/hydrophilicity correlation function, where $H_{i,j}^1$ and $H_{i,j}^2$ are given by Equation (3). Panel $(\mathbf{a1/a2})$ reflects the coupling mode between all the most contiguous residues, panel $(\mathbf{b1/b2})$ that between all the second-most contiguous residues and panel $(\mathbf{c1/c2})$ that between all the third-most contiguous residues. This figure is from Chou (2005).

Pc1.D	Pc1.C	Pc1.E
2.849582e+01	3.733935e+01	2.653059e+01
Pc1.Q	Pc1.G	Pc1.H
2.653059e+01	4.520027e+01	1.670445e+01
Pc1.I	Pc1.L	Pc1.K 2.161752e+01
1.965229e+01	4.225242e+01 Pc1.F	2.161752e+01 Pc1.P
Pc1.M 6.878302e+00	1.768706e+01	2.947844e+01
0.878302e+00 Pc1.S	1.768700e+01 Pc1.T	2.947644e+01 Pc1.W
4.913073e+01	2.456536e+01	1.277399e+01
4.3130736701 Pc1.Y	Pc1.V	Pc2.Hydrophobicity.1
2.358275e+01	2.751321e+01	2.196320e-04
Pc2.Hydrophilicity.1	Pc2.Hydrophobicity.2	Pc2.Hydrophilicity.2
1.025766e-03	-3.088876e-04	-1.834385e-04
Pc2.Hydrophobicity.3	Pc2.Hydrophilicity.3	Pc2.Hydrophobicity.4
1.174146e-03	7.400156e-04	-1.105715e-03
Pc2.Hydrophilicity.4	Pc2.Hydrophobicity.5	Pc2.Hydrophilicity.5
-4.493680e-04	1.766358e-03	1.471212e-03
Pc2.Hydrophobicity.6	Pc2.Hydrophilicity.6	Pc2.Hydrophobicity.7
-1.441572e-03	-4.913600e-03	-1.678053e-05
Pc2.Hydrophilicity.7	Pc2.Hydrophobicity.8	Pc2.Hydrophilicity.8
7.312356e-04	-1.885399e-03	-1.928708e-03
Pc2.Hydrophobicity.9	Pc2.Hydrophilicity.9	Pc2.Hydrophobicity.10
-2.931177e-03	-1.555660e-03	2.916597e-03
	Pc2.Hydrophobicity.11	
3.602591e-03	1.055082e-04	8.697920e-04
	Pc2.Hydrophilicity.12	
-9.276413e-04	-2.001384e-03	1.705044e-03
	Pc2.Hydrophobicity.14	· -
4.364007e-03	7.883453e-04	-9.441693e-04
	Pc2.Hydrophilicity.15	
-3.133437e-04	-3.599332e-03	3.689079e-05
2.483867e-03	Pc2.Hydrophobicity.17 4.832798e-04	
	Pc2.Hydrophilicity.18	
-3.142728e-04	2.021961e-03	6.421283e-05
	Pc2.Hydrophobicity.20	
-8.896690e-04	-2.986886e-04	9.304039e-04
	Pc2.Hydrophilicity.21	
-6.777458e-04	1.646818e-03	
	Pc2.Hydrophobicity.23	
3.270656e-03	2.533569e-03	2.478252e-03
	Pc2.Hydrophilicity.24	
-2.489106e-03	-1.031008e-03	-3.992322e-03
Pc2.Hydrophilicity.25	Pc2.Hydrophobicity.26	Pc2.Hydrophilicity.26
-2.596060e-03	8.690771e-04	-1.221378e-03
${\tt Pc2.Hydrophobicity.27}$	${\tt Pc2.Hydrophilicity.27}$	Pc2.Hydrophobicity.28

```
5.208649e-03 4.617400e-03 -1.088584e-03
Pc2.Hydrophilicity.28 Pc2.Hydrophobicity.29 Pc2.Hydrophilicity.29
-2.512263e-03 1.387641e-03 2.060890e-03
Pc2.Hydrophobicity.30 Pc2.Hydrophilicity.30
3.177340e-04 1.451909e-03
```

This function has the same arguments as extractPAAC().

3.10. Profile-based Descriptors

The profile-based descriptors for protein sequences are available in the **protr** package. The feature vectors of profile-based methods were based on the PSSM by running PSI-BLAST, and often show good performance. See Ye et al. (2011) and Rangwala and Karypis (2005) for details. The functions extractPSSM(), extractPSSMAcc() and extractPSSMFeature() are used to generate these descriptors. Users need to install the NCBI-BLAST+ software package first to make the functions fully functional.

4. Descriptors for Proteochemometric Modeling

Proteochemometric (PCM) modeling utilizes statistical modeling techniques to model ligand-target interaction space. The below descriptors implemented in **protr** are extensively used in Proteochemometric modeling.

- Scales-based descriptors derived by Principal Components Analysis
 - Scales-based descriptors derived by Amino Acid Properties from AAindex (Protein Fingerprint)
 - Scales-based descriptors derived by 20+ classes of 2D and 3D molecular descriptors (Topological, WHIM, VHSE, etc.)
- Scales-based descriptors derived by Factor Analysis
- Scales-based descriptors derived by Multidimensional Scaling
- BLOSUM and PAM matrix-derived descriptors

Note that each of the scales-based descriptor functions are freely to combine with the more than 20 classes of 2D and 3D molecular descriptors to construct highly customized scales-based descriptors. Of course, these functions are designed to be flexible enough that users can provide totally self-defined property matrices to construct scales-based descriptors.

For example, to compute the "topological scales" derived by PCA (using the first 5 principal components), one can use extractDescScales():

```
> 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)
```

Summary of the first 5 principal components:

```
PC1 PC2 PC3 PC4 PC5
Standard deviation 2.581537 1.754133 0.4621854 0.1918666 0.08972087
Proportion of Variance 0.666430 0.307700 0.0213600 0.0036800 0.00080000
Cumulative Proportion 0.666430 0.974130 0.9954900 0.9991700 0.99998000
```

the argument propmat involkes the AATopo dataset shipped with the **protr** package, and the argument index selects the 37 to 41 and the 43 to 47 columns (molecular descriptors) in the AATopo dataset to use, the parameter lag was set for the Auto Cross Covariance (ACC) for generating scales-based descriptors of the same length. At last, we printed the summary of the first 5 principal components (standard deviation, proportion of variance, cumulative proportion of variance).

The result is a length 175 named vector, which is consistent with the descriptors before:

> length(descscales)

[1] 175

> head(descscales, 15)

```
scl1.lag1
                 scl2.lag1
                               scl3.lag1
                                             scl4.lag1
                                                           scl5.lag1
-2.645644e-01 -1.717847e-02 1.975438e-02 -7.930659e-05 -3.710597e-05
   scl1.lag2
                 scl2.lag2
                               scl3.lag2
                                            scl4.lag2
                                                           scl5.lag2
3.548612e-01 1.343712e-01 5.699395e-03 -5.489472e-04 -6.364577e-05
   scl1.lag3
                 scl2.lag3
                               scl3.lag3
                                             scl4.lag3
                                                           scl5.lag3
2.011431e-02 -9.211136e-02 -1.461755e-03 6.747801e-04 2.386782e-04
```

For another example, to compute the descriptors derived by BLOSUM62 matrix and use the first 5 scales, one can use:

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

Relative importance of all the possible 20 scales:

```
[1] 1.204960e+01 7.982007e+00 6.254364e+00 4.533706e+00 4.326286e+00 [6] 3.850579e+00 3.752197e+00 3.538207e+00 3.139155e+00 2.546405e+00 [11] 2.373286e+00 1.666259e+00 1.553126e+00 1.263685e+00 1.024699e+00 [16] 9.630187e-01 9.225759e-01 7.221636e-01 1.020085e-01 5.868878e-16
```

The result is a length 175 named vector:

> length(blosum)

```
[1] 175
```

```
> head(blosum, 15)
```

```
scl1.lag1
                  scl2.lag1
                                 scl3.lag1
                                               scl4.lag1
                                                              scl5.lag1
0.0042370555 -0.0021502057
                              0.0005993291
                                            0.0006456375
                                                          0.0014849592
                                                              scl5.lag2
   scl1.lag2
                  scl2.lag2
                                 scl3.lag2
                                               scl4.lag2
-0.0014919096
               0.0032873726
                             0.0011734162 -0.0021758536 -0.0018127568
                  scl2.lag3
   scl1.lag3
                                 scl3.lag3
                                               scl4.lag3
                                                              scl5.lag3
-0.0029413528
               0.0001494193
                             0.0003298806 -0.0017877430 -0.0051044133
```

Dealing with gaps. In proteochemometrics, (sequence alignment) gaps can be very useful, since a gap in a certain position contains information. The **protr** package has built-in support for such gaps. We deal with the gaps by using a dummy descriptor to code for the 21st type of amino acid. The function extractScalesGap() and extractProtFPGap() can be used to deal with such gaps. See ?extractScalesGap and ?extractProtFPGap for details.

5. Similarity Calculation by Sequence Alignment

Similarity computation derived by local or global protein sequence alignment between a list of protein sequences is great need in the protein related research and applications. However, this sort of pairwise similarity computation often computationally intensive, especially when there exists many protein sequences. Luckily, this process is also highly parallelizable, the **protr** package integrates the function of parallelized similarity computation derived by local or global protein sequence alignment between a list of protein sequences.

The function twoSeqSim() calculates the alignment result between two protein sequences, and the function parSeqSim() calculates the pairwise similarity calculation with a list of protein sequences in parallel:

```
> 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 = 4, type = "local", submat = "BLOSUM62")
> print(psimmat)
           [,1]
                      [,2]
                                 [,3]
                                            [,4]
                                                        [,5]
[1,] 1.00000000 0.11825938 0.10236985 0.04921696 0.03943488
[2,] 0.11825938 1.00000000 0.18858241 0.12124217 0.06391103
[3,] 0.10236985 0.18858241 1.00000000 0.05819984 0.06175942
[4,] 0.04921696 0.12124217 0.05819984 1.00000000 0.05714638
[5,] 0.03943488 0.06391103 0.06175942 0.05714638 1.00000000
```

It should be noted that for a small number of proteins, calculating their pairwise similarity scores derived by sequence alignment in parallel may not significantly reduce the overall computation time, since each of the task only requires a relatively small time to finish, thus, computational overheads may exist and affect the performance. In testing, we used about 1,000 protein sequences on 64 CPU cores, and observed significant performance improvement comparing to the sequential computation.

Users should install the packages **foreach** and **doParallel** before using **parSeqSim()**, according to their operation system. The **protr** package will automatically decide which backend to use.

6. Similarity Calculation by GO Semantic Similarity Measures

The **protr** package also integrates the function of similarity score computation derived by Gene Ontology (GO) semantic similarity measures between a list of GO terms / Entrez Gene IDs.

The function twoGOSim() calculates the similarity derived by GO-terms semantic similarity measures between two GO terms / Entrez Gene IDs, and the function parGOSim() calculates the pairwise similarity with a list of GO terms / Entrez Gene IDs:

```
# by GO Terms
> go1 = c("G0:0005215", "G0:0005488", "G0:0005515",
          "GD:0005625", "GD:0005802", "GD:0005905")
                                                      # AP4B1
> go2 = c("G0:0005515", "G0:0005634", "G0:0005681",
          "GD:0008380", "GD:0031202")
                                                      # BCAS2
> go3 = c("G0:0003735", "G0:0005622", "G0:0005840",
          "GD:0006412")
                                                      # PDE4DIP
> glist = list(go1, go2, go3)
> gsimmat1 = parGOSim(glist, type = "go", ont = "CC")
> print(gsimmat1)
      [,1]
            [,2]
                  [,3]
[1,] 1.000 0.077 0.055
[2,] 0.077 1.000 0.220
[3,] 0.055 0.220 1.000
# by Entrez gene id
> genelist = list(c("150", "151", "152", "1814", "1815", "1816"))
> gsimmat2 = parGOSim(genelist, type = "gene")
> print(gsimmat2)
       150
             151
                   152 1814 1815 1816
    0.689 0.335 0.487 0.133 0.169 0.160
150
    0.335 0.605 0.441 0.171 0.198 0.274
152 0.487 0.441 0.591 0.151 0.178 0.198
1814 0.133 0.171 0.151 0.512 0.401 0.411
1815 0.169 0.198 0.178 0.401 0.619 0.481
1816 0.160 0.274 0.198 0.411 0.481 0.819
```

7. ProtrWeb

The web service built on **protr**, namely **ProtrWeb**, is located at:

http://protr.org

ProtrWeb (Figure 8) does not require any knowledge of programming for the users, it is a user-friendly web application for computing the protein sequence descriptors presented in the **protr** package.

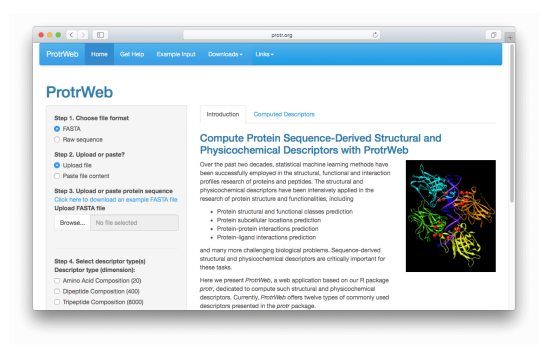


Figure 8: A screenshot of the web server **ProtrWeb**

Source code repository for this Shiny web application:

https://github.com/road2stat/protrweb

8. Miscellaneous Tools

In this section, we will briefly introduce some useful tools provided by the **protr** package.

8.1. Retrieve Protein Sequences from UniProt

This function getUniProt() gets protein sequences from uniprot.org by protein ID(s). The input ID is a character vector specifying the protein ID(s). The returned sequences are stored in a list:

```
> ids = c("P00750", "P00751", "P00752")
> prots = getUniProt(ids)
> print(prots)
```

[[1]]

[1] "MDAMKRGLCCVLLLCGAVFVSPSQEIHARFRRGARSYQVICRDEKTQMIYQQHQSWLRPVLRSNRVEYCWCN SGRAQCHSVPVKSCSEPRCFNGGTCQQALYFSDFVCQCPEGFAGKCCEIDTRATCYEDQGISYRGTWSTAESGAECT NWNSSALAQKPYSGRRPDAIRLGLGNHNYCRNPDRDSKPWCYVFKAGKYSSEFCSTPACSEGNSDCYFGNGSAYRGT HSLTESGASCLPWNSMILIGKVYTAQNPSAQALGLGKHNYCRNPDGDAKPWCHVLKNRRLTWEYCDVPSCSTCGLRQ YSQPQFRIKGGLFADIASHPWQAAIFAKHRRSPGERFLCGGILISSCWILSAAHCFQERFPPHHLTVILGRTYRVVP GEEEQKFEVEKYIVHKEFDDDTYDNDIALLQLKSDSSRCAQESSVVRTVCLPPADLQLPDWTECELSGYGKHEALSP FYSERLKEAHVRLYPSSRCTSQHLLNRTVTDNMLCAGDTRSGGPQANLHDACQGDSGGPLVCLNDGRMTLVGIISWG LGCGQKDVPGVYTKVTNYLDWIRDNMRP"

[[2]]

[1] "MGSNLSPQLCLMPFILGLLSGGVTTTPWSLARPQGSCSLEGVEIKGGSFRLLQEGQALEYVCPSGFYPYPVQ TRTCRSTGSWSTLKTQDQKTVRKAECRAIHCPRPHDFENGEYWPRSPYYNVSDEISFHCYDGYTLRGSANRTCQVNG RWSGQTAICDNGAGYCSNPGIPIGTRKVGSQYRLEDSVTYHCSRGLTLRGSQRRTCQEGGSWSGTEPSCQDSFMYDT PQEVAEAFLSSLTETIEGVDAEDGHGPGEQQKRKIVLDPSGSMNIYLVLDGSDSIGASNFTGAKKCLVNLIEKVASY GVKPRYGLVTYATYPKIWVKVSEADSSNADWVTKQLNEINYEDHKLKSGTNTKKALQAVYSMMSWPDDVPPEGWNRT RHVIILMTDGLHNMGGDPITVIDEIRDLLYIGKDRKNPREDYLDVYVFGVGPLVNQVNINALASKKDNEQHVFKVKD MENLEDVFYQMIDESQSLSLCGMVWEHRKGTDYHKQPWQAKISVIRPSKGHESCMGAVVSEYFVLTAAHCFTVDDKE HSIKVSVGGEKRDLEIEVVLFHPNYNINGKKEAGIPEFYDYDVALIKLKNKLKYGQTIRPICLPCTEGTTRALRLPP TTTCQQQKEELLPAQDIKALFVSEEEKKLTRKEVYIKNGDKKGSCERDAQYAPGYDKVKDISEVVTPRFLCTGGVSP YADPNTCRGDSGGPLIVHKRSRFIQVGVISWGVVDVCKNQKRQKQVPAHARDFHINLFQVLPWLKEKLQDEDLGFL"

[[3]]

[1] "APPIQSRIIGGRECEKNSHPWQVAIYHYSSFQCGGVLVNPKWVLTAAHCKNDNYEVWLGRHNLFENENTAQF FGVTADFPHPGFNLSLLKXHTKADGKDYSHDLMLLRLQSPAKITDAVKVLELPTQEPELGSTCEASGWGSIEPGPDB FEFPDEIQCVQLTLLQNTFCABAHPBKVTESMLCAGYLPGGKDTCMGDSGGPLICNGMWQGITSWGHTPCGSANKPS IYTKLIFYLDWINDTITENP"

8.2. Read FASTA Format files

The readFASTA() function provides a convenient way to read protein sequences stored in FASTA format files. See ?readFASTA for details. The returned sequences are stored in a named list, whose components are named with the protein sequences' names.

8.3. Read PDB Format files

The Protein Data Bank (pdb) file format is a textual file format describing the three dimensional structures of protein. The readPDB() function provides the function to read protein sequences stored in PDB format files. See ?readPDB for details.

8.4. Sanity Check of the Amino Acid Types

The protcheck() function checks if the protein sequence's amino acid types are in the 20 default types, which returns a TRUE if all the amino acids in the sequence belongs to the 20 default types:

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

```
> # A real sequence
> protcheck(x)

[1] TRUE
> # An artificial sequence
> protcheck(paste(x, "Z", sep = ""))

[1] FALSE
```

8.5. Protein Sequence Partition

The protseg() function partitions the protein sequences to create sliding windows. This is usually required when creating feature vectors for machine learning tasks. Users can specify a sequence x, and a character aa, one of the 20 amino acid types, and a positive integer k, which controls the window size (half of the window).

This function returns 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. See the example below:

```
> protseg(x, aa = "M", k = 5)

$`48`
[1] "DEKTQMIYQQH"

$`242`
[1] "LPWNSMILIGK"

$`490`
[1] "TVTDNMLCAGD"

$`525`
[1] "LNDGRMTLVGI"
```

8.6. Auto Cross Covariance (ACC) Computation

Auto Cross Covariance (ACC) is extensively used in the scales-based descriptors computation, this approach calculates the auto covariance and auto cross covariance for generating scale-based descriptors of the same length. Users can write their own scales-based descriptor functions with the help of acc() function in the **protr** package.

8.7. Pre-computed 2D and 3D Descriptor Sets for the 20 Amino Acids

The **protr** package ships with more than 20 pre-computed 2D and 3D descriptor sets for the 20 amino acids to use with the scales-based descriptors. Please use data(package = "protr") to list all the datasets included in the **protr** package.

8.8. BLOSUM and PAM Matrices for the 20 Amino Acids

The BLOSUM and PAM matrices for the 20 amino acids can be used to calculate BLOSUM and PAM matrix-derived descriptors with function extractBLOSUM(), the datasets are named in AABLOSUM45, AABLOSUM50, AABLOSUM62, AABLOSUM80, AABLOSUM100, AAPAM30, AAPAM40, AAPAM70, AAPAM120, and AAPAM250.

8.9. Meta Information of the 20 Amino Acids

As the reference, the AAMetaInfo dataset includes the meta information of the 20 amino acids used for the 2D and 3D descriptor calculation in the **protr** package. This dataset include each amino acid's name, one-letter representation, three-letter representation, SMILE representation, PubChem CID and PubChem link. See data(AAMetaInfo) for details.

9. Summary

The summary of the descriptors in the **protr** package are listed in Table 3.

Descriptor Group	Descriptor Name	Descriptor Dimension	Function Name
Amino Acid Composition	Amino Acid Composition	20	extractAAC()
	Dipeptide Composition	400	extractDC()
	Tripeptide Composition	8000	extractTC()
Autocorrelation	Normalized Moreau-Broto Auto- correlation	240^{1}	extractMoreauBroto()
	Moran Autocorrelation	240^{1}	<pre>extractMoran()</pre>
	Geary Autocorrelation	240^{1}	extractGeary()
CTD	Composition	21	<pre>extractCTDC(),</pre>
			extractCTDCClass()
	Transition	21	${\tt extractCTDT()},$
			extractCTDTClass()
	Distribution	105	${\tt extractCTDD()},$
			extractCTDDClass()
Conjoint Triad	Conjoint Triad	343	${\sf extractCTriad()},$
			extractCTriadClass()
Quasi-Sequence-Order	Sequence-Order-Coupling Number	60^{2}	extractSOCN()
	Quasi-Sequence-Order Descriptors	100^{2}	extractQSO()
Pseudo-Amino Acid Composition	Pseudo-Amino Acid Composition	50^{3}	extractPAAC()
	Amphiphilic Pseudo-Amino Acid Composition	80^{4}	extractAPAAC()

Table 3: List of commonly used descriptors in **protr**

The summary of the scales-based PCM descriptors in the **protr** package is listed in table 4. The summary of the amino acid descriptor sets used by scales-based descriptors provided in

The summary of the amino acid descriptor sets used by scales-based descriptors provided in the **protr** package is listed in table 5. Note that the non-informative descriptors (like the descriptors have only one value across all the 20 amino acids) in these datasets have already

¹The number depends on the choice of the number of properties of amino acids and the choice of the maximum values of the lag. The default is use 8 types of properties and lag = 30.

²The number depends on the maximum value of lag. By default lag = 30. And two distance matrices were used, so the descriptor dimension is $30 \times 2 = 60$ and $(20 + 30) \times 2 = 100$.

³The number depends on the choice of the number of the set of amino acid properties and the choice of the λ value. The default is use 3 types of properties proposed by Kuo-Chen Chou and $\lambda = 30$.

⁴The number depends on the choice of the λ value. The default is that $\lambda = 30$.

Table 4: List of PCM descriptors in **protr**

Derived by	Descriptor Class	Function Name
Principal Components Analysis	Scales-based descriptors derived by Principal Components Analysis Scales-based descriptors derived by amino acid properties from AAindex (a.k.a. Protein Finger- print)	<pre>extractScales(), extractScalesGap() extractProtFP(), extractProtFPGap()</pre>
	Scales-based descriptors derived by 2D and 3D molecular descriptors (Topological, WHIM, VHSE, etc.)	extractDescScales()
Factor Analysis	Scales-based descriptors derived by Factor Analysis	extractFAScales()
Multidimensional Scaling	Scales-based descriptors derived by Multidimensional Scaling	extractMDSScales()
Substitution Matrix	BLOSUM and PAM matrix-derived descriptors	<pre>extractBLOSUM()</pre>

been filtered out.

In this manual, we discussed the functions of the **protr** package, which offers a comprehensive and unique toolkit for protein sequence descriptor calculation and similarity computation.

10. How to Cite

If you feel **protr** is useful in your research, please feel free to cite our paper:

Nan Xiao, Dong-Sheng Cao, Min-Feng Zhu, and Qing-Song Xu. (2015). protr/ProtrWeb: R package and web server for generating various numerical representation schemes of protein sequences. *Bioinformatics* 31 (11), 1857–1859.

BibTeX entry:

```
author = {Xiao, Nan and Cao, Dong-Sheng and Zhu, Min-Feng and Xu, Qing-Song.},
title = {{protr/ProtrWeb: R package and web server for generating
  various numerical representation schemes of protein sequences}},
journal = {Bioinformatics},
year = {2015},
volume = {31},
number = {11},
pages = {1857--1859},
doi = {10.1093/bioinformatics/btv042},
issn = {1367-4803},
url = {http://bioinformatics.oxfordjournals.org/content/31/11/1857}
}
```

Table 5: List of the pre-calculated descriptor sets of the 20 amino acids in ${f protr}$

Dataset Name	Descriptor Set Name	Dimensionality	Calculated by
AA2DACOR	2D Autocorrelations Descriptors	92	Dragon
AA3DMoRSE	3D-MoRSE Descriptors	160	Dragon
AAACF	Atom-Centred Fragments Descriptors	6	Dragon
AABurden	Burden Eigenvalues Descriptors	62	Dragon
AAConn	Connectivity Indices Descriptors	33	Dragon
AAConst	Constitutional Descriptors	23	Dragon
AAEdgeAdj	Edge Adjacency Indices Descriptors	97	Dragon
AAEigIdx	Eigenvalue-Based Indices Descriptors	44	Dragon
AAFGC	Functional Group Counts Descriptors	5	Dragon
AAGeom	Geometrical Descriptors	41	Dragon
AAGETAWAY	GETAWAY Descriptors	194	Dragon
AAInfo	Information Indices Descriptors	47	Dragon
AAMolProp	Molecular Properties Descriptors	12	Dragon
AARandic	Randic Molecular Profiles Descriptors	41	Dragon
AARDF	RDF Descriptors	82	Dragon
AATopo	Topological Descriptors	78	Dragon
AATopoChg	Topological Charge Indices Descriptors	15	Dragon
AAWalk	Walk and Path Counts Descriptors	40	Dragon
AAWHIM	WHIM Descriptors	99	Dragon
AACPSA	CPSA Descriptors	41	Accelrys Discovery Studio
AADescAll	All the 2D Descriptors Calculated by Dragon	1171	Dragon
AAMOE2D	All the 2D Descriptors Calculated by MOE	148	$\overline{\mathrm{MOE}}$
AAMOE3D	All the 3D Descriptors Calculated by MOE	143	MOE

References

- Atchley WR, Zhao J, Fernandes AD, Drüke 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.
- Bhasin M, Raghava GPS (2004). "Classification of Nuclear Receptors Based on Amino Acid Composition and Dipeptide Composition." *Journal of Biological Chemistry*, **279**(22), 23262–6.
- Chou KC (2000). "Prediction of Protein Subcellar Locations by Incorporating Quasi-Sequence-Order Effect." Biochemical and Biophysical Research Communications, 278, 477–483.
- Chou KC (2001). "Prediction of Protein Cellular Attributes Using Pseudo-Amino Acid Composition." *PROTEINS: Structure, Function, and Genetics*, **43**, 246–255.
- Chou KC (2005). "Using Amphiphilic Pseudo Amino Acid Composition to Predict Enzyme Subfamily Classes." *Bioinformatics*, **21**, 10–19.
- Chou KC, Cai YD (2004). "Prediction of Protein Sub-cellular Locations by GO-FunD-PseAA Predictor." Biochemical and Biophysical Research Communications, 320, 1236–1239.
- Chou KC, Shen HB (2008). "Cell-PLoc: a package of Web servers for predicting subcellular localization of proteins in various organisms." Nature protocols, 3(2), 153–162.
- Damborsky J (1998). "Quantitative Structure-function and Structure-stability Relationships of Purposely Modified Proteins." *Protein Engineering*, **11**, 21–30.
- Dubchak I, Muchink I, Holbrook SR, Kim SH (1995). "Prediction of Protein Folding Class Using Global Description of Amino Acid Sequence." *Proceedings of the National Academy of Sciences*, **92**, 8700–8704.
- Dubchak I, Muchink I, Mayor C, Dralyuk I, Kim SH (1999). "Recognition of a Protein Fold in the Context of the SCOP Classification." *Proteins: Structure, Function and Genetics*, **35**, 401–407.
- Georgiev AG (2009). "Interpretable numerical descriptors of amino acid space." *Journal of Computational Biology*, **16**(5), 703–723.
- Grantham R (1974). "Amino Acid Difference Formula to Help Explain Protein Evolution." Science, 185, 862–864.
- Hellberg S, Sjoestroem M, Skagerberg B, Wold S (1987). "Peptide quantitative structure-activity relationships, a multivariate approach." *Journal of medicinal chemistry*, **30**(7), 1126–1135.
- Hopp-Woods (1981). "Prediction of Protein Antigenic Determinants from Amino Acid Sequences." Proceedings of the National Academy of Sciences, 78, 3824–3828.
- Kawashima S, Kanehisa M (2000). "AAindex: Amino Acid Index Database." Nucleic Acids Research, 28, 374.

- Kawashima S, Ogata H, Kanehisa M (1999). "AAindex: Amino Acid Index Database." *Nucleic Acids Research*, **27**, 368–369.
- Kawashima S, Pokarowski P, Pokarowska M, Kolinski A, Katayama T, Kanehisa M (2008). "AAindex: Amino Acid Index Database (Progress Report)." *Nucleic Acids Research*, **36**, D202–D205.
- Li Z, Lin H, Han Y, Jiang L, Chen X, Chen Y (2006). "PROFEAT: A Web Server for Computing Structural and Physicochemical Features of Proteins and Peptides from Amino Acid Sequence." *Nucleic Acids Research*, **34**, 32–37.
- Mei H, Liao ZH, Zhou Y, Li SZ (2005). "A new set of amino acid descriptors and its application in peptide QSARs." *Peptide Science*, **80**(6), 775–786.
- Pages H, Aboyoun P, Gentleman R, DebRoy S (2013). Biostrings: String objects representing biological sequences, and matching algorithms. R package version 2.30.1.
- Rangwala H, Karypis G (2005). "Profile-based direct kernels for remote homology detection and fold recognition." *Bioinformatics*, **21**(23), 4239–4247.
- Rao H, Zhu F, Yang G, Li Z, Chen Y (2011). "Update of PROFEAT: A Web Server for Computing Structural and Physicochemical Features of Proteins and Peptides from Amino Acid Sequence." Nucleic Acids Research, 39, 385–390.
- Sandberg M, Eriksson L, Jonsson J, Sjöström M, Wold S (1998). "New chemical descriptors relevant for the design of biologically active peptides. A multivariate characterization of 87 amino acids." *Journal of medicinal chemistry*, **41**(14), 2481–2491.
- Schneider G, Wrede P (1994). "The Rational Design of Amino Acid Sequences by Artificial Neural Networks and Simulated Molecular Evolution: Do Novo Design of an Idealized Leader Cleavage Site." Biophysical Journal, 66, 335–344.
- Shen J, Zhang J, Luo X, Zhu W, Yu K, Chen K, Li Y, Jiang H (2007). "Predicting Protein-protein Interactions Based Only on Sequences Information." *Proceedings of the National Academy of Sciences*, **104**, 4337–4341.
- Sjöström M, Rännar S, Wieslander Å (1995). "Polypeptide sequence property relationships in Escherichia coli based on auto cross covariances." Chemometrics and intelligent laboratory systems, 29(2), 295–305.
- Tian F, Zhou P, Li Z (2007). "T-scale as a novel vector of topological descriptors for amino acids and its application in QSARs of peptides." *Journal of molecular structure*, **830**(1), 106–115.
- van Westen GJ, Swier RF, Cortes-Ciriano I, Wegner JK, Overington JP, IJzerman AP, van Vlijmen HW, Bender A (2013a). "Benchmarking of protein descriptor sets in proteochemometric modeling (part 2): modeling performance of 13 amino acid descriptor sets." *Journal of cheminformatics*, **5**(1), 42.
- van Westen GJ, Swier RF, Wegner JK, IJzerman AP, van Vlijmen HW, Bender A (2013b). "Benchmarking of protein descriptor sets in proteochemometric modeling (part 1): comparative study of 13 amino acid descriptor sets." *Journal of cheminformatics*, **5**(1), 41.

- van Westen GJ, van den Hoven OO, van der Pijl R, Mulder-Krieger T, de Vries H, Wegner JK, IJzerman AP, van Vlijmen HW, Bender A (2012). "Identifying novel adenosine receptor ligands by simultaneous proteochemometric modeling of rat and human bioactivity data." *Journal of Medicinal Chemistry*, **55**(16), 7010–7020.
- van Westen GJ, Wegner JK, Geluykens P, Kwanten L, Vereycken I, Peeters A, IJzerman AP, van Vlijmen HW, Bender A (2011). "Which compound to select in lead optimization? Prospectively validated proteochemometric models guide preclinical development." *PloS one*, **6**(11), e27518.
- Venkatarajan MS, 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.
- Wikberg JE, Lapinsh M, Prusis P (2004). "Proteochemometrics: a tool for modeling the molecular interaction space." *Chemogenomics in drug discovery*, pp. 289–309.
- Xiao N, Cao DS, Zhu MF, Xu QS (2015). "protr/ProtrWeb: R package and web server for generating various numerical representation schemes of protein sequences." *Bioinformatics*, **31**(11), 1857–1859. ISSN 1367-4803. doi:10.1093/bioinformatics/btv042. URL http://bioinformatics.oxfordjournals.org/content/31/11/1857.
- Ye X, Wang G, Altschul SF (2011). "An assessment of substitution scores for protein profile—profile comparison." *Bioinformatics*, **27**(24), 3356–3363.
- Yu G, Li F, Qin Y, Bo X, Wu Y, Wang S (2010). "GOSemSim: an R package for measuring semantic similarity among GO terms and gene products." *Bioinformatics*, **26**(7), 976–978.
- Zaliani A, Gancia E (1999). "MS-WHIM scores for amino acids: a new 3D-description for peptide QSAR and QSPR studies." *Journal of chemical information and computer sciences*, **39**(3), 525–533.

Affiliation:

Nan Xiao School of Mathematics and Statistics Central South University Changsha, China E-mail: me@nanx.me

URL: https://nanx.me

Qing-Song Xu
School of Mathematics and Statistics
Central South University
Changsha, China
Dong-Sheng Cao
School of Pharmaceutical Sciences
Central South University
Changsha, China