Package ‘CORElearn’

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Title Classification, Regression and Feature Evaluation
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Description A suite of machine learning algorithms written in C++ with the R interface contains several learning techniques for classification and regression. Predictive models include e.g., classification and regression trees with optional constructive induction and models in the leaves, random forests, kNN, naïve Bayes, and locally weighted regression. All predictions obtained with these models can be explained and visualized with the ‘ExplainPrediction’ package. This package is especially strong in feature evaluation where it contains several variants of Relief algorithm and many impurity based attribute evaluation functions, e.g., Gini, information gain, MDL, and DKM. These methods can be used for feature selection or discretization of numeric attributes. The OrdEval algorithm and its visualization is used for evaluation of data sets with ordinal features and class, enabling analysis according to the Kano model of customer satisfaction. Several algorithms support parallel multithreaded execution via OpenMP. The top-level documentation is reachable through ?CORElearn.

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The package CORElearn is an R port of CORElearn data mining system. It provides various classification and regression models as well as algorithms for feature selection and evaluation. Several algorithms support parallel multithreaded execution via OpenMP (see details in function descriptions). It is possible to run many functions outside the R environment. The description and source code is available on the package web site http://lkm.fri.uni-lj.si/rmarko/software.
Details

The main functions are

- **CoreModel** which constructs classification or regression model.
  - Classification models available:
    - random forests with optional local weighing of basic models
    - decision tree with optional constructive induction in the inner nodes and/or models in the leaves
    - kNN and kNN with Gaussian kernel,
    - naive Bayes.
  - Regression models:
    - regression trees with optional constructive induction in the inner nodes and/or models in the leaves,
    - linear models with pruning techniques
    - locally weighted regression
    - kNN and kNN with Gaussian kernel.
- **predict.CoreModel** predicts with classification model labels and probabilities of new instances. For regression models it returns the predicted function value.
- **plot.CoreModel** graphically visualizes trees and random forest models
- **modelEval** computes some statistics from predictions
- **attrEval** evaluates the quality of the attributes (dependent variables) with the selected heuristic method. Feature evaluation algorithms are various variants of Relief algorithms (ReliefF, RReliefF, cost-sensitive ReliefF, etc), gain ratio, gini-index, MDL, DKM, information gain, MSE, MAE, etc.
- **ordEval** evaluates ordinal attributes with ordEval algorithm and visualizes them with **plot.ordEval**
- **infoCore** outputs certain information about CORElearn methods,
- **helpCore** prints short description of a given parameter,
- **paramCoreIO** reads/writes parameters for given model from/to file,
- **versionCore** outputs version of the package from underlying C++ library.

Some of the internal structures of the C++ part are described in **CORElearn-internal**.

For an automatically generated list of functions use **help(package=CORElearn)** or **library(help=CORElearn)**. This feature is currently not supported on all platforms and may interfere with other means of parallelization used in R, like package parallel. It is tested to work on Windows, Linux, and Mac.

For certain platforms multithreaded execution is not supported, since current set of compilers at CRAN do not fully support OpenMP. Also note that OpenMP execution may interfere with other means of parallelization on certain platforms. E.g., interference with package parallel is reported on Windows which can be prevented by setting parameter maxThreads=1. For platforms other than Linux, Windows, and OsX to support multithreading it is possible to recompile the package with appropriate tools and compilers (modify Makefile or Makefile.win in src folder, or consult authors).

**Author(s)**

Marko Robnik-Sikonja, Petr Savicky
References

Marko Robnik-Sikonja, Igor Kononenko: Theoretical and Empirical Analysis of ReliefF and RR


Majority of these references are available also from [http://lkm.fri.uni-lj.si/rmarko/papers/](http://lkm.fri.uni-lj.si/rmarko/papers/)

See Also

`CoreModel`, `predict.CoreModel`, `plot.CoreModel`, `modelEval`, `attrEval`, `ordEval`, `plot.ordEval`,
`helpCore`, `paramCoreIO`, `infoCore`, `versionCore`, `CORElearn-internal`, `classDataGen`, `regDataGen`, `ordDataGen`.

Examples

```r
# load the package
library(CORElearn)
cat(versionCore(),"\n")

# use iris data set
trainIdxs <- sample(x=nrow(iris), size=0.7*nrow(iris), replace=FALSE)
testIdxs <- c(1:nrow(iris))[!trainIdxs]

# build random forests model with certain parameters
# setting maxThreads to 0 or more than 1 forces
# utilization of several processor cores
modelRF <- CoreModel(Species ~ ., iris[trainIdxs,,], model="rf",
selectionEstimator="MDL", minNodeWeightRF=5,
rfNoTrees=100, maxThreads=1)

print(modelRF) # simple visualization, test also others with function plot

# prediction on testing set
pred <- predict(modelRF, iris[testIdxs,,], type="both")

# compute statistics
mEval <- modelEval(modelRF, iris[["Species"]][testIdxs,,], pred$class, pred$prob)
print(mEval)

## Not run:
# explain predictions on the level of model and individual instances
require(ExplainPrediction)
explainVis(modelRF, iris[trainIdxs,,], iris[testIdxs,,], method="EXPLAIN",
visLevel="model", problemName="iris", fileType="none",
classValue=1, displayColor="color")

# turn on the history in visualization window to see all instances
```
explainVis(modelRF, iris[trainIdxs[, iris[testIdxs[, method="EXPLAIN", visLevel="instance", problemName="iris", fileType="none", classValue=1, displayColor="color"]

## End(Not run)
# Clean up, otherwise the memory is still taken
destroyModels(modelRF) # clean up

# evaluate features in given data set with selected method
# instead of formula interface one can provide just
# the name or index of target variable
estReliefF <- attrEval("Species", iris,
    estimator="ReliefFexpRank", ReliefIterations=30)
print(estReliefF)

# evaluate ordered features with ordEval
profiles <- ordDataGen(200)
est <- ordEval(class ~ ., profiles, ordEvalNoRandomNormalizers=100)
# print(est)

---

attrEval

Attribute evaluation

Description

The method evaluates the quality of the features/attributes/dependent variables specified by the formula with the selected heuristic method. Feature evaluation algorithms available for classification problems are various variants of Relief and ReliefF algorithms (ReliefF, cost-sensitive ReliefF, ...), and impurity-based algorithms (information gain, gain ratio, gini-index, MDL, DKM, etc). For regression problems there are RRReliefF, MSEofMean, MSEofModel, MAEofModel, ... Parallel execution on several cores is supported for speedup.

Usage

attrEval(formula, data, estimator, costMatrix = NULL,
    outputNumericSplits=FALSE, ...)

Arguments

formula Either a formula specifying the attributes to be evaluated and the target variable, or a name of target variable, or an index of target variable.
data Data frame with evaluation data.
estimator The name of the evaluation method.
costMatrix Optional cost matrix used with certain estimators.
outputNumericSplits

Controls of the output contain the best split point for numeric attributes. This is only sensible for impurity based estimators (like information gain, gini, MDL, gain ratio, etc. in classification, and MSEofMean in regression). The default value of parameter binaryEvaluateNumericAttributes = TRUE shall not be modified in this case. If the value of outputNumericSplits = TRUE, the output is a list with attribute evaluations and numeric attributes’ splits (instead of a single vector with evaluations). See the returned value description.

Additional options used by specific evaluation methods as described in helpCore.

Details

The parameter formula can be interpreted in three ways, where the formula interface is the most elegant one, but inefficient and inappropriate for large data sets. See also examples below. As formula one can specify:

an object of class formula used as a mechanism to select features (attributes) and prediction variable (class). Only simple terms can be used and interaction expressed in formula syntax are not supported. The simplest way is to specify just response variable: class ~ . In this case all other attributes in the data set are evaluated. Note that formula interface is not appropriate for data sets with large number of variables.

a character vector specifying the name of target variable, all the other columns in data frame data are used as predictors.

an integer specifying the index of of target variable in data frame data, all the other columns are used as predictors.

The optional parameter costMatrix can provide nonuniform cost matrix to certain cost-sensitive measures (ReliefFexpC, ReliefFavgC, ReliefFpe, ReliefFpa, ReliefFsmp,GainRatioCost, DKM-cost, ReliefKukar, and MDLsmp). For other measures this parameter is ignored. The format of the matrix is costMatrix(true class, predicted class). By default a uniform costs are assumed, i.e., costMatrix(i, i) = 0, and costMatrix(i, j) = 1, for i not equal to j.

The estimator parameter selects the evaluation heuristics. For classification problem it must be one of the names returned by infoCore(what="attrEval") and for regression problem it must be one of the names returned by infoCore(what="attrEvalReg") Majority of these feature evaluation measures are described in the references given below, here only a short description is given. For classification problem they are

"ReliefEqualK" ReliefF algorithm where k nearest instances have equal weight.

"ReliefFexpRank" ReliefF algorithm where k nearest instances have weight exponentially decreasing with increasing rank. Rank of nearest instance is determined by the increasing (Manhattan) distance from the selected instance. This is a default choice for methods taking conditional dependencies among the attributes into account.

"ReliefFbestK" ReliefF algorithm where all possible k (representing k nearest instances) are tested and for each feature the highest score is returned. Nearest instances have equal weights.

"Relief" Original algorithm of Kira and Rendel (1991) working on two class problems.

"InfGain" Information gain.

"GainRatio" Gain ratio, which is normalized information gain to prevent bias to multi-valued attributes.
"MDL" Acronym for Minimum Description Length, presents method introduced in (Kononenko, 1995) with favorable bias for multi-valued and multi-class problems. Might be the best method among those not taking conditional dependencies into account.

"Gini" Gini-index.

"MyopicReliefF" Myopic version of ReliefF resulting from assumption of no local dependencies and attribute dependencies upon class.

"Accuracy" Accuracy of resulting split.

"ReliefFmerit" ReliefF algorithm where for each random instance the merit of each attribute is normalized by the sum of differences in all attributes.

"ReliefFdistance" ReliefF algorithm where k nearest instances are weighed directly with its inverse distance from the selected instance. Usually using ranks instead of distance as in ReliefFexpRank is more effective.

"ReliefFsqrDistance" ReliefF algorithm where k nearest instances are weighed with its inverse square distance from the selected instance.

"DKM" Measure named after Dietterich, Kearns, and Mansour who proposed it in 1996.

"ReliefFexpC" Cost-sensitive ReliefF algorithm with expected costs.

"ReliefFavgC" Cost-sensitive ReliefF algorithm with average costs.

"ReliefFpe" Cost-sensitive ReliefF algorithm with expected probability.

"ReliefFpa" Cost-sensitive ReliefF algorithm with average probability.

"ReliefFsmp" Cost-sensitive ReliefF algorithm with cost sensitive sampling.

"GainRatioCost" Cost-sensitive variant of GainRatio.

"DKMcost" Cost-sensitive variant of DKM.

"ReliefKukar" Cost-sensitive Relief algorithm introduced by Kukar in 1999.

"MDLsmp" Cost-sensitive variant of MDL where costs are introduced through sampling.

"ImpurityEuclid" Euclidean distance as impurity function on within node class distributions.

"ImpurityHellinger" Hellinger distance as impurity function on within node class distributions.

"UniformDKM" Dietterich-Kearns-Mansour (DKM) with uniform priors.

"UniformGini" Gini index with uniform priors.

"UniformInf" Information gain with uniform priors.

"UniformAccuracy" Accuracy with uniform priors.

"EqualDKM" Dietterich-Kearns-Mansour (DKM) with equal weights for splits.

"EqualGini" Gini index with equal weights for splits.

"EqualInf" Information gain with equal weights for splits.

"EqualHellinger" Two equally weighted splits based Hellinger distance.

"DistHellinger" Hellinger distance between class distributions in branches.

"DistAUC" AUC distance between splits.

"DistAngle" Cosine of angular distance between splits.

"DistEuclid" Euclidean distance between splits.

For regression problem the implemented measures are:
"RReliefFequalK"  RReliefF algorithm where k nearest instances have equal weight.
"RReliefFexpRank"  RReliefF algorithm where k nearest instances have weight exponentially decreasing with increasing rank. Rank of nearest instance is determined by the increasing (Manhattan) distance from the selected instance. This is a default choice for methods taking conditional dependencies among the attributes into account.
"RReliefFbestK"  RReliefF algorithm where all possible k (representing k nearest instances) are tested and for each feature the highest score is returned. Nearest instances have equal weights.
"RReliefFwithMSE"  A combination of RReliefF and MSE algorithms.
"MSEofMean"  Mean Squared Error as heuristic used to measure error by mean predicted value after split on the feature.
"MSEofModel"  Mean Squared Error of an arbitrary model used on splits resulting from the feature. The model is chosen with parameter modelTypeReg.
"MAEofModel"  Mean Absolute Error of an arbitrary model used on splits resulting from the feature. The model is chosen with parameter modelTypeReg. If we use median as the model, we get robust equivalent to MSEofMean.
"RReliefFdistance"  RReliefF algorithm where k nearest instances are weighed directly with its inverse distance from the selected instance. Usually using ranks instead of distance as in RReliefFexpRank is more effective.
"RReliefFsqrDistance"  RReliefF algorithm where k nearest instances are weighed with its inverse square distance from the selected instance.

There are some additional parameters ... available which are used by specific evaluation heuristics. Their list and short description is available by calling helpCore. See Section on attribute evaluation. The attributes can also be evaluated via random forest out-of-bag set with function rfAttrEval. Evaluation and visualization of ordered attributes is covered in function ordEval.

Value

The method returns a vector of evaluations for the features in the order specified by the formula. In case of parameter binaryEvaluateNumericAttributes=TRUE the method returns a list with two components: attrEval and splitPointNum. The attrEval contains a vector of evaluations for the features in the order specified by the formula. The splitPointNum contains the split points of numeric attributes which produced the given attribute evaluation scores.

Author(s)

Marko Robnik-Sikonja

References

Some of these references are available also from http://lkm.fri.uni-lj.si/rmarko/papers/
auxTest

See Also
CORElearn, CoreModel, rfAttrEval, ordEval, helpCore, infoCore.

Examples

# use iris data

# run method ReliefF with exponential rank distance
estReliefF <- attrEval(Species ~ ., iris,
   estimator="ReliefFexpRank", ReliefIterations=30)
print(estReliefF)

# alternatively and more appropriate for large data sets
# one can specify just the target variable
# estReliefF <- attrEval("Species", iris, estimator="ReliefFexpRank",
# ReliefIterations=30)

# print all available estimators
infoCore(what="attrEval")

Description
Test functions for the current state of the development.

Usage
testTime()
testClassPseudoRandom(s, k, m)

Arguments

s          Seed.
k          Length of required output.
m          number of streams.

Details
testTime() determines the current time. testClassPseudoRandom(s, k, m) tests the functionality of multiple streams of RNGs.

Value
Depends on the function.
Author(s)
Marko Robnik-Sikonja, Petr Savicky

See Also
CORElearn.

Examples

testTime()

calibrate C

Description
Given probability scores predictedProb as provided for example by a call to predict.CoreModel and using one of available methods given by methods the function calibrates predicted probabilities so that they match the actual probabilities of a binary class 1 provided by correctClass. The computed calibration can be applied to the scores returned by that model.

Usage

calibrate(correctClass, predictedProb, class1=1,
  method = c("isoReg","binIsoReg","binning","mdlMerge"),
  weight=NULL, noBins=10, assumeProbabilities=FALSE)

applyCalibration(predictedProb, calibration)

Arguments

correctClass     A vector of correct class labels for a binary classification problem.
predictedProb    A vector of predicted class 1 (probability) scores. In calibrate method it should be of the same length as correctClass.
class1           A class value (factor) or an index of the class value to be taken as a class to be calibrated.
method           One of isoReg, binIsoReg, binning, or mdlMerge. See details below.
weight           If specified, should be of the same length as correctClass and gives the weights for all the instances, otherwise a default weight of 1 for each instance is assumed.
noBins           The value of parameter depends on the parameter method and specifies desired or initial number of bins. See details below.
assumeProbabilities
  If assumeProbabilities=TRUE the values in predictedProb are expected to be in [0,1] range i.e., probability estimates. assumeProbabilities=FALSE the algorithm can be used as ordinary (isotonic) regression
calibration       The list resulting from a call to calibration and subsequently applied to probability scores returned by the same model.
Details

Depending on the specified method one of the following calibration methods is executed.

- "isoReg" isotonic regression calibration based on pair-adjacent violators (PAV) algorithm.
- "binning" calibration into a pre-specified number of bands given by noBins parameter, trying to make bins of equal weight.
- "binIsoReg" first binning method is executed, following by a isotonic regression calibration.
- "mdlMerge" first intervals are merged by a MDL gain criterion into a prespecified number of intervals, following by the isotonic regression calibration.

If model="binning" the parameter noBins specifies the desired number of bins i.e., calibration bands; if model="binIsoReg" the parameter noBins specifies the number of initial bins that are formed by binning before isotonic regression is applied; if model="mdlMerge" the parameter noBins specifies the number of bins formed after first applying isotonic regression. The most similar bins are merged using MDL criterion.

Value

A function returns a list with two vector components of the same length:

- interval: The boundaries of the intervals. Lower boundary 0 is not explicitly included but should be taken into account.
- calProb: The calibrated probabilities for each corresponding interval.

Author(s)

Marko Robnik-Sikonja

References


A. Niculescu-Mizil, R. Caruana: Predicting Good Probabilities With Supervised Learning. Proceedings of the 22nd International Conference on Machine Learning (ICML'05), 2005

See Also

reliabilityPlot, CORElearn, predict.CoreModel.

Examples

# generate data set separately for training the model, # calibration of probabilities and testing
train <- classDataGen(noInst=200)
cal <- classDataGen(noInst=200)
test <- classDataGen(noInst=200)

# build random forests model with default parameters
modelRF <- CoreModel(class~., train, model="rf", maxThreads=1)
# prediction
predCal <- predict(modelRF, cal, rfPredictClass=FALSE)
predTest <- predict(modelRF, test, rfPredictClass=FALSE)
destroyModels(modelRF) # clean up, model not needed anymore

# calibrate for a chosen class1 and method
class1<-1
calibration <- calibrate(cal$class, predCal$prob[,class1], class1=class1,
method="isoReg",assumeProbabilities=TRUE)

# apply the calibration to the testing set
calibratedProbs <- applyCalibration(predTest$prob[,class1], calibration)
# the calibration of probabilities can be visualized with
# reliabilityPlot function

---

classDataGen  Artificial data for testing classification algorithms

Description
The generator produces classification data with 2 classes, 7 discrete and 3 numeric attributes.

Usage

```r
classDataGen(noInst, t1=0.7, t2=0.9, t3=0.34, t4=0.32,
p1=0.5, classNoise=0)
```

Arguments

- **noInst**: Number of instances to generate.
- **t1, t2, t3**: Parameters, which control the hardness of the discrete attributes.
- **t4**: Parameter, which controls the hardness of the numeric attributes.
- **p1**: Probability of class 1.
- **classNoise**: Proportion of noise in the class variable for classification or virtual class variable for regression.

Details
Class probabilities are \( p1 \) and \( 1 - p1 \), respectively. The conditional distribution of attributes under each of the classes depends on parameters \( t1, t2, t3, t4 \) from \([0,1]\). Attributes \( a7 \) and \( x3 \) are irrelevant for all values of parameters.

Examples of extreme settings of the parameters.
- Setting satisfying \( t1 \times t2 = t3 \) implies no difference between the distributions of individual discrete attributes among the two classes. However, if \( t1 < 1 \), then the joint distribution of them is different for the two classes.
• Setting $t_1 = 1$ and $t_2 = t_3$ implies no difference between the joint distribution of the discrete attributes among the two classes.

• Setting $t_1 = 1$, $t_2 = 1$, $t_3 = 0$ implies disjoint supports of the distributions of $a_1$, $a_2$, $a_4$, $a_5$, so this allows exact classification.

• Setting $t_4 = 1$ implies no difference between the distribution of $x_1, x_2$ between the classes. Setting $t_4 = 0$ allows correct classification with probability one only using $x_1$ and $x_2$.

For class 1 the attributes have distributions

(a1, a2, a3) $D_1(t_1, t_2)$

a4, a5, a6 $D_2(t_3)$

a7 irrelevant attribute, probabilities of \{a,b,c,d\} are $\{1/2, 1/6, 1/6, 1/6\}$

x1, x2, x3 independent normal variables with mean 0 and standard deviation 1, $t_4$, 1

x4, x5 independent uniformly distributed variables on $[0,1]$

For class 2 the attributes have distributions

a1, a2, a3 $D_2(t_3)$

(a4, a5, a6) $D_1(t_1, t_2)$

a7 irrelevant attribute, probabilities of \{a,b,c,d\} are $\{1/2, 1/6, 1/6, 1/6\}$

x1, x2, x3 independent normal variables with mean 0 and st. dev. $t_4$, 1, 1

x4, x5 independent uniformly distributed variables on $[0,1]$

x3 is irrelevant for classification, since it has the same distribution under both classes.

Attributes in a bracket are mutually dependent. Otherwise, the attributes are conditionally independent for each of the two classes. This means that if we consider groups of the attributes such that the attributes in each of the two brackets form a group and each of the remaining attributes forms a group with one element, then for each class, we have 7 groups, which are conditionally independent for the given class. Note that the splitting into groups differs for class 1 and 2.

Distribution $D_1(t_1, t_2)$ consists of three dependent attributes. The distribution of individual attributes depends only on $t_1*t_2$. For a given $t_1*t_2$, the level of dependence decreases with $t_1$ and increases with $t_2$. There are two extreme settings: Setting $t_1 = 1$, $t_2 = t_1*t_2$ has the largest $t_1$ and the smallest $t_2$ and all three attributes are independent. Setting $t_1 = t_1*t_2$, $t_2 = 1$ has the smallest $t_1$ and the largest $t_2$ and also the largest dependence between attributes.

Distribution $D_2(t_3)$ is equal to $D_1(1, t_3)$, so it contains three independent attributes, whose distributions are the same as in $D_1(t_1, t_2)$ for every setting satisfying $t_1*t_2 = t_3$.

In other words, if $t_3 = t_1*t_2$, then the distributions $D_1(t_1, t_2)$ and $D_2(t_3)$ have the same distributions of individual attributes and may differ only in the dependences. There are no in $D_2(t_3)$ and there are some in $D_1(t_1, t_2)$ if $t_1 < 1$.

**Hardness of the discrete part**

Setting $t_1 = 1$ and $t_2 = t_3$ implies no difference between the discrete attributes among the two classes.

Setting satisfying $t_1*t_2 = t_3$ implies no difference between the distributions of individual discrete attributes among the two classes. However, there may be a difference in dependences.

Setting $t_1 = 1$, $t_2 = 1$, $t_3 = 0$ implies disjoint supports of the distributions of $a_1$, $a_2$, $a_4$, $a_5$, so this allows exact classification.
**Hardness of the continuous part**

Depends monotonically on t4. Setting t4 = 1 implies no difference between the classes. Setting t4 = 0 allows correct classification with probability one.

**Value**

The method `classDataGen` returns a data frame with `noInst` rows and 11 columns. Range of values of the attributes and class are

- a1: 0,1
- a2: 0,1
- a3: a,b,c,d
- a4: 0,1
- a5: 0,1
- a6: a,b,c,d
- a7: a,b,c,d
- x1: numeric
- x2: numeric
- x3: numeric
- class: 1,2

For detailed specification of attributes (columns) see details section below.

**Author(s)**

Petr Savicky

**See Also**

`regDataGen`, `ordDataGen`, `CoreModel`.

**Examples**

```r
# prepare a classification data set
classData <- classDataGen(noInst=200)

# build random forests model with certain parameters
modelRF <- CoreModel(class~., classData, model="rf",
                      selectionEstimator="MDL", minNodeWeightRF=5,
                      rfNoTrees=100, maxThreads=1)
print(modelRF)
destroyModels(modelRF) # clean up
```
classPrototypes

The typical instances of each class - class prototypes

Description

For each class the most typical instances are returned based on the highest predicted probability for each class.

Usage

classPrototypes(model, dataset, noPrototypes=10)

Arguments

model a CoreModel model.
dataset a dataset from which to get prototypes.
nPrototypes number of instances of each class to return

Details

The function uses predict.CoreModel(model, dataset) for prediction of the dataset with model. Based on the returned probabilities, it selects the nPrototypes instances with highest probabilities for each class to be typical representatives of that class, i.e., prototypes. The prototypes can be visualized by calling e.g.,
plot(model, dataset, rfGraphType="prototypes", noPrototypes = 10).

Value

A list with the most typical nPrototypes instances is returned. The list has the following attributes.

 prototopes vector with indexes of the most typical instances
clustering vector with class assignments for typical instances in vector instances
levels the names of the class values.

Author(s)

John Adeyanju Alao (as a part of his BSc thesis) and Marko Robnik-Sikonja (thesis supervisor)

References


See Also

predict.CoreModel, plot.CoreModel.
Examples

```r
dataset <- iris
md <- CoreModel(Species ~ ., dataset, model="rf", rfNoTrees=30,maxThreads=1)
typical <- classPrototypes(md, dataset, 10)
destroyModels(md) # clean up
```

Description

The package CORElearn is an R port of CORElearn data mining system. This document is a short description of the C++ part which can also serve as a standalone Linux or Windows data mining system, its organization and main classes and data structures.

Details

The C++ part is called from R functions collected in file Rinterface.R. The C++ functions called from R and providing interface to R are collected in Rfront.cpp and Rconvert.cpp. The front end for standalone version is in file frontend.cpp. For many parts of the code there are two variants, classification and regression one. Regression part usually has Reg somewhere in its name. The main classes are

- `marray`, `mmatrix` are templates for storing vectors and matrixes
- `dataStore` contains data storage and data manipulation methods, of which the most important are
  - `mmatrix<int> DiscData, DiscPredictData` contain values of discrete attributes and class for training and prediction (optional). In classification column 0 always stores class values.
  - `mmatrix<double> ContData, ContPredictData` contain values of numeric attribute and prediction values for training and prediction (optional). In regression column 0 always stores target values.
  - `marray<attribute> AttrDesc` with information about attributes’ types, number of values, min, max, column index in DiscData or ContData, ...
- `estimation`, `estimationReg` evaluate attributes with different purposes: decision/regression tree splitting, binarization, discretization, constructive induction, feature selection, etc. Because of efficiency these classes store its own data in
  - `mmatrix<int> DiscValues` containing discrete attributes and class values,
  - `mmatrix<double> ContValues` containing numeric attribute and prediction values.
- `Options` stores and handles all the parameters of the system.
- `featureTree, regressionTree` build all the models, predict with them, and create output.

Author(s)

Marko Robnik-Sikonja
CoreModel

Build a classification or regression model

Description

Builds a classification or regression model from the data and formula with given parameters. Classification models available are

- random forests, possibly with local weighing of basic models (parallel execution on several cores),
- decision tree with constructive induction in the inner nodes and/or models in the leaves,
- kNN and weighted kNN with Gaussian kernel,
- naive Bayesian classifier.

Regression models:

- regression trees with constructive induction in the inner nodes and/or models in the leaves,
- linear models with pruning techniques,
- locally weighted regression,
- kNN and weighted kNN with Gaussian kernel.

Function cvCoreModel applies cross-validation to estimate predictive performance of the model.

Usage

CoreModel(formula, data,
model=c("rf","rfNear","tree","knn","knnKernel","bayes","regTree"),
costMatrix=NULL,...)
cvCoreModel(formula, data,
model=c("rf","rfNear","tree","knn","knnKernel","bayes","regTree"),
costMatrix=NULL, folds=10, stratified=TRUE, returnModel=TRUE, ...)

Arguments

formula Either a formula specifying the attributes to be evaluated and the target variable, or a name of target variable, or an index of target variable.
data Data frame with training data.
model The type of model to be learned.
costMatrix Optional misclassification cost matrix used with certain models.
folds An integer, specifying the number of folds to use in cross-validation of model.
stratified A boolean specifying if cross-validation is to be stratified for classification problems, i.e. shall all folds have the same distribution of class values.

returnModel If TRUE the function cvCoreModel estimates predictive performance using cross-validation and returns the model build on the whole data set. If returnModel=FALSE the function only evaluates the model using cross-validation.

... Options for building the model. See helpCore.

Details

The parameter formula can be interpreted in three ways, where the formula interface is the most elegant one, but inefficient and inappropriate for large data sets. See also examples below. As formula one can specify:

an object of class formula used as a mechanism to select features (attributes) and prediction variable (class). Only simple terms can be used and interaction expressed in formula syntax are not supported. The simplest way is to specify just response variable: class ~ . In this case all other attributes in the data set are evaluated. Note that formula interface is not appropriate for data sets with large number of variables.

a character vector specifying the name of target variable, all the other columns in data frame data are used as predictors.

an integer specifying the index of target variable in data frame data, all the other columns are used as predictors.

Parameter model controls the type of the constructed model. There are several possibilities:

"rf" random forests classifier as defined by (Breiman, 2001) with some extensions,

"rfNear" random forests classifier with basic models weighted locally (Robnik-Sikonja, 2005),

"tree" decision tree with constructive induction in the inner nodes and/or models in the leaves,

"knn" k nearest neighbors classifier,

"knnKernel" weighted k nearest neighbors classifier with distance taken into account through Gaussian kernel,

"bayes" naive Bayesian classifier,

"regTree" regression trees with constructive induction in inner nodes and/or models in leaves controlled by modelTypeReg parameter. Models used in leaves of the regression tree can also be used as stand-alone regression models using option minNodeWeightTree=Inf (see examples below):

- linear models with pruning techniques
- locally weighted regression
- kNN and kNN with Gaussian kernel.

There are many additional parameters ... available which are used by different models. Their list and description is available by calling helpCore. Evaluation of attributes is covered in function attrEval.

The optional parameter costMatrix can provide nonuniform cost matrix for classification problems. For regression problem this parameter is ignored. The format of the matrix is costMatrix(true class, predicted class). By default uniform costs are assumed, i.e., costMatrix(i, i) = 0, and costMatrix(i, j) = 1, for i not equal to j.
Value

The created model is not returned as a R structure. It is stored internally in the package memory space and only its pointer (index) is returned. The maximum number of models that can be stored simultaneously is a parameter of the initialization function initCore and defaults to 16384. Models, which are not needed, may be deleted in order to free the memory using function destroyModels. By referencing the returned model, any of the stored models may be used for prediction with predict.CoreModel. What the function actually returns is a list with components:

- modelID index of internally stored model,
- terms description of prediction variables and response,
- class.lev class values for classification problem, null for regression problem,
- model the type of model used, see parameter model,
- formula the formula parameter passed.

The function cvCoreModel evaluates the model using cross-validation and function modelEval to return these additional components:

- avgs A vector with average values of each evaluation metric obtained from modelEval.
- stds A vector with standard deviations of each evaluation metric from modelEval.
- evalList A list, where each component is an evaluation metric from modelEval. Each component contains results of cross-validated runs.

In case returnModel=FALSE the function only returns the above three components are keeps no model.

Author(s)

Marko Robnik-Sikonja, Petr Savicky

References


Marko Robnik-Sikonja: CORE - a system that predicts continuous variables. Proceedings of ERK’97, Portoroz, Slovenia, 1997

Marko Robnik-Sikonja, Igor Kononenko: Discretization of continuous attributes using ReliefF. Proceedings of ERK’95, B149-152, Ljubljana, 1995

Majority of these references are available from http://lkm.fri.uni-lj.si/rmarko/papers/

See Also

CORElearn, predict.CoreModel, modelEval, attrEval, helpCore, paramCoreIO.
# use iris data set
trainIdxs <- sample(x=nrow(iris), size=0.7*nrow(iris), replace=FALSE)
testIdxs <- c(1:nrow(iris)[-trainIdxs])

# build random forests model with certain parameters
# setting maxThreads to 0 or more than 1 forces
# utilization of several processor cores
modelRF <- CoreModel(Species ~ ., iris[trainIdxs,], model="rf",
                      selectionEstimator="MDL", minNodeWeightRF=5,
                      rfNoTrees=100, maxThreads=1)
print(modelRF)  # simple visualization, test also others with function plot

# prediction on testing set
pred <- predict(modelRF, iris[testIdxs,], type="both")
meval <- modelEval(modelRF, iris[["Species"]][testIdxs], pred$class, pred$prob)
print(meval)  # evaluation of the model

# visualization of individual predictions and the model
## Not run:
require(ExplainPrediction)
explainVis(modelRF, iris[trainIdxs,], iris[testIdxs,], method="EXPLAIN",
           visLevel="model", problemName="iris", fileType="none",
           className=1, displayColor="color")

## End(Not run)
destroyModels(modelRF)  # clean up

# build decision tree with naive Bayes in the leaves
# more appropriate for large data sets one can specify just the target variable
modelDT <- CoreModel("Species", iris, model="tree", modelType=4)
print(modelDT)
destroyModels(modelDT)  # clean up

# build regression tree similar to CART
instReg <- regDataGen(200)
modelRT <- CoreModel(response~., instReg, model="regTree", modelTypeReg=1)
print(modelRT)
destroyModels(modelRT)  # clean up

# build kNN kernel regressor by preventing tree splitting
modelKernel <- CoreModel(response~., instReg, model="regTree",
                           modelTypeReg=7, minNodeWeightTree=Inf)
print(modelKernel)
destroyModels(modelKernel)  # clean up

## Not run:
# A more complex example
# Test accuracy of random forest predictor with 20 trees on iris data
# using 10-fold cross-validation.
ncases <- nrow(iris)
ind <- ceiling(10*(1:ncases)/ncases)
ind <- sample(ind,length(ind))
pred <- rep(NA,ncases)
fit <- NULL
for (i in unique(ind)) {
  # Delete the previous model, if there is one.
  fit <- CoreModel(Species ~ ., iris[ind!=i,], model="rf",
                   rfNoTrees=20, maxThreads=1)
  pred[ind==i] <- predict(fit, iris[ind==i,], type="class")
  if (!is.null(fit)) destroyModels(fit) # dispose model no longer needed
}
table(pred,iris$Species)

## End(Not run)

# a simpler way to estimate performance using cross-validation
model <- cvCoreModel(Species ~ ., iris, model="rf", rfNoTrees=20,
                      folds=10, stratified=TRUE, returnModel=TRUE,
                      maxThreads=1)
model$avgs

cvGen

cvGen

Cross-validation and stratified cross-validation

Description

Generate indices for cross-validation and stratified cross-validation

Usage

cvGen(n, k)
cvGenStratified(classVal,k)
gatherFromList(lst)

Arguments

- **n**: The number of instances in a data set.
- **k**: The number of folds in cross-validation.
- **classVal**: A vector of factors representing class values.
- **lst**: A list of lists from which we collect results of the same components.
The functions `cvGen` and `cvGenStratified` generate indices of instances from a data set which can be used in cross-validation. The function `cvGenStratified` generates the same distribution of class values in each fold. The function `gatherFromList` is an auxiliary function helping in collection of results, see the example below.

The functions `cvGen` and `cvGenStratified` return a vector of indices indicating fold membership i.e. from 1:k. The function `codegatherFromList` returns a list with components containing elements of the same name.

Marko Robnik-Sikonja

See Also

CORElearn.

Examples

```r
data <- iris
folds <- 10
foldIdx <- cvGen(nrow(data), k=folds)
evalCore<-list()
for (j in 1:folds) {
  dTrain <- data[foldIdx!=j,]
  dTest <- data[foldIdx==j,]
  modelCore <- CoreModel(Species~., dTrain, model="rf")
  predCore <- predict(modelCore, dTest)
  evalCore[[j]] <- modelEval(modelCore, correctClass=dTest$Species,
                           predictedClass=predCore$class, predictedProb=predCore$prob )
  destroyModels(modelCore)
}
results <- gatherFromList(evalCore)
sapply(results, mean)
```

```r
destroyModels
```

**Description**

Destroys internal representation of a given model or all constructed models. As side effect the memory used by the model(s) is freed.

```r
destroyModels(model=NULL)
```
Arguments

model The model structure as returned by CoreModel. The default value of NULL represents all generated models.

Details

The function destroys the model structure as returned by CoreModel. Subsequent work with this model is no longer possible. If parameter model=NULL (default value) all generated models are destroyed and memory used by their internal representation is freed.

Value

There is no return value.

Author(s)

Marko Robnik-Sikonja, Petr Savicky

See Also

CORElearn, CoreModel.

Examples

# use iris data set

# build random forests model with certain parameters
model <- CoreModel(Species ~ ., iris, model="rf",
   selectionEstimator="MDL", minNodeWeightRF=5,
   rfNoTrees=100, maxThreads=1)

# prediction
pred <- predict(model, iris, rfPredictClass=FALSE)
# print(pred)

# destruction of model's internal representation
destroyModels(model)
Description

The method `discretize` returns discretization bounds for numeric attributes and two auxiliary functions. Discretization can be obtained with one of the three discretization methods: greedy search using given feature evaluation heuristics, equal width of intervals, or equal number of instances in each interval. The attributes and target variable are specified using formula interface, target variable name or index. Feature evaluation algorithms available for classification problems are various variants of Relief and ReliefF algorithms, gain ratio, gini-index, MDL, DKM, information gain, etc. For regression problems there are RReliefF, MSEofMean, MSEofModel, MAEofMode, etc.

Usage

```r
discretize(formula, data, method=c("greedy", "equalFrequency", "equalWidth"), estimator, discretizationLookahead=3, discretizationSample=0, maxBins=0, equalDiscBins=4, ...)
applyDiscretization(data, boundsList, noDecimalsInValueName=2)
intervalMidPoint(data, boundsList, midPointMethod=c("equalFrequency", "equalWidth"))
```

Arguments

- `formula`: Either a formula specifying the attributes to be evaluated and the target variable, or a name of target variable, or an index of target variable.
- `data`: Data frame with data.
- `method`: Three discretization methods are available. With `method="greedy"` greedy search using given feature evaluation heuristics is selected, while "equalFrequency" and "equalWidth" select equal frequency (the same number of instances in each interval) and equal width discretization, respectively.
- `estimator`: The name of the evaluation method.
- `discretizationLookahead`: Discretization is performed with a greedy algorithm which adds a new boundary, until there is no improvement in evaluation function for `discretizationLookahead` number of times (0=try all possibilities). Candidate boundaries are chosen from a random sample of boundaries, whose size is `discretizationSample`.
- `discretizationSample`: Maximal number of points to try discretization (0=all sensible). Binarization of multivalued discrete features with \( k \) values is performed exhaustively, if \( 2^k - 1 \) is at most `discretizationSample`. Otherwise binarization is done greedily starting from the best separation of a single value. For ReliefF-type measures, binarization of numeric features is performed with `discretizationSample` randomly chosen splits. For other measures, the split is searched exhaustively among all possible splits.
- `maxBins`: The maximal number of discrete bins for numeric attributes used for greedy discretization (0=don’t care). This shall be an integer vector of length equal to
discretize

the number of numeric attributes or an integer which applies to all numeric attributes. The default value of 0 means that the number of bins will be determined greedily taking into account discretizationLookahead.

equalDiscBins
The number of bins used in equal frequency and equal width discretization. This shall be an integer vector of length equal to the number of numeric attributes or an integer which applies to all numeric attributes. The default value is 4.

... Additional options used by specific evaluation methods as described in helpCore.

boundsList
A list of numeric bounds which is applied to numeric attributes in data to produce discrete attributes of type factor. Numeric bounds can be obtained by calling discretize function..

noDecimalsInValueName
With how many decimal places will the numeric feature values be presented in description (i.e., levels) of feature values. The default value is 2, but will be increased if this is necessary to avoid the same description of feature values.

midPointMethod
Two methods to determine the middle points of discretization intervals are available. The “equalFrequency” method select the middle point so that each half-interval contains equal number of instances. The “equalWidth” methods sets middle point to be equally distant from the boundaries.

Details
In method discretize the parameter formula can be interpreted in three ways, where the formula interface is the most elegant one, but inefficient and inappropriate for large data sets. See CoreModel for details.

The estimator parameter selects the evaluation heuristics. For classification problem it must be one of the names returned by infoCore(what="attrEval") and for regression problem it must be one of the names returned by infoCore(what="attrEvalReg"). For details see their description in attrEval.

If the number of supplied vector in maxBins and equalDiscBins is shorter than the number of numeric attributes, the vector is coerced to the required length.

There are some additional parameters ... available which are used by specific evaluation heuristics. Their list and short description is available by calling helpCore. See Section on attribute evaluation.

The function applyDiscretization takes the discretization bounds obtain with function discretize and transforms numeric features in a data set into discrete features.

The function intervalMidPoint takes discretization bounds provided by function discretize and returns middle points of discretization intervals for numeric attributes. The middle points are computed from the data; for lowest/highest interval the minimum/maximum of the values in the data for particular attribute are implicitly taken as an additional left/right boundary point.

Value
The method discretize returns a list of discretization bounds for numeric attributes. One component of a list contains bounds for one attribute. If an attribute has all values equal, value NA is returned. If an attribute has all values equal to NA, it is skipped in the returned list.

The function applyDiscretization returns a data set where all numeric attributes are replaced with their discrete versions.
The function intervalMidPoint returns a list of vectors where each vector contains middle point of discretized intervals.

**Author(s)**

Marko Robnik-Sikonja

**References**


Some of these references are available also from [http://lkm.fri.uni-lj.si/rmarko/papers/](http://lkm.fri.uni-lj.si/rmarko/papers/)

**See Also**

CORElearn, CoreModel, attrEval, helpCore, infoCore.

**Examples**

```r
# use iris data
# run method using estimator ReliefF with exponential rank distance
discBounds <- discretize(Species ~ ., iris, method="greedy",
                        estimator="ReliefFexpRank")
print(discBounds)
discreteIris <- applyDiscretization(iris, discBounds)
prototypePoints <- intervalMidPoint(iris, discBounds,
                        midPointMethod="equalFrequency")

regData <- regDataGen(200)
discretize(response ~ ., regData, method="greedy",
            estimator="RReliefFequalK",
            maxBins=2)

# print all available estimators
#infoCore(what="attrEval")
#infoCore(what="attrEvalReg")
```

**Description**

The method display prints the tree models returned by CoreModel() function. Depending of parameter format the output is prepared for either screen or in dot format.
### Usage

```r
## S3 method for class 'CoreModel'
display(x, format=c("screen","dot"))
```

### Arguments

- `x` The model structure as returned by `CoreModel`.
- `format` The type of output, i.e., prepared for screen display or in dot language.

### Details

The tree based models returned by function `CoreModel` are visualized. Only tree based models supported, including the trees which include other prediction models in their leaves. Tree based models available are decision trees (obtained by using parameter `model="tree"` in `CoreModel`), and regression trees (with `model="regTree"`).

Models in the leaves of decision trees can be set using parameter `modelType` in `CoreModel`. At the moment naive Bayes and kNN are available, for details see `helpCore`.

Models in the leaves of regression trees can be set using parameter `modelTypeReg` in `CoreModel`. At the moment kNN, kernel regression, and several types of linear models are available, for details see `helpCore`.

The output in dot language can be used with graphViz visualization software to create model visualization in various formats.

### Value

The method invisibly returns a printed character vector.

### Author(s)

Marko Robnik-Sikonja

### See Also

`CoreModel`, `plot.CoreModel`.

### Examples

```r
# decision tree
dataset <- CO2
md <- CoreModel(Plant ~ ., dataset, model="tree")
display(md)
destroyModels(md) # clean up

# regression tree
dataset <- CO2
mdr <- CoreModel(uptake ~ ., dataset, model="regTree")
display(mdr, format="dot")
destroyModels(mdr) # clean up
```
getCoreModel  

Conversion of model to a list

Description
Function converts given model from internal structures in C++ to R’s data structures.

Usage
getCoreModel(model)

Arguments
model  The model structure as returned by CoreModel.

Details
The function converts the model referenced by model from C++ internal structures to R’s lists. Currently it is implemented only for random forests models.

Value
For random forest a resulting list contains first all the information on the forest level, followed by the list of trees. For each tree the nodes are recursively nested with indication of node type (leaf or internal node) and than required information for that data type.

Author(s)
Marko Robnik-Sikonja

See Also
CoreModel, CORElearn.

Examples
# uses iris data set
# build random forests model with certain parameters,
# do not make too many and too large trees
modelRF <- CoreModel(Species ~ ., iris, model="rf",  
    selectionEstimator="MDL", minNodeWeightRF=50, 
    rfNoTrees=5, maxThreads=1)
print(modelRF)

# get the structure of the forest
forest <- getCoreModel(modelRF)
# forest
```r
getRFsizes

destroyModels(modelRF) # clean up

getRFsizes  Get sizes of the trees in RF

Description
Get numerical characteristics of the trees in a RF model related to the size and depth.

Usage
getRFsizes(model, type=c("size", "sumdepth"))

Arguments
model The model structure as returned by CoreModel.
type The required characteristics.

Details
Size is the number of leaves. The sum of depths means the sum of the depth of all leaves.

Value
Numerical vector of the length equal to the number of trees in RF.

Author(s)
Petr Savicky

See Also
CoreModel, CORElearn.

Examples
# uses iris data set

# build random forests model with certain parameters,
# do not make too many and too large trees
modelRF <- CoreModel(Species ~ ., iris, model="rf",
  selectionEstimator="MDL", minNodeWeightRF=50,
  rfNoTrees=50, maxThreads=1)

getRFsizes(modelRF)

destroyModels(modelRF) # clean up
```
**getRpartModel**  
*Conversion of a CoreModel tree into a rpart.object*

**Description**

The function converts a given CoreModel model (decision or regression tree) into a rpart.object prepared for visualization with plot function.

**Usage**

```r
getRpartModel(model, dataset)
```

**Arguments**

- `model` A tree model produced by CoreModel
- `dataset` A data set which was used in learning of the model.

**Details**

The conversion creates rpart.object and copies CORElearn internal structures contained in memory controlled by dynamic link library written in C++.

An alternative visualization is accessible via function `display`, which outputs tree structure formatted for screen or in dot format.

**Value**

Function returns a rpart.object.

**Author(s)**

Initial version by John Adeyanju Alao, improvements by Marko Robnik-Sikonja.

**See Also**

CoreModel, plot.CoreModel, rpart.object, display

**Examples**

```
# plot a decision tree directly
dataset <- CO2
md<-CoreModel(Plant ~ ., dataset, model="tree")
plot(md, dataset)

# or indirectly
rpm <- getRpartModel(md, dataset)
# set angle to tan(0.5)=45 (degrees) and length of branches at least 5
plot(rpm, branch=0.5, minbranch=5, compress=TRUE)
# pretty=0 prints full names of attributes,
```
# numbers to 3 decimals, try to make a dendrogram more compact
text(rpm, pretty=0, digits=3)
destroyModels(md) # clean up

# an alternative is to use fancier rpart.plot package
# rpart.plot(rpm) # rpart.plot has many parameters controlling the output
# but it cannot plot models in tree leaves

---

**Description**

The behavior of CORElearn is controlled by several parameters. This is a short overview.

**Details**

There are many different parameters available. Some are general and can be used in many learning, or feature evaluation algorithms. All the values actually used by the classifier / regressor can be written to file (or read from it) using `paramCoreIO`. The parameters for the methods are split into several groups and documented below.

**Attribute/feature evaluation**

The parameters in this group may be used inside model construction via `CoreModel` and feature evaluation in `attrEval`. See `attrEval` for description of relevant evaluation methods.

Parameters `attrEvaluationInstances`, `binaryEvaluation`, `binarySplitNumericAttributes` are applicable to all attribute evaluation methods. In models which need feature evaluation (e.g., trees, random forests) they affect the selection of splits in the nodes. Other parameters may be used only in context sensitive measures, i.e., ReliefF in classification and RReliefF in regression and their variants.

`binaryEvaluation` type: logical, default value: FALSE

Shall we treat all attributes as binary and binarize them before evaluation if necessary. If TRUE, then for all multivalued discrete and all numeric features a search for the best binarization is performed. The evaluation of the best binarization found is reported. If FALSE, then multivalued discrete features are evaluated "as is" with multivalued versions of estimators. With ReliefF-type measures, numeric features are also evaluated "as is". For evaluation of numeric features with other (non-ReliefF-type) measures, they are first binarized or discretized. The choice between binarization and discretization is controlled by `binaryEvaluateNumericAttributes`. Due to performance reasons it is recommended that `binaryEvaluation`=FALSE is used. See also `discretizationSample`.

`binaryEvaluateNumericAttributes` type: logical, default value: TRUE

ReliefF like measures can evaluate numeric attributes intrinsically, others have to discretize or binarize them before evaluation; for those measures this parameter selects binarization (default) or discretization (computationally more demanding).
**multiclassEvaluation** type: integer, default value: 1, value range: 1, 4
multi-class extension for two-class-only evaluation measures (1-average of all-pairs, 2-best of all-pairs, 3-average of one-against-all, 4-best of one-against-all).

**attrEvaluationInstances** type: integer, default value: 0, value range: 0, Inf
number of instances for attribute evaluation (0=all available).

**minNodeWeightEst** type: numeric, default value: 2, value range: 0, Inf
minimal number of instances (weight) in resulting split to take it in consideration.

**ReliefIterations** type: integer, default value: 0, value range: -2, Inf
number of iterations for all variants of Relief (0=DataSize, -1=ln(DataSize) -2=sqrt(DataSize)).

**numAttrProportionEqual** type: numeric, default value: 0.04, value range: 0, 1
used in ramp function, proportion of numerical attribute’s range to consider two values equal.

**numAttrProportionDifferent** type: numeric, default value: 0.1, value range: 0, 1
used in ramp function, proportion of numerical attribute’s range to consider two values different.

**kNearestEqual** type: integer, default value: 10, value range: 0, Inf
number of neighbors to consider in equal k-nearest attribute evaluation.

**kNearestExpRank** type: integer, default value: 70, value range: 0, Inf
number of neighbors to consider in exponential rank distance attribute evaluation.

**quotientExpRankDistance** type: numeric, default value: 20, value range: 0, Inf
quotient in exponential rank distance attribute evaluation.

------

**Decision/regression tree construction**

There are several parameters controlling a construction of the tree model. Some are described here, but also attribute evaluation, stop building, model, constructive induction, discretization, and pruning options described in this document are applicable. Splits in trees are always binary, however, the option binaryEvaluation has influence on the feature selection for the split. Namely, selecting the best feature for the split is done with the given value of binaryEvaluation. If binaryEvaluation=FALSE, the features are first evaluated and the best one is finally binarized. If binaryEvaluation=TRUE, the features are binarized before selection. In this case, a search for the best binarization for all considered features is performed and the best binarizations found are used for splits. The latter option is computationally more intensive, but typically does not produce better trees.

**selectionEstimator** type: character, default value: "MDL", possible values: all from attrEval, section classification estimator for selection of attributes and binarization in classification.

**selectionEstimatorReg** type: character, default value: "RReliefFexpRank", possible values: all from attrEval, section regression estimator for selection of attributes and binarization in regression.

**minReliefEstimate** type: numeric, default value: 0, value range: -1, 1
for all variants of Relief attribute estimator: the minimal evaluation of attribute to consider the attribute useful in further processing.

**minInstanceWeight** type: numeric, default value: 0.05, value range: 0, 1
minimal weight of an instance to use it further in splitting.
Stop tree building

During tree construction the node is recursively split, until certain condition is fulfilled.

- **minNodeWeightTree** type: numeric, default value: 5, value range: 0, Inf
  minimal number of instances (weight) of a leaf in the decision or regression tree model.

- **minNodeWeightRF** type: numeric, default value: 2, value range: 0, Inf
  minimal number of instances (weight) of a leaf in the random forest tree.

- **relMinNodeWeight** type: numeric, default value: 0, value range: 0, 1
  minimal proportion of training instances in a tree node to split it further.

- **majorClassProportion** type: numeric, default value: 1, value range: 0, 1
  proportion of majority class in a classification tree node to stop splitting it.

- **rootStdDevProportion** type: numeric, default value: 0, value range: 0, 1
  proportion of root’s standard deviation in a regression tree node to stop splitting it.

Models in the tree leaves

In leaves of the tree model there can be various prediction models controlling prediction. For example instead of classification with majority of class values one can use naive Bayes in classification, or a linear model in regression, thereby expanding expressive power of the tree model.

- **modelType** type: integer, default value: 1, value range: 1, 4
  type of models used in classification tree leaves (1=majority class, 2=k-nearest neighbors, 3=k-nearest neighbors with kernel, 4=naive Bayes).

- **modelTypeReg** type: integer, default value: 5, value range: 1, 8
  type of models used in regression tree leaves (1=mean predicted value, 2=median predicted value, 3=linear by MSE, 4=linear by MDL, 5=linear reduced as in M5, 6=kNN, 7=Gaussian kernel regression, 8=locally weighted linear regression).

- **kInNN** type: integer, default value: 10, value range: 0, Inf
  number of neighbors in k-nearest neighbors models (0=all).

- **nnKernelWidth** type: numeric, default value: 2, value range: 0, Inf
  kernel width in k-nearest neighbors models.

- **bayesDiscretization** type: integer, default value: 2, value range: 1, 3
  type of discretization for naive Bayesian models (1=greedy with selection estimator, 2=equal frequency, 3=equal width).

- **discretizationIntervals** type: integer, default value: 4, value range: 1, Inf
  number of intervals in equal frequency or equal width discretizations.

Constructive induction aka. feature construction

The expressive power of tree models can be increased by incorporating additional types of splits. Operator based constructive induction is implemented in both classification and regression. The best construct is searched with beam search. At each step new constructs are evaluated with selected feature evaluation measure. With different types of operators one can control expressions in the interior tree nodes.

- **constructionMode** type: integer, default value: 15, value range: 1, 15
  sum of constructive operators (1=single attributes, 2=conjunction, 4=addition, 8=multiplication); all=1+2+4+8=15
**constructionDepth** type: integer, default value: 0, value range: 0, Inf
   maximal depth of the tree for constructive induction (0=do not do construction, 1=only at root, ...

**noCachedInNode** type: integer, default value: 5, value range: 0, Inf
   number of cached attributes in each node where construction was performed.

**constructionEstimator** type: character, default value: "MDL", possible values: all from attrEval, section classification
   estimator for constructive induction in classification.

**constructionEstimatorReg** type: character, default value: "RReliefFexpRank", possible values: all from attrEval, section regression
   estimator for constructive induction in regression.

**beamSize** type: integer, default value: 20, value range: 1, Inf
   size of the beam in search for best feature in constructive induction.

**maxConstructSize** type: integer, default value: 3, value range: 1, Inf
   maximal size of constructs in constructive induction.

**Attribute discretization and binarization**

Some algorithms cannot deal with numeric attributes directly, so we have to discretize them. Also the tree models use binary splits in nodes. The discretization algorithm evaluates split candidates and forms intervals of values. Note that setting discretizationSample=1 will force random selection of splitting point, which will speed-up the algorithm and may be perfectly acceptable for random forest ensembles.

CORElearn builds binary trees so multivalued discrete attributes have to be binarized i.e., values have to be split into twoa subset, one going left and the other going right in a node. The method used depends on the parameters and the number of attribute values. Possible methods are exhaustive (if the number of attribute values is less or equal maxValues4Exhaustive), greedy ((if the number of attribute values is less or equal maxValues4Greedy) and random ((if the number of attribute values is more than maxValues4Exhaustive). Setting maxValues4Greedy=2 will always randomly select splitting point.

**discretizationLookahead** type: integer, default value: 3, value range: 0, Inf
   Discretization is performed with a greedy algorithm which adds a new boundary, until there is no improvement in evaluation function for discretizationLookahead number of times (0=try all possibilities). Candidate boundaries are chosen from a random sample of boundaries, whose size is discretizationSample.

**discretizationSample** type: integer, default value: 50, value range: 0, Inf
   Maximal number of points to try discretization (0=all sensible). For Relieff-type measures, binarization of numeric features is performed with discretizationSample randomly chosen splits. For other measures, the split is searched among all possible splits.

**maxValues4Exhaustive** type: integer, default value: 7, value range: 2, Inf
   Maximal number of values of a discrete attribute to try finding split exhaustively. If the attribute has more values the split will be searched greedily or selected randomly based on the value of parameter maxValues4Greedy.

**maxValues4Greedy** type: integer, default value: 30, value range: 2, Inf
   Maximal number of values of a discrete attribute to try finding split greedily. If the attribute
has more values the split will be selected randomly. Setting this parameter to 2 will force random but balanced selection of splits which may be acceptable for random forest ensembles and will greatly speed-up tree construction.

Tree pruning

After the tree is constructed, to reduce noise it is beneficial to prune it.

selectedPruner type: integer, default value: 1, value range: 0, 1
decision tree pruning method used (0=none, 1=with m-estimate).

selectedPrunerReg type: integer, default value: 2, value range: 0, 4
regression tree pruning method used (0=none, 1=MDL, 2=with m-estimate, 3=as in M5, 4=error complexity as in CART (fixed alpha)).

mdlModelPrecision type: numeric, default value: 0.1, value range: 0, Inf
precision of model coefficients in MDL tree pruning.

mdlErrorPrecision type: numeric, default value: 0.01, value range: 0, Inf
precision of errors in MDL tree pruning.

mEstPruning type: numeric, default value: 2, value range: 0, Inf
m-estimate for pruning with m-estimate.

alphaErrorComplexity type: numeric, default value: 0, value range: 0, Inf
alpha for error complexity pruning.

Prediction

For some models (decision trees, random forests, naive Bayes, and regression trees) one can smoothe the output predictions. In classification models output probabilities are smoothed and in case of regression prediction value is smoothed.

smoothingType type: integer, default value: 0, value range: 0, 4
default value 0 means no smoothing (in case classification one gets relative frequencies), value 1 stands for additive smoothing, 2 is pure Laplace’s smoothing, 3 is m-estimate smoothing, and 4 means Zadrozny-Elkan type of m-estimate smoothing where smoothingValue is interpreted as \(m \cdot \hat{p}_c\) and \(\hat{p}_c\) is the prior probability of the least probable class value; for regression smoothingType has no effect, as the smoothing is controlled solely by smoothingValue.

smoothingValue type: numeric, default value: 0, value range: 0, Inf
additional parameter for some sorts of smoothing; in classification it is needed for additive, m-estimate, and Zadrozny-Elkan type of smoothing; in case of regression trees 0 means no smoothing and values larger than 0 change prediction value towards the prediction of the models in ascendant nodes.

Random forests

Random forest is quite complex model, whose construction one can control with several parameters. Momentarily only classification version of the algorithm is implemented. Besides parameters in this section one can apply majority of parameters for control of decision trees (except constructive induction and tree pruning).

rfNoTrees type: integer, default value: 100, value range: 1, Inf
number of trees in the random forest.
**rfNoSelAttr** type: integer, default value: 0, value range: -2, Inf
   number of randomly selected attributes in the node (0=\(\sqrt{\text{numOfAttr}}\), -1=\(\log_2(\text{numOfAttr})+1\), -2=all).

**rfMultipleEst** type: logical, default value: FALSE
   use multiple attribute estimators in the forest? If TRUE the algorithm uses some preselected attribute evaluation measures on different trees.

**rfkNearestEqual** type: integer, default value: 30, value range: 0, Inf
   number of nearest instances for weighted random forest classification (0=no weighing).

**rfPropWeightedTrees** type: numeric, default value: 0, value range: 0, 1
   Proportion of trees where attribute probabilities are weighted with their quality. As attribute weighting might reduce the variance between the models, the default value switches the weighing off.

**rfPredictClass** type: logical, default value: FALSE
   shall individual trees predict with majority class (otherwise with class distribution).

### General tree ensembles

In the same manner as random forests more general tree ensembles can be constructed. Additional options control sampling, tree size and regularization.

**rfSampleProp** type: numeric, default value: 0, value range: 0, 1
   proportion of the training set to be used in learning (0=bootstrap replication).

**rfNoTerminals** type: integer, default value: 0, value range: 0, Inf
   maximal number of leaves in each tree (0=build the whole tree).

**rfRegType** type: integer, default value: 2, value range: 0, 2
   type of regularization (0=no regularization, 1=global regularization, 2=local regularization).

**rfRegLambda** type: numeric, default value: 0, value range: 0, Inf
   regularization parameter lambda (0=no regularization).

### Read data directly from files

In case of very large data sets it is useful to bypass R and read data directly from files as the standalone learning system CORElearn does. Supported file formats are C4.5, M5, and native format of CORElearn. See documentation at http://lkm.fri.uni-lj.si/rmarko/software/.

**domainName** type: character,
   name of a problem to read from files with suffixes .dsc, .dat, .names, .data, .cm, and .costs

**dataDirectory** type: character,
   folder where data files are stored.

**NAString** type: character, default value: "?"
   character string which represents missing and NA values in the data files.

### Miscellaneous

**maxThreads** type: integer, default value: 0, value range: 0, Inf
   maximal number of active threads (0=allow OpenMP to set its defaults).
   As side effect, this parameter changes the number of active threads in all subsequent execution (till maxThreads is set again).
Description of certain CORElearn parameters

Description

Depending on parameter what the function prints some information on CORElearn, for example codes of available classification (or regression) attribute evaluation heuristics. For more complete description of the parameters see helpCore.

Usage

infoCore(what=c("attrEval","attrEvalReg"))

Arguments

what Selects the info to be printed.

Details

Depending on the parameter what the function some information on CORElearn.

"attrEval" Prints codes of all available classification attribute evaluation heuristics. These codes can be used as parameters for attribute evaluation methods in learning. It is internally used for validation of parameters. For more complete information see attrEval.

"attrEvalReg" prints codes of all available regression attribute evaluation heuristics. These codes can be used as parameters for attribute evaluation methods in learning. It is internally used for validation of parameters. For more complete information see attrEval.

Value

For what="attrEval" or "attrEvalReg" function returns vector of codes for all implemented classification or regression attribute evaluation heuristics, respectively.
Author(s)
Marko Robnik-Sikonja

See Also
attrEval, helpCore, CoreModel.

Examples
estClass <- infoCore(what="attrEval")
print(estClass)
infoCore(what="attrEvalReg")

modelEval  Statistical evaluation of predictions

Description
Using predictions of given model produced by predict.CoreModel and correct labels, computes some statistics evaluating the quality of the model.

Usage
modelEval(model=NULL, correctClass, predictedClass,
predictedProb=NULL, costMatrix=NULL,
priorClProb = NULL, avgTrainPrediction = NULL, beta = 1)

Arguments
model The model structure as returned by CoreModel, or NULL if some other predictions are evaluated.
correctClass A vector of correct class labels for classification problem and function values for regression problem.
predictedClass A vector of predicted class labels for classification problem and function values for regression problem.
predictedProb An optional matrix of predicted class probabilities for classification.
costMatrix Optional cost matrix can provide nonuniform costs for classification problems.
priorClProb If model=NULL a vector of prior class probabilities shall be provided in case of classification.
avgTrainPrediction If model=NULL mean of prediction values on training set shall be provided in case of regression.
beta For two class problems beta controls the relative importance of precision and recall in F-measure.
Details

The function uses the model structure as returned by `CoreModel`, `predictedClass` and `predictedProb` returned by `predict.CoreModel`. Predicted values are compared with true values and some statistics are computed measuring the quality of predictions. In classification only one of the `predictedClass` and `predictedProb` can be NULL (one of them is computed from the other under assumption that class label is assigned to the most probable class). Some of the returned statistics are defined only for two class problems, for which the confusion matrix specifying the number of instances of true/predicted class is defined as follows,

\[
\begin{array}{ccc}
\text{true/predicted class} & \text{positive} & \text{negative} \\
\text{positive} & \text{true positive (TP)} & \text{false negative (FN)} \\
\text{negative} & \text{false positive (FP)} & \text{true negative (TN)}
\end{array}
\]

Optional cost matrix can provide nonuniform costs for classification problems. For regression problem this parameter is ignored. The costs can be different from the ones used for building the model in `CoreModel` and prediction with the model in `predict.CoreModel`. If no costs are supplied, uniform costs are assumed. The format of the matrix is `costMatrix(true_class, predicted_class)`. By default a uniform costs are assumed, i.e., `costMatrix(i, i) = 0`, and `costMatrix(i, j) = 1`, for i not equal to j. See the example below.

If a non-CORElearn model is evaluated, one should set model=NULL, and a vector of prior of class probabilities `priorClProb` shall be provided in case of classification, and in case of regression `avgTrainPrediction` shall be the mean of prediction values (estimated on a e.g., training set).

Value

For classification problem function returns list with the components

- **accuracy**: classification accuracy, for two class problems this would equal
  \[
  \text{accuracy} = \frac{TP + TN}{TP + FN + FP + TN}
  \]

- **averageCost**: average classification cost

- **informationScore**: information score statistics measuring information contents in the predicted probabilities

- **AUC**: Area under the ROC curve

- **predictionMatrix**: matrix of miss-classifications also confusion matrix

- **sensitivity**: sensitivity for two class problems (also called accuracy of the positive class, i.e., acc+, or true positive rate),
  \[
  \text{rmsensitivity} = \frac{TP}{TP + FN}
  \]

- **specificity**: specificity for two class problems (also called accuracy of the negative class, i.e., acc-, or true negative rate),
  \[
  \text{specificity} = \frac{TN}{TN + FP}
  \]
brierScore Brier score of predicted probabilities (the original Brier's definition which scores all the classes not only the correct one)
kappa Cohen’s kappa statistics measuring randomness of the predictions; for perfect predictions kappa=1, for completely random predictions kappa=0
precision precision for two class problems
\[ \text{precision} = \frac{TP}{TP + FP} \]
recall recall for two class problems (the same as sensitivity)
F-measure F-measure giving a weighted score of precision and recall for two class problems
\[ F = \frac{(1 + \beta^2) \cdot \text{recall} \cdot \text{precision}}{\beta^2 \cdot \text{recall} + \text{precision}} \]
G-mean geometric mean of positive and negative accuracy,
\[ G = \sqrt{\text{senstivity} \cdot \text{specificity}} \]
KS Kolmogorov-Smirnov statistics defined for binary classification problems, reports the distance between the probability distributions of positive class for positive and negative instances, see (Hand, 2005), value 0 means no separation, and value 1 means perfect separation,
\[ KS = \max_t |TPR(t) - FPR(t)| \]
see definitions of TPR and FPR below
TPR true positive rate \( TPR = \frac{TP}{TP + FN} \) at maximal value of KS statistics
FPR false positive rate \( FPR = \frac{FP}{FP + TN} \) at maximal value of KS statistics
For regression problem the returned list has components
MSE square root of Mean Squared Error
RMSE Relative Mean Squared Error
MAE Mean Absolute Error
RMAE Relative Mean Absolute Error

Author(s)
Marko Robnik-Sikonja

References
noEqualRows

See Also

CORElearn, CoreModel, predict.CoreModel.

Examples

# use iris data

# build random forests model with certain parameters
model <- CoreModel(Species ~ ., iris, model="rf",
  selectionEstimator="MDL", minNodeWeightRF=5,
  rfNoTrees=100, maxThreads=1)

# prediction with node distribution
pred <- predict(model, iris, rfPredictClass=FALSE)

# Model evaluation
mEval <- modelEval(model, iris[["Species"]], pred$class, pred$prob)
print(mEval)

# use nonuniform cost matrix
noClasses <- length(levels(iris[["Species"]]))
costMatrix <- 1 - diag(noClasses)
costMatrix[3,1] <- costMatrix[3,2] <- 5 # assume class 3 is more valuable
mEvalCost <- modelEval(model, iris[["Species"]], pred$class, pred$prob,
  costMatrix=costMatrix)
print(mEvalCost)

destroyModels(model) # clean up

noEqualRows

Number of equal rows in two data sets

Description

Counts number of equal rows in two data sets. The two data sets shall have equal number of columns.

Usage

noEqualRows(data1, data2, tolerance=1e-5, countOnce=TRUE)

Arguments

data1 The first data set.
data2 The second data set.
tolerance Tolerated difference between two rows.
countOnce Shall each equal row in data1 be counted just once, or number of rows it is equal to in data2.
Details

Rows are compared using column-wise comparisons. The sum of differences up to a given tolerance are tolerated.

Value

Integer value giving the count of equal instances.

Author(s)

Marko Robnik-Sikonja

See Also

CORElearn.

Examples

```r
# uses two randomly generated data sets
set.seed(12345)
d1 <- classDataGen(100)
d2 <- classDataGen(100)
noEqualRows(d1, d2, tolerance=1e-4)
```

---

ordDataGen | Artificial data for testing ordEval algorithms

Description

The generator produces ordinal data simulating different profiles of attributes: basic, performance, excitement and irrelevant.

Usage

```r
ordDataGen(noInst, classNoise=0)
```

Arguments

- `noInst` Number of instances to generate.
- `classNoise` Proportion of randomly determined values in the class variable.
Details

Problem is described by six important and two irrelevant features. The important features correspond to different feature types from the marketing theory: two basic features ($B_{\text{weak}}$ and $B_{\text{strong}}$), two performance features ($P_{\text{weak}}$ and $P_{\text{strong}}$), two excitement features ($E_{\text{weak}}$ and $E_{\text{strong}}$), and two irrelevant features ($I_{\text{uniform}}$ and $I_{\text{normal}}$). The values of all features are randomly generated integer values from 1 to 5, indicating for example score assigned to each of the features by the survey’s respondent. The dependent variable for each instance (class) is the sum of its features’ effects, which we scale to the uniform distribution of integers 1-5, indicating, for example, an overall score assigned by the respondent.

$$C = b_w(B_{\text{weak}}) + b_s(B_{\text{strong}}) + p_w(P_{\text{weak}}) + p_s(P_{\text{strong}}) + e_w(E_{\text{weak}}) + e_s(E_{\text{strong}})$$

Value

The method returns a data.frame with noInst rows and 9 columns. Range of values of the attributes and class are integers in [1,5]

Author(s)

Marko Robnik-Sikonja

See Also

classDataGen, regDataGen, ordEval,

Examples

```r
#prepare a data set
dat <- ordDataGen(200)

# evaluate ordered features with ordEval
est <- ordEval(class ~ ., dat, ordEvalNoRandomNormalizers=100)
# print(est)
plot(est)
```

Description

The method evaluates the quality of ordered attributes specified by the formula with ordEval algorithm.

Usage

```r
ordEval(formula, data, file=NULL, rndFile=NULL,
        variant=c("allNear","attrDist1","classDist1"), ...)```
Arguments

- **formula**: Either a formula specifying the attributes to be evaluated and the target variable, or a name of target variable, or an index of target variable.
- **data**: Data frame with evaluation data.
- **file**: Name of file where evaluation results will be written to.
- **rndFile**: Name of file where evaluation of random normalizing attributes will be written to.
- **variant**: Name of the variant of ordEval algorithm. Can be any of "allNear", "attrDist1", or "classDist1".
- **...**: Other options specific to ordEval or common to other context-sensitive evaluation methods (e.g., ReliefF).

Details

The parameter **formula** can be interpreted in three ways, where the formula interface is the most elegant one, but inefficient and inappropriate for large data sets. See also examples below. As **formula** one can specify:

- **an object of class** `formula` used as a mechanism to select features (attributes) and prediction variable (class). Only simple terms can be used and interaction expressed in formula syntax are not supported. The simplest way is to specify just response variable: `class ~ .` In this case all other attributes in the data set are evaluated. Note that formula interface is not appropriate for data sets with large number of variables.

- **a character vector** specifying the name of target variable, all the other columns in data frame `data` are used as predictors.

- **an integer** specifying the index of target variable in data frame `data`, all the other columns are used as predictors.

In the data frame `data` take care to supply the ordinal data as factors and to provide equal levels for them (this is not necessary what one gets with `read.table`). See example below.

The output can be optionally written to files `file` and `rndFile`, in a format used by visualization methods in `plotOrdEval`.

The variant of the algorithm actually used is controlled with **variant** parameter which can have values "allNear", "attrDist1", and "classDist1". The default value is "allNear" which takes all nearest neighbors into account in evaluation of attributes. Variant "attrDist1" takes only neighbors with attribute value at most 1 different from current case into account (for each attribute separately). This makes sense when we want to see the thresholds of reinforcement, and therefore observe just small change up or down (it makes sense to combine this with `equalUpDown=TRUE` in `plot.ordEval` function). The "classDist1" variant takes only neighbors with class value at most 1 different from current case into account. This makes sense if we want to observe strictly small changes in upward/downward reinforcement and has little effect in practical applications.

There are some additional parameters (note ...) some of which are common with other context-sensitive evaluation methods (e.g., ReliefF). Their list of common parameters is available in `helpCore` (see subsection on attribute evaluation therein). The parameters specific to **ordEval** are:
ordEval

ordEvalNoRandomNormalizers  type: integer, default value: 0, value range: 0, Inf.
number of randomly shuffled attributes for normalization of each attribute (0=no normalization). This parameter should be set to a reasonably high value (e.g., 200) in order to produce reliable confidence intervals with plot.ordEval. The parameters ordEvalBootstrapNormalize and ordEvalNormalizingPercentile only make sense if this parameter is larger than 0.

ordEvalBootstrapNormalize  type: logical, default value: FALSE
are features used for normalization constructed with bootstrap sampling or random permutation.

ordEvalNormalizingPercentile  type: numeric, default value: 0.025, value range: 0, 0.5
percentile defines the length of confidence interval obtained with random normalization. Percentile t forms interval by taking the \( n \cdot t \) and \( n(1 - t) \) random evaluation as the confidence interval boundaries, thereby forming \( 100(1 - 2t)\% \) confidence interval (t=0.025 gives 95% confidence interval). The value \( n \) is set by ordEvalNoRandomNormalizers parameter.

attrWeights  type: character,
a character vector representing a list of attribute weights in the ordEval distance measure.

Evaluation of attributes without specifics of ordered attributes is covered in function attrEval.

Value

The method returns a list with following components:

- **reinfPosAV**  a matrix of positive reinforcement for attributes’ values,
- **reinfNegAV**  a matrix of negative reinforcement for attributes’ values,
- **anchorAV**  a matrix of anchoring for attributes’ values,
- **noAV**  a matrix containing count for each value of each attribute,
- **reinfPosAttr**  a vector of positive reinforcement for attributes,
- **reinfNegAttr**  a matrix of negative reinforcement for attributes,
- **anchorAttr**  a matrix of anchoring for attributes,
- **noAVattr**  a vector containing count of valid values of each attribute,
- **rndReinfPosAV**  a three dimensional array of statistics for random normalizing attributes’ positive reinforcement for attributes’ values,
- **rndReinfPosAV**  a three dimensional array of statistics for random normalizing attributes’ negative reinforcement for attributes’ values,
- **rndAnchorAV**  a three dimensional array of statistics for random normalizing attributes’ anchoring for attributes’ values,
- **rndReinfPosAttr**  a three dimensional array of statistics for random normalizing attributes’ positive reinforcement for attributes,
- **rndReinfPosAttr**  a three dimensional array of statistics for random normalizing attributes’ negative reinforcement for attributes,
- **rndAnchorAttr**  a three dimensional array of statistics for random normalizing attributes’ anchoring for attributes.
attrNames the names of attributes
valueNames the values of attributes
noAttr number of attributes
ordVal maximal number of attribute values
variant the variant of the algorithm used
file the file to store the results
rndFile the file to store random normalizations

The statistics used are median, 1st quartile, 3rd quartile, low and high percentile selected by ordEvalNormalizingPercentile, mean, standard deviation, and expected probability according to value distribution. With these statistics we can visualize significance of reinforcements using adapted box and whiskers plot.

Author(s)
Marko Robnik-Sikonja

References
Some of the references are available also from http://lkm.fri.uni-lj.si/rmarko/papers/

See Also
plot.ordEval, CORElearn, CoreModel, helpCore, infoCore.

Examples
#prepare a data set
dat <- ordDataGen(200)

# evaluate ordered features with ordEval
est <- ordEval(class ~ ., dat, ordEvalNoRandomNormalizers=100)
# print(est)
printOrdEval(est)
plot(est)
Description

All the parameters of the given model are written directly to file, or read from file into model.

Usage

```r
paramCoreIO(model, fileName, io=c("read","write"))
```

Arguments

- **model**: The model structure as returned by `CoreModel`.
- **fileName**: Name of the parameter file.
- **io**: Controls whether the parameters will be read or written.

Details

The function uses the `model` structure as returned by `CoreModel` and reads or writes all its parameters from/to file. If parameter `io="read"` parameters are read from file `filename`. If parameter `io="write"` parameters are written to file `filename`.

Value

Returns invisible list with parameters passed to C function: `list(modelID, fileName, io)`.

Author(s)

Marko Robnik-Sikonja

See Also

`CORElearn`, `helpCore`.

Examples

```r
# use iris data
# build random forests model with certain parameters
modelRF <- CoreModel(Species ~ ., iris, model="rf",
                     selectionEstimator="MDL", minNodeWeightRF=5,
                     rfNoTrees=50, maxThreads=1)

# writes all the used parameters to file
paramCoreIO(modelRF, "parameters.par", io="write")
# and reads them back into the model
paramCoreIO(modelRF, "parameters.par", io="read")
```
The method `plot` visualizes the models returned by `CoreModel()` function or summaries obtained by applying these models to data. Different plots can be produced depending on the type of the model.

**Usage**

```r
## S3 method for class 'CoreModel'
plot(x, trainSet, rfGraphType=c("attrEval", "outliers", "scaling", "prototypes", "attrEvalCluster"), clustering=NULL, ...)
```

**Arguments**

- `x`: The model structure as returned by `CoreModel`.
- `trainSet`: The data frame containing training data which produced the model `x`.
- `rfGraphType`: The type of the graph to produce for random forest models. See details.
- `clustering`: The clustering of the training instances used in some model types. See details.
- `...`: Other options controlling graphical output passed to additional graphical functions.

**Details**

The output of function `CoreModel` is visualized. Depending on the model type, different visualizations are produced. Currently, classification tree, regression tree, and random forests are supported (models "tree", "regTree", "rf", and "rfNear").

For classification and regression trees (models "tree" and "regTree") the visualization produces a graph representing structure of classification and regression tree, respectively. This process exploits graphical capabilities of `rpart.plot` package. Internal structures of `CoreModel` are converted to `rpart.object` and then visualized by calling `rpart.plot` using default parameters. Any additional parameters are passed on to this function. For further control use the `getRpartModel` function and call the function `rpart.plot` or `plot.rpart` with different parameters. Note that `rpart.plot` can only display a single value in a leaf, which is not appropriate for model trees using e.g., linear regression in the leaves. For these cases function `display` is a better alternative.

For random forest models (models "rf" and "rfNear") different types of visualizations can be produced depending on the `graphType` parameter:
- "attrEval" the attributes are evaluated with random forest model and the importance scores are then visualized. For details see rfAttrEval.
- "attrEvalClustering" similarly to the "attrEval" the attributes are evaluated with random forest model and the importance scores are then visualized, but the importance scores are generated for each cluster separately. The parameter clustering provides clustering information on the trainSet. If clustering parameter is set to NULL, the class values are used as clustering information and visualization of attribute importance for each class separately is generated. For details see rfAttrEvalClustering.
- 'outliers' the random forest proximity measure of training instances in trainSet is visualized and outliers for each class separately can be detected. For details see rfProximity and rfOutliers.
- 'prototypes' typical instances are found based on predicted class probabilities and their values are visualized (see classPrototypes).
- "scaling" returns a scaling plot of training instances in a two dimensional space using random forest based proximity as the distance (see rfProximity and a scaling function cmdscale).

Value

The method returns no value.

Author(s)

John Adeyanju Alao (initial implementation) and Marko Robnik-Sikonja (integration, improvements)

References


See Also

CoreModel, rfProximity, pam, rfClustering, rfAttrEvalClustering, rfOutliers, classPrototypes, cmdscale

Examples

# decision tree
dataset <- iris
md <- CoreModel(Species ~ ., dataset, model="tree")
plot(md, dataset) # additional parameters are passed directly to rpart.plot

# Additional visualizations can be obtained by explicit conversion to rpart.object
#rpm <- getRpartModel(md,dataset)
# and than setting graphical parameters in plot.rpart and text.rpart
#require(rpart)
# E.g., set angle to tan(0.5)=45 (degrees) and length of branches at least 5,
# try to make a dendrogram more compact
#plot(rpm, branch=0.5, minbranch=5, compress=TRUE)
plot.ordEval

Visualization of ordEval results

Description

The method plot visualizes the results of ordEval algorithm with an adapted box-and-whiskers plots. The method printOrdEval prints summary of the results in a text format.

Usage

plotOrdEval(file, rndFile, ...)

## S3 method for class 'ordEval'
plot(x, graphType=c("avBar", "attrBar", "avSlope"), ...)

printOrdEval(x)

Arguments

x The object containing results of ordEval algorithm obtained by calling ordEval. If this object is not given, it has to be constructed from files file and rndFile.
file Name of file where evaluation results of ordEval algorithm were written to.
rndFile Name of file where evaluation of random normalizing attributes by ordEval algorithm were written to.
graphType The type of the graph to produce. Can be any of "avBar", "attrBar", "avSlope".
...
Other options controlling graphical output, used by specific graphical methods. See details.
Details

The output of function `ordEval` either returned directly or stored in files `file` and `rndFile` is read and visualized. The type of graph produced is controlled by `graphType` parameter:

- **avBar** the positive and negative reinforcement of each value of each attribute is visualized as the length of the bar. For each value also a normalizing modified box and whiskers plot is produced above it, showing the confidence interval of the same attribute value under the assumption that the attribute contains no information. If the length of the bar is outside the normalizing whiskers, this is a statistically significant indication that the value is important.
- **attrBar** the positive and negative reinforcement for each attribute is visualized as the length of the bar. This reinforcement is weighted sum of contributions of individual values visualized with `avBar` graph type.
- **avSlope** the positive and negative reinforcement of each value of each attribute is visualized as the slope of the line segment connecting consequent values

The `avBar` and `avSlope` produce several graphs (one for each attribute). In order to see them all on an interactive device use `devAskNewPage`. On some platforms or in RStudio environment the graphical window stores the history and one can browse through recent pages. Alternatively use any of non-interactive devices such as `pdf` or `postscript`. Some support for opening and handling of these devices is provided by function `preparePlot`. The user should take care to call `dev.off` after completion of the operations.

There are some additional optional parameters ... which are important to all or for some graph types.

- **ciType** The type of the confidence interval in "avBar" and "attrBar" graph types. Can be "two.sided", "upper", "lower", or "none". Together with `ordEvalNormalizingPercentile` parameter in `ordEval`, `ciType`, `ciDisplay`, and `ciDecorate` controls the type, length and display of confidence intervals for each value.
- **ciDisplay** The way how confidence intervals are displayed. Can be "box" or "color". The value "box" displays confidence interval as box and whiskers plot above the actual value with whiskers representing confidence percentiles. The value "color" displays only the upper limit of confidence interval, namely the value (represented with a length of the bar) beyond the confidence interval is displayed with more intensive color or shade.
- **ciDecorate** controls if the reinforcement factors stretching outside the confidence intervals of possible random effects are decorated by being circled with an ellipse. The default value NULL means that there are no decorations, other values are interpreted as colors in the function `draw.elipse`, e.g., `ciDecorate="red"` draws red ellipses around statistically significant reinforcemnets.
- **equalUpDown** a boolean specifying if upward and downward reinforcement of the same value are to be displayed side by side on the same level; it usually makes sense to set this parameter to TRUE when specifying a single value differences by setting `variant="attrDist1"` in `ordEval` function.
- **graphTitle** specifies text to incorporate into the title.
- **attrIdx** displays plot for a single attribute with specified index.
- **xlabel** label of lower horizontal axis.
- **ylabLeft** label of the left-hand vertical axis.
• ylabRight label of the right-hand vertical axis.
• colors a vector with four colors specifying colors of reinforcement bars for down, down_beyond, up, and up_beyond, respectively. If set to NULL this produces black and white graph with shades of gray. The colors down_beyond and up_beyond depict the confidence interval if parameter ciDisplay="color".
   The default values are colors=c("green","lightgreen","blue","lightblue").

Value

The method returns no value.

Author(s)

Marko Robnik-Sikonja

References


Some of the references are available also from http://lkm.fri.uni-lj.si/rmarko/papers/

See Also

ordEval, helpCore, preparePlot, CORElearn

Examples

# prepare a data set
dat <- ordDataGen(200)

# evaluate ordered features with ordEval
oe <- ordEval(class ~ ., dat, ordEvalNoRandomNormalizers=200)
plot(oe)
# printOrdEval(oe)

# the same effect we achieve by storing results to files
tmp <- ordEval(class ~ ., dat, file="profiles.oe",
    rndFile="profiles.oer", ordEvalNoRandomNormalizers=200)
plotOrdEval(file="profiles.oe", rndFile="profiles.oer",
    graphType="attrBar")
# clean up for the sake of R package checks
file.remove("profiles.oe")
file.remove("profiles.oer")
predict.CoreModel

Prediction using constructed model

Description

Using a previously built model and new data, predicts the class value and probabilities for classification problem and function value for regression problem.

Usage

```r
## S3 method for class 'CoreModel'
predict(object, newdata, ..., costMatrix=NULL,
        type=c("both","class","probability"))
```

Arguments

- `object`: The model structure as returned by `CoreModel`.
- `newdata`: Data frame with fresh data.
- `costMatrix`: Optional cost matrix can provide nonuniform costs for classification problems.
- `type`: Controls what will be return value in case of classification.
- `...`: Other model dependent options for prediction. See `helpCore`.

Details

The function uses the `object` structure as returned by `CoreModel` and applies it on the data frame `newdata`. The `newdata` must be transformable using the formula specified for building the model (with dependent variable removed). If the dependent variable is present in `newdata`, it is ignored.

Optional cost matrix can provide nonuniform costs for classification problems. For regression problem this parameter is ignored. The costs can be different from the ones used for building the model in `CoreModel`.

Value

For regression model a vector of predicted values for given input instances. For classification problem the parameter `type` controls what is returned. With default value "both" function returns a list with two components class and probabilities containing predicted class values and probabilities for all class values, respectively. With type set to "class" or "probability" the function returns only the selected component as vector or matrix.

Author(s)

Marko Robnik-Sikonja, Petr Savicky

See Also

- `CORElearn`
- `CoreModel`
- `modelEval`
- `helpCore`
- `paramCoreIO`
Examples

# use iris data set

# build random forests model with certain parameters
modelRF <- CoreModel(Species ~ ., iris, model="rf",
                      selectionEstimator="MDL",minNodeWeightRF=5,rfNoTrees=100)
print(modelRF)

# prediction with node distribution
pred <- predict(modelRF, iris, rfPredictClass=FALSE, type="both")
# print(pred)

destroyModels(modelRF) # clean up

preparePlot

Prepare graphics device

Description

Based on provided fileName opens and sets appropriate graphical device: pdf, postscript, interactive graphical window, or (only on windows) windows metafile.

Usage

preparePlot(fileName="Rplot",...)

Arguments

fileName Name of the file to store the output to.
...

Further parameters passed to device.

Details

The function opens the graphical output device based on fileName extension. The extensions .pdf, .ps, .jpg, .bmp, .tif, .png, .tiff or none select pdf, postscript, jpeg, bmp, tiff, png, bitmap or a default (interactive) graphical device.

On Windows also .emf extension is supported which opens win.metafile and creates vector graphics in windows enhanced metafile format.

The extension .tiff opens bitmap device which produces bitmap via postscript device. Therefore it requires Ghostscript to be installed and on the executable path.

Some sensible default values are passed to created devices, but further options can be passed via ....

Value

A plot device is opened and nothing is returned to the R interpreter.
# prepare a data set
dat <- ordDataGen(200)
# evaluate ordered features with ordEval
oe <- ordEval(class ~ ., dat, ordEvalNoRandomNormalizers=200)

# the following line if uncommented will create a separate
# postscript file with given filename for each attribute
# preparePlot("myGraph%03d.ps")

plot(oe)
dev.off()

regDataGen  Artificial data for testing regression algorithms

Description
The generator produces regression data data with 4 discrete and 7 numeric attributes.

Usage
regDataGen(noInst, t1=0.8, t2=0.5, noise=0.1)

Arguments
noInst  Number of instances to generate.
t1, t2  Parameters controlling the shape of the distribution.
noise  Parameter controlling the amount of noise. If noise=0, there is no noise. If noise = 1, then the level of the signal and noise are the same.

Details
The response variable is derived from x4, x5, x6 using two different functions. The choice depends on a hidden variable, which determines weather the response value would follow a linear dependency \( f = x_4 - 2x_5 + 3x_6 \), or a nonlinear one \( f = \cos(4\pi x_4)(2x_5 - 3x_6) \).

Attributes a1, a2, x1, x2 carry some information on the hidden variables depending on parameters t1, t2. Extreme values of the parameters are t1=0.5 and t2=1, when there is no information. On the
other hand, if \( t1=0 \) or \( t1=1 \) then each of the attributes \( a1, a2 \) carries full information. If \( t2=0 \), then each of \( x1, x2 \) carries full information on the hidden variable.

The attributes \( x4, x5, x6 \) are available with a noise level depending on parameter \( \text{noise} \). If \( \text{noise}=0 \), there is no noise. If \( \text{noise}=1 \), then the level of the signal and noise are the same.

**Value**

Returns a `data.frame` with \( \text{noInst} \) rows and 11 columns. Range of values of the attributes and response are:

- \( a1 \): 0,1
- \( a2 \): a,b,c,d
- \( a3 \): 0,1 (irrelevant)
- \( a4 \): a,b,c,d (irrelevant)
- \( x1 \): numeric (gaussian with different sd for each class)
- \( x2 \): numeric (gaussian with different sd for each class)
- \( x3 \): numeric (gaussian, irrelevant)
- \( x4 \): numeric from [0,1]
- \( x5 \): numeric from [0,1]
- \( x6 \): numeric from [0,1]
- \( \text{response} \): numeric

**Author(s)**

Petr Savicky

**See Also**

`classDataGen`, `ordDataGen`, `CoreModel`.

**Examples**

```r
#prepare a regression data set
regData <- regDataGen(noInst=200)

# build regression tree similar to CART
modelRT <- CoreModel(response ~ ., regData, model="regTree", modelTypeReg=1)
print(modelRT)

destroyModels(modelRT) # clean up
```
reliabilityPlot  

Plots reliability plot of probabilities

Description

Given probability scores `probScore` and true probabilities `trueProb` the methods plots one against the other using a selected boxing method which groups scores and probabilities to show calibration of probabilities in given probability bands.

Usage

```r
reliabilityPlot(probScore, trueProb, titleText='', boxing="equipotent", noBins=10, classValue = 1, printWeight=FALSE)
```

Arguments

- `probScore`: A vector of predicted probabilities for a given class `classValue`.
- `trueProb`: A vector of true probabilities for a given `classValue`, should be of the same length as `probScore`.
- `titleText`: The text of the graph title.
- `boxing`: One of "unique", "equidistant" or "equipotent", determines the grouping of probabilities. See details below.
- `noBins`: The value of parameter depends on the parameter `boxing` and specifies the number of bins. See details below.
- `classValue`: A class value (factor) or an index of the class value (integer) for which reliability plot is made.
- `printWeight`: A boolean specifying if box weights are to be printed.

Details

Depending on the specified `boxing` the probability scores are grouped in one of three possible ways

- "unique" each unique probability score forms its own box.
- "equidistant" forms `noBins` equally wide boxes.
- "equipotent" forms `noBins` boxes with equal number of scores in each box.

The parameter `trueProb` can represent either probabilities (in [0, 1] range, in most cases these will be 0s or 1s), or the true class values from which the method will form 0 and 1 values corresponding to probabilities for class value `classValue`.

Value

A function returns a graph containing reliability plot on a current graphical device.

Author(s)

Marko Robnik-Sikonja
rfAttrEval

Attribute evaluation with random forest

Description

The method evaluates the quality of the features/attributes/dependent variables used in the given random forest model.

Usage

rfAttrEval(model)
rfAttrEvalClustering(model, dataset, clustering=NULL)
**Arguments**

- **model**: The model of type rf or rfNear as returned by `CoreModel`.
- **dataset**: Training instances that produced random forest model.
- **clustering**: A clustering vector of dataset training instances used in model.

**Details**

The attributes are evaluated via provided random forest’s out-of-bag sets. Values for each attribute in turn are randomly shuffled and classified with random forest. The difference between average margin of non-shuffled and shuffled instances serves as a quality estimate of the attribute. The function `rfAttrEvalClustering` uses a clustering of the training instances to produce importance score of attributes for each cluster separately. If parameter `clustering` is set to NULL the actual class values of the instances are used as clusters thereby producing the evaluation of attributes specific for each of the class values.

**Value**

In case of `rfAttrEval` a vector of evaluations for the features in the order specified by the formula used to generate the provided model. In case of `rfAttrEvalClustering` a matrix is returned, where each row contains evaluations for one of the clusters.

**Author(s)**

Marko Robnik-Sikonja (thesis supervisor) and John Adeyanju Alao (as a part of his BSc thesis)

**References**


**See Also**

`CORElearn`, `CoreModel`, `attrEval`.

**Examples**

```r
# build random forests model with certain parameters
modelRF <- CoreModel(Species ~ ., iris, model="rf",
  selectionEstimator="MDL", minNodeWeightRF=5,
  rfNoTrees=100, maxThreads=1)
rfAttrEval(modelRF) # feature evaluations

x <- rfAttrEval(modelRF) # feature evaluations for each class
print(x)

destroyModels(modelRF) # clean up
```
rfClustering

Random forest based clustering

Description

Creates a clustering of random forest training instances. Random forest provides proximity of its training instances based on their out-of-bag classification. This information is usually passed to visualizations (e.g., scaling) and attribute importance measures.

Usage

rfClustering(model, noClusters=4)

Arguments

- model: a random forest model returned by `CoreModel`
- noClusters: number of clusters

Details

The method calls `pam` function for clustering, initializing its distance matrix with random forest based similarity by calling `rfProximity` with argument `model`.

Value

An object of class `pam` representing the clustering (see `?pam.object` for details), the most important being a vector of cluster assignments (named `cluster`) to training instances used to generate the model.

Author(s)

John Adeyanju Alao (as a part of his BSc thesis) and Marko Robnik-Sikonja (thesis supervisor)

References


See Also

`CoreModel rfProximity pam`

Examples

```r
set<-iris
md<-CoreModel(Species ~ ., set, model="rf", rfNoTrees=30, maxThreads=1)
mdCluster<-rfClustering(md, 5)
destroyModels(md) # clean up
```
Description

The method returns internal out-of-bag performance evaluation for given random forests model.

Usage

rfOOB(model)

Arguments

model The model of type rf or rfNear as returned by CoreModel.

Details

The method returns random forest performance estimations obtained via its out-of-bag sets. The performance measures returned are classification accuracy, average classification margin, and correlation between trees in the forest. The classification margin is defined as the difference between probability of the correct class and probability of the most probable incorrect class. The correlation between models is estimated as the ratio between classification margin variance and variance of the forest as defined in (Breiman, 2001).

Value

The list containing three performance measures computed with out-of-bag instances is returned:

accuracy the classification accuracy of the forest,
margin the average margin of classification with the forest,
correlation the correlation between trees in the forest.

Author(s)

Marko Robnik-Sikonja.

References


See Also

CORElearn, CoreModel.
Examples

# build random forests model with certain parameters
modelRF <- CoreModel(Species ~ ., iris, model="rf",
    selectionEstimator="MDL", minNodeWeightRF=5,
    rfNoTrees=100, maxThreads=1)
rfOOB(modelRF)

destroyModels(modelRF) # clean up

rfOutliers

Random forest based outlier detection

Description

Based on random forest instance proximity measure detects training cases which are different to all other cases.

Usage

rfOutliers(model, dataset)

Arguments

model a random forest model returned by CoreModel
dataset a training set used to generate the model

Details

Strangeness is defined using the random forest model via a proximity matrix (see rfProximity). If the number is greater than 10, the case can be considered an outlier according to Breiman 2001.

Value

For each instance from a dataset the function returns a numeric score of its strangeness to other cases.

Author(s)

John Adeyanju Alao (as a part of his BSc thesis) and Marko Robnik-Sikonja (thesis supervisor)

References


See Also

CoreModel, rfProximity, rfClustering.
**rfProximity**

A random forest based proximity function

**Description**

Random forest computes similarity between instances with classification of out-of-bag instances. If two out-of-bag cases are classified in the same tree leaf the proximity between them is incremented.

**Usage**

rfProximity(model, outProximity=TRUE)

**Arguments**

- **model**: a CORElearn model of type random forest.
- **outProximity**: if TRUE, function returns a proximity matrix, else it returns a distance matrix.

**Details**

A proximity is transformed into distance with expression distance=sqrt(1-proximity).

**Value**

Function returns an M by M matrix where M is the number of training instances. Returned matrix is used as an input to other function (see rfOutliers and rfClustering).

**Author(s)**

John Adeyanju Alao (as a part of his BSc thesis) and Marko Robnik-Sikonja (thesis supervisor)

**References**

See Also

CoreModel, rfOutliers, cmdscale, rfClustering.

Examples

md <- CoreModel(Species ~ ., iris, model="rf", rfNoTrees=30, maxThreads=1)
pr <- rfProximity(md, outProximity=TRUE)
# visualization
require(lattice)
levelplot(pr)
destroyModels(md) # clean up

saveRF

Saves/loads random forests model to/from file

Description

saveRF: the internal structure of given random forests model is saved to file. loadRF: the internal structure of random forests model is loaded from given file and a model is created and returned.

Usage

saveRF(model, fileName)
loadRF(fileName)

Arguments

model The model structure as returned by CoreModel.
fileName Name of the file to save/load the model to/from.

Details

The function saveRF saves the internal structure of given random forests model to file. The structures from C++ code are stored to the file with specified file, while internal structures from R are stored to file named fileName.Rda. The model must be a valid structure returned by CoreModel.

The function loadRF loads the internal structure of random forests saved in a specified files and returns access to it.

Value

saveRF invisibly returns some debugging information, while loadRF returns a loaded model as a list, similarly to CoreModel.
Author(s)

Marko Robnik-Sikonja

See Also

CORElearn, CoreModel.

Examples

# use iris data set

# build random forests model with certain parameters
modelRF <- CoreModel(Species ~ ., iris, model="rf",
                       selectionEstimator="MDL", minNodeWeightRF=5,
                       rfNoTrees=100, maxThreads=1)
print(modelRF)

# prediction with node distribution
pred <- predict(modelRF, iris, rfPredictClass=FALSE, type="both")
# print(pred)

# saves the random forests model to file
saveRF(modelRF, "tempRF.txt")

# restore the model to another model
loadedRF = loadRF("tempRF.txt")

# prediction should be the same
predLoaded <- predict(loadedRF, iris, rfPredictClass=FALSE, type="both")
# print(predLoaded)
# sum of differences should be zero subject to numeric imprecision
sum(pred$probabilities - predLoaded$probabilities)
cat("Are predicted classes of original and retrieved models equal? ",
     all(pred$class == predLoaded$class), "\n")
cat("Are predicted probabilities of original and retrieved model equal? ",
     all(pred$probabilities == predLoaded$probabilities), "\n")

# clean up the models when no longer needed
destroyModels(modelRF)
destroyModels(loadedRF)

# clean up for the sake of R package checks
file.remove("tempRF.txt")
file.remove("tempRF.txt.Rda")
Verification of the CORElearn installation

Description

Performs a partial check of the classification part of CORElearn.

Usage

testCoreClass(continue=TRUE)
testCoreAttrEval(continue=TRUE)
testCoreReg(continue=TRUE)
testCoreOrdEval(continue=TRUE)
testCoreNA(continue=TRUE)
testCoreRPORT(continue=TRUE)
testCoreRand(continue=TRUE)
allTests(continue=TRUE, timed=FALSE)

Arguments

continue Logical. Whether a warning or an error should be generated when a test fails.
timed Logical. Whether the time usage should be printed.

Details

Functions testCoreClass(), testCoreAttrEval(), testCoreReg() evaluate functions CoreModel(), predict.CoreModel(), modelEval(), and attrEval() and perform a partial check of the obtained results.

Function testNA() performs a test of consistency NA and NaN between R and CORElearn.

Functions testCoreRPORT() and testCoreRand() test, whether the R_PORT directive is defined in C code and whether R random number generator is used. These tests are mostly used for debugging.

Function allTests() calls all the above functions and prints a table of the results. If an error is found, a more detailed information is printed and the continuation of the tests depends on the argument continue.

Value

The functions have no output value. The result OK or FAILED is printed.

Author(s)

Marko Robnik-Sikonja, Petr Savicky

See Also

CORElearn.
Examples

allTests() # run all tests and generate an error, if any of the tests fails

<table>
<thead>
<tr>
<th>versionCore</th>
<th>Package version</th>
</tr>
</thead>
</table>

Description

Prints package version obtained from C code.

Usage

versionCore()

Arguments

None.

Details

The function returns the information about the current version obtained from underlying C library `CORElearn`.

Value

Character string with information about the version.

Author(s)

Marko Robnik-Sikonja, Petr Savicky

See Also

`CORElearn`.

Examples

# load the package
library(CORElearn)

# print its version
versionCore()
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