Package 'LedPred'

March 6, 2025

Title Learning from DNA to Predict Enhancers

Description This package aims at creating a predictive model of regulatory sequences used to score unknown sequences based on the content of DNA motifs, next-generation sequencing (NGS) peaks and signals and other numerical scores of the sequences using supervised classification. The package contains a workflow based on the support vector machine (SVM) algorithm that maps features to sequences, optimize SVM parameters and feature number and creates a model that can be stored and used to score the regulatory potential of unknown sequences.

Version 1.40.0

Date 2016-08-13

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Depends R (>= 3.2.0), e1071 (>= 1.6)

- **Imports** akima, ggplot2, irr, jsonlite, parallel, plot3D, plyr, RCurl, ROCR, testthat
- License MIT | file LICENSE

LazyData true

biocViews SupportVectorMachine, Software, MotifAnnotation, ChIPSeq, Sequencing, Classification

NeedsCompilation no

BugReports https://github.com/aitgon/LedPred/issues

RoxygenNote 5.0.1

git_url https://git.bioconductor.org/packages/LedPred

git_branch RELEASE_3_20

git_last_commit ee0237e

git_last_commit_date 2024-10-29

Repository Bioconductor 3.20

Date/Publication 2025-03-06

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createModel

Create the model with the optimal features

Description

createModel function creates a SVM model from the training data set with the selected features.

Usage

```
createModel(data, cl = 1, kernel = "radial", cost = 1, gamma = 1,
valid.times = 10, feature.ranking = NULL, feature.nb = NULL,
file.prefix = NULL)
```

Arguments

| data | data.frame containing the training set | | | | |
|-----------------|--|--|--|--|--|
| cl | integer indicating the column number corresponding to the response vector that classify positive and negative regions (default = 1) | | | | |
| kernel | SVM kernel, a character string: "linear" or "radial". (default = "radial") | | | | |
| cost | The SVM cost parameter for both linear and radial kernels. If NULL (default), the function $mcTune$ is run. | | | | |
| gamma | The SVM gamma parameter for radial kernel. If radial kernel and NULL (default), the function ${\tt mcTune}$ is run. | | | | |
| valid.times | Integer indicating how many times the training set will be split for the cross validation step (default = 10). This number must be smaller than positive and negative sets sizes. | | | | |
| feature.ranking | | | | | |
| | List of ordered features. | | | | |
| feature.nb | the optimal number of feature to use from the list of ordered features. | | | | |
| file.prefix | A character string that will be used as a prefix followed by "_model.RData" for the resulting model file, if it is NULL (default), no model is saved | | | | |

Value

the best SVM model

crm.features

Examples

```
crm.features
```

This is data to be included in my package

Description

This is data to be included in my package

evaluateModelPerformance

Evaluate model performances

Description

evaluateModelPerformance function computes the precision and recall measures to evaluate the model through cross validation steps using ROCR package.

Usage

```
evaluateModelPerformance(data, cl = 1, valid.times = 10,
feature.ranking = NULL, feature.nb = NULL,
numcores = ifelse(.Platform$0S.type == "windows", 1, parallel::detectCores()
- 1), file.prefix = NULL, kernel = "linear", cost = NULL,
gamma = NULL)
```

Arguments

| data | data.frame containing the training set | | | |
|-----------------|--|--|--|--|
| cl | integer indicating the column number corresponding to the response vector that classify positive and negative regions (default = 1) | | | |
| valid.times | Integer indicating how many times the training set will be split for the cross validation step (default = 10). This number must be smaller than positive and negative sets sizes. | | | |
| feature.ranking | | | | |
| | List of ordered features. | | | |
| feature.nb | the optimal number of feature to use from the list of ordered features. | | | |
| numcores | Number of cores to use for parallel computing (default: the number of available cores in the machine - 1) | | | |

| file.prefix | A character string that will be used as a prefix followed by "_ROCR_perf.png" for the result plot file, if it is NULL (default), no plot is returned |
|-------------|--|
| kernel | SVM kernel, a character string: "linear" or "radial". (default = "radial") |
| cost | The SVM cost parameter for both linear and radial kernels. If NULL (default), the function mcTune is run. |
| gamma | The SVM gamma parameter for radial kernel. If radial kernel and NULL (default), the function mcTune is run. |

Value

A list with two objects.

| probs | The predictions computed by the model for each subset during the cross-validation |
|--------|---|
| labels | The actual class for each subset |

Examples

feature.ranking This is data to be included in my package

Description

This is data to be included in my package

LedPred

Creates an SVM model given a feature matrix

Description

The LedPred function computes the best SVM parameters, defines the optimal features for creating the SVM model by running sequentially mcTune, rankFeatures, tuneFeatureNb and createModel. The performances of this model are then computed usong evaluateModelPerformance.

Usage

```
LedPred(data = NULL, cl = 1, ranges = list(gamma = c(1, 10), cost = c(1,
10)), cost = NULL, gamma = NULL, kernel = "linear", valid.times = 10,
file.prefix = NULL, numcores = ifelse(.Platform$0S.type == "windows", 1,
parallel::detectCores() - 1), step.nb = 10, halve.above = 100)
```

LedPred

Arguments

| data.frame containing the training set |
|---|
| integer indicating the column number corresponding to the response vector that classify positive and negative regions (default = 1) |
| list object containing one (linear kernel) or two (radial kernel) vectors of integers corresponding to SVM cost and SVM gamma parameters to test. |
| The SVM cost parameter for both linear and radial kernels. If NULL (default), the function mcTune is run. |
| The SVM gamma parameter for radial kernel. If radial kernel and NULL (default), the function mcTune is run. |
| SVM kernel, a character string: "linear" or "radial". (default = "radial") |
| Integer indicating how many times the training set will be split for the cross validation step (default = 10). This number must be smaller than positive and negative sets sizes. |
| A character string that will be used as a prefix for the result files. If it is NULL (default), no plot is returned |
| Number of cores to use for parallel computing (default: the number of available cores in the machine - 1) |
| Number of features to add at each step (default = 10) |
| During RFE, all the features are ranked at the first round and the half lowest ranked features (that contribute the least in the model) are removed for the next round. When the number of feature is lower or equal to halve.above, the features are removed one by one. (default=100) |
| |

Value

A list of the object produced at each step

| best.params | A list of the parameters giving the lowest misclassification error |
|-----------------|--|
| feature.ranking | |
| | List of ordered features from rankFeatures |
| feature.nb | he optimal number of feature to use from the list of ordered features from ${\tt tuneFeatureNb}$ |
| model.svm | The best SVM model createModel |
| probs.label.lis | t |
| | |

 $The\ cross-validation\ results\ from\ evaluate {\tt ModelPerformance}$

```
data(crm.features)
#cost_vector <- c(1,3,10)
#gamma_vector <- c(1,3,10)
#ledpred.list=LedPred(data.granges=crm.features, cl=1, ranges = list(cost=cost_vector,
# gamma=gamma_vector), kernel="linear", halve.above=50)
#names(ledpred.list)</pre>
```

```
mapFeaturesToCRMs
```

Description

The mapFeaturesToCRMs function allows the user to create a training set matrix to build a predictive model. The training set is composed of positive regions (known to be involved in the pathway of interest) and negative regions (randomly picked or known to not be involved in the pathway of interest) that will be described (scored) by features. Three types of features file format are accepted: Position specific scoring matrices modeling motifs recognised by transcription factors, bed files containing region coordinates for any discrete feature (NGS peaks, conservation blocks) and wig/bigWig files containing signal data. This script has been tested with version 0.99 of the online server. Go here to see current version of the server http://ifbprod.aitorgonzalezlab.org/map_features_to_crms.php

Usage

```
mapFeaturesToCRMs(URL = "http://ifbprod.aitorgonzalezlab.org/map_features_to_crms.php",
positive.bed = NULL, genome = NULL, negative.bed = NULL,
shuffling = NULL, background.seqs = NULL, genome.info = NULL,
pssm = NULL, background.freqs = NULL, ngs = NULL, bed.overlap = NULL,
my.values = NULL, feature.ranking = NULL, feature.nb = NULL,
crm.feature.file = NULL, stderr.log.file = NULL, stdout.log.file = NULL)
```

Arguments

| URL | URL of the server REST target |
|--|---|
| positive.bed | Positive bed file path. Compulsory |
| genome | Genome code, eg. dm3 for Drosophila Melanogaster. Compulsory |
| negative.bed | Negative bed file path. |
| shuffling | Integer with number of time shuffle background sequences (background.seqs). If negative.bed is NULL and shuffling is set at 0, the feature matrix does not contain negative sequences. It is useful to produce a test set matrix. |
| background.seqs | |
| | Background sequences used for shuffling. If shuffling = 0 , set this parameter at 0 . |
| genome.info | File require for shuffling bed. If shuffling = 0 , set this parameter at 0 . |
| pssm | Position specific scoring matrices |
| background from | S |
| backgi bullu. H eq | |
| backgi ounu. n eq | Background frequencies of nucleotides in genome |
| ngs | Background frequencies of nucleotides in genome NGS (bed and wig) files |
| ngs bed.overlap | Background frequencies of nucleotides in genome NGS (bed and wig) files Minimal overlap as a fraction of query sequence with NGS bed peak. Equivalent with intersectBed -f argument. Default 1bp. |
| ngs bed.overlap my.values | Background frequencies of nucleotides in genome NGS (bed and wig) files Minimal overlap as a fraction of query sequence with NGS bed peak. Equivalent with intersectBed -f argument. Default 1bp. Bed file where fourth column are values to append to the SVM matrix |
| ngs bed.overlap my.values feature.ranking | Background frequencies of nucleotides in genome NGS (bed and wig) files Minimal overlap as a fraction of query sequence with NGS bed peak. Equivalent with intersectBed -f argument. Default 1bp. Bed file where fourth column are values to append to the SVM matrix |
| ngs bed.overlap my.values feature.ranking | Background frequencies of nucleotides in genome NGS (bed and wig) files Minimal overlap as a fraction of query sequence with NGS bed peak. Equivalent with intersectBed -f argument. Default 1bp. Bed file where fourth column are values to append to the SVM matrix File with ranked features (Output of rankFeatures). It is used for scoring a query bed file |

mcTune

```
crm.feature.file
Path to feature matrix file
stderr.log.file
Path to error log
stdout.log.file
Path to standard output log
```

Value

| feature.matrix | a data frame where each row is a region and each column a feature, each cell |
|----------------|--|
| | carry a score, the first column is the response vector |
| stdout.log | Standard output log of mapFeaturesToCRMs script in server |
| stderr.log | Standard error log of mapFeaturesToCRMs script in server |
| | |

Examples

```
## Not run:
dirPath <- system.file("extdata", package="LedPred")</pre>
 file.list <- list.files(dirPath, full.names=TRUE)</pre>
 background.freqs <- file.list[grep("freq", file.list)]</pre>
 positive.regions <- file.list[grep("positive", file.list)]</pre>
 negative.regions <- file.list[grep("negative", file.list)]</pre>
 TF.matrices <- file.list[grep("tf", file.list)]</pre>
 ngs.path <- system.file("extdata/ngs", package="LedPred")</pre>
 ngs.files=list.files(ngs.path, full.names=TRUE)
 crm.features.list <- mapFeaturesToCRMs(positive.bed=positive.regions,</pre>
     negative.bed=negative.regions, background.freqs=background.freqs,
     pssm=TF.matrices, genome="dm3", ngs=ngs.files,
     crm.feature.file = "crm.features.tab",
     stderr.log.file = "stderr.log", stdout.log.file = "stdout.log")
 names(crm.features.list)
 class(crm.features.list$crm.features)
 crm.features.list$stdout.log
 crm.features.list$stderr.log
```

End(Not run)

```
mcTune
```

Tuning the SVM parameters

Description

The mcTune function is a modified version of the function tune from package e1071 [6]. It tests the different combinations of C and gamma parameters given as vectors in a list and will return the prediction error computed during the cross-validation step.

Usage

```
mcTune(data, cl = 1, ranges = list(gamma = c(1, 10), cost = c(1, 10)),
kernel = "linear", valid.times = 10, file.prefix = NULL,
numcores = ifelse(.Platform$0S.type == "windows", 1, parallel::detectCores()
- 1))
```

Arguments

| data | data.frame containing the training set |
|-------------|--|
| cl | integer indicating the column number corresponding to the response vector that classify positive and negative regions (default = 1) |
| ranges | list object containing one (linear kernel) or two (radial kernel) vectors of integers corresponding to SVM cost and SVM gamma parameters to test. |
| kernel | SVM kernel, a character string: "linear" or "radial". (default = "radial") |
| valid.times | Integer indicating how many times the training set will be split for the cross validation step (default = 10). This number must be smaller than positive and negative sets sizes. |
| file.prefix | A character string that will be used as a prefix followed by "_c_g_eval.png" for result plot files, if it is NULL (default), no plot is returned |
| numcores | Number of cores to use for parallel computing (default: the number of available cores in the machine - 1) |

Value

A list of class tune

| best.parameters | 3 | |
|------------------|---|--|
| | A list of the parameters giving the lowest misclassification error | |
| best.performance | | |
| | The lowest misclassification error | |
| method | The method used | |
| nparcomb | the number of tested parameter combinations | |
| train.ind | The indexes used to produce subsets during the cross validation step | |
| sampling | The cross-validation fold number | |
| performances | A matrix summarizing the cross-validation step with the error for each tested parameter at each round and the dispersion of these errors (regarding to the average error) | |
| best.model | The model produced by the best parameters | |

```
data(crm.features)
cost.vector <- c(1,3,10,30)
gamma.vector <- c(1,3,10,30)
#c.g.obj <- mcTune(data.granges= crm.features, ranges = list(cost=cost.vector,
# gamma=gamma.vector), kernel='linear', file.prefix = "test")
#names(c.g.obj)
# cost <- c.g.obj$best.parameters$cost
# gamma <- c.g.obj$best.parameters$gamma</pre>
```

rankFeatures

Description

The rankFeatures function performs a Recursive Feature Elimination (RFE) on subsets of the feature matrix. For each subset the features are ranked according to the weight attributed by SVM at each round of elimination and the average rank of each feature over the subsets is returned. We recommand to save the object containing the ranked features for the following steps.

Usage

```
rankFeatures(data, cl = 1, halve.above = 100, valid.times = 10,
kernel = "linear", cost = 1, gamma = 1,
numcores = ifelse(.Platform$0S.type == "windows", 1, parallel::detectCores()
- 1), file.prefix = NULL)
```

Arguments

| data.frame containing the training set |
|---|
| integer indicating the column number corresponding to the response vector that classify positive and negative regions (default = 1) |
| During RFE, all the features are ranked at the first round and the half lowest ranked features (that contribute the least in the model) are removed for the next round. When the number of feature is lower or equal to halve.above, the features are removed one by one. (default=100) |
| Integer indicating how many times the training set will be split (default = 10). This number must be smaller than positive and negative sets sizes. |
| SVM kernel, a character string: "linear" or "radial". (default = "radial") |
| The SVM cost parameter for both linear and radial kernels. If NULL (default), the function mcTune is run. |
| The SVM gamma parameter for radial kernel. If radial kernel and NULL (default), the function mcTune is run. |
| Number of cores to use for parallel computing (default: the number of available cores in the machine - 1) |
| A character string that will be used as a prefix for output file, if it is NULL (default), no file is writen. |
| |

Value

A 3-columns data frame with ranked features. First column contains the feature names, the second the original position of the feature in the feature.matrix and the third the average rank over the subsets.

scoreData

Description

scoreData function predict new regulatory regions using SVM model from a test data set

Usage

```
scoreData(data, ledpred = NULL, model = NULL, score.file = NULL)
```

Arguments

| data | data.frame containing the test set. This test set must have the same descriptive features as the one that were used to build the model. |
|------------|--|
| ledpred | Returned object from the LedPred function |
| model | Returned object of the createModel function |
| score.file | A character string that will be used as the file name for the output file, if it is NULL (default), no file is writen. The output file takes the form of two columns with object names and scores. |

Value

A 2-columns dataframe. First column containg the SVM model prediction probabilities and the second containing the corresponding regions

Examples

```
data(crm.features)
data(svm.model)
#pred.test <- scoreData(data.granges=crm.features, model=svm.model,
# score.file="test_prediction.tab")</pre>
```

tuneFeatureNb

Selecting the optimal number of features

Description

tuneFeatureNb iterates through increasing feature numbers to calculate kappa values which represents the performance of the model computed with the given features. We recommand to save the object containing the optimal number of features for the following steps.

Usage

```
tuneFeatureNb(data, cl = 1, feature.ranking, step.nb = 10,
valid.times = 10, cost = NULL, gamma = NULL, kernel = "linear",
numcores = ifelse(.Platform$0S.type == "windows", 1, parallel::detectCores()
- 1), file.prefix = NULL)
```

tuneFeatureNb

Arguments

| data | data.frame containing the training set |
|-----------------|--|
| cl | integer indicating the column number corresponding to the response vector that classify positive and negative regions (default = 1) |
| feature.ranking | |
| | List of ordered features. |
| step.nb | Number of features to add at each step (default = 10) |
| valid.times | Integer indicating how many times the training set will be split for the cross validation step (default = 10). This number must be smaller than positive and negative sets sizes. |
| cost | The SVM cost parameter for both linear and radial kernels. If NULL (default), the function mcTune is run. |
| gamma | The SVM gamma parameter for radial kernel. If radial kernel and NULL (de-fault), the function mcTune is run. |
| kernel | SVM kernel, a character string: "linear" or "radial". (default = "radial") |
| numcores | Number of cores to use for parallel computing (default: the number of available cores in the machine - 1) |
| file.prefix | A character string that will be used as a prefix followed by "_kappa_measures.png" for the result plot file. If it is NULL (default), no plot is returned |

Value

A list with two objects.

| performance | 2-columns data frame. first column correspond to the number of tested features, second column contains the corresponding kappa value |
|-----------------|--|
| best.feature.nb | |
| | Integer corresponding to the number of features producing the model with the highest kappa value |

```
data(crm.features)
data(feature.ranking)
cost <- 1
gamma <- 1
#feature.nb.obj <- tuneFeatureNb(data.granges=crm.features,
# feature.ranking=feature.ranking, kernel='linear', cost=cost,gamma=gamma,
# file.prefix = "test")
#names(feature.nb.obj)</pre>
```

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