Package 'EPX'

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Type Package

Title Ensemble of Phalanxes

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Description An ensemble method for the statistical detection of a rare class in two-class classification problems. The method uses an ensemble of classifiers where the constituent models of the ensemble use disjoint subsets (phalanxes) of explanatory variables. We provide an implementation of the phalanx-formation algorithm. Please see Tomal et al. (2015) <doi:10.1214/14-AOAS778>, Tomal et al. (2016) <doi:10.1021/acs.jcim.5b00663>, and Tomal et al. (2019) <arXiv:1706.06971> for more details.

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Encoding UTF-8

LazyData TRUE

Depends R (>= 3.6.0), foreach, randomForest, doRNG

Imports nnet, doParallel, rngtools

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2 AHR

R topics documented:

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Description

Calculates average hitrate (AHR), which is equivalent to average precision. When there are ties in ranking (not all values in phat are unique), the result is the expection of AHR. The algorithm that produces this analytic result assumes that the items in any tied group are in an arbitrary order within the group.

Usage

```
AHR(y, phat, ...)
```

Arguments

y True (binary) response vector where 1 is the rare/relevant class.

Numeric vector of estimated probabilities of relevance.

Further arguments passed to or from other methods.

Details

Implementation adapted from Wang (2005, Chapter 3). Please also see Chapter 3 of Wang (2005) for AHR and expected AHR formulas.

Value

Numeric value of average hitrate; expected average hitrate when there are ties.

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References

Wang, M. (2005). Statistical Methods for High Throughput Screening Drug Discovery Data (Doctoral thesis). University of Waterloo, Waterloo, Ontario, Canada.

Examples

BNhold

AID348 hold-out data using Burden Numbers for testing the EPX package

Description

AID348 hold-out data with 24 burden numbers as explanatory variables. Demonstrates in a timely manner epx, the phalanx-formation algorithm in **EPX** and associated functions summary.epx, predict.epx, plot.epx, hit.curve.

Usage

BNhold

Format

A dataframe with 3946 rows and 25 variables:

WBN Burden numbers descriptor set with 24 variables.

y The response variable where 1 denotes active and 0 inactive.

References

Tomal, J. H., Welch, W. J., & Zamar, R. H. (2015). Ensembling classification models based on phalanxes of variables with applications in drug discovery. *The Annals of Applied Statistics*, *9*(1), 69-93. doi: 10.1214/14AOAS778

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BNsample

AID348 sample (training) data with Burden Numbers for testing the EPX package

Description

AID348 sample (training) dataset with 24 burden numbers as explanatory variables. Demonstrates in a timely manner epx, the phalanx-formation algorithm in **EPX** and associated functions summary.epx, predict.epx, plot.epx, cv.epx, hit.curve.

Usage

BNsample

Format

A dataframe with 1000 rows and 25 variables:

WBN Burden numbers descriptor set with 24 variables.

y The response variable where 1 denotes active and 0 inactive.

References

Tomal, J. H., Welch, W. J., & Zamar, R. H. (2015). Ensembling classification models based on phalanxes of variables with applications in drug discovery. *The Annals of Applied Statistics*, *9*(1), 69-93. doi: 10.1214/14AOAS778

cv.epx

Balanced K-fold cross-validation for an "epx" object

Description

Balanced K-fold cross-validation based on an "epx" object. Hence, we have biased cross-validation as we do not re-run the phalanx-formation algorithm for each fold.

Usage

```
cv.epx(
  epx,
  folds = NULL,
  K = 10,
  folds.out = FALSE,
  classifier.args = list(),
  performance.args = list(),
  ...
)
```

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Arguments

epx Object of class "epx".

folds Optional vector specifying to which fold each observation belongs. Must be an

n-length vector (*n* being the number of observations) with integer values only

in the range from 1 to K.

K Number of folds; default is 10.

folds.out Indicates whether a vector indicating fold membership for each of the observa-

tions will be output; default is FALSE.

classifier.args

Arguments for the base classifier specified by epx; default is that used in epx

formation.

performance.args

Arguments for the performance measure specified by epx; default is that used in

epx formation.

. . . Further arguments passed to or from other methods.

Value

An (n+1) by (p+1) matrix, where n is the number of observations used to train epx and p is the number of (final) phalanxes. Column p+1 of the matrix contains the predicted probabilities of relevance from the ensemble of phalanxes, and row n+1 is the performance (choice of performance measure determined by the "epx" object) of the corresponding column.

Setting folds.out as TRUE changes the output of cv.epx into a list of two elements:

EPX.CV The (n+1) by (p+1) matrix returned by default when folds.out = FALSE.

FOLDS . USED A vector of length n with integer values only in the range from 1 to K indicating

to which fold each observation was randomly assigned for cross-validation.

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```
cv.folds <- cv.epx(model, folds.out = TRUE)
tail(cv.folds[[1]])  # same as first example
table(cv.folds[[2]])  # number of observations in each of the 10 folds

## 10 runs of 10-fold balanced cross-validation (using default settings)
set.seed(761)
cv.ahr <- NULL  # store AHR of each ensemble
for (i in 1:10) {
    cv.i <- cv.epx(model)
    cv.ahr <- c(cv.ahr, cv.i[nrow(cv.i), ncol(cv.i)])
}
boxplot(cv.ahr)  # to see variation in AHR

## End(Not run)</pre>
```

ерх

Fitting an Ensemble of Phalanxes

Description

epx forms phalanxes of variables from training data for binary classification with a rare class. The phalanxes are disjoint subsets of variables, each of which is fit with a base classifier. Together they form an ensemble.

Usage

```
epx(
    x,
    y,
    phalanxes.initial = c(1:ncol(x)),
    alpha = 0.95,
    nsim = 1000,
    rmin.target = 1,
    classifier = "random forest",
    classifier.args = list(),
    performance = "AHR",
    performance.args = list(),
    computing = "sequential",
    ...
)
```

Arguments

x Explanatory variables (predictors, features) contained in a data frame.

y Binary response variable vector (numeric or integer): 1 for the rare class, 0 for the majority class.

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phalanxes.initial

Initial variable group indices; default one group per variable. Example: vector c(1, 1, 2, 2, 3, ...) puts variables 1 and 2 in group 1, variables 3 and 4 in group, 2, etc. Indices cannot be skipped, e.g., c(1, 3, 3, 4, 4, 3, 1) skips group 2 and is

invalid.

alpha Lower-tail probability for the critical quantile of the reference distribution of the

performance measure for a classifier that ranks at random (i.e., the predictors

have no explanatory power); default is 0.95.

nsim Number of simulations for the reference empirical distribution of the perfor-

mance measure; default is 1000.

rmin.target To merge the pair of groups with the minimum ratio of performance measures

(ensemble of models to single model) into a single group their ratio must be less

than rmin.target, otherwise merging stops; default is 1.

classifier Base classifier, one of c("random forest", "logistic regression", "neural

network"); default is "random forest", which uses randomForest.

classifier.args

Arguments for the base classifier specified in a list as follows: list(argName1 = value1, argName2 = value2,...). If the list is empty, the classifier will use

its defaults. For "random forest", user may specify replace, cutoff, nodesize, maxnodes.

For "logistic regression" there are no options. For "neural network", user may

specify size, trace.

performance Performance assessment metric, one of c("AHR", "IE", "TOP1", "RKL"); de-

fault is AHR.

performance.args

Arguments for the performance measure specified in a list as follows: list(argName1

= value1, argName2 = value2,...). If the list is empty, the performance measure will use its defaults. Currently, only IE takes an argument list, and its only

argument is cutoff.

computing Whether to compute sequentially or in parallel. Input is one of c("sequential", "parallel");

default is "sequential".

... Further arguments passed to or from other methods.

Details

Please see Tomal et al. (2015) for more description of phalanx formation.

Value

Returns an object of class epx, which is a list containing the following components:

PHALANXES

List of four vectors, each the same length as the number of explanatory variables (columns in x): phalanxes.initial, phalanxes.filtered, phalanxes.merged, phalanxes.final. Each vector contains the phalanx membership indices of all explanatory variables at one of the four stages of phalanx-formation. Element i of a vector is the index of the phalanx to which variable i belongs. Phalanx 0 does not exist and so membership in phalanx 0 indicates that the variable does not belong to any phalanx; it has been screened out.

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PHALANXES.FINAL.PERFORMANCE

Vector of performance measures of the final phalanxes: the first element is for phalanx 1, etc.

PHALANXES.FINAL.FITS

A matrix with number of rows equal to the number of observations in the training data and number of columns equal to the number of final phalanxes. Column i contains the predicted probabilities of class 1 from fitting the base classifier to the variables in phalanx i.

ENSEMBLED.FITS The predicted probabilities of class 1 from the ensemble of phalanxes based on phalanxes.final.

BASE.CLASSIFIER.ARGS

(Parsed) record of user-specified arguments for classifier.

PERFORMANCE.ARGS

(Parsed) record of user-specified arguments for performance.

X User-provided data frame of explanatory variables.

Y User-provided binary response vector.

References

Tomal, J. H., Welch, W. J., & Zamar, R. H. (2015). Ensembling classification models based on phalanxes of variables with applications in drug discovery. *The Annals of Applied Statistics*, *9*(1), 69-93. doi: 10.1214/14AOAS778

See Also

summary.epx prints a summary of the results, and cv.epx assesses performance via cross-validation.

```
# Example with data(harvest)
## Phalanx-formation using a base classifier with 50 trees (default = 500)
set.seed(761)
model \leftarrow epx(x = harvest[, -4], y = harvest[, 4],
             classifier.args = list(ntree = 50))
## Phalanx-membership of explanatory variables at the four stages
## of phalanx formation (0 means not in a phalanx)
model $PHALANXES
## Summary of the final phalanxes (matches above)
summary(model)
## Not run:
## Parallel computing
clusters <- parallel::detectCores()</pre>
cl <- parallel::makeCluster(clusters)</pre>
doParallel::registerDoParallel(cl)
set.seed(761)
model.par \leftarrow epx(x = harvest[, -4], y = harvest[, 4],
```

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```
computing = "parallel")
parallel::stopCluster(cl)
## End(Not run)
```

harvest

Simulated dataset for testing the EPX package

Description

A simulated dataset from Yuan et al. (2012) with three explanatory variables. Demonstrates in a timely manner epx, the phalanx-formation algorithm in **EPX** and associated functions summary.epx, predict.epx, plot.epx, cv.epx, hit.curve.

Usage

harvest

Format

A dataframe with 190 rows and 4 variables:

LogP Octanol/water partition coefficient (-2 to 7).

MeltPt Melting point (120 to 280 degrees Celsius).

MolWt Molecular weight (200 to 800).

y The response variable where 1 denotes active and 0 inactive.

References

Yuan, Y., Chipman, H. A., & Welch, W. J. (2012). Harvesting Classification Trees for Drug Discovery. *Journal of Chemical Information and Modeling*, *52*(12), 3169-3180. doi: 10.1021/ci3000216

hit.curve

Plot hit curve

Description

Plots the hit curve corresponding to phat and y.

Usage

```
hit.curve(y, phat, max.cutoff = min(100, length(y)), plot.hc = T, ...)
```

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Arguments

У	True binary response vector where 1 denotes the relevant rare class.
phat	Vector of estimated probabilities of relevance.
max.cutoff	Maximum number of observations selected, equivalently the maximum shortlist cutoff; default is $min(100, length(y))$.
plot.hc	Whether to return a plot of the hit curve; default is TRUE.
	Further arguments passed to or from other methods.

Details

Order the cases by decreasing phat (predicted probabilities of relevance) values, and plot the expected number and actual number of hits as cases are selected. Cases with tied phat values are grouped together. See plot.epx for plotting the hit curve for an "epx" object.

Value

Plot of the hit curve (if plot.hc = TRUE) and a list with the following vectors:

select	Number of observations in each tied phat group; select[1], select[2], are the numbers of observations with the largest predicted probability of relevance (max(phat)), the second largest value in phat, etc.
p	Unique phat values; $p[1]$, $p[2]$, are the largest value in phat, the second largest value in phat, etc.
nhits	Number of hits (truly relevant observations) in each tied phat group.
nhitlast	Number of hits after max.cutoff observations selected.

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IE Calculate Initial Enhancement

Description

Calculates initial enhancement (IE), which is the precision at one specific shortlist length (cutoff) normalised by the proportion of relevants in the total sample size (Tomal et al. 2015). Since IE is a rescaling of precision, we expect IE and AHR to lead to similar conclusions as an assessment metric for the EPX algorithm.

Usage

```
IE(y, phat, cutoff = length(y)/2, ...)
```

Arguments

y True (binary) response vector where 1 is the rare/relevant class.

Numeric vector of estimated probabilities of relevance.

Shortlist cutoff length, and so must not exceed length of y; default is half the sample size.

Further arguments passed to or from other methods.

Details

Let c be the cutoff and h(c) be the hitrate at c. Let also A be the total number of relevants and N be the total number of observations. IE is defined as

$$IE = h(c)/(A/N)$$

IE calculation does not change whether there are ties in phat or not.

Value

Numeric value of IE.

References

Tomal, J. H., Welch, W. J., & Zamar, R. H. (2015). Ensembling classification models based on phalanxes of variables with applications in drug discovery. *The Annals of Applied Statistics*, *9*(1), 69-93. doi: 10.1214/14AOAS778

```
## IE when there are no ties in phat:
resp <- c(1, 1, 0, 0, 0, 0, 0, 1, 0, 0)
prob <- (10:1) * 0.1
IE(y = resp, phat = prob, cutoff = 3)</pre>
```

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```
# expect answer: (2/3) / (3/10)
## IE when there are ties
resp <- c(1, 1, 0, 0, 0, 0, 0, 1, 0, 0)
prob <- c(1, 1, 1, 0.4, 0.4, 0.3, 0.2, 0.15, 0.1, 0)
IE(y = resp, phat = prob, cutoff = 3)
# expect answer: same as above</pre>
```

plot.epx

Plot hit curve for an "epx" object

Description

Plots the hit curve for the fitted values of an "epx" object.

Usage

```
## S3 method for class 'epx'
plot(x, max.cutoff = min(100, length(x$Y)), plot.hc = TRUE, ...)
```

Arguments

X	Object of class "epx".
max.cutoff	Maximum number of observations selected, equivalently the maximum shortlist cutoff; default is $min(100, length(x$Y))$.
plot.hc	Whether to make a plot of the hit curve; default is TRUE.
	Further arguments passed to or from other methods.

Details

Order the cases by decreasing phat (predicted probabilities of relevance) values, and plot the expected number and actual number of hits as cases are selected. Cases with tied phat values are grouped together. See hit.curve in order to plot a hit curve in general.

Value

Plot of the hit curve (if plot.hc = TRUE) and a list with the following vectors:

select	Number of observations in each tied phat group; select[1], select[2], are the numbers of observations with the largest predicted probability of relevance (max(phat)), the second largest value in phat, etc.
p	Unique phat values; $p[1]$, $p[2]$, are the largest value in phat, the second largest value in phat, etc.
nhits	Number of hits (truly relevant observations) in each tied phat group.
nhitlast	Number of hits after max cut off observations selected

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Examples

predict.epx

Predict with an "epx" object

Description

Predicted values based on an "epx" object; may specify different base classifier arguments than those used for phalanx-formation.

Usage

```
## S3 method for class 'epx'
predict(object, newdata, classifier.args = list(), ...)
```

Arguments

object Object of class "epx".

newdata An optional data frame specifiying variables with which to predict; if omitted

and classifier.args are not specified, the fitted (ensembled) values are used.

classifier.args

Additional arguments for the base classifier; same base classifier as that used for

phalanx-formation (specified in epx).

. . . Further arguments passed to or from other methods.

Value

Numeric vector of predicted values (double).

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Examples

RKL

Calculate rank last

Description

The performance measure rank last (RKL) is calculated as follows: after ranking the observations in decreasing order via phat, RKL is the rank of the last truly relevant observation. Hence, RKL can take on integer values from 1 to n, where n is the total number of observations. If there are ties, the last object in the tied group determines RKL. That is, if all n objects are tied at the first rank but only one object is truly relevant, RKL will have a value of n.

Usage

```
RKL(y, phat, ...)
```

Arguments

y True (binary) response vector where 1 is the rare/relevant class.

Numeric vector of estimated probabilities of relevance.

Further arguments passed to or from other methods.

Value

Numeric value of RKL.

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Examples

```
## without ties in phat
resp <- c(rep(1, 50), rep(0, 50))
prob <- (1:100)*0.01
RKL(y = resp, phat = prob) # expect 100
resp <- c(rep(0, 50), rep(1, 50))
RKL(y = resp, phat = prob) # expect 50
## with ties in phat
resp <- sample(c(1, 0), 100, replace = TRUE)
prob <- rep(1, 100)
RKL(y = resp, phat = prob) # expect 100</pre>
```

summary.epx

Summarising an "epx" object

Description

```
summary method for class "epx".
```

Usage

```
## S3 method for class 'epx'
summary(object, ...)
```

Arguments

```
objectObject of class "epx" returned by epx....Further arguments passed to or from other methods.
```

Value

Prints a summary of the object returned by the phalanx-formation algorithm epx.

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```
## The summary agrees with
(model$PHALANXES)[[4]]
```

TOP1

Calculate TOP1

Description

The performance measure TOP1 is calculated as follows: after sorting the observations by their predicted probabilities of relevance (phat) in decreasing order so the first ranked observation has the highest probability of relevance, if the first ranked observation is truly relevant, TOP1 has a value of 1. Otherwise TOP1 is 0. If there are ties for the first rank, all the corresponding observations must be truly relevant for TOP1 to score 1.

Usage

```
TOP1(y, phat, ...)
```

Arguments

y True (binary) response vector where 1 is the rare/relevant class.

Numeric vector of estimated probabilities of relevance.

Further arguments passed to or from other methods.

Value

Numeric value of TOP1.

```
## with ties in phat

resp <- c(0, rep(1, 99))
prob <- rep(1, 100)
TOP1(y = resp, phat = prob) # expect 0

resp <- c(1, 1, 1, rep(0, 95), 1, 1)
prob <- c(1, 1, 1, rep(0, 97))
TOP1(y = resp, phat = prob) # expect 1

## no ties in phat
resp <- c(0, rep(1, 99))
prob <- (100:1)*0.01
TOP1(y = resp, phat = prob) # expect 0

resp <- c(1, rep(0, 99))
TOP1(y = resp, phat = prob) # expect 1</pre>
```

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