

R topics documented:

arulesCBA-package ......................................................... 2
CBA .................................................................................. 2
CBA_helpers ................................................................. 5
CBA_ruleset ................................................................. 6
discretizeDF:supervised ...................................................... 9
FOIL .............................................................................. 10
LUCS_KDD_CBA .............................................................. 12
Lymphography ............................................................. 14
mineCARs ...................................................................... 15
Mushroom ................................................................. 18
predict.CBA ................................................................. 19
prepareTransactions ....................................................... 20
RCAR ........................................................................... 21
RWeka_CBA ................................................................. 23
transactions2DF ............................................................ 25

Index 27

arulesCBA-package  

arulesCBA: Classification Based on Association Rules

Description

Provides the infrastructure for association rule-based classification including the algorithms CBA, 
CMAR, CPAR, C4.5, FOIL, PART, PRM, RCAR, and RIPPER to build associative classifiers. 

Author(s)

Michael Hahsler

CBA

Classification Based on Association Rules Algorithm (CBA)

Description

Build a classifier based on association rules using the ranking, pruning and classification strategy of 
the CBA algorithm by Liu, et al. (1998).
CBA

Usage

```r
CBA(
  formula,
  data,
  pruning = "M1",
  parameter = NULL,
  control = NULL,
  balanceSupport = FALSE,
  disc.method = "mdlp",
  verbose = FALSE,
  ...
)

pruneCBA_M1(formula, rules, transactions, verbose = FALSE)

pruneCBA_M2(formula, rules, transactions, verbose = FALSE)
```

Arguments

- **formula**: A symbolic description of the model to be fitted. Has to be of form `class ~ .` or `class ~ predictor1 + predictor2`.
- **data**: `arules::transactions` containing the training data or a `data.frame` which is automatically discretized and converted to transactions with `prepareTransactions()`.
- **pruning**: Pruning strategy used: "M1" or "M2".
- **parameter, control**: Optional parameter and control lists for `apriori`.
- **balanceSupport**: BalanceSupport parameter passed to `mineCARs()` function.
- **disc.method**: Discretization method used to discretize continuous variables if data is a `data.frame` (default: "mdlp"). See `discretizeDF.supervised()` for more supervised discretization methods.
- **verbose**: Show progress?
- **...**: For convenience, additional parameters are used to create the parameter control list for `apriori` (e.g., to specify the support and confidence thresholds).
- **rules, transactions**: prune a set of rules using a transaction set.

Details

Implementation the CBA algorithm with the M1 or M2 pruning strategy introduced by Liu, et al. (1998).

Candidate classification association rules (CARs) are mined with the `APRIORI` algorithm but minimum support is only checked for the LHS (rule coverage) and not the whole rule. Rules are ranked by confidence, support and size. Then either the M1 or M2 algorithm are used to perform database coverage pruning and default rule pruning.
Value

Returns an object of class CBA representing the trained classifier.

Author(s)

Ian Johnson and Michael Hahsler

References


See Also

CBA, mineCARs().

Examples

data("iris")

# 1. Learn a classifier using automatic default discretization
classifier <- CBA(Species ~ ., data = iris, supp = 0.05, conf = 0.9)
classifier

# inspect the rule base
inspect(classifier$rules)

# make predictions
predict(classifier, head(iris))
table(pred = predict(classifier, iris), true = iris$Species)

# 2. Learn classifier from transactions (and use verbose)
iris_trans <- prepareTransactions(Species ~ ., iris, disc.method = "mdlp")
iris_trans
classifier <- CBA(Species ~ ., data = iris_trans, supp = 0.05, conf = 0.9, verbose = TRUE)
classifier

# make predictions. Note: response extracts class information from transactions.
predict(classifier, head(iris_trans))
table(pred = predict(classifier, iris_trans), true = response(Species ~ ., iris_trans))
**CBA_helpers**

**Helper Functions For Dealing with Classes**

**Description**

Helper functions to extract the response from transactions or rules, determine the class frequency, majority class, transaction coverage and the uncovered examples per class.

**Usage**

- `classes(formula, x)`
- `response(formula, x)`
- `classFrequency(formula, x, type = "relative")`
- `majorityClass(formula, transactions)`
- `transactionCoverage(transactions, rules)`
- `uncoveredClassExamples(formula, transactions, rules)`
- `uncoveredMajorityClass(formula, transactions, rules)`

**Arguments**

- `formula`: A symbolic description of the model to be fitted.
- `x, transactions`: An object of class `arules::transactions` or `rules`.
- `type`: "relative" or "absolute" to return proportions or absolute counts.

**Value**

- `response`: returns the response label as a factor.
- `classFrequency`: returns the item frequency for each class label as a vector.
- `majorityClass`: returns the most frequent class label in the transactions.

**Author(s)**

Michael Hahsler

**See Also**

`itemFrequency()`, `rules`, `arules::transactions`. 
Examples

data("iris")

iris.disc <- discretizeDF.supervised(Species ~ ., iris)
iris.trans <- as(iris.disc, "transactions")
inspect(head(iris.trans, n = 3))

# convert the class items back to a class label
response(Species ~ ., head(iris.trans, n = 3))

# Class labels
classes(Species ~ ., iris.trans)

# Class distribution. The iris dataset is perfectly balanced.
classFrequency(Species ~ ., iris.trans)

# Majority class
# (Note: since all class frequencies for iris are the same, the first one is returned)
majorityClass(Species ~ ., iris.trans)

# Use for CARs
cars <- mineCARs(Species ~ ., iris.trans, parameter = list(support = 0.3))

#’ # Class labels
classes(Species ~ ., cars)

# Number of rules for each class
classFrequency(Species ~ ., cars, type = "absolute")

# conclusion (item in the RHS) of the rule as a class label
response(Species ~ ., cars)

# How many rules (using the first three rules) cover each transactions?
transactionCoverage(iris.trans, cars[1:3])

# Number of transactions per class not covered by the first three rules
uncoveredClassExamples(Species ~ ., iris.trans, cars[1:3])

# Majority class of the uncovered examples
uncoveredMajorityClass(Species ~ ., iris.trans, cars[1:3])

---

**CBA_ruleset**

*Constructor for Objects for Classifiers Based on Association Rules*

**Description**

Objects for classifiers based on association rules have class CBA. A creator function `CBA_ruleset()` and several methods are provided.
Usage

CBA_ruleset(
  formula,
  rules,
  default,
  method = "first",
  weights = NULL,
  bias = NULL,
  model = NULL,
  discretization = NULL,
  description = "Custom rule set",
  ...
)

Arguments

formula A symbolic description of the model to be fitted. Has to be of form class ~ .. The class is the variable name (part of the item label before =).
rules A set of class association rules mined with mineCARs() or apriori() (from arules).
default Default class. If not specified then objects that are not matched by rules are classified as NA.
method Classification method "first" found rule or "majority".
weights Rule weights for the majority voting method. Either a quality measure available in the classification rule set or a numeric vector of the same length are the classification rule set can be specified. If missing, then equal weights are used
bias Class bias vector.
model An optional list with model information (e.g., parameters).
discretization A list with discretization information used by predict() to discretize data supplied as a data.frame.
description Description field used when the classifier is printed.
... Additional arguments added as list elements to the CBA object.

Details

CBA_ruleset() creates a new object of class CBA using the provides rules as the rule base. For method "first", the user needs to make sure that the rules are predictive and sorted from most to least predictive.

Value

A object of class CBA representing the trained classifier with fields:

formula used formula.
rules the classifier rule base.
default default class label (uses partial matching against the class labels).
method  classification method.
weights  rule weights.
bias  class bias vector if available.
model  list with model description.
discretization  discretization information.
description  description in human readable form.

rules returns the rule base.

Author(s)

Michael Hahsler

See Also

mineCARs()

Examples

```r
## Example 1: create a first-matching-rule classifier with non-redundant rules
## sorted by confidence.
data("iris")

# discretize and create transactions
iris.disc <- discretizeDF.supervised(Species ~ ., iris)
trans <- as(iris.disc, "transactions")

# create rule base with CARs
cars <- mineCARs(Species ~ ., trans, parameter = list(support = .01, confidence = .8))
cars <- cars[!is.redundant(cars)]
cars <- sort(cars, by = "conf")

# create classifier and use the majority class as the default if no rule matches.
cl <- CBA_ruleset(Species ~ .,
  rules = cars,
  default = uncoveredMajorityClass(Species ~ ., trans, cars),
  method = "first")

# look at the rule base
inspect(cl$rules)

# make predictions
prediction <- predict(cl, trans)
table(prediction, response(Species ~ ., trans))
accuracy(prediction, response(Species ~ ., trans))

# Example 2: use weighted majority voting.
cl <- CBA_ruleset(Species ~ .,
  rules = cars,
  method = "first")
```
discretizeDF.supervised

Supervised Methods to Convert Continuous Variables into Categorical Variables

Description

This function implements several supervised methods to convert continuous variables into a categorical variables (factor) suitable for association rule mining and building associative classifiers. A whole data.frame is discretized (i.e., all numeric columns are discretized).

Usage

discretizeDF.supervised(formula, data, method = "mdlp", dig.lab = 3, ...)

Arguments

formula a formula object to specify the class variable for supervised discretization and the predictors to be discretized in the form class ~ . or class ~ predictor1 + predictor2.
data a data.frame containing continuous variables to be discretized
method discretization method. Available are: "mdlp", "caim", "cacc", "ameva", "chi2", "chimerge", "extendedchi2", and "modchi2".
dig.lab integer; number of digits used to create labels.
... Additional parameters are passed on to the implementation of the chosen discretization method.
Details

discretizeDF.supervised() only implements supervised discretization. See discretizeDF() in package arules for unsupervised discretization.

Value

discretizeDF() returns a discretized data.frame. Discretized columns have an attribute "discretized:breaks" indicating the used breaks or and "discretized:method" giving the used method.

Author(s)

Michael Hahsler

See Also

Unsupervised discretization from arules: discretize(), discretizeDF().
Details about the available supervised discretization methods from discretization: discretization::mdlp, discretization::caim, discretization::cace, discretization::ameva, discretization::chi2, discretization::chiM, discretization::extendChi2, discretization::modChi2.

Examples

data("iris")
summary(iris)

# supervised discretization using Species
iris.disc <- discretizeDF.supervised(Species ~ ., iris)
summary(iris.disc)

attributes(iris.disc$Sepal.Length)

# discretize the first few instances of iris using the same breaks as iris.disc
discretizeDF(head(iris), methods = iris.disc)

# only discretize predictors Sepal.Length and Petal.Length
iris.disc2 <- discretizeDF.supervised(Species ~ Sepal.Length + Petal.Length, iris)
head(iris.disc2)

———

FOIL

Use FOIL to learn a rule set for classification

———

Description

Build a classifier rule base using FOIL (First Order Inductive Learner), a greedy algorithm that learns rules to distinguish positive from negative examples.
Usage

FOIL(
  formula,
  data,
  max_len = 3,
  min_gain = 0.7,
  best_k = 5,
  disc.method = "mdlp"
)

Arguments

  formula  A symbolic description of the model to be fitted. Has to be of form class ~ .
            or class ~ predictor1 + predictor2.
  data     A data.frame or arules::transactions containing the training data. Data frames
            are automatically discretized and converted to transactions with prepareTransactions().
  max_len  maximal length of the LHS of the created rules.
  min_gain minimal gain required to expand a rule.
  best_k   use the average expected accuracy (laplace) of the best k rules per class for
            prediction.
  disc.method Discretization method used to discretize continuous variables if data is a data.frame
            (default: "mdlp"). See discretizeDF.supervised() for more supervised discretization methods.

Details

Implements FOIL (Quinlan and Cameron-Jones, 1995) to learn rules and then use them as a classifier following Xiaoxin and Han (2003).

For each class, we find the positive and negative examples and learn the rules using FOIL. Then the rules for all classes are combined and sorted by Laplace accuracy on the training data.

Following Xiaoxin and Han (2003), we classify new examples by

1. select all the rules whose bodies are satisfied by the example;
2. from the rules select the best k rules per class (highest expected Laplace accuracy);
3. average the expected Laplace accuracy per class and choose the class with the highest average.

Value

Returns an object of class CBA representing the trained classifier.

Author(s)

Michael Hahsler
References


Yin, Xiaoxin and Jiawei Han. CPAR: Classification based on Predictive Association Rules, SDM, 2003. doi:10.1137/1.9781611972733.40

Examples

data("iris")

# learn a classifier using automatic default discretization
classifier <- FOIL(Species ~ ., data = iris)
classifier

# inspect the rule base
inspect(classifier$rules)

# make predictions for the first few instances of iris
predict(classifier, head(iris))

---

LUCS_KDD_CBA

Interface to the LUCS-KDD Implementations of CMAR, PRM and CPAR

Description

Interface for the LUCS-KDD Software Library Java implementations of CMAR (Li, Han and Pei, 2001), PRM, and CPAR (Yin and Han, 2003). \textbf{Note:} The Java implementations is not part of \texttt{arulesCBA} and is only free for \texttt{non-commercial use}.

Usage

\begin{verbatim}
FOIL2(formula, data, best_k = 5, disc.method = "mdlp", verbose = FALSE)
CPAR(formula, data, best_k = 5, disc.method = "mdlp", verbose = FALSE)
PRM(formula, data, best_k = 5, disc.method = "mdlp", verbose = FALSE)
CMAR(
    formula,
    data,
    support = 0.1,
    confidence = 0.5,
    disc.method = "mdlp",
    verbose = FALSE
)
\end{verbatim}
Arguments

- **formula**: a symbolic description of the model to be fitted. Has to be of form `class ~ .` or `class ~ predictor1 + predictor2`.
- **data**: A data.frame or `arules::transactions` containing the training data. Data frames are automatically discretized and converted to transactions with `prepareTransactions()`.
- **best_k**: use average expected accuracy of the best k rules per class for prediction.
- **disc.method**: Discretization method used to discretize continuous variables if data is a data.frame (default: "mdlp"). See `discretizeDF.supervised()` for more supervised discretization methods.
- **verbose**: Show verbose output?
- **support, confidence**: minimum support and minimum confidence thresholds for CMAR (range [0, 1]).

Details

- **Requirement**: The code needs a **JDK (Java Software Development Kit) Version 1.8 (or higher)** installation. On some systems (Windows), you may need to set the `JAVA_HOME` environment variable so the system finds the compiler.
- **Memory**: The memory for Java can be increased via R options. For example: `options(java.parameters = "-Xmx1024m")`
- **Note**: The implementation does not expose the min. gain parameter for CPAR, PRM and FOIL2. It is fixed at 0.7 (the value used by Yin and Han, 2001). FOIL2 is an alternative Java implementation to the native implementation of FOIL already provided in the `arulesCBA`. FOIL exposes min. gain.

Value

Returns an object of class **CBA** representing the trained classifier.

References

- Li W., Han, J. and Pei, J. CMAR: Accurate and Efficient Classification Based on Multiple Class-Association Rules, ICDM, 2001, pp. 369-376.
- Yin, Xiaoxin and Jiawei Han. CPAR: Classification based on Predictive Association Rules, SDM, 2003. doi:10.1137/1.9781611972733.40
- Frans Coenen et al. The LUCS-KDD Software Library, [https://cgi.csc.liv.ac.uk/~frans/KDD/Software/](https://cgi.csc.liv.ac.uk/~frans/KDD/Software/)

Examples

```r
# make sure you have a Java SDK Version 1.4.0+ and not a headless installation.
system("java -version")

data("iris")

# build a classifier, inspect rules and make predictions
cl <- CMAR(Species ~ ., iris, support = .2, confidence = .8, verbose = TRUE)
c1
```
inspect(cl$rules)
predict(cl, head(iris))

cl <- CPAR(Species ~ ., iris)
cl

c1 <- PRM(Species ~ ., iris)
cl

c1 <- FOIL2(Species ~ ., iris)
cl

---

**Lymphography**

---

**The Lymphography Domain Data Set (UCI)**

---

**Description**

This is lymphography domain obtained from the University Medical Centre, Institute of Oncology, Ljubljana, Yugoslavia. It was repeatedly used in the machine learning literature.

**Format**

A data frame with 147 observations on the following 19 variables.

- **class** a factor with levels normalfind metastases malignlymph fibrosis
- **lymphatics** a factor with levels normal arched deformed displaced
- **blockofaffere** a factor with levels no yes
- **bloflymphc** a factor with levels no yes
- **bloflymphs** a factor with levels no yes
- **bypass** a factor with levels no yes
- **extravasates** a factor with levels no yes
- **regenerationof** a factor with levels no yes
- **earlyuptakein** a factor with levels no yes
- **lymnodesdimin** a factor with levels 0 1 2 3
- **lymnodesenlar** a factor with levels 1 2 3 4
- **changesinlym** a factor with levels bean oval round
- **defectinnode** a factor with levels no lacunar lacmarginal laccentral
- **changesinnode** a factor with levels no lacunar lacmarginal laccentral
- **changesinstru** a factor with levels no grainy droplike coarse diluted reticular stripped
- **faint**
- **specialforms** a factor with levels no chalices vesicles
- **dislocationof** a factor with levels no yes
- **exclusionofno** a factor with levels no yes
- **noofnodesin** a factor with levels 0-9 10-19 20-29 30-39 40-49 50-59 60-69 >=70
mineCARs

Source

The data set was obtained from the UCI Machine Learning Repository at [http://archive.ics.uci.edu/ml/datasets/Lymphography](http://archive.ics.uci.edu/ml/datasets/Lymphography).

References

This lymphography domain was obtained from the University Medical Centre, Institute of Oncology, Ljubljana, Yugoslavia. Thanks go to M. Zwitter and M. Soklic for providing the data. Please include this citation if you plan to use this database.

Examples

data("Lymphography")
summary(Lymphography)

mineCARs

Mine Class Association Rules

Description

Class Association Rules (CARs) are association rules that have only items with class values in the RHS as introduced for the CBA algorithm by Liu et al., 1998.

Usage

mineCARs(
  formula,
  transactions,
  parameter = NULL,
  control = NULL,
  balanceSupport = FALSE,
  verbose = TRUE,
  ...
)

Arguments

- **formula**: A symbolic description of the model to be fitted.
- **transactions**: An object of class `arules::transactions` containing the training data.
- **parameter, control**: Optional parameter and control lists for `apriori()`.
- **balanceSupport**: logical; if TRUE, class imbalance is counteracted by using class specific minimum support values. Alternatively, a support value for each class can be specified (see Details section).
mineCARs

verbose logical; report progress?

For convenience, the mining parameters for `apriori()` can be specified as.... Examples are the support and confidence thresholds, and the maxlen of rules.

Details

Class association rules (CARs) are of the form

\[ P \Rightarrow c_i, \]

where the LHS \( P \) is a pattern (i.e., an itemset) and \( c_i \) is a single items representing the class label.

Mining parameters. Mining parameters for `apriori()` can be either specified as a list (or object of `arules::APparameter`) as argument `parameter` or, for convenience, as arguments in ... Note: `mineCARs()` uses by default a minimum support of 0.1 (for the LHS of the rules via parameter `originalSupport = FALSE`), a minimum confidence of 0.5 and a maxlen (rule length including items in the LHS and RHS) of 5.

Balancing minimum support. Using a single minimum support threshold for a highly class imbalanced dataset will lead to the problem, that minority classes will only be presented in very few rules. To address this issue, `balanceSupport = TRUE` can be used to adjust minimum support for each class dependent on the prevalence of the class (i.e., the frequency of the \( c_i \) in the transactions) similar to the minimum class support suggested for CBA by Liu et al (2000) we use

\[ \text{minsupp}_i = \text{minsupp}_t \frac{\text{supp}(c_i)}{\max(\text{supp}(C))}, \]

where \( \max(\text{supp}(C)) \) is the support of the majority class. Therefore, the defined minimum support is used for the majority class and then minimum support is scaled down for classes which are less prevalent, giving them a chance to also produce a reasonable amount of rules. In addition, a named numerical vector with a support values for each class can be specified.

Value

Returns an object of class `rules`.

Author(s)

Michael Hahsler

References


Examples

data("iris")

# discretize and convert to transactions
iris.trans <- prepareTransactions(Species ~ ., iris)

# mine CARs with items for "Species" in the RHS.
# Note: mineCars uses a default a minimum coverage (lhs support) of 0.1, a
# minimum confidence of .5 and maxlen of 5
cars <- mineCars(Species ~ ., iris.trans)
inspect(head(cars))

# specify minimum support and confidence

cars <- mineCars(Species ~ ., iris.trans,
                  parameter = list(support = 0.3, confidence = 0.9, maxlen = 3))
inspect(head(cars))

# for convenience this can also be written without a list for parameter using ...
cars <- mineCars(Species ~ ., iris.trans, support = 0.3, confidence = 0.9, maxlen = 3)

# restrict the predictors to items starting with "Sepal"
cars <- mineCars(Species ~ Sepal.Length + Sepal.Width, iris.trans)
inspect(cars)

# using different support for each class
cars <- mineCars(Species ~ ., iris.trans, balanceSupport = c(
    "Species=setosa" = 0.1,
    "Species=versicolor" = 0.5,
    "Species=virginica" = 0.01), confidence = 0.9)
cars

# balance support for class imbalance
data("Lymphography")
Lymphography.trans <- as(Lymphography, "transactions")

classFrequency(class ~ ., Lymphography.trans)

# mining does not produce CARs for the minority classes
cars <- mineCars(class ~ ., Lymphography.trans, support = .3, maxlen = 3)

classFrequency(class ~ ., cars, type = "absolute")

# Balance support by reducing the minimum support for minority classes
cars <- mineCars(class ~ ., Lymphography.trans, support = .3, maxlen = 3,
                 balanceSupport = TRUE)

classFrequency(class ~ ., cars, type = "absolute")

# Mine CARs from regular transactions (a negative class item is automatically added)
data(Groceries)
cars <- mineCars("whole milk" ~ ., Groceries,
                 balanceSupport = TRUE, support = 0.01, confidence = 0.8)
inspect(sort(cars, by = "lift"))
The Mushroom data set includes descriptions of hypothetical samples corresponding to 23 species of gilled mushrooms in the Agaricus and Lepiota Family. It contains information about 8123 mushrooms. 4208 (51.8\% edible and 3916 (48.2\% features plus the class attribute (edible or not).

Format

A data frame with 8123 observations on the following 23 variables.

- **Class** a factor with levels edible poisonous
- **CapShape** a factor with levels bell conical flat knobbed sunken convex
- **CapSurf** a factor with levels fibrous grooves smooth scaly
- **CapColor** a factor with levels buff cinnamon red gray brown pink green purple white yellow
- **Bruises** a factor with levels no bruises
- **Odor** a factor with levels almond creosote foul anise musty none pungent spicy fishy
- **GillAttached** a factor with levels attached free
- **GillSpace** a factor with levels close crowded
- **GillSize** a factor with levels broad narrow
- **GillColor** a factor with levels buff red gray chocolate black brown orange pink green purple white yellow
- **StalkShape** a factor with levels enlarging tapering
- **StalkRoot** a factor with levels bulbous club equal rooted
- **SurfaceAboveRing** a factor with levels fibrous silky smooth scaly
- **SurfaceBelowRing** a factor with levels fibrous silky smooth scaly
- **ColorAboveRing** a factor with levels buff cinnamon red gray brown orange pink white yellow
- **ColorBelowRing** a factor with levels buff cinnamon red gray brown orange pink white yellow
- **VeilType** a factor with levels partial
- **VeilColor** a factor with levels brown orange white yellow
- **RingNumber** a factor with levels none one two
- **RingType** a factor with levels evanescent flaring large none pendant
- **Spore** a factor with levels buff chocolate black brown orange green purple white yellow
- **Population** a factor with levels brown yellow
- **Habitat** a factor with levels woods grasses leaves meadows paths urban waste

Source

The data set was obtained from the UCI Machine Learning Repository at [http://archive.ics.uci.edu/ml/datasets/Mushroom](http://archive.ics.uci.edu/ml/datasets/Mushroom).
References


Examples

data(Mushroom)

summary(Mushroom)

predict.CBA

Model Prediction for Classifiers Based on Association Rules

Description

Predicts classes for new data using a CBA classifier.

Usage

## S3 method for class 'CBA'
predict(object, newdata, type = c("class", "score"), ...)

accuracy(pred, true)

Arguments

object An object of class CBA.
newdata A data.frame or arules::transactions containing rows of new entries to be classified.
type Predict "class" labels. Some classifiers can also return "scores".
... Additional arguments are ignored.
pred, true two factors with the same level representing the predictions and the ground truth (e.g., obtained with response()).

Value

A factor vector with the classification result.

Author(s)

Michael Hahsler
**Examples**

```r
data("iris")

train_id <- sample(seq_len(nrow(iris)), 130)
iris_train <- iris[train_id, ]
iris_test <- iris[-train_id, ]

c1 <- CBA(Species ~., iris_train)
pr <- predict(c1, iris_test)
pr

accuracy(pr, response(Species ~., iris_test))
```

---

**prepareTransactions**  
*Prepare Data for Associative Classification*

**Description**

Converts data.frame into transactions suitable for classification based on association rules.

**Usage**

```r
prepareTransactions(formula, data, disc.method = "mdlp", match = NULL)
```

**Arguments**

- `formula`: the formula.
- `data`: a data.frame with the data.
- `disc.method`: Discretization method used to discretize continuous variables if data is a data.frame (default: "mdlp"). See `discretizeDF.supervised()` for more supervised discretization methods.
- `match`: typically NULL. Only used internally if data is a already a set of transactions.

**Details**

To convert a data.frame into items in a transaction dataset for classification, the following steps are performed:

1. All continuous features are discretized using class-based discretization (default is MDLP) and each range is represented as an item.
2. Factors are converted into items, one item for each level.
3. Each logical is converted into an item.
4. If the class variable is a logical, then a negative class item is added.

Steps 1-3 are skipped if data is already as `arules::transactions` object.
Value

An object of class `arules::transactions` from `arules` with an attribute called "disc_info" that contains information on the used discretization for each column.

Author(s)

Michael Hahsler

See Also

`arules::transactions`, `transactions2DF()`.

Examples

```r
# Perform discretization and convert to transactions
data("iris")
iris_trans <- prepareTransactions(Species ~ ., iris)
inspect(head(iris_trans))

# A negative class item is added for regular transaction data (here "!canned beer")
# Note: backticks are needed in formulas with item labels that contain a space or special character.
data("Groceries")
g2 <- prepareTransactions(`canned beer` ~ ., Groceries)
inspect(head(g2))
```

---

**RCAR**

*Regularized Class Association Rules for Multi-class Problems (RCAR+)*

Description

Build a classifier based on association rules mined for an input dataset and weighted with LASSO regularized logistic regression following RCAR (Azmi, et al., 2019). RCAR+ extends RCAR from a binary classifier to a multi-class classifier and can use support-balanced CARs.

Usage

```r
RCAR(
  formula,
  data,
  lambda = NULL,
  alpha = 1,
  glmnet.args = NULL,
  cv.glmnet.args = NULL,
  parameter = NULL,
  control = NULL,
  balanceSupport = FALSE,
)```
Arguments

formula A symbolic description of the model to be fitted. Has to be of form class ~ . or class ~ predictor1 + predictor2.
data A data.frame or arules::transactions containing the training data. Data frames are automatically discretized and converted to transactions with prepareTransactions().
lambda The amount of weight given to regularization during the logistic regression learning process. If not specified (NULL) then cross-validation is used to determine the best value (see Details section).
alpha The elastic net mixing parameter. alpha = 1 is the lasso penalty (default RCAR), and alpha = 0 the ridge penalty.
cv.glmnet.args, glmnet.args A list of arguments passed on to cv.glmnet() and glmnet(), respectively. See Example section.
parameter, control Optional parameter and control lists for apriori().
balanceSupport balanceSupport parameter passed to mineCARs().
disc.method Discretization method for factorizing numeric input (default: "mdlp"). See discretizeDF.supervised() for more supervised discretization methods.
verbose Report progress?
...

Details

RCAR+ extends RCAR from a binary classifier to a multi-class classifier using regularized multinomial logistic regression via glmnet.

If lambda is not specified (NULL) then cross-validation with the largest value of lambda such that error is within 1 standard error of the minimum is used to determine the best value (see cv.glmnet() also for how to perform cross-validation in parallel).

Value

Returns an object of class CBA representing the trained classifier with the additional field model containing a list with the following elements:

all_rules all rules used to build the classifier, including the rules with a weight of zero.
reg_model them multinomial logistic regression model as an object of class glmnet().
cv contains the results for the cross-validation used determine lambda.
Author(s)

Tyler Giallanza and Michael Hahsler

References


Examples

data(“iris”)

classifier <- RCAR(Species~., iris)
classifier

# inspect the rule base sorted by the larges class weight
inspect(sort(classifier$rules, by = “weight”))

# make predictions for the first few instances of iris
predict(classifier, head(iris))

# inspecting the regression model, plot the regularization path, and
# plot the cross-validation results to determine lambda
str(classifier$model$reg_model)
plot(classifier$model$reg_model)
plot(classifier$model$cv)

# show progress report and use 5 instead of the default 10 cross-validation folds.
classifier <- RCAR(Species~., iris, cv.glmnet.args = list(nfolds = 5), verbose = TRUE)

RWeka_CBA

**CBA classifiers based on rule-based classifiers in RWeka**

Description

Provides CBA-type classifiers based on RIPPER (Cohen, 1995), C4.5 (Quinlan, 1993) and PART (Frank and Witten, 1998) using the implementation in Weka via RWeka (Hornik et al, 2009).

Usage

RIPPER_CBA(formula, data, control = NULL, disc.method = “mdlp”)

PART_CBA(formula, data, control = NULL, disc.method = “mdlp”)

C4.5_CBA(formula, data, control = NULL, disc.method = “mdlp”)
Arguments

- **formula**: A symbolic description of the model to be fitted. Has to be of form `class ~ .` or `class ~ predictor1 + predictor2`.
- **data**: A data.frame or `arules::transactions` containing the training data. Data frames are automatically discretized and converted to transactions with `prepareTransactions()`.
- **control**: Algorithmic control options for R/Weka Rule learners (see Details Section).
- **disc.method**: Discretization method used to discretize continuous variables if data is a data.frame (default: "mdlp"). See `discretizeDF.supervised()` for more supervised discretization methods.

Details

You need to install package **RWeka** to use these classifiers.

See R/Weka functions **RWeka::JRip()** (RIPPER), **RWeka::J48()** (C4.5 rules), **RWeka::PART()** for algorithm details and how control options can be passed on via `control`. An example is given in the Examples Section below.

Memory for **RWeka** can be increased using the R options (e.g., `options(java.parameters = "-Xmx1024m")`) before **RWeka** or **rJava** is loaded or any RWeka-based classigier in this package is used.

Value

Returns an object of class **CBA** representing the trained classifier.

Author(s)

Michael Hahsler

References


Examples

# Java and RWeka need to be installed

## Not run:
data(iris)
# learn a classifier using automatic default discretization
classifier <- RIPPER_CBA(Species ~ ., data = iris)
classifier

# inspect the rule base
inspect(classifier$rules)

# make predictions for the first few instances of iris
predict(classifier, head(iris))

table(predict(classifier, iris), iris$Species)

# C4.5
classifier <- C4.5_CBA(Species ~ ., iris)
inspect(classifier$rules)

# To use algorithmic options (here for PART), you need to load RWeka
library(RWeka)

# control options can be found using the Weka Option Wizard (WOW)
WOW(PART)

# build PART with control option U (Generate unpruned decision list) set to TRUE
classifier <- PART_CBA(Species ~ ., data = iris, control = RWeka::Weka_control(U = TRUE))
classifier
inspect(classifier$rules)
predict(classifier, head(iris))

## End(Not run)

---

**transactions2DF**  
*Convert Transactions to a Data.Frame*

**Description**

Convert transactions back into data.frames by combining the items for the same variable into a single column.

**Usage**

transactions2DF(transactions, itemLabels = FALSE)

**Arguments**

- `transactions`: an object of class `transactions`.
- `itemLabels`: logical; use the complete item labels (variable=level) as the levels in the data.frame? By default, only the levels are used.
Value

Returns a data.frame.

Author(s)

Michael Hahsler

Examples

data(“iris”)
iris_trans <- prepareTransactions(Species ~ ., iris)
iris_trans

# standard conversion
iris_df <- transactions2DF(iris_trans)
head(iris_df)

# use item labels in the data.frame
iris_df2 <- transactions2DF(iris_trans, itemLabels = TRUE)
head(iris_df2)

# Conversion of transactions without variables in itemInfo
data(“Groceries”)
head(transactions2DF(Groceries), 2)

# Conversion of transactions prepared for classification
g2 <- prepareTransactions(‘shopping bags’ ~ ., Groceries)
head(transactions2DF(g2), 2)
Index

* datasets
  - Lymphography, 14
  - Mushroom, 18
* manip
  - discretizeDF.supervised, 9
  - accuracy (predict.CBA), 19
  - apriori(), 7, 15, 16, 22
  - arules::AParameter, 16
  - arules::transactions, 3, 5, 11, 13, 15, 19–22, 24
  - arulesCBA-package, 2
  - CBA, 2, 4, 11, 13, 19, 22, 24
  - CBA_helpers, 5
  - CBA_ruleset, 6
  - classes (CBA_helpers), 5
  - classFrequency (CBA_helpers), 5
  - CMAR (LUCS_KDD_CBA), 12
  - CPAR (LUCS_KDD_CBA), 12
  - cv.glmnet(), 22
  - discretization::ameva, 10
  - discretization::cacc, 10
  - discretization::caim, 10
  - discretization::chi2, 10
  - discretization::chiM, 10
  - discretization::extendChi2, 10
  - discretization::mdlp, 10
  - discretization::modChi2, 10
  - discretize (discretizeDF.supervised), 9
  - discretize(), 10
  - discretizeDF(), 10
  - discretizeDF.supervised, 9
  - discretizeDF.supervised(), 3, 11, 13, 19–22, 24
  - FOIL, 10, 13
  - foil (FOIL), 10
  - FOIL2 (LUCS_KDD_CBA), 12
  - glmnet(), 22
  - itemFrequency(), 5
  - LUCS_KDD_CBA, 12
  - Lymphography, 14
  - majorityClass (CBA_helpers), 5
  - mineCARs, 15
  - mineCARs(), 3, 4, 7, 8, 16, 22
  - Mushroom, 18
  - PART_CBA (RWeka_CBA), 23
  - predict (predict.CBA), 19
  - predict(), 7
  - predict.CBA, 19
  - prepareTransactions, 20
  - prepareTransactions(), 3, 11, 13, 22, 24
  - PRM (LUCS_KDD_CBA), 12
  - pruneCBA_M1 (CBA), 2
  - pruneCBA_M2 (CBA), 2
  - RCAR, 21
  - rcar (RCAR), 21
  - response (CBA_helpers), 5
  - response(), 19
  - RIPPER_CBA (RWeka_CBA), 23
  - rules, 5, 16
  - RWeka::C4.5_CBA, 23
  - RWeka::JRip(), 24
  - RWeka::PART(), 24
  - RWeka_CBA, 23
  - transactionCoverage (CBA_helpers), 5
  - transactions, 25
  - transactions2DF, 25
  - uncoveredClassExamples (CBA_helpers), 5
  - uncoveredMajorityClass (CBA_helpers), 5

27