Package ‘superml’

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Type Package

Title Build Machine Learning Models Like Using Python's Scikit-Learn Library in R

Version 0.5.7

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Description The idea is to provide a standard interface to users who use both R and Python for building machine learning models. This package provides a scikit-learn's fit, predict interface to train machine learning models in R.

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Encoding UTF-8

LazyData true

URL https://github.com/saraswatmks/superml

BugReports https://github.com/saraswatmks/superml/issues

Depends R(>= 3.6), R6(>= 2.2)

Imports data.table (>= 1.10), Rcpp (>= 1.0), assertthat (>= 0.2), Metrics (>= 0.1)

LinkingTo Rcpp, BH, RcppArmadillo

Suggests knitr, rlang, testthat, rmarkdown, naivebayes(>= 0.9), ClusterR(>= 1.1), FNN(>= 1.1), ranger(>= 0.10), caret(>= 6.0), xgboost(>= 0.6), glmnet(>= 2.0), e1071(>= 1.7)

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VignetteBuilder knitr

NeedsCompilation yes

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BM25 Matching

BM25 stands for Best Matching 25. It is widely used for ranking documents and a preferred method than TF*IDF scores. It is used to find the similar documents from a corpus, given a new document. It is popularly used in information retrieval systems. This implementation is based on c++ functions hence quite optimised as well.

Usage

bm_25(document, corpus, top_n)

Arguments

document     a string for which to find similar documents
corpus       a vector of strings against which document is to be matched
top_n        top n similar documents to find
Value

a vector containing similar documents and their scores

Examples

docs <- c("chimpanzees are found in jungle",
"chimps are jungle animals",
"Mercedes automobiles are best",
"merc is made in germany",
"chimps are intelligent animals")

sentence <- "automobiles are"
s <- bm_25(document=sentence, corpus=docs, top_n=2)

cla_train

Description

Training Dataset used for classification examples. This is classic titanic dataset used to predict if a passenger will survive or not in titanic ship disaster.

Usage

cla_train

Format

An object of class data.table (inherits from data.frame) with 891 rows and 12 columns.

Source

https://www.kaggle.com/c/titanic/data

Counter

Calculate count of values in a list or vector

Description

Handy function to calculate count of values given in a list or vector

Usage

Counter(data, sort = TRUE, decreasing = FALSE)
CountVectorizer

Arguments

- data: should be a vector or list of input values
- sort: a logical value, to sort the result or not
- decreasing: a logical value, the order of sorting to be followed

Value

count of values in a list

Examples

d <- list(c('i','am','bad'),c('you','are','also','bad'))
counts <- Counter(d, sort=TRUE, decreasing=TRUE)

Description

Creates CountVectorizer Model.

Details

Given a list of text, it generates a bag of words model and returns a sparse matrix consisting of token counts.

Public fields

- sentences: a list containing sentences
- max_df: When building the vocabulary ignore terms that have a document frequency strictly higher than the given threshold, value lies between 0 and 1.
- min_df: When building the vocabulary ignore terms that have a document frequency strictly lower than the given threshold, value lies between 0 and 1.
- max_features: Build a vocabulary that only consider the top max_features ordered by term frequency across the corpus.
- ngram_range: The lower and upper boundary of the range of n-values for different word n-grams or char n-grams to be extracted. All values of n such such that min_n <= n <= max_n will be used. For example an ngram_range of c(1, 1) means only unigrams, c(1, 2) means unigrams and bigrams, and c(2, 2) means only bigrams.
- split: splitting criteria for strings, default: " "
- lowercase: convert all characters to lowercase before tokenizing
- regex: regex expression to use for text cleaning.
- remove_stopwords: a list of stopwords to use, by default it uses its inbuilt list of standard stopwords
- model: internal attribute which stores the count model
Methods

Public methods:

• `CountVectorizer$new()`
• `CountVectorizer$fit()`
• `CountVectorizer$fit_transform()`
• `CountVectorizer$transform()`
• `CountVectorizer$clone()`

Method `new()`:

Usage:
`CountVectorizer$new(  
  min_df,
  max_df,
  max_features,
  ngram_range,
  regex,
  remove_stopwords,
  split,
  lowercase
)`

Arguments:

min_df numeric, When building the vocabulary ignore terms that have a document frequency strictly lower than the given threshold, value lies between 0 and 1.
max_df numeric, When building the vocabulary ignore terms that have a document frequency strictly higher than the given threshold, value lies between 0 and 1.
max_features integer, Build a vocabulary that only consider the top max_features ordered by term frequency across the corpus.
ngram_range vector, The lower and upper boundary of the range of n-values for different word n-grams or char n-grams to be extracted. All values of n such such that min_n <= n <= max_n will be used. For example an ngram_range of c(1, 1) means only unigrams, c(1, 2) means unigrams and bigrams, and c(2, 2) means only bigrams.
regex character, regex expression to use for text cleaning.
remove_stopwords list, a list of stopwords to use, by default it uses its inbuilt list of standard english stopwords
split character, splitting criteria for strings, default: " "
lowercase logical, convert all characters to lowercase before tokenizing, default: TRUE

Details: Create a new ‘CountVectorizer’ object.

Returns: A ‘CountVectorizer’ object.

Examples:
`cv = CountVectorizer$new(min_df=0.1)`

Method `fit()`:

Usage:
CountVectorizer$fit(sentences)

*Arguments:*
sentences  a list of text sentences

*Details:*  Fits the countvectorizer model on sentences

*Returns:*  NULL

*Examples:*
sents = c('i am alone in dark.','mother_mary a lot',
        'alone in the dark?','many mothers in the lot....')
cv = CountVectorizer$new(min_df=0.1)
cv$fit(sents)

**Method** fit_transform():

*Usage:*
CountVectorizer$fit_transform(sentences)

*Arguments:*
sentences  a list of text sentences

*Details:*  Fits the countvectorizer model and returns a sparse matrix of count of tokens

*Returns:*  a sparse matrix containing count of tokens in each given sentence

*Examples:*
sents = c('i am alone in dark.','mother_mary a lot',
        'alone in the dark?','many mothers in the lot....')
cv <- CountVectorizer$new(min_df=0.1)
cv_count_matrix <- cv$fit_transform(sents)

cv_count_matrix

**Method** transform():

*Usage:*
CountVectorizer$transform(sentences)

*Arguments:*
sentences  a list of new text sentences

*Details:*  Returns a matrix of count of tokens

*Returns:*  a sparse matrix containing count of tokens in each given sentence

*Examples:*
sents = c('i am alone in dark.','mother_mary a lot',
        'alone in the dark?','many mothers in the lot....')

new_sents <- c("dark at night","mothers day")
cv = CountVectorizer$new(min_df=0.1)
cv$fit(sents)
cv_count_matrix <- cv$transform(new_sents)

**Method** clone():  The objects of this class are cloneable with this method.

*Usage:*
CountVectorizer$clone(deep = FALSE)

*Arguments:*
deep  Whether to make a deep clone.
**Examples**

```r
## Method `CountVectorizer$new`
## ------------------------------------------------

cv = CountVectorizer$new(min_df=0.1)

## Method `CountVectorizer$fit`
## ------------------------------------------------

sents = c('i am alone in dark.', 'mother_mary a lot', 'alone in the dark?', 'many mothers in the lot....')
cv = CountVectorizer$new(min_df=0.1)
cv$fit(sents)

## Method `CountVectorizer$fit_transform`
## ------------------------------------------------

sents = c('i am alone in dark.', 'mother_mary a lot', 'alone in the dark?', 'many mothers in the lot....')
cv <- CountVectorizer$new(min_df=0.1)
cv_count_matrix <- cv$fit_transform(sents)

## Method `CountVectorizer$transform`
## ------------------------------------------------

sents = c('i am alone in dark.', 'mother_mary a lot', 'alone in the dark?', 'many mothers in the lot....')
new_sents <- c('dark at night', 'mothers day')
cv = CountVectorizer$new(min_df=0.1)
cv$fit(sents)
cv_count_matrix <- cv$transform(new_sents)
```

---

**dot**  
*Dot product similarity in vectors*

---

**Description**

Computes the dot product between two given vectors.

**Usage**

```r
dot(a, b, norm = TRUE)
```
**Arguments**

- **a**: numeric vector
- **b**: numeric vector
- **norm**: logical, compute normalised dot product, default=True

**Value**

numeric vector containing sdot product score

**Examples**

```r
a <- runif(5)
b <- runif(5)
s <- dot(a, b)
```

---

**dotmat**

*Dot product similarity between a vector and matrix*

**Description**

Computes the dot product between a vector and a given matrix. The vector returned has a dot product similarity value for each row in the matrix.

**Usage**

```r
dotmat(a, b, norm = TRUE)
```

**Arguments**

- **a**: numeric vector
- **b**: numeric matrix
- **norm**: logical, compute normalised dot product, default=True

**Value**

numeric vector containing dot product scores
**Description**

Runs grid search cross validation scheme to find best model training parameters.

**Details**

Grid search CV is used to train a machine learning model with multiple combinations of training hyper-parameters and finds the best combination of parameters which optimizes the evaluation metric. It creates an exhaustive set of hyperparameter combinations and train model on each combination.

**Public fields**

- trainer: superml trainer object, could be either XGBTrainer, RFTrainer, NBTrainer etc.
- parameters: a list of parameters to tune
- n_folds: number of folds to use to split the train data
- scoring: scoring metric used to evaluate the best model, multiple values can be provided. currently supports: auc, accuracy, mse, rmse, logloss, mae, f1, precision, recall
- evaluation_scores: parameter for internal use

**Methods**

**Public methods:**

- `GridSearchCV$new()`
- `GridSearchCV$fit()`
- `GridSearchCV$best_iteration()`
- `GridSearchCV$clone()`

**Method new():**

*Usage:*

```
GridSearchCV$new(trainer = NA, parameters = NA, n_folds = NA, scoring = NA)
```

*Arguments:*

- trainer: superml trainer object, could be either XGBTrainer, RFTrainer, NBTrainer etc.
- parameters: a list of parameters to tune
- n_folds: number of folds to use to split the train data
- scoring: character, scoring metric used to evaluate the best model, multiple values can be provided. currently supports: auc, accuracy, mse, rmse, logloss, mae, f1, precision, recall

*Details:* Create a new ‘GridSearchCV’ object.

*Returns:* A ‘GridSearchCV’ object.

*Examples:*
rf <- RFTrainer$new()
gst <- GridSearchCV$new(trainer = rf,
  parameters = list(n_estimators = c(100),
    max_depth = c(5,2,10)),
    n_folds = 3,
    scoring = c('accuracy','auc'))

Method fit():

Usage:
GridSearchCV$fit(X, y)

Arguments:
X data.frame or data.table
y character, name of target variable

Details: Trains the model using grid search

Returns: NULL

Examples:
rf <- RFTrainer$new()
gst <- GridSearchCV$new(trainer = rf,
  parameters = list(n_estimators = c(100),
    max_depth = c(5,2,10)),
    n_folds = 3,
    scoring = c('accuracy','auc'))
data("iris")
gst$fit(iris, "Species")

Method best_iteration():

Usage:
GridSearchCV$best_iteration(metric = NULL)

Arguments:
metric character, which metric to use for evaluation

Details: Returns the best parameters

Returns: a list of best parameters

Examples:
rf <- RFTrainer$new()
gst <- GridSearchCV$new(trainer = rf,
  parameters = list(n_estimators = c(100),
    max_depth = c(5,2,10)),
    n_folds = 3,
    scoring = c('accuracy','auc'))
data("iris")
gst$fit(iris, "Species")
gst$best_iteration()

Method clone(): The objects of this class are cloneable with this method.
**Usage:**
GridSearchCV$clone(deep = FALSE)

**Arguments:**
deep Whether to make a deep clone.

**Examples**

```r
## Method `GridSearchCV$new`
rf <- RFTrainer$new()
gst <- GridSearchCV$new(trainer = rf,
parameters = list(n_estimators = c(100),
max_depth = c(5,2,10)),
n_folds = 3,
scoring = c('accuracy','auc'))

## Method `GridSearchCV$fit`
rf <- RFTrainer$new()
gst <- GridSearchCV$new(trainer = rf,
parameters = list(n_estimators = c(100),
max_depth = c(5,2,10)),
n_folds = 3,
scoring = c('accuracy','auc'))
data("iris")
gst$fit(iris, "Species")

## Method `GridSearchCV$best_iteration`
rf <- RFTrainer$new()
gst <- GridSearchCV$new(trainer = rf,
parameters = list(n_estimators = c(100),
max_depth = c(5,2,10)),
n_folds = 3,
scoring = c('accuracy','auc'))
data("iris")
gst$fit(iris, "Species")
gst$best_iteration()
```
Description

Calculates out-of-fold mean features (also known as target encoding) for train and test data. This strategy is widely used to avoid overfitting or causing leakage while creating features using the target variable. This method is experimental. If the results you get are unexpected, please report them in github issues.

Usage

kFoldMean(train_df, test_df, colname, target, n_fold = 5, seed = 42)

Arguments

train_df          train dataset
test_df           test dataset
colname           name of categorical column
target            the target or dependent variable, should be a string.
n_fold            the number of folds to use for doing kfold computation, default=5
seed              the seed value, to ensure reproducibility, it could be any positive value, default=42

Value

a train and test data table with out-of-fold mean value of the target for the given categorical variable

Examples

train <- data.frame(region=c('del', 'csk', 'rcb', 'del', 'csk', 'pune', 'guj', 'del'),
                     win = c(0,1,1,0,0,0,0,1))
test <- data.frame(region=c('rcb', 'csk', 'rcb', 'del', 'guj', 'pune', 'csk', 'kol'))
train_result <- kFoldMean(train_df = train,
                          test_df = test,
                          colname = 'region',
                          target = 'win',
                          seed = 1220)$train

test_result <- kFoldMean(train_df = train,
                          test_df = test,
                          colname = 'region',
                          target = 'win',
                          seed = 1220)$test
Description

Trains a k-means machine learning model in R

Details

Trains an unsupervised K-Means clustering algorithm. It borrows mini-batch k-means function from ClusterR package written in c++, hence it is quite fast.

Public fields

- clusters: the number of clusters
- batch.size: the size of the mini batches
- num.init: number of times the algorithm will be run with different centroid seeds
- max.iters: the maximum number of clustering iterations
- init.fraction: percentage of data to use for the initialization centroids (applies if initializer is kmeans++ or optimal.init). Should be a float number between 0.0 and 1.0.
- initializer: the method of initialization. One of, optimal_init, quantile_init, kmeans++ and random.
- early.stop.iter: continue that many iterations after calculation of the best within-cluster-sum-of-squared-error
- verbose: either TRUE or FALSE, indicating whether progress is printed during clustering
- centroids: a matrix of initial cluster centroids. The rows of the CENTROIDS matrix should be equal to the number of clusters and the columns should be equal to the columns of the data
- tol: a float number. If, in case of an iteration (iteration > 1 and iteration < max.iters) "tol" is greater than the squared norm of the centroids, then kmeans has converged
- tol.optimal.init: tolerance value for the 'optimal_init' initializer. The higher this value is, the far apart from each other the centroids are.
- seed: integer value for random number generator (RNG)
- model: use for internal purpose
- max.clusters: either a numeric value, a contiguous or non-contiguous numeric vector specifying the cluster search space

Methods

Public methods:

- KMeansTrainer$new()
- KMeansTrainer$fit()
- KMeansTrainer$predict()
• KMeansTrainer$clone()

**Method** `new()`:

**Usage:**
KMeansTrainer$new(
  clusters,
  batch_size = 10,
  num_init = 1,
  max_iters = 100,
  init_fraction = 1,
  initializer = "kmeans++",
  early_stop_iter = 10,
  verbose = FALSE,
  centroids = NULL,
  tol = 1e-04,
  tol_optimal_init = 0.3,
  seed = 1,
  max_clusters = NA
)

**Arguments:**
- `clusters` numeric, When building the vocabulary ignore terms that have a document frequency strictly lower than the given threshold, value lies between 0 and 1.
- `batch_size` numeric, When building the vocabulary ignore terms that have a document frequency strictly higher than the given threshold, value lies between 0 and 1.
- `num_init` integer, use top features sorted by count to be used in bag of words matrix.
- `max_iters` character, regex expression to use for text cleaning.
- `init_fraction` list, a list of stopwords to use, by default it uses its inbuilt list of standard stopwords.
- `initializer` character, splitting criteria for strings, default: " ".
- `early_stop_iter` continue that many iterations after calculation of the best within-cluster-sum-of-squared-error.
- `verbose` either TRUE or FALSE, indicating whether progress is printed during clustering.
- `centroids` a matrix of initial cluster centroids. The rows of the CENTROIDS matrix should be equal to the number of clusters and the columns should be equal to the columns of the data.
- `tol` a float number. If, in case of an iteration (iteration > 1 and iteration < max_iters) "tol" is greater than the squared norm of the centroids, then kmeans has converged.
- `tol_optimal_init` tolerance value for the 'optimal_init' initializer. The higher this value is,
  the far apart from each other the centroids are.
- `seed` integer value for random number generator (RNG).
- `max_clusters` either a numeric value, a contiguous or non-contiguous numeric vector specifying the cluster search space.

**Details:** Create a new ‘KMeansTrainer’ object.

**Returns:** A ‘KMeansTrainer’ object.

**Examples:**
data <- rbind(replicate(20, rnorm(1e4, 2)),
              replicate(20, rnorm(1e4, -1)),
              replicate(20, rnorm(1e4, 5)))
km_model <- KMeansTrainer$new(clusters=2, batch_size=30, max_clusters=6)

Method `fit()`:

Usage:
KMeansTrainer$fit(X, y = NULL, find_optimal = FALSE)

Arguments:
X data.frame or matrix containing features
y NULL only kept here for superml's standard way
find_optimal logical, to find the optimal clusters automatically

Details: Trains the KMeansTrainer model

Returns: NULL

Examples:
data <- rbind(replicate(20, rnorm(1e4, 2)),
              replicate(20, rnorm(1e4, -1)),
              replicate(20, rnorm(1e4, 5)))
km_model <- KMeansTrainer$new(clusters=2, batch_size=30, max_clusters=6)
km_model$fit(data, find_optimal = FALSE)

Method `predict()`:

Usage:
KMeansTrainer$predict(X)

Arguments:
X data.frame or matrix

Details: Returns the prediction on test data

Returns: a vector of predictions

Examples:
data <- rbind(replicate(20, rnorm(1e4, 2)),
              replicate(20, rnorm(1e4, -1)),
              replicate(20, rnorm(1e4, 5)))
km_model <- KMeansTrainer$new(clusters=2, batch_size=30, max_clusters=6)
km_model$fit(data, find_optimal = FALSE)
predictions <- km_model$predict(data)

Method `clone()`: The objects of this class are cloneable with this method.

Usage:
KMeansTrainer$clone(deep = FALSE)

Arguments:
depth Whether to make a deep clone.
Examples

```r
# Method `KMeansTrainer$new`

data <- rbind(replicate(20, rnorm(1e4, 2)),
              replicate(20, rnorm(1e4, -1)),
              replicate(20, rnorm(1e4, 5)))
km_model <- KMeansTrainer$new(clusters=2, batch_size=30, max_clusters=6)

# Method `KMeansTrainer$fit`

km_model$fit(data, find_optimal = FALSE)

# Method `KMeansTrainer$predict`

predictions <- km_model$predict(data)
```

KNNTrainer

*K Nearest Neighbours Trainer*

**Description**

Trains a k nearest neighbour model using fast search algorithms. KNN is a supervised learning algorithm which is used for both regression and classification problems.

**Format**

*R6Class* object.

**Usage**

For usage details see Methods, Arguments and Examples sections.

```r
bst = KNNTrainer$new(k=1, prob=FALSE, algorithm=NULL, type="class")
bst$fit(X_train, X_test, "target")
bst$predict(type)
```
Methods

$new()$ Initialise the instance of the trainer
$fit()$ trains the knn model and stores the test prediction
$predict()$ returns predictions

Arguments

- **k** number of neighbours to predict
- **prob** if probability should be computed, default=FALSE
- **algorithm** algorithm used to train the model, possible values are 'kd_tree', 'cover_tree', 'brute'
- **type** type of problem to solve i.e. regression or classification, possible values are 'reg' or 'class'

Public fields

- **k** number of neighbours to predict
- **prob** if probability should be computed, default=FALSE
- **algorithm** algorithm used to train the model, possible values are 'kd_tree', 'cover_tree', 'brute'
- **type** type of problem to solve i.e. regression or classification, possible values are 'reg' or 'class'
- **model** for internal use

Methods

Public methods:

- KNNTrainer$new()
- KNNTrainer$fit()
- KNNTrainer$predict()
- KNNTrainer$clone()

Method **new()**:

Usage:

KNNTrainer$new(k, prob, algorithm, type)

Arguments:

- **k** number of neighbours to predict
- **prob** if probability should be computed, default=FALSE
- **algorithm** algorithm used to train the model, possible values are 'kd_tree', 'cover_tree', 'brute'
- **type** type of problem to solve i.e. regression or classification, possible values are 'reg' or 'class'

Details: Create a new ‘KNNTrainer’ object.

Returns: A ‘KNNTrainer’ object.

Examples:
data("iris")

iris$Species <- as.integer(as.factor(iris$Species))

xtrain <- iris[1:100,]
xtest <- iris[101:150,]

bst <- KNNTrainer$new(k=3, prob=TRUE, type="class")
bst$fit(xtrain, xtest, 'Species')
pred <- bst$predict(type="raw")

Method fit():
Usage:
KNNTrainer$fit(train, test, y)
Arguments:
  train  data.frame or matrix
  test   data.frame or matrix
  y      character, name of target variable
Details: Trains the KNNTrainer model
Returns: NULL
Examples:
data("iris")

iris$Species <- as.integer(as.factor(iris$Species))

xtrain <- iris[1:100,]
xtest <- iris[101:150,]

bst <- KNNTrainer$new(k=3, prob=TRUE, type="class")
bst$fit(xtrain, xtest, 'Species')

Method predict():
Usage:
KNNTrainer$predict(type = "raw")
Arguments:
  type   character, 'raw' for labels else 'prob'
Details: Predicts the nearest neighbours for test data
Returns: a list of predicted neighbours
Examples:
data("iris")

iris$Species <- as.integer(as.factor(iris$Species))

xtrain <- iris[1:100,]
xtest <- iris[101:150,]

bst <- KNNTrainer$new(k=3, prob=TRUE, type="class")
bst$fit(xtrain, xtest, 'Species')
pred <- bst$predict(type="raw")

**Method** clone(): The objects of this class are cloneable with this method.

**Usage:**
KNNTrainer$clone(deep = FALSE)

**Arguments:**
deep Whether to make a deep clone.

**Examples**

data("iris")

iris$Species <- as.integer(as.factor(iris$Species))

xtrain <- iris[1:100,]
xtest <- iris[101:150,]

bst <- KNNTrainer$new(k=3, prob=TRUE, type="class")
bst$fit(xtrain, xtest, 'Species')
pred <- bst$predict(type="raw")

## ------------------------------------------------
## Method 'KNNTrainer$new'
## ------------------------------------------------

data("iris")

iris$Species <- as.integer(as.factor(iris$Species))

xtrain <- iris[1:100,]
xtest <- iris[101:150,]

bst <- KNNTrainer$new(k=3, prob=TRUE, type="class")
bst$fit(xtrain, xtest, 'Species')
pred <- bst$predict(type="raw")

## ------------------------------------------------
## Method 'KNNTrainer$fit'
## ------------------------------------------------

data("iris")

iris$Species <- as.integer(as.factor(iris$Species))

xtrain <- iris[1:100,]
xtest <- iris[101:150,]
bst <- KNNTrainer$new(k=3, prob=TRUE, type="class")
bst$fit(xtrain, xtest, 'Species')

## Method 'KNNTrainer$predict'
## ------------------------------------------------
data("iris")
iris$Species <- as.integer(as.factor(iris$Species))

xtrain <- iris[1:100,]
xtest <- iris[101:150,]

bst <- KNNTrainer$new(k=3, prob=TRUE, type="class")
bst$fit(xtrain, xtest, 'Species')
pred <- bst$predict(type="raw")

---

**LabelEncoder**

**Label Encoder**

**Description**

Encodes and decodes categorical variables into integer values and vice versa. This is a commonly performed task in data preparation during model training, because all machine learning models require the data to be encoded into numerical format. It takes a vector of character or factor values and encodes them into numeric.

**Format**

*R6Class* object.

**Usage**

For usage details see Methods, Arguments and Examples sections.

```r
lbl = LabelEncoder$new()
lbl$fit(x)
lbl$fit_transform(x)
lbl$transform(x)
```

**Methods**

$new() Initialise the instance of the encoder
$fit() creates a memory of encodings but doesn’t return anything
$transform() based on encodings learned in fit method is applies the transformation
$fit_transform() encodes the data and keep a memory of encodings simultaneously
$inverse_transform() encodes the data and keep a memory of encodings simultaneously
Arguments

data a vector or list containing the character / factor values

Public fields

input_data internal use
encodings internal use
decodings internal use
fit_model internal use

Methods

Public methods:

• LabelEncoder$fit()
• LabelEncoder$fit_transform()
• LabelEncoder$transform()
• LabelEncoder$inverse_transform()
• LabelEncoder$clone()

Method fit():

Usage:
LabelEncoder$fit(data_col)

Arguments:
data_col a vector containing non-null values

Details: Fits the labelencoder model on given data

Returns: NULL, calculates the encoding and save in memory

Examples:
data_ex <- data.frame(Score = c(10,20,30,4), Name=c('Ao','Bo','Bo','Co'))
lbl <- LabelEncoder$new()
lbl$fit(data_ex$Name)
data_ex$Name <- lbl$fit_transform(data_ex$Name)
decode_names <- lbl$inverse_transform(data_ex$Name)

Method fit_transform():

Usage:
LabelEncoder$fit_transform(data_col)

Arguments:
data_col a vector containing non-null values

Details: Fits and returns the encoding

Returns: encoding values for the given input data

Examples:
data_ex <- data.frame(Score = c(10,20,30,4), Name=c('Ao','Bo','Bo','Co'))
lbl <- LabelEncoder$new()
lbl$fit(data_ex$Name)
data_ex$Name <- lbl$fit_transform(data_ex$Name)

Method transform():
Usage:
LabelEncoder$transform(data_col)
Arguments:
data_col a vector containing non-null values
Details: Returns the encodings from the fitted model
Returns: encoding values for the given input data
Examples:
data_ex <- data.frame(Score = c(10,20,30,4), Name=c('Ao','Bo','Bo','Co'))
lbl <- LabelEncoder$new()
lbl$fit(data_ex$Name)
data_ex$Name <- lbl$transform(data_ex$Name)

Method inverse_transform():
Usage:
LabelEncoder$inverse_transform(coded_col)
Arguments:
coded_col a vector containing label encoded values
Details: Gives back the original values from a encoded values
Returns: original values from the label encoded data
Examples:
data_ex <- data.frame(Score = c(10,20,30,4), Name=c('Ao','Bo','Bo','Co'))
lbl <- LabelEncoder$new()
lbl$fit(data_ex$Name)
data_ex$Name <- lbl$fit_transform(data_ex$Name)
decode_names <- lbl$inverse_transform(data_ex$Name)

Method clone(): The objects of this class are cloneable with this method.
Usage:
LabelEncoder$clone(deep = FALSE)
Arguments:
dep Whether to make a deep clone.

Examples

data_ex <- data.frame(Score = c(10,20,30,4), Name=c('Ao','Bo','Bo','Co'))
lbl <- LabelEncoder$new()
data_ex$Name <- lbl$fit_transform(data_ex$Name)
decode_names <- lbl$inverse_transform(data_ex$Name)
## Method \`LabelEncoder$fit\`
## -----------------------------------------------

data_ex <- data.frame(Score = c(10, 20, 30, 4), Name=c('Ao', 'Bo', 'Bo', 'Co'))
lbl <- LabelEncoder$new()
lbl$fit(data_ex$Name)
data_ex$Name <- lbl$fit_transform(data_ex$Name)
decode_names <- lbl$inverse_transform(data_ex$Name)

## Method \`LabelEncoder$fit_transform\`
## -----------------------------------------------

data_ex <- data.frame(Score = c(10, 20, 30, 4), Name=c('Ao', 'Bo', 'Bo', 'Co'))
lbl <- LabelEncoder$new()
lbl$fit(data_ex$Name)
data_ex$Name <- lbl$fit_transform(data_ex$Name)

## Method \`LabelEncoder$transform\`
## -----------------------------------------------

data_ex <- data.frame(Score = c(10, 20, 30, 4), Name=c('Ao', 'Bo', 'Bo', 'Co'))
lbl <- LabelEncoder$new()
lbl$fit(data_ex$Name)
data_ex$Name <- lbl$transform(data_ex$Name)

decode_names <- lbl$inverse_transform(data_ex$Name)

---

**LMTrainer**

*Linear Models Trainer*

**Description**

Trains regression, lasso, ridge model in R

**Details**

Trains linear models such as Logistic, Lasso or Ridge regression model. It is built on glmnet R package. This class provides fit, predict, cross validation functions.
Public fields

family  type of regression to perform, values can be "gaussian" , "binomial" , "multinomial" , "mgaussian"
weights  observation weights. Can be total counts if responses are proportion matrices. Default is 1 for each observation
alpha  The elasticnet mixing parameter, alpha=1 is the lasso penalty, alpha=0 the ridge penalty, alpha=NULL is simple regression
lambda  the number of lambda values - default is 100
standardize  normalise the features in the given data
standardize.response  normalise the dependent variable between 0 and 1, default = FALSE
model  internal use
cvmodel  internal use
Flag  internal use
is_lasso  internal use
iid_names  internal use

Methods

Public methods:

•  LMTrainer$new()
•  LMTrainer$fit()
•  LMTrainer$predict()
•  LMTrainer$cv_model()
•  LMTrainer$cv_predict()
•  LMTrainer$get_importance()
•  LMTrainer$clone()

Method new():

Usage:
LMTrainer$new(family, weights, alpha, lambda, standardize.response)

Arguments:

family  character, type of regression to perform, values can be "gaussian" , "binomial" , "multinomial" , "mgaussian"
weights  numeric, observation weights. Can be total counts if responses are proportion matrices. Default is 1 for each observation
alpha  integer, The elasticnet mixing parameter, alpha=1 is the lasso penalty, alpha=0 the ridge penalty, alpha=NULL is simple regression
lambda  integer, the number of lambda values - default is 100
standardize.response  logical, normalise the dependent variable between 0 and 1, default = FALSE

Details:  Create a new ‘LMTrainer’ object.

Returns:  A ‘LMTrainer’ object.
Examples:
\dontrun{
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
        "RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
}

Method \textbf{fit}():

\textbf{Usage}: 
LMTrainer$fit(X, y)

\textbf{Arguments}:
\begin{itemize}
  \item \textbf{X} data.frame containing train featuers
  \item \textbf{y} character, name of target variable
\end{itemize}

\textbf{Details}: Fits the LMTrainer model on given data

\textbf{Returns}: NULL, train the model and saves internally

\textbf{Examples}:
\dontrun{
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
        "RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$fit(X = housing, y = 'MEDV')
}\n
Method \textbf{predict}():

\textbf{Usage}:
LMTrainer$predict(df, lambda = NULL)

\textbf{Arguments}:
\begin{itemize}
  \item \textbf{df} data.frame containing test features
  \item \textbf{lambda} integer, the number of lambda values - default is 100. By default it picks the best value from the model.
\end{itemize}

\textbf{Details}: Returns predictions for test data

\textbf{Returns}: vector, a vector containing predictions

\textbf{Examples}:
\dontrun{
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
        "RAD","TAX","PTRATIO","B","LSTAT","MEDV")
housing$MEDV <- NULL
housing$MEDV[1:10] <- NA
housing <- na.omit(housing)
housing <- housing[1:100,]
}

\textbf{Examples}:
\dontrun{
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
        "RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$fit(X = housing, y = 'MEDV')
lf$predict(df = housing, lambda = NULL)
}
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$fit(X = housing, y = 'MEDV')
predictions <- lf$cv_predict(df = housing)

Method cv_model():

Usage:
LMTrainer$cv_model(X, y, nfolds, parallel, type.measure = "deviance")

Arguments:
X data.frame containing test features
y character, name of target variable
nfolds integer, number of folds
parallel logical, if do parallel computation. Default=FALSE
type.measure character, evaluation metric type. Default = deviance

Details: Train regression model using cross validation

Returns: NULL, trains the model and saves it in memory

Examples:
\dontrun{
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$cv_model(X = housing, y = 'MEDV', nfolds = 5, parallel = FALSE)
}

Method cv_predict():

Usage:
LMTrainer$cv_predict(df, lambda = NULL)

Arguments:
df data.frame containing test features
lambda integer, the number of lambda values - default is 100. By default it picks the best value from the model.

Details: Get predictions from the cross validated regression model

Returns: vector a vector containing predicted values

Examples:
\dontrun{
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)

\dontrun{
LMTrainer

"RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$cv_model(X = housing, y = 'MEDV', nfolds = 5, parallel = FALSE)
predictions <- lf$cv_predict(df = housing)
}

Method \texttt{get\_importance}():

\textit{Usage:}
LMTrainer$\texttt{get\_importance}()

\textit{Details:} Get feature importance using model coefficients

\textit{Returns:} a matrix containing feature coefficients

\textit{Examples:}
\begin{verbatim}
\dontrun{
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
"RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$cv_model(X = housing, y = 'MEDV', nfolds = 5, parallel = FALSE)
predictions <- lf$cv_predict(df = housing)
coefs <- lf$get\_importance()
}
\end{verbatim}

Method \texttt{clone}(): The objects of this class are cloneable with this method.

\textit{Usage:}
LMTrainer$\texttt{clone}(deep = FALSE)

\textit{Arguments:}
depth \hspace{1cm} Whether to make a deep clone.

\textit{Examples}

\begin{verbatim}
## ---------------------
## Method \texttt{LMTrainer$\texttt{new}}
## ---------------------

## Not run:
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
"RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)

## End(Not run)
\end{verbatim}
## Method `LMTrainer$fit`

### Not run:
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
            "RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$fit(X = housing, y = 'MEDV')

### Method `LMTrainer$predict`

### Not run:
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
            "RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$fit(X = housing, y = 'MEDV')
predictions <- lf$cv_predict(df = housing)

### Method `LMTrainer$cv_model`

### Not run:
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names <- c("CRIM","ZN","INDUS","CHAS","NOX","RM","AGE","DIS",
            "RAD","TAX","PTRATIO","B","LSTAT","MEDV")
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$cv_model(X = housing, y = 'MEDV', nfolds = 5, parallel = FALSE)

### Method `LMTrainer$cv_predict`
NBTrainer <- read.table(LINK)
names(housing) <- names
lf <- LMTrainer$new(family = "gaussian", alpha=1)
lf$cv_model(X = housing, y = "MEDV", nfolds = 5, parallel = FALSE)
predictions <- lf$cv_predict(df = housing)

## End(Not run)

## Method `LMTrainer$get_importance`
## --------------------------------

## Not run:
LINK <- "http://archive.ics.uci.edu/ml/machine-learning-databases/housing/housing.data"
housing <- read.table(LINK)
names(housing) <- names
lf <- LMTrainer$new(family = 'gaussian', alpha=1)
lf$cv_model(X = housing, y = 'MEDV', nfolds = 5, parallel = FALSE)
predictions <- lf$cv_predict(df = housing)
coefs <- lf$get_importance()
## End(Not run)

---

NBTrainer

Naive Bayes Trainer

Description

Trains a probabilistic naive bayes model

Details

Trains a naive bayes model. It is built on top high performance naivebayes R package.

Public fields

prior numeric vector with prior probabilities. vector with prior probabilities of the classes. If unspecified, the class proportions for the training set are used. If present, the probabilities should be specified in the order of the factor levels.

laplace value used for Laplace smoothing. Defaults to 0 (no Laplace smoothing)

usekernel if TRUE, density is used to estimate the densities of metric predictors

model for internal use
Methods

Public methods:

- `NBTrainer$new()`
- `NBTrainer$fit()`
- `NBTrainer$predict()`
- `NBTrainer$clone()`

Method `new()`:

Usage:
```r
NBTrainer$new(prior, laplace, usekernel)
```

Arguments:
- `prior` numeric, prior numeric vector with prior probabilities. vector with prior probabilities of the classes. If unspecified, the class proportions for the training set are used. If present, the probabilities should be specified in the order of the factor levels.
- `laplace` numeric, value used for Laplace smoothing. Defaults to 0 (no Laplace smoothing)
- `usekernel` logical, if TRUE, density is used to estimate the densities of metric predictors

Details: Create a new `NBTrainer` object.

Returns: A `NBTrainer` object.

Examples:
```r
data(iris)
nb <- NBTrainer$new()
```

Method `fit()`:

Usage:
```r
NBTrainer$fit(X, y)
```

Arguments:
- `X` data.frame containing train features
- `y` character, name of target variable

Details: Fits the naive bayes model

Returns: NULL, trains and saves the model in memory

Examples:
```r
data(iris)
nb <- NBTrainer$new()
nb$fit(iris, 'Species')
```

Method `predict()`:

Usage:
```r
NBTrainer$predict(X, type = "class")
```

Arguments:
- `X` data.frame containing test features
- `type` character, if the predictions should be labels or probability
Details: Returns predictions from the model

Returns: NULL, trains and saves the model in memory

Examples:

data(iris)
nb <- NBTrainer$new()
nb$fit(iris, 'Species')
y <- nb$predict(iris)

Method clone(): The objects of this class are cloneable with this method.

Usage:

NBTrainer$clone(deep = FALSE)

Arguments:

depth Whether to make a deep clone.

Examples

## ------------------------------------------------
## Method `NBTrainer$new`
## ------------------------------------------------

data(iris)
nb <- NBTrainer$new()

## ------------------------------------------------
## Method `NBTrainer$fit`
## ------------------------------------------------

data(iris)
nb <- NBTrainer$new()
nb$fit(iris, 'Species')

## ------------------------------------------------
## Method `NBTrainer$predict`
## ------------------------------------------------

data(iris)
nb <- NBTrainer$new()
nb$fit(iris, 'Species')
y <- nb$predict(iris)

normalise1d

Description

Normalises a 1 dimensional vector towards unit p norm. By default, p = 2 is used. For a given vector, eg: c(1,2,3), norm value is calculated as `|x| / lxl` where `lxl` is calculated as the square root of sum of square of values in the given vector.
Usage

normalise1d(vec, pnorm = 2L)

Arguments

vec vector containing integers or numeric values.

pnorm integer, default: 2

Value

a vector containing normalised values

Examples

val <- c(1,10,5,3,8)
norm_val <- normalise1d(val)

Description

Normalises a matrix towards unit p norm row wise or column wise. By default, p = 2 is used. To normalise row wise, use axis=0. To normalise column wise, use axis=1. as the square root of sum of square of values in the given vector.

Usage

normalise2d(mat, pnorm = 2L, axis = 1L)

Arguments

mat numeric matrix

pnorm integer value, default value=2

axis integer (0 or 1), row wise = 0, column wise = 1

Value

normalised numeric matrix
RandomSearchCV

Examples

```r
mat <- matrix(runif(12), 3, 4)

## normalise matrix row wise
r <- normalise2d(mat, axis=0)

## normalise matrix column wise
r <- normalise2d(mat, axis=1)
```

---

RandomSearchCV  
Random Search CV

Description

Hyperparameter tuning using random search scheme.

Details

Given a set of hyper parameters, random search trainer provides a faster way of hyper parameter tuning. Here, the number of models to be trained can be defined by the user.

Super class

```
superml::GridSearchCV -> RandomSearchTrainer
```

Public fields

- `n_iter` number of models to be trained

Methods

Public methods:

- `RandomSearchCV$new()`
- `RandomSearchCV$fit()`
- `RandomSearchCV$clone()`

Method `new()`:

*Usage:*

```r
RandomSearchCV$new(
  trainer = NA,
  parameters = NA,
  n_folds = NA,
  scoring = NA,
  n_iter = NA
)
```

*Arguments:*

RandomSearchCV

trainer: superml trainer object, must be either XGBTTrainer, LMTrainer, RFTrainer, NBTrainer
parameters: list, list containing parameters
n_folds: integer, number of folds to use to split the train data
scoring: character, scoring metric used to evaluate the best model, multiple values can be provided. Currently supports: auc, accuracy, mse, rmse, logloss, mae, f1, precision, recall
n_iter: integer, number of models to be trained

Details: Create a new ‘RandomSearchTrainer’ object.

Returns: A ‘RandomSearchTrainer’ object.

Examples:
rf <- RFTrainer$new()
rst <- RandomSearchCV$new(trainer = rf,
parameters = list(n_estimators = c(100,500),
max_depth = c(5,2,10,14)),
n_folds = 3,
scoring = c('accuracy','auc'),
n_iter = 4)

trainer superml trainer object, must be either XGBTTrainer, LMTrainer, RFTrainer, NBTrainer
parameters: list, list containing parameters
n_folds: integer, number of folds to use to split the train data
scoring: character, scoring metric used to evaluate the best model, multiple values can be provided. Currently supports: auc, accuracy, mse, rmse, logloss, mae, f1, precision, recall
n_iter: integer, number of models to be trained

Details: Create a new ‘RandomSearchTrainer’ object.

Returns: A ‘RandomSearchTrainer’ object.

Examples:
rf <- RFTrainer$new()
rst <- RandomSearchCV$new(trainer = rf,
parameters = list(n_estimators = c(100,500),
max_depth = c(5,2,10,14)),
n_folds = 3,
scoring = c('accuracy','auc'),
n_iter = 4)

data("iris")
rst$fit(iris, "Species")
rst$best_iteration()

Method clone(): The objects of this class are cloneable with this method.

Usage:
RandomSearchCV$clone(deep = FALSE)

Arguments:
depth: Whether to make a deep clone.
Examples

```r
## Method `RandomSearchCV$new`
rf <- RFTrainer$new()
rst <- RandomSearchCV$new(trainer = rf,
                         parameters = list(n_estimators = c(100,500),
                              max_depth = c(5,2,10,14)),
                         n_folds = 3,
                         scoring = c('accuracy','auc'),
                         n_iter = 4)
```

```r
data(“iris”)
rst$fit(iris, “Species”) rst$best_iteration()
```

Description

Training Dataset used for regression examples. In this data set, we have to predict the sale price of the houses.

Usage

reg_train

Format

An object of class `data.table` (inherits from `data.frame`) with 1460 rows and 81 columns.

Source

**Random Forest Trainer**

**Description**

Trains a random forest model.

**Details**

Trains a Random Forest model. A random forest is a meta estimator that fits a number of decision tree classifiers on various sub-samples of the dataset and use averaging to improve the predictive accuracy and control over-fitting. This implementation uses ranger R package which provides faster model training.

**Public fields**

- `n_estimators` the number of trees in the forest, default= 100
- `max_features` the number of features to consider when looking for the best split. Possible values are `auto` (default) takes sqrt(num_of_features), `sqrt` same as auto, `log` takes log(num_of_features), `none` takes all features
- `max_depth` the maximum depth of each tree
- `min_node_size` the minimum number of samples required to split an internal node
- `criterion` the function to measure the quality of split. For classification, `gini` is used which is a measure of gini index. For regression, the `variance` of responses is used.
- `classification` whether to train for classification (1) or regression (0)
- `verbose` show computation status and estimated runtime
- `seed` seed value
- `class_weights` weights associated with the classes for sampling of training observation
- `always_split` vector of feature names to be always used for splitting
- `importance` Variable importance mode, one of 'none', 'impurity', 'impurity_corrected', 'permutation'. The 'impurity' measure is the Gini index for classification, the variance of the responses for regression. Defaults to “impurity”

**Methods**

**Public methods:**

- `RFTrainer$new()`
- `RFTrainer$fit()`
- `RFTrainer$predict()`
- `RFTrainer$get_importance()`
- `RFTrainer$clone()`

**Method new():**
Usage:
RFTrainer$new(
  n_estimators,
  max_depth,
  max_features,
  min_node_size,
  classification,
  class_weights,
  always_split,
  verbose,
  save_model,
  seed,
  importance
)

Arguments:
n_estimators integer, the number of trees in the forest, default= 100
max_depth integer, the maximum depth of each tree
max_features integer, the number of features to consider when looking for the best split. Possible values are auto (default) takes sqrt(num_of_features), sqrt same as auto, log takes log(num_of_features), none takes all features
min_node_size integer, the minimum number of samples required to split an internal node
classification integer, whether to train for classification (1) or regression (0)
class_weights weights associated with the classes for sampling of training observation
always_split vector of feature names to be always used for splitting
verbose logical, show computation status and estimated runtime
save_model logical, whether to save model
seed integer, seed value
importance Variable importance mode, one of 'none', 'impurity', 'impurity_corrected', 'permutation'. The 'impurity' measure is the Gini index for classification, the variance of the responses for regression. Defaults to "impurity"

Details: Create a new ‘RFTrainer’ object.

Returns: A ‘RFTrainer’ object.

Examples:
data("iris")
bst <- RFTrainer$new(n_estimators=10,
  max_depth=4,
  classification=1,
  seed=42,
  verbose=TRUE)

Method fit():

Usage:
RFTrainer$fit(X, y)

Arguments:
RFTrainer

**Details:** Trains the random forest model

**Returns:** NULL, trains and saves the model in memory

**Examples:**
```r
data("iris")
bst <- RFTrainer$new(n_estimators=10,
                       max_depth=4,
                       classification=1,
                       seed=42,
                       verbose=TRUE)
bst$fit(iris, 'Species')
```

**Method predict():**

**Usage:**
```r
RFTrainer$predict(df)
```

**Arguments:**
- `df` data.frame containing test features

**Details:** Return predictions from random forest model

**Returns:** a vector containing predictions

**Examples:**
```r
data("iris")
bst <- RFTrainer$new(n_estimators=10,
                       max_depth=4,
                       classification=1,
                       seed=42,
                       verbose=TRUE)
bst$fit(iris, 'Species')
predictions <- bst$predict(iris)
```

**Method get_importance():**

**Usage:**
```r
RFTrainer$get_importance()
```

**Details:** Returns feature importance from the model

**Returns:** a data frame containing feature predictions

**Examples:**
```r
data("iris")
bst <- RFTrainer$new(n_estimators=50,
                       max_depth=4,
                       classification=1,
                       seed=42,
                       verbose=TRUE)
bst$fit(iris, 'Species')
predictions <- bst$predict(iris)
bst)get_importance()
```
Method `clone()`: The objects of this class are cloneable with this method.

Usage:
RFTrainer$clone(deep = FALSE)

Arguments:
- deep Whether to make a deep clone.

Examples

```r
## Method `RFTrainer$new`
## ------------------------------
data("iris")
bst <- RFTrainer$new(n_estimators=10,  
                     max_depth=4,  
                     classification=1,  
                     seed=42,  
                     verbose=TRUE)
```

```r
## Method `RFTrainer$fit`
## ------------------------------
data("iris")
bst <- RFTrainer$new(n_estimators=10,  
                     max_depth=4,  
                     classification=1,  
                     seed=42,  
                     verbose=TRUE)
bst$fit(iris, 'Species')
```

```r
## Method `RFTrainer$predict`
## ------------------------------
data("iris")
bst <- RFTrainer$new(n_estimators=10,  
                     max_depth=4,  
                     classification=1,  
                     seed=42,  
                     verbose=TRUE)
bst$fit(iris, 'Species')
predictions <- bst$predict(iris)
```

```r
## Method `RFTrainer$get_importance`
## ------------------------------
data("iris")
bst <- RFTrainer$new(n_estimators=50,
```
smoothMean

smoothMean Calculator

Description

Calculates target encodings using a smoothing parameter and count of categorical variables. This approach is more robust to possibility of leakage and avoid overfitting.

Usage

```
smoothMean(
  train_df, test_df, colname, target,
  min_samples_leaf = 1, smoothing = 1, noise_level = 0
)
```

Arguments

- `train_df`: train dataset
- `test_df`: test dataset
- `colname`: name of categorical column
- `target`: name of target column
- `min_samples_leaf`: minimum samples to take category average into account
- `smoothing`: smoothing effect to balance categorical average vs prior
- `noise_level`: random noise to add, optional

Value

a train and test data table with mean encodings of the target for the given categorical variable
Examples

```r
train <- data.frame(region=c('del','csk','rcb','del','csk','pune','guj','del'),
                    win = c(0,1,1,0,0,1,0,1))
test <- data.frame(region=c('rcb','csk','rcb','del','guj','pune','csk','kol'))

# calculate encodings
all_means <- smoothMean(train_df = train,
                        test_df = test,
                        colname = 'region',
                        target = 'win')
train_mean <- all_means$train
test_mean <- all_means$test
```

Description

For a given vector, return the indexes of the sorted array and not the sorted array itself.

Usage

```r
sort_index(vec, ascending = TRUE)
```

Arguments

- `vec`: numeric vector
- `ascending`: logical, order to return (ascending or descending), default = True

Value

numeric vector containing sorted indexes

Examples

```r
v <- c(10,3,1,4)
j <- sort_index(v)
```
**TfidfVectorizer**

**TfidfVectorizer**

**Description**

Creates a tf-idf matrix

**Details**

Given a list of text, it creates a sparse matrix consisting of tf-idf score for tokens from the text.

**Super class**

`superml::CountVectorizer` -> `TfidfVectorizer`

**Public fields**

- `sentences` a list containing sentences
- `max_df` When building the vocabulary ignore terms that have a document frequency strictly higher than the given threshold, value lies between 0 and 1.
- `min_df` When building the vocabulary ignore terms that have a document frequency strictly lower than the given threshold, value lies between 0 and 1.
- `max_features` use top features sorted by count to be used in bag of words matrix.
- `ngram_range` The lower and upper boundary of the range of n-values for different word n-grams or char n-grams to be extracted. All values of n such such that min_n <= n <= max_n will be used. For example an ngram_range of c(1, 1) means only unigrams, c(1, 2) means unigrams and bigrams, and c(2, 2) means only bigrams.
- `split` splitting criteria for strings, default: " "
- `lowercase` convert all characters to lowercase before tokenizing
- `regex` regex expression to use for text cleaning.
- `remove_stopwords` a list of stopwords to use, by default it uses its inbuilt list of standard stopwords
- `smooth_idf` logical, to prevent zero division, adds one to document frequencies, as if an extra document was seen containing every term in the collection exactly once
- `norm` logical, if TRUE, each output row will have unit norm ‘l2’: Sum of squares of vector elements is 1. if FALSE returns non-normalized vectors, default: TRUE

**Methods**

**Public methods:**

- `TfidfVectorizer$new()`
- `TfidfVectorizer$fit()`
- `TfidfVectorizer$fit_transform()`
- `TfidfVectorizer$transform()`
• TfIdfVectorizer$clone()

Method new():

Usage:
TfIdfVectorizer$new(
  min_df,
  max_df,
  max_features,
  ngram_range,
  regex,
  remove_stopwords,
  split,
  lowercase,
  smooth_idf,
  norm
)

Arguments:

min_df numeric, When building the vocabulary ignore terms that have a document frequency strictly lower than the given threshold, value lies between 0 and 1.

max_df numeric, When building the vocabulary ignore terms that have a document frequency strictly higher than the given threshold, value lies between 0 and 1.

max_features integer, Build a vocabulary that only consider the top max_features ordered by term frequency across the corpus.

ngram_range vector, The lower and upper boundary of the range of n-values for different word n-grams or char n-grams to be extracted. All values of n such such that min_n <= n <= max_n will be used. For example an ngram_range of c(1, 1) means only unigrams, c(1, 2) means unigrams and bigrams, and c(2, 2) means only bigrams.

regex character, regex expression to use for text cleaning.

remove_stopwords list, a list of stopwords to use, by default it uses its inbuilt list of standard english stopwords

split character, splitting criteria for strings, default: " "

lowercase logical, convert all characters to lowercase before tokenizing, default: TRUE

smooth_idf logical, to prevent zero division, adds one to document frequencies, as if an extra document was seen containing every term in the collection exactly once

norm logical, if TRUE, each output row will have unit norm ‘l2’: Sum of squares of vector elements is 1. if FALSE returns non-normalized vectors, default: TRUE

Details: Create a new ‘TfIdfVectorizer’ object.

Returns: A ‘TfIdfVectorizer’ object.

Examples:
TfIdfVectorizer$new()

Method fit():

Usage:
TfIdfVectorizer$fit(sentences)
**TfidfVectorizer**

*Arguments:*
sentences a list of text sentences

*Details:*  Fits the TfidfVectorizer model on sentences

*Returns:*  NULL

*Examples:*
sents = c('i am alone in dark.','mother_mary a lot','alone in the dark?','many mothers in the lot....')
tf = TfidfVectorizer$new(smooth_idf = TRUE, min_df = 0.3)
tf$fit(sents)

**Method fit_transform():**

*Usage:*
TfidfVectorizer$fit_transform(sentences)

*Arguments:*
sentences  a list of text sentences

*Details:*  Fits the TfidfVectorizer model and returns a sparse matrix of count of tokens

*Returns:*  a sparse matrix containing tf-idf score for tokens in each given sentence

*Examples:*
\dontrun{
sents <- c('i am alone in dark.','mother_mary a lot','alone in the dark?','many mothers in the lot....')
tf <- TfidfVectorizer$new(smooth_idf = TRUE, min_df = 0.1)
tf_matrix <- tf$fit_transform(sents)
}

**Method transform():**

*Usage:*
TfidfVectorizer$transform(sentences)

*Arguments:*
sentences  a list of new text sentences

*Details:*  Returns a matrix of tf-idf score of tokens

*Returns:*  a sparse matrix containing tf-idf score for tokens in each given sentence

*Examples:*
\dontrun{
sents = c('i am alone in dark.','mother_mary a lot','alone in the dark?','many mothers in the lot....')
new_sents <- c("dark at night","mothers day")
tf = TfidfVectorizer$new(min_df=0.1)
tf$fit(sents)
tf_matrix <- tf$transform(new_sents)
}

**Method clone():**  The objects of this class are cloneable with this method.
Usage:
TfIdfVectorizer$clone(deep = FALSE)

Arguments:
deep  Whether to make a deep clone.

Examples

```
## Method `TfIdfVectorizer$new'
TfIdfVectorizer$new()

## Method `TfIdfVectorizer$fit'
sents = c('i am alone in dark.', 'mother_mary a lot',
  'alone in the dark?', 'many mothers in the lot....')
tf = TfIdfVectorizer$new(smooth_idf = TRUE, min_df = 0.3)
tf$fit(sents)

## Method `TfIdfVectorizer$fit_transform'
sents <- c('i am alone in dark.', 'mother_mary a lot',
  'alone in the dark?', 'many mothers in the lot....')
tf <- TfIdfVectorizer$new(smooth_idf = TRUE, min_df = 0.1)
tf_matrix <- tf$fit_transform(sents)

## Method `TfIdfVectorizer$transform'
new_sents <- c("dark at night", "mothers day")
tf = TfIdfVectorizer$new(min_df=0.1)
tf$fit(sents)
tf_matrix <- tf$transform(new_sents)
```
XGBTrainer  

Extreme Gradient Boosting Trainer

Description

Trains a XGBoost model in R

Details

Trains a Extreme Gradient Boosting Model. XGBoost belongs to a family of boosting algorithms that creates an ensemble of weak learner to learn about data. It is a wrapper for original xgboost R package, you can find the documentation here: http://xgboost.readthedocs.io/en/latest/parameter.html

Public fields

booster  the trainer type, the values are gbtree(default), gblinear, dart:gbtree
objective  specify the learning task. Check the link above for all possible values.
nthread  number of parallel threads used to run, default is to run using all threads available
silent  0 means printing running messages, 1 means silent mode
n_estimators  number of trees to grow, default = 100
learning_rate  Step size shrinkage used in update to prevents overfitting. Lower the learning rate, more time it takes in training, value lies between between 0 and 1. Default = 0.3
gamma Minimum loss reduction required to make a further partition on a leaf node of the tree. The larger gamma is, the more conservative the algorithm will be. Value lies between 0 and infinity. Default = 0
max_depth  the maximum depth of each tree, default = 6
min_child_weight Minimum sum of instance weight (hessian) needed in a child. If the tree partition step results in a leaf node with the sum of instance weight less than min_child_weight, then the building process will give up further partitioning. In linear regression task, this simply corresponds to minimum number of instances needed to be in each node. The larger min_child_weight is, the more conservative the algorithm will be. Value lies between 0 and infinity. Default = 1
subsample  Subsample ratio of the training instances. Setting it to 0.5 means that XGBoost would randomly sample half of the training data prior to growing trees, and this will prevent overfitting. Subsampling will occur once in every boosting iteration. Value lies between 0 and 1. Default = 1
colsample_bytree  Subsample ratio of columns when constructing each tree. Subsampling will occur once in every boosting iteration. Value lies between 0 and 1. Default = 1
lambda  L2 regularization term on weights. Increasing this value will make model more conservative. Default = 1
alpha  L1 regularization term on weights. Increasing this value will make model more conservative. Default = 0
eval_metric Evaluation metrics for validation data, a default metric will be assigned according to objective
print_every print training log after n iterations. Default = 50
feval custom evaluation function
early_stopping Used to prevent overfitting, stops model training after this number of iterations if there is no improvement seen
maximize If feval and early_stopping_rounds are set, then this parameter must be set as well. When it is TRUE, it means the larger the evaluation score the better.
custom_objective custom objective function
save_period when it is non-NULL, model is saved to disk after every save_period rounds, 0 means save at the end.
save_name the name or path for periodically saved model file.
xgb_model a previously built model to continue the training from. Could be either an object of class xgb.Booster, or its raw data, or the name of a file with a previously saved model.
callbacks a list of callback functions to perform various task during boosting. See callbacks. Some of the callbacks are automatically created depending on the parameters’ values. User can provide either existing or their own callback methods in order to customize the training process.
verbose If 0, xgboost will stay silent. If 1, xgboost will print information of performance. If 2, xgboost will print some additional information. Setting verbose > 0 automatically engages the cb.evaluation.log and cb.print.evaluation callback functions.
watchlist what information should be printed when verbose=1 or verbose=2. Watchlist is used to specify validation set monitoring during training. For example user can specify watchlist=list(validation1=mat1, validation2=mat2) to watch the performance of each round’s model on mat1 and mat2
num_class set number of classes in case of multiclassification problem
weight a vector indicating the weight for each row of the input.
na_missing by default is set to NA, which means that NA values should be considered as 'missing' by the algorithm. Sometimes, 0 or other extreme value might be used to represent missing values. This parameter is only used when input is a dense matrix.
feature_names internal use, stores the feature names for model importance
cv_model internal use

Methods

Public methods:
- XGBTrainer$new()
- XGBTrainer$cross_val()
- XGBTrainer$fit()
- XGBTrainer$predict()
- XGBTrainer$show_importance()
- XGBTrainer$clone()
Method new():

Usage:
XGBTrainer$new(
  booster,
  objective,
  nthread,
  silent,
  n_estimators,
  learning_rate,
  gamma,
  max_depth,
  min_child_weight,
  subsample,
  colsample_bytree,
  lambda,
  alpha,
  eval_metric,
  print_every,
  feval,
  early_stopping,
  maximize,
  custom_objective,
  save_period,
  save_name,
  xgb_model,
  callbacks,
  verbose,
  num_class,
  weight,
  na_missing
)

Arguments:

booster the trainer type, the values are gbtree(default), gblinear, dart:gbtree
objective specify the learning task. Check the link above for all possible values.
nthread number of parallel threads used to run, default is to run using all threads available
silent 0 means printing running messages, 1 means silent mode
n_estimators number of trees to grow, default = 100
learning_rate Step size shrinkage used in update to prevents overfitting. Lower the learning
rate, more time it takes in training, value lies between between 0 and 1. Default = 0.3
gamma Minimum loss reduction required to make a further partition on a leaf node of the tree.
The larger gamma is, the more conservative the algorithm will be. Value lies between 0 and
infinity, Default = 0
max_depth the maximum depth of each tree, default = 6
min_child_weight Minimum sum of instance weight (hessian) needed in a child. If the tree
partition step results in a leaf node with the sum of instance weight less than min_child_weight,
then the building process will give up further partitioning. In linear regression task, this
simply corresponds to minimum number of instances needed to be in each node. The larger min_child_weight is, the more conservative the algorithm will be. Value lies between 0 and infinity. Default = 1

subsample Subsample ratio of the training instances. Setting it to 0.5 means that XGBoost would randomly sample half of the training data prior to growing trees. and this will prevent overfitting. Subsampling will occur once in every boosting iteration. Value lies between 0 and 1. Default = 1

colsample_bytree Subsample ratio of columns when constructing each tree. Subsampling will occur once in every boosting iteration. Value lies between 0 and 1. Default = 1

lambda L2 regularization term on weights. Increasing this value will make model more conservative. Default = 1

alpha L1 regularization term on weights. Increasing this value will make model more conservative. Default = 0

eval_metric Evaluation metrics for validation data, a default metric will be assigned according to objective

print_every print training log after n iterations. Default = 50

eval custom evaluation function

early_stopping Used to prevent overfitting, stops model training after this number of iterations if there is no improvement seen

maximize If feval and early_stopping_rounds are set, then this parameter must be set as well.

When it is TRUE, it means the larger the evaluation score the better.

custom_objective custom objective function

save_period when it is non-NULL, model is saved to disk after every save_period rounds, 0 means save at the end.

save_name the name or path for periodically saved model file.

xgb_model a previously built model to continue the training from. Could be either an object of class xgb.Booster, or its raw data, or the name of a file with a previously saved model.

callbacks a list of callback functions to perform various task during boosting. See callbacks.

Some of the callbacks are automatically created depending on the parameters’ values. User can provide either existing or their own callback methods in order to customize the training process.

verbose If 0, xgboost will stay silent. If 1, xgboost will print information of performance. If 2, xgboost will print some additional information. Setting verbose > 0 automatically engages the cb.evaluation.log and cb.print.evaluation callback functions.

num_class set number of classes in case of multiclassification problem

weight a vector indicating the weight for each row of the input.

na_missing by default is set to NA, which means that NA values should be considered as ‘missing’ by the algorithm. Sometimes, 0 or other extreme value might be used to represent missing values. This parameter is only used when input is a dense matrix.

Details: Create a new ‘XGBTrainer’ object.

Returns: A ‘XGBTrainer’ object.

Examples:

library(data.table)
df <- copy(iris)
# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species))-1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
  maximize = FALSE,
  eval_metric = 'merror',
  num_class=3,
  n_estimators = 2)

Method cross_val():

Usage:
XGBTrainer$cross_val(X, y, nfolds = 5, stratified = TRUE, folds = NULL)

Arguments:
X data.frame
y character, name of target variable
nfolds integer, number of folds
stratified logical, whether to use stratified sampling
folds the list of CV folds’ indices - either those passed through the folds parameter or randomly generated.

Details: Trains the xgboost model using cross validation scheme

Returns: NULL, trains a model and saves it in memory

Examples:
\dontrun{
library(data.table)
df <- copy(iris)

# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species))-1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
  maximize = FALSE,
  eval_metric = 'merror',
  num_class=3,
  n_estimators = 2)

# do cross validation to find optimal value for n_estimators
xgb$cross_val(X = df, y = 'Species',nfolds = 3, stratified = TRUE)
}

Method fit():

Usage:
XGBTrainer$fit(X, y, valid = NULL)
**Arguments:**
- `X` data.frame, training data
- `y` character, name of target variable
- `valid` data.frame, validation data

**Details:** Fits the xgboost model on given data

**Returns:** NULL, trains a model and keeps it in memory

**Examples:**

```r
library(data.table)
df <- copy(iris)

# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species))-1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
    maximize = FALSE,
    eval_metric = 'merror',
    num_class=3,
    n_estimators = 2)

xgb$fit(df, 'Species')
```

**Method predict():**

**Usage:**

```r
XGBTrainer$predict(df)
```

**Arguments:**
- `df` data.frame, test data set

**Details:** Returns predicted values for a given test data

**Returns:** xgboost predictions

**Examples:**

```r
# library(data.table)
df <- copy(iris)

# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species))-1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
    maximize = FALSE,
    eval_metric = 'merror',
    num_class=3,
    n_estimators = 2)

xgb$fit(df, 'Species')

# make predictions
preds <- xgb$predict(as.matrix(iris[,1:4]))
```
**Method** show_importance():

*Usage:*

XGBTrainer$show_importance(type = "plot", topn = 10)

*Arguments:*

- **type** character, could be 'plot' or 'table'
- **topn** integer, top n features to display

*Details:* Shows feature importance plot

*Returns:* a table or a plot of feature importance

*Examples:*

```r
\dontrun{
library(data.table)
df <- copy(iris)

# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species))-1

# initialise model
xgb <- XGBTrainer$new(objective = "multi:softmax",
                         maximize = FALSE,
                         eval_metric = "merror",
                         num_class=3,
                         n_estimators = 2)
xgb$fit(df, 'Species')
xgb$show_importance()
}
```

**Method** clone(): The objects of this class are cloneable with this method.

*Usage:*

XGBTrainer$clone(deep = FALSE)

*Arguments:*

- **deep** Whether to make a deep clone.

*Examples*

```r
## ------------------------------------------------
## Method XGBTrainer$new
## ------------------------------------------------
library(data.table)
df <- copy(iris)

# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species))-1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
                       maximize = FALSE,
                       eval_metric = 'merror',
                       num_class=3,
                       n_estimators = 2)
xgb$fit(df, 'Species')
xgb$show_importance()
```
```r
xgb <- XGBTrainer$new(objective = 'multi:softmax',
                       maximize = FALSE,
                       eval_metric = 'merror',
                       num_class=3,
                       n_estimators = 2)

## Method `XGBTrainer$cross_val`

library(data.table)
df <- copy(iris)
# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species)) - 1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
                       maximize = FALSE,
                       eval_metric = 'merror',
                       num_class=3,
                       n_estimators = 2)

# do cross validation to find optimal value for n_estimators
xgb$cross_val(X = df, y = 'Species', nfolds = 3, stratified = TRUE)

## Method `XGBTrainer$fit`

library(data.table)
df <- copy(iris)
# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species)) - 1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
                       maximize = FALSE,
                       eval_metric = 'merror',
                       num_class=3,
                       n_estimators = 2)
xgb$fit(df, 'Species')

## Method `XGBTrainer$predict`

```

# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species))-1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
                      maximize = FALSE,
                      eval_metric = 'merror',
                      num_class=3,
                      n_estimators = 2)

xgb$fit(df, 'Species')

# make predictions
preds <- xgb$predict(as.matrix(iris[,1:4]))

# Method `XGBTrainer$show_importance`

library(data.table)
df <- copy(iris)

# convert characters/factors to numeric
df$Species <- as.numeric(as.factor(df$Species))-1

# initialise model
xgb <- XGBTrainer$new(objective = 'multi:softmax',
                      maximize = FALSE,
                      eval_metric = 'merror',
                      num_class=3,
                      n_estimators = 2)

xgb$fit(df, 'Species')
xgb$show_importance()

# End(Not run)
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