Using the BioSeqClass Package

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1 Overview

There are now 863 completely sequenced genomes of cellular organisms in NCBI genome database. Nevertheless, functional annotation drops far behind sequencing because functional validation experiments are time-consuming and costly. Taken model organism Homo

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sapiens, Mus musculus and Saccharomyces cere-visiae as examples, only 16and 18annotations in Gene Ontology), respectively. Thus computational methods for predicting function is still a fun-damental complement. The most common com-putation approach is biological sequence based classification, since sequence information is still the most abundant and reliable. Se-quence based classification has been used in: discovering new microRNA candidates, predicting transcription factor binding sites, detecting protein post-translational modification sites, and so on.

Features and models are two basic factors for classification. Features generally are numerical values that can be used to distinguish different classes. Therefore it is preferable to select features that can achieve better and faster classification. Classification models are built from features by various algorithms, and it is necessary to evaluate its prediction ability by cross validation or jackknife test. For biological sequences, there are additional steps: one is to reduce homolog sequences which might result in overestimation of prediction accuracy, and then another most important step is to convert sequences into numerical features. Thus, the general workflow for sequence-based classifications includes (Figure 1): reduce homolog sequences; extract features from sequences and code them to numerical values; evaluate and select features; build classification model and evaluate its performance.

Here we present an R package (BioSeqClass) to carry out the general workflow for biological sequence based classification. It contains diverse fearure coding schemas for RNA, DNA and proteins, supports feature seletion, and integrates multiple classification methods.

2 Installation

2.1 Requirements

BioSeqClass employs some external programs to extract biological properties and use other R packages to build classification model:

- 1. BioSeqClass imported R packages are listed in table 1. These packages will be automatically installed when Biocalss is firstly loaded.
- 2. External programs are used to assist the performance of BioSeqClass (see table 2). Some programs are invoked via their web service, and some ones are needed to be installed at the local computer.

Note: You do not need to install programs listed in table 2, unless you will use the related function in BioSeqClass.

2.2 Installation

The biocLite script is used to install PAnnBuilder from within R:

- > source("http://bioconductor.org/biocLite.R")
- > biocLite("BioSeqClass")

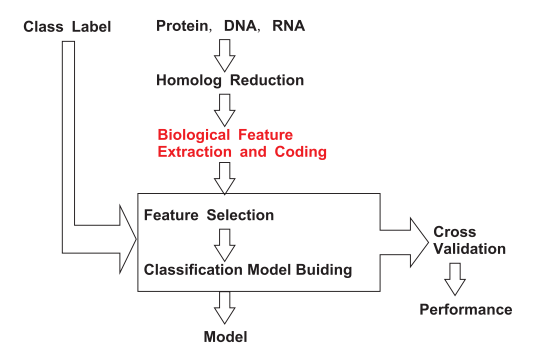


Figure 1: Workflow for Biological Sequence based Classification.

Users also can use the installation script "BioSeqClass.R" to download and install BioSeqClass package.

- > source("http://www.biosino.org/download/BioSeqClass/BioSeqClass.R")
- > BioSeqClass()

Load package:

> library(BioSeqClass)

Note: Web Connection is needed to install BioSeqClass and its required packages. All the codes in this vignette were tested in R 2.8.0 and 2.9.0, thus the latest R version is recommended.

3 Function description

3.1 Homolog Reduction

Homologous sequences in training/testing data may lead to overestimation of prediction accuracy. Therefore, the first step for sequence based classification and prediction is homolog reduction based on sequence similarity. Taking computation complexity and similarity restriction into consideration, homolog reductions for full-length sequences and fragment sequences are different. We have designed different functions to deal with them, respectively (see table 3).

- hr It employs cdhitHR and aligndisHR to filter homolog sequences by sequence similarity. cdhitHR is designed to filter full-length protein or gene sequences. aligndisHR is designed for aligned sequences with equal length.
- cdhithrall It uses cd-hit program to do homolog reduction ("formatdb" and "blastall" are required for running cd-hit program). CD-HIT is a program for clustering large protein database at high sequence identity threshold (Li and Godzik, 2006).
- aligndisHR It uses the number of different residues to do homolog reduction (D et al., 2008). The algorithm proceeds in a stepwise manner by first eliminating sequences that were different from another in exactly 1 position. Elimination proceeds one peptide at a time; Re-evaluate after each peptide is removed. Once no further homologs of distance 1 remain, homologs of distance 2 are eliminated, and so forth until identity between all peptides are less than given cutoff.

3.2 Feature Extraction and Numerical Coding

3.2.1 Biological sequences

RNA, DNA and protein are three kinds of basic biological sequences. RNA and DNA are composed of bases, while proteins are composed of amino acids. Furthermore, amino acids

have different physical-chemical properties, which were used to divide amino acids into different groups. These elements and groups are basic objects for feature extraction (see table 4).

3.2.2 Feature Coding

Feature coding means to extract features from sequences and convert them into numerical values. The frequently used features are the basic elements of sequence (bases, amino acids), physical-chemical properties, secondary structures, and so on. There are many methods to convert features to numerical values. The simplest is the composition of element. But more sophisticated conversions are preferrable for achieving better distinguishing power. Previous studies have shown that feature coding is the key point for the accuracy of classification and prediction. Here we have summarized various feature coding methods used in published papers and carried out them in BioSeqClass (see table 5). These functions will allow more diverse choices of coding strategies and accelerate the feature coding process. We also provide a function featureEvaluate to test the performance of models with different feature coding schemes and different classification algorithms.

3.3 Feature Selection

Features are important for the accuracy of prediction model. However, it does not mean that the more the better. Computation time is usually increased with the increase of number of features. Conflictive features would even reduce the accuracy. Therefore, suitable feature selection is needed for better prediction performance and less computation cost. We provided two functions for feature selection (see table 6).

3.4 Model Building and Performance Evaluation

Besides features, classification method is another factor that influences classification. Different cases may have different perference over classification methods. Multiple classification methods are integrated and available in BioSeqClass (see table 7). To evaluate and compare classification models, performance assessment is done for each model, including precision, sensitivity, specificity, accuracy, and matthews correlation coefficient.

Table 1: Imported R packages.

Existing R Package	Functions used by BioSeqClass
Biostrings	readAAStringSet, writeXStringSet
e1071	svm
ipred	bagging
klaR	svmlight, NaiveBayes
random Forest	randomForest
class	knn
tree	tree
nnet	nnet
rpart	rpart
party	ctree
foreign	write.arff
Biobase	addVigs2WinMenu

4 Examples

To illustrate the use of BioSeqClass, lysine acetylation site prediction is taken as an example.

- 1. Suppose the original data are protein FASTA sequences and lysine acetylation sites. You can use getTrain to extract the flanking peptides of acetylation sites as positive dataset, and filter these peptides based on sequence identity. Lysine without acetylation annotation are regarded as negative dataset, and are filtered like the positive dataset. Considering the computational time, only 20 positive data are used as examples in the following codes.

 - > # Only a part of lysine acetylation sites are used for demo.
 - > posfile1=tempfile()
 - > write.table(read.table(posfile,sep='\t',header=F)[1:20,], posfile1,
 - + sep='\t', quote=F, row.names=F, col.names=F)
 - > seqList = getTrain(file, posfile1, aa="K", w=7, identity=0.4)
 - [1] "Positive Site: 18"
 - [1] "Positive Protein: 9"
 - [1] "Positive Site After Homolog Reduction: 16"
 - [1] "Positive Protein After Homolog Reduction: 9"

Table 2: Invoked External Programs.

External	Description	Related BioSeqClass	Need In-	Ref
Program		Function	stalled?	
cd-hit	a program for clustering large	cdhitHR	Yes	(Li and Godzik, 2006)
	protein database at high se-			
	quence identity threshold			
blastpgp	PSI-BLAST (Position-Specific	featurePSSM	Yes	(Altschul et al., 1997)
	Iterated BLAST) for capturing			
	the conservation pattern			
SVMlight	support vector machine	classifyModelSVMLIGHT	No	(T, 1999)
DSSP	a database of secondary struc-	getDSSP	No	(Kabsch and Sander, 1983)
	ture assignments for protein en-			
	tries in the Protein Data Bank			
	(PDB)			
Proteus2	predict secondary structure	predictPROTEUS	No	(S et al., 2006)
HMMER	predict domains with hmmpfam	predictPFAM	No	(Eddy, 1998)
	using models of Pfam database			

Table 3: Summary Table for Homolog Reduction Functions.

Function	Description	Ref
hr	employ cdhitHR and aligndisHR to do homolog reduction	
cdhitHR	invoke cd-hit to cluster sequences	(Li and Godzik, 2006)
aligndisHR	calculated identity of aligned sequences	(D et al., 2008)

Table 4: Summary Table for Base and Amino Acid Groups.

Function	Description	Ref
elements	basic elements of biological sequence	
aaClass	amino acids groups depend on their physical-chemical properties:	(CS et al., 2006)
	hydrophobicity, normalized Van der Waals volume, polarizability,	
	polarity, and so on	

Table 5: Summary table for Feature Coding Functions.

Type	Function	Feature Coding Scheme	Ref
	featureBinary	use 0-1 vector to code each element	(Jia et al., 2006; Chen
DNA,			et al., 2006)
RNA, or	featureCTD	numeric vector for the composition, transition	(Cai et al., 2003)
protein		and distribution of properties	
	featureFragmentComposition	numeric vector for the frequency of k-mer se-	(CS et al., 2004; RY
		quence fragment	et al., 2002)
	featureGapPairComposition	numeric vector for the frequency of g-spaced ele-	(CS et al., 2006)
		ment pair	
	featureCKSAAP	integer vector for the number of k-spaced element	(Chen et al., 2008)
		pair (k cycled from 0 to g)	
	featureHydro	hydrophobic effect	(A and R, 1988; MJ
			et al., 2000)
	featureACH	average cumulative hydrophobicity over a sliding	(Zhang et al., 2008;
protein		window	Kurgan et al., 2007)
protein	featureAAindex	numeric vector measuring the physicochemical	(S et al., 1999; Cai
		and biochemical properties based on AAindex	and Lu, 2008)
		database	
	featureACI	numeric vector measuring the average cumulative	
		properties in AAindex	/771
	featureACF	numeric vector measuring the Auto Correlation	(Zhang et al., 1998;
	C	Function (ACF) of properties in AAindex	Liu and Chou, 1998)
	featurePseudoAAComp	numeric vector for the pseudo amino acid com-	(Chou, 2001)
	f t DGGM	position proposed by Chou,K.C.	(71
	featurePSSM	numeric vector for the normalized position-	(Zhang et al., 2005;
		specific score of PSSM generated by PSI-BLAST	Chung-Tsai Su and
			Ou, 2006; S and A, 2005; Zhang et al.,
			2008; Zhang et al.,
	featureDOMAIN	vector for the number of domain. Domains can	(Jia et al., 2007)
	TeatureDoMAIN	be obtained by 'predictPFAM' function.	(31a Ct al., 2001)
	featureSSC	coding for secondary structure of protein. Sec-	(Zheng and Kurgan,
	10dbd1cbb0	ondary structure can be got by 'predictPRO-	2008)
		TEUS' or 'getDSSP'.	2000)
D. 1. 1	featureBDNAVIDEO	Conformational and physicochemical DNA fea-	(v. Ponomarenko
DNA or		tures from B-DNA-VIDEO database	et al., 1999)
RNA	featureDIPRODB	conformational and thermodynamic dinu-	(Friedel et al., 2009)
		cleotide properties from DiProDB database	
		(http://diprodb.fli-leibniz.de)	

Table 6: Summary Table for Feature Selection Functions.

Function	Description	Ref
selectWeka	feature selction by methods in WEKA	(Witten and Fran, 2005)
selectFFS	feature forward selction based on the performance of	(Cai and Lu, 2008)
	classification model	
classify	build and test model with cross validation, also support	
	feature selection by envoking WEKA	

Table 7: Summary Table for Classification Methods.

Function	Description	Depended R package
classifyModelLIBSVM	support vector machine by LIBSVM	e1071
classifyModelSVMLIGHT	support vector machine by SVM-light	klaR
classifyModelNB	naive bayes	klaR
classifyModelRF	random forest	random Forest
classifyModelKNN	k-nearest neighbor	class
classifyModelTree	tree model	tree
classifyModelNNET	neural net algorithm	VR
classifyModelRPART	recursive partitioning trees	rpart
classifyModelCTREE	conditional inference trees	party
classifyModelCTREELIBSVM	combine conditional inference trees and sup-	party, e1071
	port vector machine	
classifyModelBAG	bagging method	ipred

```
[1] "Negative Site: 215"
```

- [1] "Negative Protein: 9"
- [1] "Negative Site After Homolog Reduction and Random Selection: 16"
- [1] "Negative Protein After Homolog Reduction and Random Selection: 8"
- 2. If the original data are non-redundant positive/negative peptides. We directly read the data into R and assign class labels for them.

```
> tmpDir=file.path(path.package('BioSeqClass'), 'example')
> tmpFile1=file.path(tmpDir, 'acetylation_K.pos40.pep')
> tmpFile2=file.path(tmpDir, 'acetylation_K.neg40.pep')
> posSeq=as.matrix(read.csv(tmpFile1,header=F,sep='\t',row.names=1))[,1]
> negSeq=as.matrix(read.csv(tmpFile2,header=F,sep='\t',row.names=1))[,1]
> seq=c(posSeq,negSeq)
> classLable=c(rep("+1",length(posSeq)),rep("-1",length(negSeq)))
> length(seq)
[1] 1200
```

- 3. Once you have positive/negative datasets, you can code them to numeric vectors by functions listed in table 5. Function featureBinary and featureGapPairComposition are taken as examples of different coding methods, which use binary 0-1 coding and the composition of gapped amino acid pair, respectively. Other functions can be used in the same way.
 - > # Use 0-1 binary coding.
 > feature1 = featureBinary(seq,elements("aminoacid"))
 > dim(feature1)
 [1] 1200 300
 - > # Use the compostion of paired amino acids.
 > feature2 = featureGapPairComposition(seq,0,elements("aminoacid"))
 - > feature2 = featureGapPairComposition(seq,0,elements("aminoacid"),
 > dim(feature2)
 - [1] 1200 400
- 4. classify is used to build classification model under cross validation. It also supports feature selection by invoking WEKA. Models built with selected features usually can obtain higher accuracy. In the following codes, two models are built by classify. The 1st classification model 'LIBSVM_CV5' is built by support vector machine with linear kernel and get an accuracy of 0.56 under 5-fold cross validation. The 2nd classification model 'FS_LIBSVM_CV5' is also built by support vector machine with linear kernel, but a feature selection method called "CfsSubsetEval" is used before building model. Thus the 2nd model using feature selection achieves an higher accuracy of 0.62 than the 1st model using all features.

```
data = data.frame(feature1,classLable)
>
    # Use support vector machine and 5 fold cross validation to do classification
>
    LIBSVM_CV5 = classify(data, classifyMethod='libsvm',
                 cv=5, svm.kernel='linear',svm.scale=F)
    LIBSVM_CV5[["totalPerformance"]]
                                                         prc
337.0000000 337.0000000 263.0000000 263.0000000
                                                   0.5599397
                                                               0.5616667
         sp
                    acc
                                mcc
                                              рс
 0.5616667
              0.5616667
                          0.1243355
                                      0.3918323
    # Features selection is done by envoking "CfsSubsetEval" method in WEKA.
    FS_LIBSVM_CV5 = classify(data, classifyMethod='libsvm',
>
                    cv=5, evaluator='CfsSubsetEval', search='BestFirst',
                    svm.kernel='linear', svm.scale=F)
    FS_LIBSVM_CV5[["totalPerformance"]] ## Accuracy is increased by feature select
         tp
                     tn
                                 fp
                                              fn
                                                         prc
                                                                      sn
258.0000000 488.0000000 112.0000000 342.0000000
                                                   0.6924855
                                                               0.4300000
                                \mathtt{mcc}
 0.8133333
              0.6216667
                          0.2621407
                                      0.3651539
    # Selected features:
    colnames(data)[FS_LIBSVM_CV5$features[[1]]]
                "BIN.2_R"
 [1] "BIN.1_A"
                           "BIN.3_K"
                                      "BIN.4_K"
                                                 "BIN.5_K"
                                                             "BIN.5_N"
 [7] "BIN.6_N"
                "BIN.6_G"
                           "BIN.6_T"
                                      "BIN.6_P" "BIN.7_P"
                                                             "BIN.9_W"
[13] "BIN.9_H" "BIN.9_Y"
                           "BIN.9_L"
                                      "BIN.10_Q" "BIN.11_W" "BIN.11_V"
```

5. Different feature coding methods usually might result in different prediction performance. featureEvaluate can be used to test multiple feature coding methods. Figure 2 shows the 3D plot of prediction accuracy varied with feature coding functions and parameters. It can be generated by employing featureEvaluate as follows (Note: It may be time consuming!):

[19] "BIN.12_K" "BIN.12_F" "BIN.13_K" "BIN.13_I" "BIN.13_M" "BIN.14_A"

[25] "BIN.15_K" "BIN.15_E" "BIN.15_A" "BIN.15_T"

fileName = tempfile() # Note: It may be time consuming. testFeatureSet = featureEvaluate(seq, classLable, fileName, cv=5, ele.type='aminoacid', featureMethod=c('Binary', 'GapPairComposition'), classifyMethod='libsvm', group=c('aaH', 'aaV', 'aaZ', 'aaP', 'aaF', 'aaS', 'aaE'), g=0, +

```
summary = read.csv(fileName,sep="\t",header=T)
>
    # Plot the result of 'featureEvaluate'
    require("scatterplot3d")
>
    tmp1 = summary[,"Feature_Function"]
    tmp2 = as.factor(sapply(as.vector(summary[, 'Feature_Parameter']),
           function(x){unlist(strsplit(x,split='; '))[1]}))
    testResult = data.frame(as.integer(tmp2), as.integer(tmp1), summary[, "acc"])
>
    s3d=scatterplot3d(testResult,
>
        color=c('red','blue')[testResult[,2]], pch=19, xlab='Parameter',
        ylab='Feature Coding',
        zlab='Accuracy', lab=c(9,3,7),
        x.ticklabs=gsub('class: ','',sort(unique(tmp2))),
        type='h',ylim=c(0,3),y.margin.add=2.5,
        y.ticklabs=c('',gsub('feature','',sort(unique(tmp1))),'') )
```

6. Features from multiple functions can be combined and re-selected to increase the prediction accuracy. In the following code chunk, the first three feature sets from 'testFeature-Set' are combined together ('testFeatureSet' is generated in the aforementioned codes by featureEvaluate). Then feature selection functions (classify and selectFFS) can be employed to selecte features. (classify has been illustrated in the aforementioned code chunk. Thus selectFFS is used here to do feature forward selection to select a subset with maximum prediction accuracy (Note: It may be time consuming!). The process of feature selection and the increasing accuracy are shown in Figure 3.

```
feature.index = 1:3
>
    tmp <- testFeatureSet[[1]]$data</pre>
>
    tmp1 <- testFeatureSet[[feature.index[1]]]$model</pre>
    colnames(tmp) <- paste(</pre>
      tmp1["Feature_Function"],
      tmp1["Feature_Parameter"],
      colnames(tmp),sep=" ; ")
    data <- tmp[,-ncol(tmp)]</pre>
>
    for(i in 2:length(feature.index) ){
      tmp <- testFeatureSet[[feature.index[i]]]$data</pre>
      tmp1 <- testFeatureSet[[feature.index[i]]]$model</pre>
      colnames(tmp) <- paste(</pre>
         tmp1["Feature_Function"],
         tmp1["Feature_Parameter"],
         colnames(tmp),sep=" ; ")
      data <- data.frame(data, tmp[,-ncol(tmp)] )</pre>
    name <- colnames(data)</pre>
>
    data <- data.frame(data, tmp[,ncol(tmp)] ) ## Combined features</pre>
```

```
# Use 'selectFFS' to do feature forward selection.

# Note: It may be time consuming.

combineFeatureResult = selectFFS(data,accCutoff=0.005,

classifyMethod="knn",cv=5) ## It is time consuming.

tmp = sapply(combineFeatureResult,function(x){

c(length(x$features),x$performance["acc"])})

plot(tmp[1,],tmp[2,],xlab="Feature Number",ylab="Accuracy",

pch=19)

lines(tmp[1,],tmp[2,])
```

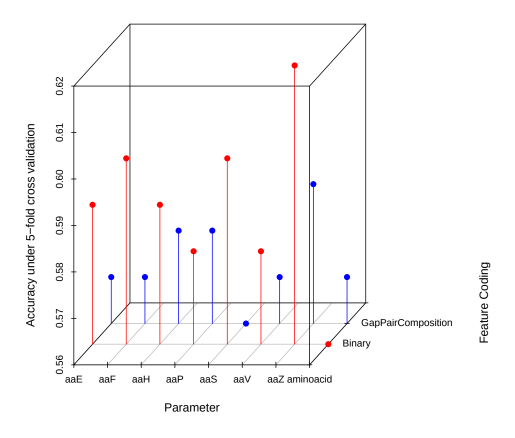


Figure 2: Result of featureEvaluate.

5 Session Information

This vignette was generated using the following package versions:

R version 3.0.0 (2013-04-03)

Platform: x86_64-unknown-linux-gnu (64-bit)

locale:

[1] LC_CTYPE=en_US.UTF-8 LC_NUMERIC=C
[3] LC_TIME=en_US.UTF-8 LC_COLLATE=C
[5] LC_MONETARY=en_US.UTF-8 LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=C LC_NAME=C

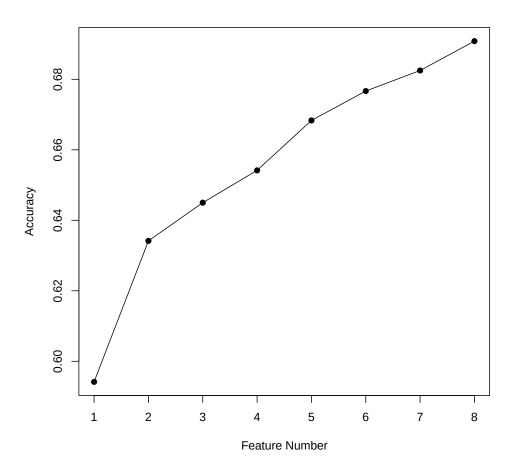


Figure 3: Result of selectFFS.

```
[9] LC_ADDRESS=C LC_TELEPHONE=C
```

[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:

[1] stats graphics grDevices utils datasets methods base

other attached packages:

[1] BioSeqClass_1.18.0 scatterplot3d_0.3-33

loaded via a namespace (and not attached):

[1]	Biobase_2.20.0	BiocGenerics_0.6.0	Biostrings_2.28.0	IRanges_1.18.0
[5]	MASS_7.3-26	class_7.3-7	coin_1.0-21	colorspace_1.2-1
[9]	e1071_1.6-1	foreign_0.8-53	ipred_0.9-1	klaR_0.6-8
[13]	modeltools_0.2-19	nnet_7.3-6	parallel_3.0.0	party_1.0-6
[17]	randomForest_4.6-7	rpart_4.1-1	splines_3.0.0	stats4_3.0.0
[21]	survival_2.37-4	tools_3.0.0	tree_1.0-33	

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