Package 'FeaLect'

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Title	Scores Features for Feature Selection	
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Depends lars, rms

Author Habil Zare

Type Package

Description For each feature, a score is computed that can be useful for feature selection. Several random subsets are sampled from the input data and for each random subset, various linear models are fitted using lars method. A score is assigned to each feature based on the tendency of LASSO in including that feature in the models. Finally, the average score and the models are returned as the output. The features with relatively low scores are recommended to be ignored because they can lead to overfitting of the model to the training data. Moreover, for each random subset, the best set of features in terms of global error is returned. They are useful for applying Bolasso, the alternative feature selection method that recommends the intersection of features subsets.

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Description

Suppose you have a feature matrix with 200 features and only 20 samples and your goal is to build a classifier. You can run the FeaLect() function to compute the scores for your features. Only the relatively high score features (say the top 20) are recommended for further analysis. In this way, one can prevent overfitting by reducing the number of features significantly.

Details

The DESCRIPTION file:

FeaLect Package: Type: Package

Title: Scores Features for Feature Selection

Version: 1.20 Date: 2020-02-25 Author: Habil Zare

Maintainer: Habil Zare <zare@u.washington.edu>

Depends:

Description: For each feature, a score is computed that can be useful for feature selection. Several random subsets an

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linear scores

doctor.validate Validates a model using validating samples.

ignore.redundant Refines a feature matrix

input.check.FeaLect Checks the inputs to Fealect() function.

mcl_sll MCL and SLL lymphoma subtypes

random.subset Selects a random subset of the input.

train.doctor Fits various models based on a combination on

penalized linear models and logistic

regression.

Author(s)

Habil Zare

Maintainer: Habil Zare <zare@u.washington.edu>

References

Zare, Habil, et al. "Scoring relevancy of features based on combinatorial analysis of Lasso with application to lymphoma diagnosis." *BMC genomics*. Vol. 14. No. 1. BioMed Central, 2013.

See Also

FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect, lars-package, and SparseLearner-package

Examples

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
L <- as.numeric(mcl_sll[ ,1]) # The labels
names(L) <- rownames(F)
message(dim(F)[1], " samples and ",dim(F)[2], " features.")

## For this data, total.num.of.models is suggested to be at least 100.
FeaLect.result.1 <-FeaLect(F=F,L=L,maximum.features.num=10,total.num.of.models=20,talk=TRUE)</pre>
```

compute.balanced

Balances between negative and positive samples by oversampling.

Description

If negative samples are less than positive ones, more copies of the negative cases are added and vice versa.

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Usage

```
compute.balanced(F_, L_)
```

Arguments

F_ The feature matrix, each column is a feature.

L_ The vector of labels named according to the rows of F.

Details

Considerably unbalanced classes may be probabilistic for fitting some models.

Value

Returns a list of:

F_ The feature matrix, each column is a feature.

L_ The vector of labels named according to the rows of F.

Author(s)

Habil Zare

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
L <- as.numeric(mcl_sll[ ,1]) # The labels
names(L) <- rownames(F)
message(L)

balanced <- compute.balanced(F_=F, L_=L)
message(balanced$L_)</pre>
```

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```
compute.logistic.score
```

Fits a logistic regression model using the linear scores

Description

A logistic regression model is fitted to the linear scores using lrm() function and the logistic scores are computed using the formula: 1/(1+exp(-(a+bX))) where a and b are the logistic coefficients.

Usage

```
compute.logistic.score(F_-, L_-, considered.features, training.samples, validating.samples, linear.scores, report.fitting.failure = TRUE)
```

Arguments

F_ The feature matrix, each column is a feature.

__ The vector of labels named according to the rows of F.

training.samples

The names of rows of F that should be considered as training samples.

validating.samples

The names of rows of F that should be considered as validating samples.

considered.features

The names of columns of F that determine the features of interest.

linear.scores A vector that contains for each training or validating sample, a linear score predicted by the linear method.

report.fitting.failure

If TRUE, any failure in fitting the linear of logistic models will be printed.

Details

The logistic regression will be fitted to all training and validating samples.

Value

Returns a list of:

logistic.scores

A vector of predicted logistic values for all samples.

logistic.cofs The coefficients that are computed by logistic regression.

Note

Logistic regression is also done on top of fitting the linear models.

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Author(s)

Habil Zare

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

```
FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect
```

Examples

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
L <- as.numeric(mcl_sll[ ,1]) # The labels
names(L) <- rownames(F)
all.samples <- rownames(F); ts <- all.samples[5:10]; vs <- all.samples[c(1,22)]
L <- L[c(ts,vs)]
L
asymptotic.scores <- c(1,0.9,0.8,0.2,0.1,0.1,0.7,0.2)
compute.logistic.score(F_=F, L_=L, training.samples=ts, validating.samples=vs, considered.features=colnames(F),linear.scores= asymptotic.scores)</pre>
```

doctor.validate

Validates a model using validating samples.

Description

A model fitted on the training samples, can be validated on a separate validating set. The recall, precision, and accuracy of the model are computed.

Usage

```
doctor.validate(true.labels, predictions)
```

Arguments

```
true.labels A vector of 0 and 1. predictions A vector of 0 and 1.
```

Details

F-measure is equal to: 2 times precision times recall / (precision+recall).

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Value

F-measure, precision, and recall are calculated. Also, the mis-labeled cases are reported.

Author(s)

Habil Zare

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect

Examples

```
tls <- c(1,1,1,0,0)
ps <- c(1,1,0,1,0)
names(tls) <- 1:5; names(ps) <- 1:5
doctor.validate(true.labels=tls, predictions=ps)</pre>
```

FeaLect

Computes the scores of the features.

Description

Several random subsets are sampled from the input data and for each random subset, various linear models are fitted using lars method. A score is assigned to each feature based on the tendency of LASSO in including that feature in the models. Finally, the average score and the models are returned as the output.

Usage

```
FeaLect(F, L, maximum.features.num = dim(F)[2], total.num.of.models, gamma = 3/4,
    persistence = 1000, talk = FALSE, minimum.class.size = 2,
    report.fitting.failure = FALSE, return_linear.models = TRUE, balance = TRUE,
    replace = TRUE, plot.scores = TRUE)
```

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Arguments

F The feature matrix, each column is a feature.

The vector of labels named according to the rows of F.

maximum.features.num

Upto this number of features are allowed to contribute to each linear model.

total.num.of.models

The total number of models that are fitted.

gamma A value in range 0-1 that determines the relative size of sample subsets.

persistence Maximum number of tries for randomly choosing samples, If we try this many

times and the obtained labels are all the same, we give up (maybe the whole labels are the same) with the error message: " Not enough variation in the la-

bels...".

talk If TRUE, some messages are printed during the computations.

minimum.class.size

The size of both positive and negative classes should be greater than this thresh-

old after sampling.

report.fitting.failure

If TRUE, any failure in fitting the linear of logistic models will be printed.

return_linear.models

The models are memory intensive, so for if they more than 1000, we may decide

to ignore them to prevent memory outage.

balance If TRUE, the cases will be balanced for the same number of positive vs. nega-

tives by oversampling before fitting the linear model.

replace If TRUE, the subsets are sampled with replacement.

plot. scores If TRUE, the scores are plotted in logarithmic scale after each iteration.

Details

See the reference for more details.

Value

Returns a list of:

log. scores A vector containing the logarithm of final scores.

feature.matrix The input feature matrix.

labels The input labels

total.num.of.models

The total number of models that are fitted.

maximum.features.num

Upto this number of features are allowed to contribute to each linear model.

feature.scores.history

The matrix of history of feature scores where column i contains the scores after i runs.

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num.of.features.score

A vector, entry i contains the number of times that i has been the best number of features.

best.feature.num

The i'th value of this vector is the best number of features for the i'th model.

mislabeling.record

A vector that keeps track of the frequency of mislabelling for each cases.

doctors List of all models which are created by train.doctor() function. best.features.intersection

Best features are computed for each sampling and their intersection is reported as this vector of features names

features.with.best.global.error

A list containing the sets of features. The set i was the best for i'th sampling.

time.taken Total time used for executing this function.

Note

Logistic regression is also done on top of fitting the linear models.

Author(s)

Habil Zare

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
L <- as.numeric(mcl_sll[ ,1]) # The labels
names(L) <- rownames(F)
message(dim(F)[1], " samples and ",dim(F)[2], " features.")
## For this data, total.num.of.models is suggested to be at least 100.
FeaLect.result <-FeaLect(F=F,L=L,maximum.features.num=10,total.num.of.models=20,talk=TRUE)</pre>
```

ignore.redundant

ignore.redundant

Refines a feature matrix

Description

If the value a feature is the same for all points (e.g. =0), it can be ignored.

Usage

```
ignore.redundant(F, num.of.values = 1)
```

Arguments

F The feature matrix, each column is a feature.

num.of.values A feature should have more than this threshold non-zero values not to be ignored.

Value

The refined feature matrix.

Author(s)

Habil Zare

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
#F <- cbind(F, rep(1, times=dim(F)[1]))
message(dim(F)[1], " samples and ",dim(F)[2], " features.")

G <- ignore.redundant(F)
message("for ",dim(G)[1], " samples, ",dim(G)[2], " features are left.")</pre>
```

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input.check.FeaLect

Checks the inputs to Fealect() function.

Description

We should have: F as a matrix, L as a vector, and length of L be equal to number of rows of F. They should have names accordingly.

Usage

```
input.check.FeaLect(F_, L_, maximum.features.num, gamma)
```

Arguments

F_ The feature matrix, each column is a feature.

L_ The vector of labels named according to the rows of F.

maximum.features.num

Upto this number of features are allowed to contribute to each linear model.

gamma A value in range 0-1 that determines the relative size of sample subsets.

Details

If the input is not appropriate, error or warning message will be produced.

Value

Returns a list of:

F_ The feature matrix, each column is a feature.

The vector of labels named according to the rows of F.

maximum.features.num

Upto this number of features are allowed to contribute to each linear model.

Author(s)

Habil Zare

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect

mcl_sll

Examples

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
L <- as.numeric(mcl_sll[ ,1]) # The labels
names(L) <- rownames(F)

checked <- input.check.FeaLect(F_=F, L_=L, maximum.features.num=10, gamma=3/4)</pre>
```

 mcl_sll

MCL and SLL lymphoma subtypes

Description

A total of 237 features are identified for 22 lymphoma patients.

Usage

```
data(mcl_sll)
```

Format

A matrix. Each of the 237 columns represents a features except the first column which contains the label vector. Each of the 22 rows represents a patients.

Details

7 cases diagnosed with Mantel Cell Lymphoma (MCL) and 15 cases with Small Lymphocytic Lymphoma (SLL). The presented features are computed based on flow cytometry data The fist column contains the label vector which has value 1 for MCL cases and 0 for SLL cases.

Source

British Columbia Cancer Agency

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect

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Examples

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
L <- as.numeric(mcl_sll[ ,1]) # The labels
names(L) <- rownames(F)
message(dim(F)[1], " samples and ",dim(F)[2], " features.")
L</pre>
```

random.subset

Selects a random subset of the input.

Description

If a subset of samples are selected randomly, the navigate of positive classes might be too sparse or even empty. This function will repeat sampling until the classes are appropriate in this sense.

Usage

```
random.subset(F_, L_, gamma, persistence = 1000, minimum.class.size=2, replace)
```

Arguments

F_ The feature matrix, each column is a feature.

L_ The vector of labels named according to the rows of F.

gamma A value in range 0-1 that determines the relative size of sample subsets.

persistence Maximum number of tries for randomly choosing samples, If we try this many

times and the obtained labels are all the same, we give up (maybe the whole labels are the same) with the error message: " Not enough variation in the la-

bels...".

minimum.class.size

A lower bound on the number of samples in each class.

replace If TRUE, sampling is done by replacement.

Details

The function also returns a refined feature matrix by ignoring too sparse features after sampling.

Value

Returns a list of:

X_ The sampled feature matrix, each column is a feature after ignoring the redun-

dant ones.

Y_ The vector of labels named according to the rows of X_.

remainder.samples

The names of the rows of F_ which do not appear in X_, later on can be used for validation.

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Author(s)

Habil Zare

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

```
FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect
```

Examples

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
L <- as.numeric(mcl_sll[ ,1]) # The labels
names(L) <- rownames(F)
message(dim(F)[1], " samples and ",dim(F)[2], " features.")

XY <- random.subset(F_=F, L_=L, gamma=3/4,replace=TRUE)
XY$remainder.samples</pre>
```

train.doctor

Fits various models based on a combination on penalized linear models and logistic regression.

Description

Various linear models are fitted to the training samples using lars method. The models differ in the number of features and each is validated by validating samples. A score is also assigned to each feature based on the tendency of LASSO in including that feature in the models.

Usage

```
train.doctor(F_, L_, training.samples, validating.samples, considered.features,
maximum.features.num, balance = TRUE, return_linear.models = TRUE,
report.fitting.failure = FALSE)
```

Arguments

F_ The feature matrix, each column is a feature.

L_ The vector of labels named according to the rows of F.

training.samples

The names of rows of F that should be considered as training samples.

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validating.samples

The names of rows of F that should be considered as validating samples.

considered.features

The names of columns of F that determine the features of interest.

maximum.features.num

Upto this number of features are allowed to contribute to each linear model.

balance If TRUE, the cases will be balanced for the same number of positive vs. nega-

tives by oversampling before fitting the linear model.

return_linear.models

The models are memory intensive, so for if they more than 1000, we may decide

to ignore them to prevent memory outage.

report.fitting.failure

If TRUE, any failure in fitting the linear of logistic models will be printed.

Details

See the reference for more details.

Value

Returns a list of:

linear.models The result of model fitting computed by lars().

best.number.of.features

According to best accuracy.

probabilities The best computed logistic score.

accuracy The best F-measure.

best.logistic.cof

According to best accuracy.

contribution.to.feature.scores

This vector should be added to the total feature scores.

 $\verb|contribution.to.feature.scores.frequency| \\$

This vector should be added to the total frequency of features.

training.samples

Input, the names of rows of F that should be considered as training samples.

validating.samples

Input, the names of rows of F that should be considered as validating samples.

precision Ratio of number of true positives to predicted positives.

recall Ratio of number of true positives to real positives.

selected.features.sequence

A list of sets of features which are selected in different models.

global.errors A vector of global error of the linear fits.

features.with.best.global.error

A vector of names of good features in terms of global error of linear fits.

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Note

Logistic regression is also done on top of fitting the linear models.

Author(s)

Habil Zare

References

"Statistical Analysis of Overfitting Features", manuscript in preparation.

See Also

FeaLect, train.doctor, doctor.validate, random.subset, compute.balanced,compute.logistic.score, ignore.redundant, input.check.FeaLect

```
library(FeaLect)
data(mcl_sll)
F <- as.matrix(mcl_sll[ ,-1]) # The Feature matrix
L <- as.numeric(mcl_sll[ ,1]) # The labels
names(L) <- rownames(F)
message(dim(F)[1], " samples and ",dim(F)[2], " features.")

all.samples <- rownames(F); ts <- all.samples[5:10]; vs <- all.samples[c(1,22)]

doctor <- train.doctor(F_=F, L_=L, training.samples=ts, validating.samples=vs, considered.features=colnames(F), maximum.features.num=10)</pre>
```

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