**Pse-in-One 2.0:** a web server for generating comprehensive modes of pseudo components of DNA, RNA, and protein sequences

# Manual of stand-alone program of Pse-in-One-Analysis

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**Home-page**: <a href="http://bioinformatics.hitsz.edu.cn/Pse-in-One2.0/">http://bioinformatics.hitsz.edu.cn/Pse-in-One2.0/</a>







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# 1. Introduction

The **Pse-in-One 2.0** web server is an update version of **Pse-in-One** (1). **Pse-in-One 2.0** is able to generate totally 51 different modes of pseudo components for DNA, RNA, and protein sequences, including 20 modes for DNA sequences (**Table 1**), 14 modes for RNA sequences (**Table 2**), and 17 modes for protein sequences (**Table 3**). Compared with the old one, a total of 23 new pseudo component modes were added. In order to handle large dataset, the stand-alone program of **Pse-in-One 2.0** is given. Compared with the old version, a new facility called **Pse-in-One-Analysis** has been added, by which all the tedious jobs in developing a predictor, such as selecting optimal features and parameters as well as evaluating anticipated prediction quality, can be automatically fulfilled by the computer. Besides, to make the program lightweight and easy to use, all the 23 newly-added modes were added to the standalone program. More details will be introduced in the following parts of this manual.

#### 2. Installation

The **Pse-in-One 2.0** package can be run on Linux (64-bit) and Windows (64-bit) operating system. The full package and documents of **Pse-in-One 2.0** are available at <a href="http://bioinformatics.hitsz.edu.cn/Pse-in-One2.0/download">http://bioinformatics.hitsz.edu.cn/Pse-in-One2.0/download</a>. Before using **Pse-in-One 2.0**, the Python software should be first installed and configured. Python 2.7 64-bit is recommended, which can be downloaded from <a href="https://www.python.org">https://www.python.org</a>.

After Python installed, the Python package Numpy (2) should be downloaded and installed from here: <a href="http://www.numpy.org/">http://www.numpy.org/</a>, or use the following command if Internet is accessible:

> pip install numpy

For Windows operating system, the Windows 7 or later versions are supported. The next step is the installation and configuration of LIBSVM (3). Extract the package to a directory. After un-zip the downloaded **Pse-in-One 2.0** package, make sure that the "libsym.dll" is available in the directory "...\libsym\windows"

For Linux operating system, the LIBSVM should be configured firstly. Un-zip the Pse-Analysis package to a folder, for example, "~/usr". Navigate to "~/usr/Pse-in-One 2.0/libsvm" directory, and type the command: > make

After executing successfully, then navigate to "~/usr/ Pse-in-One 2.0/libsvm/python" directory, and type the command:

> make

If gnuplot has not been installed, use the following command lines to install gnuplot: > sudo apt-get install gnuplot

Now, **Pse-in-One 2.0** is ready to use.

## 3. Function description

### 3.1 Directory structure

The main directory contains several Python files and folders. "nac.py", "acc.py", "pse.py", "sc.py" and "profile.py" are five executive Python scripts used for generating feature vectors based on the input sequence files and the selected feature extraction methods. "train.py" and "predict.py" are two executive scripts used for doing the analysis. The details of their functions will be introduced in the following sections. "const.py" contains the constants used in the scripts. "util.py" provides the useful functions used in the scripts and "util\_sc.py" provides some specific functions used for "sc.py". "libsvm" folder contains the LIBSVM package. The tool for drawing ROC curve is in the "gnuplot" folder. "acc\_pssm" folder contains the tools used for ACC-PSSM, AC-PSSM and CC-PSSM methods. "pdt" folder contains the tools used for PDT and PDT-Profile methods. "psiblast" folder contains the tools used for generating frequency profiles of protein sequences. "docs" folder contains the related documents of Pse-in-One 2.0. In "data" folder, there are four subfolders: "example" folder contains the dataset files used in the example; "final\_results" folder is used for storing the generated model file while the "gen\_files" folder is used for storing the generated data files in the parameter selection process. The other files in the "data" folder are used for feature extraction methods. Modifications of these files are not suggested.

#### 3.2 Feature extraction

#### **3.2.1 Scripts**

"nac.py", "acc.py", "pse.py", "sc.py" and "profile.py" are five executive Python scripts used for generating feature vectors based on the input sequence files and the selected feature extraction methods.

The "nac.py" is used for calculating the modes in the category nucleic acid composition or amino acid composition; the "acc.py" is used for calculating the modes in autocorrelation category. The "pse.py" is used for calculating the modes in the category pseudo nucleotide composition or pseudo amino acid composition. The "sc.py" is used for calculating the modes in predicted structure composition category. The "profile.py" is used for calculating the modes in profile-based features category.

## 3.2.2 Input and output

The input file for "nac.py", "acc.py", "pse.py" and "profile.py" should be in a valid FASTA format that consists of a single initial line beginning with a greater-than symbol (">") in the first column, followed by lines of sequence data. The words right after the ">" symbol in the single initial line are optional and only used for the purpose of identification and description.

For "sc.py", the input file should be in a valid FASTA format with the secondary structure as follows:

>example

GCAUCCGGGUUGAGGUAGGUUGUAUGGUUUAGAGUUACACCCUGGG AGUUAACUGUACAACCUUCUAGCUUUCCUUGGAGC

The output file formats support three choices that are suitable for downstream computational analyses, such as machine learning. The first and the default choice is the tab format. In this format, all data is separated by TABs. The second one is the

LIBSVM's sparse data format. For this format, each line contains an instance and is ended by a '\n' character, like <label> <index1>:<value1> <index2>:<value2> ... . The <label> is a category label of the sequence. The pair <index>:<value> gives a feature (attribute) value: <index> is an integer starting from 1 and <value> is a real number. The third output format is the csv format. This format is similar to the tab format. The only difference is the separation characters between data are commas.

#### 3.2.3 Physicochemical Properties Selection

The Physicochemical Properties Selection file is a text file that contains a list of property names used for generating the modes in categories: autocorrelation, pseudo nucleotide composition/ pseudo amino acid composition. For example, if you want to use the "Rise", "Tilt" and "Shift" of DNA dinucleotide for calculating, the Physicochemical Properties Selection file should be written as follows:



After saving this file as "propChosen.txt" and specifying it using the command "-i propChosen.txt", or just "I propChosen.txt", the above three properties will be used in calculations. Meanwhile, you can also use the command "-a True" to select all the built-in physicochemical properties for the corresponding sequence type, which can be selected by using parameter DNA, RNA or PROTEIN.

The complete lists of physicochemical properties for DNA, RNA and protein sequences used in the stand-alone program are provided in **Table 4-12**.

### 3.2.4 User-defined Physicochemical Properties

In the user-defined physicochemical index files, each index should be represented in three lines. The first line must start with a greater-than symbol (">") in the first column. The words right after the ">" symbol in the single initial line are optional and only used for the purpose of identification and description of the index. The second line lists the names of the sequence compositions (i.e. amino acids, nucleotides, dinucleotides, or trinucleotides, etc), which should be sorted in the alphabet order, such as 'A' 'C' ... 'AA' 'AC'. All the elements in this line should be separated by TAB. The corresponding values of these sequence compositions are listed in the third line, which are separated by TAB.

For example, if you defined a physicochemical property "user\_property", the user-defined physicochemical index file should be written as follows:

After saving this file as "user\_defined.txt" and specifying it using the command "-e user\_defined.txt", or just "E user\_defined.txt", the properties defined by user will be used in calculations.

## 3.3 Pse-in-One-Analysis

The facility **Pse-in-One-Analysis** includes two main scripts: "train.py" and "predict.py".

#### **3.3.1 train.py**

#### **Basic functions**

The "train.py" is used for training SVM-based predictors and evaluating their performance based on the input benchmark datasets. Both binary classification and multiclass classification are supported. There are three main processes of "train.py", including parameter selection, model training and cross validation. In the parameter selection process, the parameters of LIBSVM are optimized on the validation sets. In this process, the multiprocessing technique is employed to significantly reduce the computational cost. In the model training process, the LIBSVM package is employed to train the prediction models. Finally, in the cross validation process, the performance of the constructed predictors is evaluated by k-fold cross-validation, jackknife or independent dataset test which can be selected by users. For more details of these three processes, please refer to the "Methods description" section.

#### Input and output

The input files of "train.py" are at least two files of feature vectors in LIBSVM format generated by the feature extraction methods in "nac.py", "acc.py", "pse.py" and "sc.py". For binary classification problem, two files need to be input, storing the positive samples and the negative samples, respectively. For multiclass classification, at least three files are needed. The output file is the trained SVM model listing the parameters used in the training process and the log information, for example:

```
c,128,g,0.5,b,0,bi_or_multi,0
svm_type c_svc
kernel_type rbf
gamma 0.5
nr_class 2
total_sv 2871
rho 33.5904
label 1 -1
nr_sv 1441 1430
SV
128 1:0.00108139 2:0.00108139 3:0.00108139 .....
```

### 3.3.2 predict.py

#### **Basic functions**

The "predict.py" predicts the unseen samples independent from the benchmark dataset based on the trained model generated by using "train.py". For binary classification, the performance of the constructed predictors is evaluated by five common performance measures, and the corresponding ROC curves can also be generated. For multiclass classification, only one measure is calculated. For more information of these functions, please refer to the "**Methods description**" section.

#### **Input and output**

The input file of "predict.py" is an independent file of feature vectors in LIBSVM format generated by feature extraction methods. If the label information of the samples

is available, the performance measures of the predictors will be calculated based on the predicted labels and the input real labels, otherwise, the performance will not be evaluated. One label should be listed in each line in the label file, for example:



The output of "predict.py" is a file containing the predicted labels in the same format as the input label file.

## 4. Commands

## 4.1 "nac.py" usage

Command line arguments for "nac.py":

| Required            | description   |
|---------------------|---|
| inputfiles          | The input files in FASTA format. More than one file could be input. |
| {DNA, RNA, Protein} | The sequence type.  |
| method              | The method name of nucleic acid composition.                        |

| Optional           | description  |
|--------------------|--|
| -h,help            | Show this help message and exit.   |
| -out               | The output files used for storing results. The number of output files should be the same as that of input files.   |
| -k K               | The k value of kmer.   |
| -m M               | For mismatch. The max value inexact matching. $(m < k)$ . $(default = 1)$  |
| -delta             | For subsequence method. The value of penalized factor. (0<=delta<=1). (default = 1)  |
| -r {0,1}           | Whether consider the reverse complement or not. 1  |
| -f {tab, svm, csv} | means True, 0 means False. (default = 0) The output format (default = tab).  |
|                    | tab Simple format, delimited by TAB. svm -   |
|                    | - The LIBSVM training data format.   |
|                    | csv The format that can be loaded into a spreadsheet program.  |
| -labels            | The libSVM output file label. If the argument "-f" is set as "svm", this argument is required. And the number of labels should be the same as that of the input files. For binary classification problem, the labels should be '+1' or '-1'; For multiclass classification problem, the labels can be set as |
|                    | integers.  |

| -ps      | The input positive source file in FASTA format for IDKmer. Only for IDKmer method.                              |
|----------|---|
| -ns      | The input negative source file in FASTA format for IDKmer. Only for IDKmer method.                              |
| -max_dis | The max distance value of DR and Distance Pair. Only for DR and Distance Pair methods(default = 3).             |
| -cp      | The reduced alphabet scheme. Choose one of the four: cp_13, cp_14, cp_19, cp_20. Only for Distance Pair method. |

# 4.2 "acc.py" usage

Command line arguments for "acc.py":

| Required            | description   |
|---------------------|---|
| inputfiles          | The input files in FASTA format. More than one file could be input. |
| {DNA, RNA, Protein} | The sequence type.  |
| method              | The method name of autocorrelation.                                 |

| Optional           | description   |
|--------------------|---|
| -h,help            | Show this help message and exit.                      |
| -out               | The output files used for storing results. The number |
|                    | of output files should be the same as that of input   |
|                    | files.  |
| -lag LAG           | The value of lag.                                     |
| -i I               | The index file user chosen.                           |
| -e E               | The user-defined index file.                          |
| -all_index         | Choose all physicochemical indices.                   |
| -no_all_index      | Do not choose all physicochemical indices, default.   |
| -f {tab, svm, csv} | The output format ( $default = tab$ ).                |
|                    | tab Simple format, delimited by TAB.                  |
|                    | svm The LIBSVM training data format.                  |
|                    | csv The format that can be loaded into a spreadsheet  |
|                    | program.  |
| -labels            | The libSVM output file label. If the argument "-f" is |
|                    | set as "svm", this argument is required. And the      |
|                    | number of labels should be the same as that of the    |
|                    | input files. For binary classification problem, the   |
|                    | labels should be '+1' or '-1'; For multiclass         |
|                    | classification problem, the labels can be set as      |
|                    | integers.   |
| -lamada            | The value of lamada. Only for MAC, GAC, NMBAC         |
|                    | methods (default=1).                                  |
| -oli               | Choose one kind of Oligonucleotide:                   |
|                    | 0 represents dinucleotide, default;                   |
|                    | 1 represents trinucleotide.                           |
|                    |   |

# 4.3 "pse.py" usage

Command line arguments for "pse.py":

| Required            | description   |
|---------------------|---|
| inputfiles          | The input files in FASTA format. More than one file could be input. |
| {DNA, RNA, Protein} | The sequence type.  |
| method              | The method name of pseudo components.                               |

| Optional           | description  |
|--------------------|--|
| -h,help            | Show this help message and exit.                             |
| -out               | The output files used for storing results. The number        |
|                    | of output files should be the same as that of input          |
|                    | files.   |
| -lamada            | The value of lamada (default=2).                             |
| -w W               | The value of weight (default=0.1).                           |
| -k K               | The value of kmer, it works only with PseKNC method.         |
| -e Е               | The user-defined index file, this parameter only needs to be |
|                    | set for PC-PseDNC-General, PC-PseTNC-General, SC-            |
|                    | PseDNC-General, SC-PseTNC-General, PC-PseAAC-                |
|                    | General or SC-PseAAC-General.                                |
| -all_index         | Choose all physicochemical indices.                          |
| -no_all_index      | Do not choose all physicochemical indices, default.          |
| -f {tab, svm, csv} | The output format (default $=$ tab).                         |
|                    | tab Simple format, delimited by TAB.                         |
|                    | svm The LIBSVM training data format.                         |
|                    | csv The format that can be loaded into a spreadsheet         |
|                    | program.   |
| -labels            | The libSVM output file label. If the argument "-f" is        |
|                    | set as "svm", this argument is required. And the             |
|                    | number of labels should be the same as that of the           |
|                    | input files. For binary classification problem, the          |
|                    | labels should be '+1' or '-1'; For multiclass                |
|                    | classification problem, the labels can be set as             |
|                    | integers.  |

# 4.4 "sc.py" usage

Command line arguments for "sc.py":

| Required            | description   |
|---------------------|---|
| inputfiles          | The input files in FASTA format. More than one file could be input. |
| {DNA, RNA, Protein} | The sequence type.  |
| method              | The method name of pseudo components.                               |
| Optional            | description   |

-h, --help Show this help message and exit.

|                    | 9  |
|--------------------|--|
| -out               | The output files used for storing results. The number of output files should be the same as that of input files  |
| -k K               | The number of k adjacent structure statuses (default=2). It works only with PseSSC method.   |
| -n N               | The maximum distance between structure statuses (default=0). It works only with PseDPC method.   |
| -r R               | The value of lambda, represents the highest counted rank (or tier) of the structural correlation along a RNA chain (default=2).  |
| -w W               | The weight factor used to adjust the effect of the correlation factors (default=0.1).  |
| -f {tab, svm, csv} | The output format (default = tab). tab Simple format, delimited by TAB.  |
|                    | svm The LIBSVM training data format. csv The format that can be loaded into a spreadsheet  |
| -labels            | program.  The libSVM output file label. If the argument "-f" is set as "svm", this argument is required. And the number of labels should be the same as that of the input files. For binary classification problem, the labels should be '+1' or '-1'; For multiclass classification problem, the labels can be set as integers. |
|                    |  |

# 4.5 "profile.py" usage

Command line arguments for "profile.py":

| Required   | description   |
|------------|---|
| inputfiles | The input files in FASTA format. More than one file could be input. |
| method     | The method name of pseudo components.                               |

| memou              | The method hame of poeddo components.   |
|--------------------|---|
| Optional           | description   |
| -h,help            | Show this help message and exit.  |
| -out               | The output files used for storing results. The number of output files should be the same as that of input files.  |
| -n N               | For Top-n-gram, PDT-Profile methods. The value of top-n-gram. The value cam only be 1, 2 or 3.  |
| -lamada            | For PDT, PDT-Profile methods. The value of lamada   |
| -max_dis           | For DT methods. The max distance value of residues (default = $3$ ).  |
| -lag LAG           | For ACC-PSSM, AC-PSSM and CC-PSSM methods. The value of lag (default = 2).  |
| -f {tab, svm, csv} | The output format (default = tab).  tab Simple format, delimited by TAB.  svm The LIBSVM training data format.  csv The format that can be loaded into a spreadsheet program. |
|                    |   |

| -labels  | The libSVM output file label. If the argument "-f" is set as "svm", this argument is required. And the number of labels should be the same as that of the input files. For binary classification problem, the labels should be '+1' or '-1'; For multiclass classification problem, the labels can be set as integers. | 10 |
|----------|--|----|
| -cpu CPU | The maximum number of CPU cores used for multiprocessing in generating frequency profile. Default value is 1.  |    |

# 4.5 "train.py" usage

Command line arguments for "train.py":

| requir | red description  |
|--------|--|
| files  | The input files in LIBSVM format, generated by feature extraction methods mentioned above or <b>Pse-in-One 2.0</b> |
|        | webserver.   |
|        | For binary classification, two files needed.   |
| -m M   | For multiclass classification, at least three files needed. The name of the trained SVM model.                     |

| Optional         | description   |
|------------------|---|
| -h,help          | Show this help message and exit.  |
| -p {ACC,MCC,AUC} | The performance metric used for parameter selection. Default value is "ACC".  |
| -v V             | The cross validation mode.  |
|                  | n: (an integer larger than 0) n-fold cross validation.  |
|                  | j: (character "j") jackknife cross validation.  |
|                  | i: (character 'i') independent test set method.   |
| -i_files         | Set the range of parameters to be optimized.<br>0: small range set c from -5 to 10, step is 2; g<br>from -10 to 5, step is 2.   |
| -opt             | 1: large range set c from -5 to 10, step is 1; g from -10 to 5, step is 1.  Default value is 0.  Set the range of parameters to be optimized.  0: small range set c from -5 to 10, step is 2; g                         |
| -b {0,1}         | from -10 to 5, step is 2.  1: large range set c from -5 to 10, step is 1; g from -10 to 5, step is 1.  Default value is 0.  Whether to train a SVC or SVR model for probability estimates, 0 or 1.  Default value is 0. |

The maximum number of CPU cores used for multiprocessing during parameter selection process. Default value is the number of all available CPU cores.

## 4.6 "predict.py" usage

Command line arguments for "predict.py":

| required   | description   |
|------------|---|
| inputfiles | The input files in LIBSVM format, generated by feature extraction methods mentioned above or <b>Pse-in-One 2.0</b> webserver. |
| -m M       | The name of the trained SVM model.  |

| optional       | description   |  |
|----------------|---|--|
| -h,help        | Show this help message and exit.  |  |
| -labels LABELS | The real label file. Optional.  |  |
| -o O           | The output file name listing the predicted labels. The default name is "output_labels.txt". |  |

## 4.7 Example

An example of using **Pse-in-One 2.0** to construct machine learning predictor for solving a specific task in bioinformatics is given.

**Example:** Reconstructing the predictor PseDNA-Pro for DNA binding protein identification based on the benchmark dataset (4), and evaluating its performance on an independent dataset (5) by using **Pse-in-One 2.0**.

The benchmark dataset contains 525 positive samples and 550 negative samples. There are 93 positive samples and 93 negative samples in the independent dataset. The benchmark dataset and independent dataset are available at

http://bioinformatics.hitsz.edu.cn/PseDNA-Pro/Resources/benchmark\_dataset.pdf, and, http://journals.plos.org/plosone/article/asset?unique&id=info:doi/10.1371/journal.pone.0 086703.s002, respectively.

In this example, the files "protein\_pos.txt" and "protein\_neg.txt" contain the positive dataset and negative dataset of the benchmark dataset, respectively. The samples of the independent dataset and their labels are stored in the files "protein\_test.txt" and "labels.txt", respectively. All these four files are available in the "/data/example" folder.

Firstly, to construct the predictor PseDNA-Pro, extract features based on the benchmark dataset.

Convert the positive dataset "protein\_pos.txt" and negative dataset "protein\_neg.txt" into feature vectors in LIBSVM format, and save as output files "pos\_svm.txt" and "neg\_svm.txt":

```
neg_svm.txt
```

The content of the output file "pos\_svm.txt" is as follows:

```
+1 1:0.04075964 2:0.01358655 3:0.0 4:0.08151929 5:0.04075964 6:0.0271731...
+1 1:0.29181499 2:0.19233261 3:0.0 4:0.0 5:0.0 6:0.19233261 ...
+1 1:0.12355132 2:0.0 3:0.05615969 4:0.07862357 5:0.01123194 6:0.04492775...
```

The content of the output file "neg\_svm.txt" is as follows:

```
-1 1:0.06394096 2:0.06394096 3:0.05195203 4:0.02397786 5:0.04395941 ...
-1 1:0.07606996 2:0.0 3:0.05071331 4:0.03728919 5:0.02535665 ...
-1 1:0.04873717 2:0.01624572 3:0.03858359 4:0.05076788 5:0.0263993 ...
.....
```

After feature extraction, construct the predictor PseDNA-Pro based on "pos\_svm.txt" and "neg\_svm.txt":

```
python train.py pos_svm.txt neg_svm.txt -m protein.model -opt 0 -v 5
```

The output information is as follows:

ACC = 0.7451

```
Processing...
Parameter selection is in processing...
Iteration c = 7 g = -10 finished.
Iteration c = -5 g = -1 finished.
Iteration c = -5 g = -10 finished.
Iteration c = -5 g = -7 finished.
Iteration c = -2 g = -10 finished.
Iteration c = -5 g = 5 finished.
Iteration c = -2 g = 5 finished.
Iteration c = 4 g = -7 finished.
Iteration c = 1 g = 2 finished.
. . . . . .
. . . . . .
Iteration c = 10 g = -1 finished.
Iteration c = 10 g = -10 finished.
Iteration c = 4 g = -10 finished.
Iteration c = -2 g = -7 finished.
Iteration c = 10 g = 2 finished.
Iteration c = 7 g = 5 finished.
Iteration c = 10 g = 5 finished.
The time cost for parameter selection is 4.48s
Parameter selection completed.
The optimal parameters for the dataset are: C = 0.25 gamma = 32
Model training is in processing...
The cross validation results are as follows:
```

MCC = 0.4946 AUC = 0.8212Sn = 0.7392

Sp = 0.7525

The ROC curve has been saved. You can check it here: /Pse-in-One 2.0/data/final\_results/cross\_validation.png

Model training completed.

The model has been saved. You can check it here:

/Pse-in-One 2.0/data/final\_results/protein.model

Done.

Used time: 6.57s

The generated ROC curve is shown in **Fig. 1**.

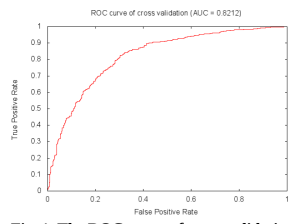


Fig .1. The ROC curve of cross validation.

Then, the performance of PseDNA-Pro can be further evaluated by using the independent dataset:

Firstly, convert the independent dataset "protein\_test.txt" into feature vectors in LIBSVM format, and save as an output file "test\_svm.txt":

python pse.py ./data/example/protein\_test.txt Protein PC-PseAAC -lamada 2 -w 0.05 -f svm -labels 0 -out test\_svm.txt

The content of the output file "test\_svm.txt" is as follows:

```
+1 1:0.09717009 2:0.0 3:0.04318671 4:0.04318671 5:0.02159335 ...

+1 1:0.04954724 2:0.00707818 3:0.05662542 4:0.07078177 5:0.05662542 ...

+1 1:0.05782654 2:0.01084248 3:0.05421239 4:0.06866902 5:0.04336991 ...

.....
```

After feature extraction, predict the independent dataset using the following commad:

python predict.py test\_svm.txt -m protein.model -labels ./data/example/labels.txt

The output information is as follows:

Processing...

The parameters of RBF kernel:

c = 0.25 g = 32

The performance evaluations are as follows:

ACC = 0.6720

MCC = 0.3487

AUC = 0.7294

Sn = 0.7527

Sp = 0.5914

The ROC curve has been saved. You can check it here:

/Pse-in-One 2.0/data/final\_results/predicted\_roc.png

The predicted labels have been saved. You can check it here:

/Pse-in-One 2.0/data/final\_results/output\_labels.txt

Done.

Used time: 3.16s

As shown in this example, the PseDNA-Pro can be easily constructed based on the benchmark dataset by using the script "train.py", and then evaluated on the independent dataset by using "predict.py".

## 5. Methods description

#### **5.1 Feature extraction**

The **Pse-in-One 2.0** web server is able to generate totally 51 different modes of pseudo components for DNA, RNA, and protein sequences, including 20 modes for DNA sequences (**Table 1**), 14 modes for RNA sequences (**Table 2**), and 17 modes for protein sequences (**Table 3**). The detailed information of the 51 methods will be introduced in Pse-in-One 2.0 description document which can be downloaded from here: <a href="http://bioinformatics.hitsz.edu.cn/Pse-in-One2.0/static/download/Pse-in-One%202.0\_description.pdf">http://bioinformatics.hitsz.edu.cn/Pse-in-One2.0/static/download/Pse-in-One%202.0\_description.pdf</a>.

#### **5.2 Parameter selection**

In LIBSVM there are two parameters c and g which can determine the performance of the predictor. **Pse-in-One 2.0** is able to automatically optimize these parameters based on the best performance on the validation set by using the new facility **Pse-in-One-Analysis**. Users can choose a range of the two parameters for optimizing. For more information of the input format, please refer to "**Commands**" section. To improve the efficiency of this procedure, multiprocessing technique is applied, which significantly reduces the computational cost. One of the three performance measures, including Accuracy (ACC), Mathew's Correlation Coefficient (MCC) and Area Under roc Curve (AUC) can be used as the golden standard to optimize the parameters.

## **5.3 Model training**

In the model training process, this model is trained based on LIBSVM with RBF kernel. The trained SVM model and all the parameters are saved in a separate file, which will be used as the input for "predict.py".

#### **5.4 Cross validation**

**Pse-in-One 2.0** provides three types of cross validation options, including k-fold cross validation, jackknife (leave-one-out cross validation) and independent dataset test, which can be chosen by the argument "-v". Please refer to "**Commands**" section for more details.

For binary classification, the performance of the predictor is measured by five common performance measures, including the accuracy (Acc), Mathew's Correlation Coefficient (MCC), Area Under roc Curve (AUC), sensitivity (Sn), and specificity (Sp). Furthermore, the ROC (Receiver Operating Characteristic) (6) curve will also be generated and saved in a PNG file.

For multiclass classification, only the performance measure of Acc is calculated since the other measures are not suitable for multiclass classification.

Besides, if the parameter "-b" of libsvm is set, the prediction probability values will be output and save as a file, thus users can do further analysis with these data.

### **5.5** Sequence prediction

The "predict.py" is used to predict the unseen samples based on the model trained by using "train.py". The performance of the predictors can be further evaluated on the independent datasets. If the label information of the independent dataset is not available, the performance of the predictor will not be evaluated, and only the predicted labels are given. Otherwise, this script will output the predicted labels. For binary classification, the five performance measures (Acc, MCC, AUC, Sn, and Sp) will be calculated along with the corresponding ROC curve saved as a PNG file; for multiclass classification, only the performance measure Acc will be calculated.

**Table 1.** 20 modes of DNA sequences.

| Category                 | Mode          | Description  |
|--------------------------|---------------|--|
|                          | Kmer          | Basic kmer (7)   |
|                          | RevKmer       | Reverse complementary kmer(8,9)                                    |
|                          | <b>IDKmer</b> | increment of diversity (10-12)                                     |
| Nucleic acid Composition | Mismatch      | The occurrences of kmers, allowing at most m mismatches (13-15)    |
|                          | Subsequence   | The occurences of kmers, allowing non-contiguous maches (13,15,16) |
|                          | DAC           | Dinucleotide-based auto covariance (17,18)                         |
|                          | DCC           | Dinucleotide-based cross covariance (17,18)                        |
|                          | DACC          | Dinucleotide-based auto-cross covariance (17,18)                   |
|                          | TAC           | Trinucleotide-based auto covariance (17)                           |
| Autocorrelation          | TCC           | Trinucleotide-based cross covariance (17)                          |
|                          | TACC          | Trinucleotide-based auto-cross covariance (17)                     |
|                          | MAC           | Moran autocorrelation (19,20)                                      |
|                          | GAC           | Geary autocorrelation (20,21)                                      |
|                          | NMBAC         | Normalized Moreau-Broto autocorrelation (20,22)                    |
|                          | PseDNC        | Pseudo dinucleotide composition (23)                               |
|                          | PseKNC        | Pseudo k-tuple nucleotide composition (24,25)                      |
|                          | PC-PseDNC-    | General parallel correlation                                       |
|                          | General       | pseudo dinucleotide composition (26)                               |
| Pseudo nucleotide        | PC-PseTNC-    | General parallel correlation                                       |
| composition              | General       | pseudo trinucleotide<br>composition (26)                           |
|                          | SC-PseDNC-    | General series correlation   |
|                          | General       | pseudo dinucleotide composition (26)                               |
|                          | SC-PseTNC-    | General series correlation   |
|                          | General       | pseudo trinucleotide composition (26)                              |

**Table 2.** 14 modes of RNA sequences.

| Category                 | Mode               | Description                        |
|--------------------------|--------------------|------------------------------------|
|                          | Kmer               | Basic kmer (27)                    |
|                          | Mismatch           | The occurrences of kmers,          |
|                          |                    | allowing at most m                 |
| Nucleic acid Composition |                    | mismatches (13-15)                 |
|                          | Subsequence        | The occurences of kmers,           |
|                          |                    | allowing non-contiguous            |
|                          |                    | maches (13,15,16)                  |
|                          | DAC                | Dinucleotide-based auto            |
|                          |                    | covariance (17,18,28)              |
|                          | DCC                | Dinucleotide-based cross           |
|                          |                    | covariance (17,18,28)              |
|                          | DACC               | Dinucleotide-based auto-           |
|                          |                    | cross covariance                   |
| Autocorrelation          |                    | (17,18,28)                         |
| ratocorrelation          | MAC                | Moran autocorrelation              |
|                          |                    | (19,20)                            |
|                          | GAC                | Geary autocorrelation              |
|                          |                    | (20,21)                            |
|                          | NMBAC              | Normalized Moreau-                 |
|                          |                    | Broto autocorrelation              |
|                          | DOD DVG G          | (20,22)                            |
|                          | PC-PseDNC- General | General parallel                   |
|                          |                    | correlation pseudo                 |
| Pseudo nucleotide        |                    | dinucleotide composition           |
| composition              | SC-PseDNC-General  | (18,20) General series correlation |
|                          | SC-PseDNC-General  | pseudo dinucleotide                |
|                          |                    | composition (18,20)                |
|                          | Triplet            | Local structure-sequence           |
|                          | Tipici             | triplet element (29)               |
|                          | PseSSC             | Pseudo-structure status            |
| Predicted Structure      | 1 50000            | composition (30)                   |
| composition              | PseDPC             | Pseudo-distance structure          |
|                          | 150010             | status pair composition            |
|                          |                    | (31)                               |

**Table 3.** 17 modes of protein sequences.

| Category               | Mode              | Description                |
|------------------------|-------------------|----------------------------|
|                        | Kmer              | Basic kmer (32)            |
| Amino acid composition | DR                | Distance-based Residue     |
|                        |                   | (33)                       |
|                        | Distance Pair     | PseAAC of Distance-        |
|                        |                   | Pairs and Reduced          |
|                        |                   | Alphabet (34)              |
|                        | AC                | Auto covariance (17,28)    |
|                        | CC                | Cross covariance (17,28)   |
| A vyta a a malati a m  | ACC               | Auto-cross covariance      |
| Autocorrelation        |                   | (17,28)                    |
|                        | PDT               | Physicochemical distance   |
|                        |                   | transformation (35)        |
|                        | PC-PseAAC         | Parallel correlation       |
|                        |                   | pseudo amino acid          |
|                        |                   | composition (36)           |
|                        | SC-PseAAC         | Series correlation pseudo  |
|                        |                   | amino acid composition     |
| Pseudo amino acid      |                   | (37)                       |
| composition            | PC-PseAAC-General | General parallel           |
|                        |                   | correlation pseudo amino   |
|                        |                   | acid composition (36,38)   |
|                        | SC-PseAAC-General | General series correlation |
|                        |                   | pseudo amino acid          |
|                        |                   | composition (37,38)        |
|                        | Top-n-gram        | Select and combine the n   |
|                        |                   | most frequenct amino       |
|                        |                   | acids according to their   |
|                        |                   | frequencies. (32)          |
|                        | PDT-Pofile        | Profile-based              |
|                        |                   | Physicochemical distance   |
|                        |                   | transformation (35)        |
| Profile-based features | DT                | Distance-based Top-n-      |
|                        |                   | gram (33)                  |
|                        | AC-PSSM           | Profile-based Auto         |
|                        |                   | covariance (17)            |
|                        | CC-PSSM           | Profile-based Cross        |
|                        |                   | covariance (17)            |
|                        | ACC-PSSM          | Profile-based Auto-cross   |
|                        |                   | covariance (17)            |

**Table 4.** The names of the 148 physicochemical indices for dinucleotides.

| Base stacking               | Protein                       | B-DNA twist                    |
|-----------------------------|-------------------------------|--------------------------------|
|                             | induced deformability         |                                |
| Propeller twist             | Duplex stability:(freeenergy) | Duplex tability(disruptenergy) |
| Protein DNA twist           | Stabilising energy of Z-DNA   | Aida_BA_transition             |
| Breslauer_dS                | Electron_interaction          | Hartman_trans_free_energy      |
| Lisser_BZ_transition        | Polar_interaction             | SantaLucia_dG                  |
| Sarai_flexibility           | Stability                     | Stacking_energy                |
| Sugimoto_dS                 | Watson-                       | Twist                          |
| 8 =                         | Crick_interaction             |                                |
| Shift                       | Slide                         | Rise                           |
| Twist stiffness             | Tilt stiffness                | Shift_rise                     |
| Twist_shift                 | Enthalpy1                     | Twist_twist                    |
| Shift2                      | Tilt3                         | Tilt1                          |
| Slide (DNA-protein          | Tilt_shift                    | Twist_tilt                     |
| complex)1                   |                               |                                |
| Roll_rise                   | Stacking energy               | Stacking energy1               |
| Propeller Twist             | Roll11                        | Rise (DNA-protein complex)     |
| Roll2                       | Roll3                         | Roll1                          |
| Slide_slide                 | Enthalpy                      | Shift_shift                    |
| Flexibility_slide           | Minor Groove Distance         | Rise (DNA-protein complex)1    |
| Roll (DNA-protein complex)1 | Entropy                       | Cytosine content               |
| Major Groove Distance       | Twist (DNA-protein complex)   | Purine (AG) content            |
| Tilt slide                  | Major Groove Width            | Major Groove Depth             |
| Free energy6                | Free energy7                  | Free energy4                   |
| Free energy3                | Free energy1                  | Twist roll                     |
| Flexibility_shift           | Shift (DNA-protein complex)1  | Thymine content                |
| Tip                         | Keto (GT) content             | Roll stiffness                 |
| Entropy1                    | Roll_slide                    | Slide (DNA-protein complex)    |
| Twist2                      | Twist5                        | Twist4                         |
| Tilt (DNA-protein           | Twist_slide                   | Minor Groove Depth             |
| complex)1                   | I WISt_SHGC                   | Willion Groove Depth           |
| Persistance Length          | Rise3                         | Shift stiffness                |
| Slide3                      | Slide2                        | Slide1                         |
| Rise1                       | Rise stiffness                | Mobility to bend towards minor |
|                             |                               | groove                         |
| Dinucleotide GC Content     | A-philicity                   | Wedge                          |
| DNA denaturation            | Bending stiffness             | Free energy5                   |
| Breslauer_dG                | Breslauer_dH                  | Shift (DNA-protein complex)    |
| Helix-Coil_transition       | Ivanov_BA_transition          | Slide_rise                     |
| SantaLucia_dH               | SantaLucia_dS                 | Minor Groove Width             |
| Sugimoto_dG                 | Sugimoto_dH                   | Twist1                         |
| Tilt                        | Roll                          | Twist7                         |
| Clash Strength              | Roll_roll                     | Roll (DNA-protein complex)     |
| Adenine content             | Direction                     | Probability contacting         |
|                             |                               | nucleosome core                |
| Roll_shift                  | Shift_slide                   | Shift1                         |
| Tilt4                       | Tilt2                         | Free energy8                   |
| Twist (DNA-protein          | Tilt_rise                     | Free energy2                   |
|                             |                               |                                |

|                            |                      | 20                                    |
|----------------------------|----------------------|---------------------------------------|
| complex)1                  |                      |                                       |
| Stacking energy2           | Stacking energy3     | Rise_rise                             |
| Tilt_tilt                  | Roll4                | Tilt_roll                             |
| Minor Groove Size          | GC content           | Inclination                           |
| Slide stiffness            | Melting Temperature1 | Twist3                                |
| Tilt (DNA-protein complex) | Guanine content      | Twist6                                |
| Major Groove Size          | Twist_rise           | Rise2                                 |
| Melting Temperature        | Free energy          | Mobility to bend towards major groove |
| Bend                       |                      |                                       |

**Table 5.** The names of the 12 physicochemical indices for trinucleotides.

| Bendability (DNAse)    | Bendability (consensus) | Trinucleotide GC Content |
|------------------------|-------------------------|--------------------------|
| Consensus_roll         | Consensus-Rigid         | Dnase I                  |
| MW-Daltons             | MW-kg                   | Nucleosome               |
| Nucleosome positioning | Dnase I-Rigid           | Nucleosome-Rigid         |

**Table 6.** The names of the 90 physicochemical indices for dinucleotides.

| Base stacking         | Protein induced deformability   | B-DNA twist                     |
|-----------------------|---------------------------------|---------------------------------|
| Dinucleotide GC       | A-philicity                     | Propeller twist                 |
| Content               | 1 2                             | 1                               |
| Duplex stability-free | Duplex stability-disrupt energy | DNA denaturation                |
| energy                | 1 7 1 27                        |                                 |
| Bending stiffness     | Protein DNA twist               | Stabilising energy of Z-<br>DNA |
| Aida_BA_transition    | Breslauer_dG                    | Breslauer_dH                    |
| Breslauer_dS          | Electron_interaction            | Hartman_trans_free_ener         |
|                       |                                 | gy                              |
| Helix-                | Ivanov_BA_transition            | Lisser_BZ_transition            |
| Coil_transition       |                                 |                                 |
| Polar_interaction     | SantaLucia_dG                   | SantaLucia_dH                   |
| SantaLucia_dS         | Sarai_flexibility               | Stability                       |
| Stacking_energy       | Sugimoto_dG                     | Sugimoto_dH                     |
| Sugimoto_dS           | Watson-Crick_interaction        | Twist                           |
| Tilt                  | Roll                            | Shift                           |
| Slide                 | Rise                            | Stacking energy                 |
| Bend                  | Tip                             | Inclination                     |
| Major Groove          | Major Groove Depth              | Major Groove Size               |
| Width                 |                                 |                                 |
| Major Groove          | Minor Groove Width              | Minor Groove Depth              |
| Distance              |                                 |                                 |
| Minor Groove Size     | Minor Groove Distance           | Persistance Length              |
| Melting               | Mobility to bend towards major  | Mobility to bend towards        |
| Temperature           | groove                          | minor groove                    |
| Propeller Twist       | Clash Strength                  | Enthalpy                        |
| Free energy           | Twist_twist                     | Tilt_tilt                       |
| Roll_roll             | Twist_tilt                      | Twist_roll                      |
| Tilt_roll             | Shift_shift                     | Slide_slide                     |
| Rise_rise             | Shift_slide                     | Shift_rise                      |
| Slide_rise            | Twist_shift                     | Twist_slide                     |
| Twist_rise            | Tilt_shift                      | Tilt_slide                      |
| Tilt_rise             | Roll_shift                      | Roll_slide                      |
|                       |                                 |                                 |

| Roll_rise         | Slide stiffness   | Shift stiffness |
|-------------------|-------------------|-----------------|
| Roll stiffness    | Rise stiffness    | Tilt stiffness  |
| Twist stiffness   | Wedge             | Direction       |
| Flexibility_slide | Flexibility_shift | Entropy         |

**Table 7.** The names of the 6 physicochemical indices for dinucleotides.

| Twist | Tilt  | Roll |  |
|-------|-------|------|--|
| Shift | Slide | Rise |  |

**Table 8.** The names of the 22 physicochemical indices for dinucleotides.

| Shift (RNA)          | Hydrophilicity (RNA)  |
|----------------------|-----------------------|
| Hydrophilicity (RNA) | GC content            |
| Purine (AG) content  | Keto (GT) content     |
| Adenine content      | Guanine content       |
| Cytosine content     | Thymine content       |
| Slide (RNA)          | Rise (RNA)            |
| Tilt (RNA)           | Roll (RNA)            |
| Twist (RNA)          | Stacking energy (RNA) |
| Enthalpy (RNA)       | Entropy (RNA)         |
| Free energy (RNA)    | Free energy (RNA)     |
| Enthalpy (RNA)       | Entropy (RNA)         |

**Table 9.** The names of the 11 physicochemical indices for dinucleotides.

| Shift           | Slide          | Rise    |
|-----------------|----------------|---------|
| Tilt            | Roll           | Twist   |
| Stacking energy | Enthalpy       | Entropy |
| Free energy     | Hydrophilicity |         |

**Table 10.** The names of the 547 physicochemical indices for amino acids.

| Hydrophobicity | Hydrophilicity | Mass       |
|----------------|----------------|------------|
| ARGP820102     | ARGP820103     | BEGF750101 |
| BHAR880101     | BIGC670101     | BIOV880101 |
| BROC820102     | BULH740101     | BULH740102 |
| BUNA790103     | BURA740101     | BURA740102 |
| CHAM820102     | CHAM830101     | CHAM830102 |
| CHAM830105     | CHAM830106     | CHAM830107 |
| CHOC760101     | CHOC760102     | CHOC760103 |
| CHOP780201     | CHOP780202     | CHOP780203 |
| CHOP780206     | CHOP780207     | CHOP780208 |
| CHOP780211     | CHOP780212     | CHOP780213 |
| CHOP780216     | CIDH920101     | CIDH920102 |
| CIDH920105     | СОНЕ430101     | CRAJ730101 |
| DAWD720101     | DAYM780101     | DAYM780201 |
| EISD840101     | EISD860101     | EISD860102 |
| FASG760102     | FASG760103     | FASG760104 |
| FAUJ880101     | FAUJ880102     | FAUJ880103 |
| FAUJ880106     | FAUJ880107     | FAUJ880108 |
| FAUJ880111     | FAUJ880112     | FAUJ880113 |
| FINA910102     | FINA910103     | FINA910104 |
| GEIM800102     | GEIM800103     | GEIM800104 |
| GEIM800107     | GEIM800108     | GEIM800109 |
| GOLD730101     | GOLD730102     | GRAR740101 |
|                |                |            |

|            |            | 22                       |
|------------|------------|--------------------------|
| GUYH850101 | HOPA770101 | HOPT810101               |
| HUTJ700103 | ISOY800101 | ISOY800102               |
| ISOY800105 | ISOY800106 | ISOY800107               |
| JANJ780102 | JANJ780103 | JANJ790101               |
| JOND750102 | JOND920101 | JOND920102               |
| KANM800101 | KANM800102 | KANM800103               |
| KARP850102 | KARP850103 | KHAG800101               |
| KRIW790101 | KRIW790102 | KRIW790103               |
| LEVM760101 | LEVM760102 | LEVM760103               |
| LEVM760106 | LEVM760107 | LEVM780101               |
| LEVM780104 | LEVM780105 | LEVM780106               |
| LIFS790102 | LIFS790103 | MANP780101               |
| MAXF760103 | MAXF760104 | MAXF760105               |
| MEEJ800101 | MEEJ800102 | MEEJ810101               |
| MEIH800102 | MEIH800103 | MIYS850101               |
| NAGK730103 | NAKH900101 | NAKH900102               |
| NAKH900105 | NAKH900106 | NAKH900107               |
| NAKH900110 | NAKH900111 | NAKH900112               |
| NAKH920102 | NAKH920103 | NAKH920104               |
| NAKH920107 | NAKH920108 | NISK800101               |
| OOBM770101 | OOBM770102 | OOBM770103               |
| OOBM850101 | OOBM850102 | OOBM850103               |
| PALJ810101 | PALJ810102 | PALJ810103               |
| PALJ810106 | PALJ810107 | PALJ810108               |
| PALJ810111 | PALJ810112 | PALJ810113               |
| PALJ810116 | PARJ860101 | PLIV810101               |
| PONP800103 | PONP800104 | PONP800105               |
| PONP800108 | PRAM820101 | PRAM820102               |
| PRAM900102 | PRAM900103 | PRAM900104               |
| QIAN880101 | QIAN880102 | QIAN880103               |
| QIAN880106 | QIAN880107 | QIAN880108               |
| QIAN880111 | QIAN880112 | QIAN880113               |
| QIAN880116 | QIAN880117 | QIAN880118               |
| QIAN880121 | QIAN880122 | QIAN880123               |
| QIAN880126 | QIAN880127 | QIAN880128               |
| QIAN880131 | QIAN880132 | QIAN880133               |
| QIAN880136 | QIAN880137 | QIAN880138               |
| RACS770102 | RACS770103 | RACS820101               |
| RACS820104 | RACS820105 | RACS820106               |
| RACS820109 | RACS820110 | RACS820111               |
| RACS820107 | RADA880101 | RAC5820111<br>RADA880102 |
| RADA880105 | RADA880106 | RADA880102               |
| RICJ880102 | RICJ880103 | RICJ880104               |
| RICJ880102 | RICJ880103 | RICJ880104<br>RICJ880109 |
| RICJ880112 | RICJ880113 | RICJ880114               |
| RICJ880117 | ROBB760101 | ROBB760102               |
| ROBB760105 | ROBB760101 | ROBB760102<br>ROBB760107 |
|            |            |                          |
| ROBB760110 | ROBB760111 | ROBB760112               |
| ROSG850101 | ROSG850102 | ROSM880101               |
| SIMZ760101 | SNEP660101 | SNEP660102               |
| SUEM840101 | SUEM840102 | SWER830101               |
| TANS770103 | TANS770104 | TANS770105               |
| TANS770108 | TANS770109 | TANS770110               |
| VASM830103 | VELV850101 | VENT840101               |

|            |                          | 23           |
|------------|--------------------------|--------------|
| WEBA780101 | WERD780101               | WERD780102   |
| WOEC730101 | WOLR810101               | WOLS870101   |
| YUTK870101 | YUTK870102               | YUTK870103   |
| ZIMJ680101 | ZIMJ680102               | ZIMJ680103   |
| AURR980101 | AURR980102               | AURR980103   |
| AURR980106 | AURR980107               | AURR980108   |
| AURR980111 | AURR980112               | AURR980113   |
| AURR980116 | AURR980117               | AURR980118   |
| ONEK900101 | ONEK900102               | VINM940101   |
| VINM940104 | MUNV940101               | MUNV940102   |
| MUNV940105 | WIMW960101               | KIMC930101   |
| PARS000101 | PARS000102               | KUMS000101   |
| KUMS000104 | TAKK010101               | FODM020101   |
| NADH010103 | NADH010104               | NADH010105   |
| MONM990201 | KOEP990101               | KOEP990102   |
| CEDJ970103 | CEDJ970104               | CEDJ970105   |
| FUKS010103 | FUKS010104               | FUKS010105   |
| FUKS010108 | FUKS010109               | FUKS010110   |
| AVBF000101 | AVBF000102               | AVBF000103   |
| AVBF000106 | AVBF000107               | AVBF000108   |
| MITS020101 | TSAJ990101               | TSAJ990102   |
| WILM950101 | WILM950102               | WILM950103   |
| GUOD860101 | JURD980101               | BASU050101   |
| SUYM030101 | PUNT030101               | PUNT030102   |
| GEOR030103 | GEOR030104               | GEOR030105   |
| GEOR030108 | GEOR030104<br>GEOR030109 | ZHOH040101   |
| BAEK050101 | HARY940101               | PONJ960101   |
| OLSK800101 | KIDA850101               | GUYH850102   |
| GUYH850105 | ROSM880104               | ROSM880105   |
| BLAS910101 | CASG920101               | CORJ870101   |
| CORJ870104 | CORJ870105               | CORJ870106   |
| MIYS990101 | MIYS990102               | MIYS990103   |
| ENGD860101 | FASG890101               | TANS770101   |
| ANDN920101 | ARGP820101               | TANS770106   |
| BEGF750102 | BEGF750103               | VASM830101   |
| BIOV880102 | BROC820101               | VHEG790101   |
| BUNA790101 | BUNA790102               | WERD780103   |
| CHAM810101 | CHAM820101               | WOLS870102   |
| CHAM830103 | CHAM830104               | YUTK870104   |
| CHAM830108 | CHOC750101               | ZIMJ680104   |
| CHOC760104 | CHOP780101               | AURR980104   |
| CHOP780204 | CHOP780205               | AURR980109   |
| CHOP780209 | CHOP780210               | AURR980114   |
| CHOP780214 | CHOP780215               | AURR980119   |
| CIDH920103 | CIDH920104               | VINM940102   |
| CRAJ730102 | CRAJ730103               | MUNV940103   |
| DESM900101 | DESM900102               | MONM990101   |
| EISD860103 | FASG760101               | KUMS000102   |
| FASG760105 | FAUJ830101               | NADH010101   |
| FAUJ880104 | FAUJ880105               | NADH010101   |
| FAUJ880109 | FAUJ880110               | CEDJ970101   |
| FINA770101 | FINA910101               | FUKS010101   |
| GARJ730101 | GEIM800101               | FUKS010106   |
| GEIM800105 | GEIM800106               | FUKS010111   |
| OLIMOU103  | OLIMOUTUU                | 1 0120010111 |

|            |            | 24         |
|------------|------------|------------|
| GEIM800110 | GEIM800111 | AVBF000104 |
| GRAR740102 | GRAR740103 | AVBF000109 |
| HUTJ700101 | HUTJ700102 | COSI940101 |
| ISOY800103 | ISOY800104 | WILM950104 |
| ISOY800108 | JANJ780101 | BASU050102 |
| JANJ790102 | JOND750101 | GEOR030101 |
| JUKT750101 | JUNJ780101 | GEOR030106 |
| KANM800104 | KARP850101 | ZHOH040102 |
| KLEP840101 | KRIW710101 | DIGM050101 |
| KYTJ820101 | LAWE840101 | GUYH850103 |
| LEVM760104 | LEVM760105 | JACR890101 |
| LEVM780102 | LEVM780103 | CORJ870102 |
| LEWP710101 | LIFS790101 | CORJ870107 |
| MAXF760101 | MAXF760102 | MIYS990104 |
| MAXF760106 | MCMT640101 | TANS770102 |
| MEEJ810102 | MEIH800101 | TANS770107 |
| NAGK730101 | NAGK730102 | VASM830102 |
| NAKH900103 | NAKH900104 | WARP780101 |
| NAKH900108 | NAKH900109 | WERD780104 |
| NAKH900113 | NAKH920101 | WOLS870103 |
| NAKH920105 | NAKH920106 | ZASB820101 |
| NISK860101 | NOZY710101 | ZIMJ680105 |
| OOBM770104 | OOBM770105 | AURR980105 |
| OOBM850104 | OOBM850105 | AURR980110 |
| PALJ810104 | PALJ810105 | AURR980115 |
| PALJ810109 | PALJ810110 | AURR980120 |
| PALJ810114 | PALJ810115 | VINM940103 |
| PONP800101 | PONP800102 | MUNV940104 |
| PONP800106 | PONP800107 | BLAM930101 |
| PRAM820103 | PRAM900101 | KUMS000103 |
| PTIO830101 | PTIO830102 | NADH010102 |
| QIAN880104 | QIAN880105 | NADH010107 |
| QIAN880109 | QIAN880110 | CEDJ970102 |
| QIAN880114 | QIAN880115 | FUKS010102 |
| QIAN880119 | QIAN880120 | FUKS010107 |
| QIAN880124 | QIAN880125 | FUKS010112 |
| QIAN880129 | QIAN880130 | AVBF000105 |
| QIAN880134 | QIAN880135 | YANJ020101 |
| QIAN880139 | RACS770101 | PONP930101 |
| RACS820102 | RACS820103 | KUHL950101 |
| RACS820107 | RACS820108 | BASU050103 |
| RACS820112 | RACS820113 | GEOR030102 |
| RADA880103 | RADA880104 | GEOR030107 |
| RADA880108 | RICJ880101 | ZHOH040103 |
| RICJ880105 | RICJ880106 | WOLR790101 |
| RICJ880110 | RICJ880111 | GUYH850104 |
| RICJ880115 | RICJ880116 | COWR900101 |
| ROBB760103 | ROBB760104 | CORJ870103 |
| ROBB760108 | ROBB760109 | CORJ870108 |
| ROBB760113 | ROBB790101 | MIYS990105 |
| ROSM880102 | ROSM880103 | SNEP660104 |
| SNEP660103 |            |            |

**Table 11.** The names of the 3 physicochemical indices for amino acids.

| Hydrophobicity | hydrophilicity | mass |
|----------------|----------------|------|
|----------------|----------------|------|

**Table 12.** The names of the 2 physicochemical indices for amino acids.

| Hydrophobicity | hydrophilicity |  |
|----------------|----------------|--|

#### References

- 1. Liu, B., Liu, F., Wang, X., Chen, J., Fang, L. and Chou, K.-C. (2015) Pse-in-One: a web server for generating various modes of pseudo components of DNA, RNA, and protein sequences. Nucleic acids research, 43, W65-W71.
- 2. Van Der Walt, S., Colbert, S.C. and Varoquaux, G. (2011) The NumPy array: a structure for efficient numerical computation. Computing in Science & Engineering, 13, 22-30.
- 3. Chang, C.C. and Lin, C.J. (2011) LIBSVM: A Library for Support Vector Machines. Acm T Intel Syst Tec, 2, 1-27.
- 4. Liu, B., Xu, J., Fan, S., Xu, R., Zhou, J. and Wang, X. (2015) PseDNA-Pro: DNA-Binding Protein Identification by Combining Chou's PseAAC and Physicochemical Distance Transformation. Molecular Informatics, 34, 8-17.
- 5. Lou, W., Wang, X., Chen, F., Chen, Y., Jiang, B. and Zhang, H. (2014) Sequence based prediction of DNA-binding proteins based on hybrid feature selection using random forest and Gaussian naive Bayes. PLoS One, 9, e86703.
- 6. Fawcett, T. (2006) An introduction to ROC analysis. Pattern recognition letters, 27, 861-874.
- 7. Lee, D., Karchin, R. and Beer, M.A. (2011) Discriminative prediction of mammalian enhancers from DNA sequence. Genome research, 21, 2167-2180.
- 8. Gupta, S., Dennis, J., Thurman, R.E., Kingston, R., Stamatoyannopoulos, J.A. and Noble, W.S. (2008) Predicting human nucleosome occupancy from primary sequence. PLoS computational biology, 4, e1000134.
- 9. Noble, W.S., Kuehn, S., Thurman, R., Yu, M. and Stamatoyannopoulos, J. (2005) Predicting the in vivo signature of human gene regulatory sequences. Bioinformatics, 21 Suppl 1, i338-343.
- 10. Chen, W., Luo, L. and Zhang, L. (2010) The organization of nucleosomes around splice sites. Nucleic acids research, 38, 2788-2798.
- 11. Liu, G., Liu, J., Cui, X. and Cai, L. (2012) Sequence-dependent prediction of recombination hotspots in Saccharomyces cerevisiae. Journal of theoretical biology, 293, 49-54.
- 12. Liu, B., Liu, F., Fang, L., Wang, X. and Chou, K.-C. (2015) repDNA: a Python package to generate various modes of feature vectors for DNA sequences by incorporating user-defined physicochemical properties and sequence-order effects. Bioinformatics, 31, 1307-1309.
- 13. El-Manzalawy, Y., Dobbs, D. and Honavar, V. (2008) Predicting flexible length linear B-cell epitopes. Computational Systems Bioinformatics, 7, 121-132.
- 14. Leslie, C.S., Eskin, E., Cohen, A., Weston, J. and Noble, W.S. (2004) Mismatch

- string kernels for discriminative protein classification. Bioinformatics, 20, 467-476.
- 15. Luo, L., Li, D., Zhang, W., Tu, S., Zhu, X. and Tian, G. (2016) Accurate Prediction of Transposon-Derived piRNAs by Integrating Various Sequential and Physicochemical Features. PLoS ONE, 11, e0153268.
- 16. Lodhi, H., Saunders, C., Shawe-Taylor, J., Cristianini, N. and Watkins, C. (2002) Text classification using string kernels. Journal of Machine Learning Research, 2, 419-444.
- 17. Dong, Q., Zhou, S. and Guan, J. (2009) A new taxonomy-based protein fold recognition approach based on autocross-covariance transformation. Bioinformatics, 25, 2655-2662.
- 18. Friedel, M., Nikolajewa, S., Sühnel, J. and Wilhelm, T. (2009) DiProDB: a database for dinucleotide properties. Nucleic acids research, 37, D37-D40.
- 19. Horne, D.S. (1988) Prediction of protein helix content from an autocorrelation analysis of sequence hydrophobicities. Biopolymers, 27, 451-477.
- 20. Chen, W., Zhang, X., Brooker, J., Lin, H., Zhang, L. and Chou, K.-C. (2015b) PseKNC-General: a cross-platform package for generating various modes of pseudo nucleotide compositions. Bioinformatics, 31, 119-120.
- 21. Sokal, R.R. and Thomson, B.A. (2006) Population structure inferred by local spatial autocorrelation: an example from an Amerindian tribal population. American journal of physical anthropology, 129, 121-131.
- 22. 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.
- 23. Chen, W., Feng, P.M., Lin, H. and Chou, K.C. (2013) iRSpot-PseDNC: identify recombination spots with pseudo dinucleotide composition. Nucleic Acids Res, 41, e68.
- 24. Guo, S.-H., Deng, E.-Z., Xu, L.-Q., Ding, H., Lin, H., Chen, W. and Chou, K.-C. (2014) iNuc-PseKNC: a sequence-based predictor for predicting nucleosome positioning in genomes with pseudo k-tuple nucleotide composition. Bioinformatics, btu083.
- 25. Lin, H., Deng, E.-Z., Ding, H., Chen, W. and Chou, K.-C. (2014) iPro54-PseKNC: a sequence-based predictor for identifying sigma-54 promoters in prokaryote with pseudo k-tuple nucleotide composition. Nucleic acids research, 42, 12961-12972.
- 26. Liu, B., Zhang, D., Xu, R., Xu, J., Wang, X., Chen, Q., Dong, Q. and Chou, K.-C. (2014) Combining evolutionary information extracted from frequency profiles with sequence-based kernels for protein remote homology detection. Bioinformatics, 30, 472-479.
- 27. Wei, L., Liao, M., Gao, Y., Ji, R., He, Z. and Zou, Q. (2014) Improved and promising identification of human microRNAs by incorporating a high-quality negative set. IEEE/ACM Transactions on Computational Biology and Bioinformatics, 11, 192-201.
- 28. Guo, Y., Yu, L., Wen, Z. and Li, M. (2008) Using support vector machine combined with auto covariance to predict protein—protein interactions from protein sequences. Nucleic acids research, 36, 3025-3030.
- 29. Xue, C., Li, F., He, T., Liu, G.-P., Li, Y. and Zhang, X. (2005) Classification of real and pseudo microRNA precursors using local structure-sequence features and support vector machine. BMC bioinformatics, 6, 1.
- 30. Liu, B., Fang, L., Liu, F., Wang, X., Chen, J. and Chou, K.-C. (2015) Identification of real microRNA precursors with a pseudo structure status composition approach. PloS one, 10, e0121501.
- 31. Liu, B., Fang, L., Liu, F., Wang, X. and Chou, K.-C. (2016) iMiRNA-PseDPC: microRNA precursor identification with a pseudo distance-pair composition approach. Journal of Biomolecular Structure and Dynamics, 34, 223-235.
- 32. Liu, B., Wang, X., Lin, L., Dong, Q. and Wang, X. (2008) A discriminative method for protein remote homology detection and fold recognition combining Top-ngrams and latent semantic analysis. BMC bioinformatics, 9, 1.
- 33. Liu, B., Xu, J., Zou, Q., Xu, R., Wang, X. and Chen, Q. (2014) Using distances

- between Top-n-gram and residue pairs for protein remote homology detection. Bmc Bioinformatics, 15, 1.
- 34. Liu, B., Xu, J., Lan, X., Xu, R., Zhou, J., Wang, X. and Chou, K.-C. (2014) iDNA-Prot| dis: identifying DNA-binding proteins by incorporating amino acid distance-pairs and reduced alphabet profile into the general pseudo amino acid composition. PloS one, 9, e106691.
- 35. Liu, B., Wang, X., Chen, Q., Dong, Q. and Lan, X. (2012) Using amino acid physicochemical distance transformation for fast protein remote homology detection. PLoS One, 7, e46633.
- 36. Chou, K.C. (2001) Prediction of protein cellular attributes using pseudo-amino acid composition. Proteins: Structure, Function, and Bioinformatics, 43, 246-255.
- 37. Chou, K.-C. (2005) Using amphiphilic pseudo amino acid composition to predict enzyme subfamily classes. Bioinformatics, 21, 10-19.
- 38. Kawashima, S., Pokarowski, P., Pokarowska, M., Kolinski, A., Katayama, T. and Kanehisa, M. (2008) AAindex: amino acid index database, progress report 2008. Nucleic acids research, 36, D202-D205.