

Package: daltoolbox (via r-universe)

September 13, 2024

Title Leveraging Experiment Lines to Data Analytics

Version 1.0.767

Description The natural increase in the complexity of current research experiments and data demands better tools to enhance productivity in Data Analytics. The package is a framework designed to address the modern challenges in data analytics workflows. The package is inspired by Experiment Line concepts. It aims to provide seamless support for users in developing their data mining workflows by offering a uniform data model and method API. It enables the integration of various data mining activities, including data preprocessing, classification, regression, clustering, and time series prediction. It also offers options for hyper-parameter tuning and supports integration with existing libraries and languages. Overall, the package provides researchers with a comprehensive set of functionalities for data science, promoting ease of use, extensibility, and integration with various tools and libraries. Information on Experiment Line is based on Ogasawara et al. (2009) <[doi:10.1007/978-3-642-02279-1_20](https://doi.org/10.1007/978-3-642-02279-1_20)>.

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URL <https://github.com/cefet-rj-dal/daltoolbox>

Encoding UTF-8

Roxygen list(markdown = TRUE)

RoxygenNote 7.3.1

Imports FNN, MLmetrics, caret, class, cluster, dbscan, dplyr, e1071, elmNNRcpp, forecast, ggplot2, nnet, randomForest, reshape, tree, reticulate

Config/reticulate list(packages = list(list(package = ``scipy"), list(package = ``torch"), list(package = ``pandas"), list(package = ``numpy"), list(package = ``matplotlib"), list(package = ``scikit-learn"), list(package = ``functools"), list(package = ``operator"), list(package = ``sys")))

Repository <https://cefet-rj-dal.r-universe.dev>

RemoteUrl <https://github.com/cefet-rj-dal/daltoolbox>

RemoteRef HEAD

RemoteSha c6b64bac989905da343412a0c3e6488272338449

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action	<i>Action</i>
--------	---------------

Description

Executes the action of model applied in provided data

Usage

```
action(obj, ...)
```

Arguments

obj	object: a dal_base object to apply the transformation on the input dataset.
...	optional arguments.

Value

The result of an action of the model applied in provided data

Examples

```
data(iris)
# an example is minmax normalization
trans <- minmax()
trans <- fit(trans, iris)
tiris <- action(trans, iris)
```

action.dal_transform *Action implementation for transform*

Description

A default function that defines the action to proxy transform method

Usage

```
## S3 method for class 'dal_transform'  
action(obj, ...)
```

Arguments

obj	object
...	optional arguments

Value

Transformed data

Examples

```
#See ?minmax for an example of transformation
```

adjust_class_label *adjust categorical mapping*

Description

vector value is adjusted to a categorical mapping

Usage

```
adjust_class_label(x, valTrue = 1, valFalse = 0)
```

Arguments

x	vector to be categorized
valTrue	value to represent true
valFalse	value to represent false

Value

an adjusted categorical mapping

adjust_data.frame *Adjust to data frame*

Description

dataset data is adjusted to a data.frame

Usage

```
adjust_data.frame(data)
```

Arguments

data dataset

Value

The data argument

Examples

```
data(iris)
df <- adjust_data.frame(iris)
```

adjust_factor *adjust factors*

Description

vector value is adjusted to a factor

Usage

```
adjust_factor(value, ilevels, slevels)
```

Arguments

value vector to be converted into factor
ilevels order for categorical values
slevels labels for categorical values

Value

an adjusted factor

adjust_matrix	<i>adjust to matrix</i>
---------------	-------------------------

Description

dataset data is adjusted to a matrix

Usage

```
adjust_matrix(data)
```

Arguments

data dataset

Value

an adjusted matrix

Examples

```
data(iris)
mat <- adjust_matrix(iris)
```

adjust_ts_data	<i>adjust ts_data</i>
----------------	-----------------------

Description

dataset data is adjusted to a ts_data

Usage

```
adjust_ts_data(data)
```

Arguments

data dataset

Value

an adjusted ts_data

autoenc_encode *Autoencoder - Encode*

Description

Creates an deep learning autoencoder to encode a sequence of observations. It wraps the pytorch library.

Usage

```
autoenc_encode(  
    input_size,  
    encoding_size,  
    batch_size = 32,  
    num_epochs = 1000,  
    learning_rate = 0.001  
)
```

Arguments

input_size	input size
encoding_size	encoding size
batch_size	size for batch learning
num_epochs	number of epochs for training
learning_rate	learning rate

Value

a autoenc_encode object.

Examples

```
#See example at https://nbviewer.org/github/cefet-rj-dal/daltoolbox-examples
```

autoenc_encode_decode *Autoencoder - Encode*

Description

Creates an deep learning autoencoder to encode a sequence of observations. It wraps the pytorch library.

Usage

```

autoenc_encode_decode(
  input_size,
  encoding_size,
  batch_size = 32,
  num_epochs = 1000,
  learning_rate = 0.001
)

```

Arguments

input_size	input size
encoding_size	encoding size
batch_size	size for batch learning
num_epochs	number of epochs for training
learning_rate	learning rate

Value

a autoenc_encode_decode object.

Examples

#See example at <https://nbviewer.org/github/cefet-rj-dal/daltoolbox-examples>

Boston

Boston Housing Data (Regression)

Description

housing values in suburbs of Boston.

- crim: per capita crime rate by town.
- zn: proportion of residential land zoned for lots over 25,000 sq.ft.
- indus: proportion of non-retail business acres per town
- chas: Charles River dummy variable (= 1 if tract bounds)
- nox: nitric oxides concentration (parts per 10 million)
- rm: average number of rooms per dwelling
- age: proportion of owner-occupied units built prior to 1940
- dis: weighted distances to five Boston employment centres
- rad: index of accessibility to radial highways
- tax: full-value property-tax rate per \$10,000
- ptratio: pupil-teacher ratio by town
- black: $1000(Bk - 0.63)^2$ where Bk is the proportion of blacks by town
- lstat: percentage of lower status of the population
- medv: Median value of owner-occupied homes in \$1000's

Usage

```
data(Boston)
```

Format

Regression Dataset.

Source

This dataset was obtained from the MASS library.

References

Creator: Harrison, D. and Rubinfeld, D.L. Hedonic prices and the demand for clean air, J. Environ. Economics & Management, vol.5, 81-102, 1978.

Examples

```
data(Boston)
head(Boston)
```

categ_mapping

Categorical mapping

Description

Categorical mapping provides a way to map the levels of a categorical variable to new values. Each possible value is converted to a binary attribute.

Usage

```
categ_mapping(attribute)
```

Arguments

attribute attribute to be categorized.

Value

A data frame with binary attributes, one for each possible category.

Examples

```
cm <- categ_mapping("Species")
iris_cm <- transform(cm, iris)

# can be made in a single column
species <- iris[, "Species", drop=FALSE]
iris_cm <- transform(cm, species)
```

classification	<i>classification</i>
----------------	-----------------------

Description

Ancestor class for classification problems

Usage

```
classification(attribute, slevels)
```

Arguments

attribute	attribute target to model building
slevels	• possible values for the target classification

Value

classification object

Examples

```
#See ?cla_dtree for a classification example using a decision tree
```

cla_dtree	<i>Decision Tree for classification</i>
-----------	-----------------------------------------

Description

Creates a classification object that uses the Decision Tree algorithm for classification. It wraps the tree library.

Usage

```
cla_dtree(attribute, slevels)
```

Arguments

attribute	attribute target to model building.
slevels	The possible values for the target classification.

Value

A classification object that uses the Decision Tree algorithm for classification.

Examples

```
data(iris)
slevels <- levels(iris$Species)
model <- cla_dtree("Species", slevels)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, iris)
train <- sr$train
test <- sr$test

model <- fit(model, train)

prediction <- predict(model, test)
predictand <- adjust_class_label(test[, "Species"])
test_eval <- evaluate(model, predictand, prediction)
test_eval$metrics
```

cla_knn

K Nearest Neighbor Classification

Description

Classifies using the K-Nearest Neighbor algorithm. It wraps the class library.

Usage

```
cla_knn(attribute, slevels, k = 1)
```

Arguments

attribute	attribute target to model building.
slevels	Possible values for the target classification.
k	A vector of integers indicating the number of neighbors to be considered.

Value

A knn object.

Examples

```
data(iris)
slevels <- levels(iris$Species)
model <- cla_knn("Species", slevels, k=3)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, iris)
```

```
train <- sr$train
test <- sr$test

model <- fit(model, train)

prediction <- predict(model, test)
predictand <- adjust_class_label(test[, "Species"])
test_eval <- evaluate(model, predictand, prediction)
test_eval$metrics
```

cla_majority

Majority Classification

Description

This function creates a classification object that uses the majority vote strategy to predict the target attribute. Given a target attribute, the function counts the number of occurrences of each value in the dataset and selects the one that appears most often.

Usage

```
cla_majority(attribute, slevels)
```

Arguments

attribute	attribute target to model building.
slevels	Possible values for the target classification.

Value

Returns a classification object.

Examples

```
data(iris)
slevels <- levels(iris$Species)
model <- cla_majority("Species", slevels)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, iris)
train <- sr$train
test <- sr$test

model <- fit(model, train)

prediction <- predict(model, test)
predictand <- adjust_class_label(test[, "Species"])
test_eval <- evaluate(model, predictand, prediction)
test_eval$metrics
```

`cla_mlp`*MLP for classification*

Description

Creates a classification object that uses the Multi-Layer Perceptron (MLP) method. It wraps the nnet library.

Usage

```
cla_mlp(attribute, slevels, size = NULL, decay = 0.1, maxit = 1000)
```

Arguments

<code>attribute</code>	attribute target to model building
<code>slevels</code>	possible values for the target classification
<code>size</code>	number of nodes that will be used in the hidden layer
<code>decay</code>	how quickly it decreases in gradient descent
<code>maxit</code>	maximum iterations

Value

a classification object

Examples

```
data(iris)
slevels <- levels(iris$Species)
model <- cla_mlp("Species", slevels, size=3, decay=0.03)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, iris)
train <- sr$train
test <- sr$test

model <- fit(model, train)

prediction <- predict(model, test)
predictand <- adjust_class_label(test[, "Species"])
test_eval <- evaluate(model, predictand, prediction)
test_eval$metrics
```

cla_nb	<i>Naive Bayes Classifier</i>
--------	-------------------------------

Description

Classification using the Naive Bayes algorithm It wraps the e1071 library.

Usage

```
cla_nb(attribute, slevels)
```

Arguments

attribute	attribute target to model building.
slevels	Possible values for the target classification.

Value

A classification object.

Examples

```
data(iris)
slevels <- levels(iris$Species)
model <- cla_nb("Species", slevels)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, iris)
train <- sr$train
test <- sr$test

model <- fit(model, train)

prediction <- predict(model, test)
predictand <- adjust_class_label(test[, "Species"])
test_eval <- evaluate(model, predictand, prediction)
test_eval$metrics
```

cla_rf	<i>Random Forest for classification</i>
--------	-----------------------------------------

Description

Creates a classification object that uses the Random Forest method It wraps the randomForest library.

Usage

```
cla_rf(attribute, slevels, nodesize = 5, ntree = 10, mtry = NULL)
```

Arguments

attribute	attribute target to model building
slevels	possible values for the target classification
nodesize	node size
ntree	number of trees
mtry	number of attributes to build tree

Value

obj

Examples

```
data(iris)
slevels <- levels(iris$Species)
model <- cla_rf("Species", slevels, ntree=5)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, iris)
train <- sr$train
test <- sr$test

model <- fit(model, train)

prediction <- predict(model, test)
predictand <- adjust_class_label(test[, "Species"])
test_eval <- evaluate(model, predictand, prediction)
test_eval$metrics
```

cla_svm

SVM for classification

Description

Creates a classification object that uses the Support Vector Machine (SVM) method for classification. It wraps the e1071 library.

Usage

```
cla_svm(attribute, slevels, epsilon = 0.1, cost = 10, kernel = "radial")
```


Arguments

attribute	attribute target to model building
slevels	possible values for the target classification
epsilon	parameter that controls the width of the margin around the separating hyperplane
cost	parameter that controls the trade-off between having a wide margin and correctly classifying training data points
kernel	the type of kernel function to be used in the SVM algorithm (linear, radial, polynomial, sigmoid)

Value

A SVM classification object

Examples

```
data(iris)
slevels <- levels(iris$Species)
model <- cla_svm("Species", slevels, epsilon=0.0, cost=20.000)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, iris)
train <- sr$train
test <- sr$test

model <- fit(model, train)

prediction <- predict(model, test)
predictand <- adjust_class_label(test[, "Species"])
test_eval <- evaluate(model, predictand, prediction)
test_eval$metrics
```

 cla_tune

Classification Tune

Description

Classification Tune

Usage

```
cla_tune(base_model, folds = 10, metric = "accuracy")
```

Arguments

base_model	base model for tuning
folds	number of folds for cross-validation
metric	metric used to optimize

Value

a `cla_tune` object.

Examples

```
# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, iris)
train <- sr$train
test <- sr$test

# hyper parameter setup
tune <- cla_tune(cla_mlp("Species", levels(iris$Species)))
ranges <- list(size=c(3:5), decay=c(0.1))

# hyper parameter optimization
model <- fit(tune, train, ranges)

# testing optimization
test_prediction <- predict(model, test)
test_predictand <- adjust_class_label(test[, "Species"])
test_eval <- evaluate(model, test_predictand, test_prediction)
test_eval$metrics
```

cluster

Cluster

Description

Defines a cluster method.

Usage

```
cluster(obj, ...)
```

Arguments

`obj` a clusterer object.
`...` optional arguments.

Value

clustered data.

Examples

```
#See ?cluster_kmeans for an example of transformation
```

clusterer	<i>Clusterer</i>
-----------	------------------

Description

Ancestor class for clustering problems

Usage

```
clusterer()
```

Value

a clusterer object

Examples

```
#See ?cluster_kmeans for an example of transformation
```

cluster_dbscan	<i>DBSCAN</i>
----------------	---------------

Description

Creates a clusterer object that uses the DBSCAN method It wraps the dbscan library.

Usage

```
cluster_dbscan(minPts = 3, eps = NULL)
```

Arguments

minPts	minimum number of points
eps	distance value

Value

A dbscan object.

Examples

```
# setup clustering
model <- cluster_dbscan(minPts = 3)

#load dataset
data(iris)

# build model
model <- fit(model, iris[,1:4])
clu <- cluster(model, iris[,1:4])
table(clu)

# evaluate model using external metric
eval <- evaluate(model, clu, iris$Species)
eval
```

cluster_kmeans	<i>k-means</i>
----------------	----------------

Description

Creates a clusterer object that uses the k-means method It wraps the stats library.

Usage

```
cluster_kmeans(k = 1)
```

Arguments

k The number of clusters to form.

Value

A k-means object.

Examples

```
# setup clustering
model <- cluster_kmeans(k=3)

#load dataset
data(iris)

# build model
model <- fit(model, iris[,1:4])
clu <- cluster(model, iris[,1:4])
table(clu)

# evaluate model using external metric
eval <- evaluate(model, clu, iris$Species)
eval
```

cluster_pam	<i>PAM</i>
-------------	------------

Description

Creates a clusterer object that uses the Partition Around Medoids (PAM) method It wraps the cluster library.

Usage

```
cluster_pam(k = 1)
```

Arguments

k The number of clusters to generate.

Value

A PAM object.

Examples

```
# setup clustering
model <- cluster_pam(k = 3)

#load dataset
data(iris)

# build model
model <- fit(model, iris[,1:4])
clu <- cluster(model, iris[,1:4])
table(clu)

# evaluate model using external metric
eval <- evaluate(model, clu, iris$Species)
eval
```

clu_tune	<i>Clustering Tune</i>
----------	------------------------

Description

Clustering Tune

Usage

```
clu_tune(base_model)
```

Arguments

base_model base model for tuning

Value

a clu_tune object.

Examples

```
data(iris)

# fit model
model <- clu_tune(cluster_kmeans(k = 0))
ranges <- list(k = 1:10)
model <- fit(model, iris[,1:4], ranges)
model$k
```

dal_base

Class dal_base

Description

The dal_base class is an abstract class for all dal descendants classes. It provides both fit() and action() functions

Usage

```
dal_base()
```

Value

A dal_base object

Examples

```
trans <- dal_base()
```

`dal_learner`*DAL Learner*

Description

A ancestor class for clustering, classification, regression, and time series regression. It also provides the basis for specialized evaluation of learning performance.

An example of a learner is a decision tree (`cla_dtree`)

Usage

```
dal_learner()
```

Value

a learner

Examples

```
#See ?cla_dtree for a classification example using a decision tree
```

`dal_transform`*DAL Transform*

Description

A transformation method applied to a dataset. If needed, the fit can be called to adjust the transform.

Usage

```
dal_transform()
```

Value

a `dal_transform` object.

Examples

```
#See ?minmax for an example of transformation
```

`dal_tune`*DAL Tune*

Description

Ancestor class for hyper parameter optimization

Usage

```
dal_tune(base_model, folds = 10)
```

Arguments

<code>base_model</code>	base model for tuning
<code>folds</code>	number of folds for cross-validation

Value

a `dal_tune` object.

Examples

```
#See ?cla_tune for classification tuning  
#See ?reg_tune for regression tuning  
#See ?ts_tune for time series tuning
```

`data_sample`*Data Sample*

Description

The `data_sample` function in R is used to randomly sample data from a given data frame. It can be used to obtain a subset of data for further analysis or modeling.

Two basic specializations of `data_sample` are `sample_random` and `sample_stratified`. They provide random sampling and stratified sampling, respectively.

Data sample provides both training and testing partitioning (`train_test`) and k-fold partitioning (`k_fold`) of data.

Usage

```
data_sample()
```

Value

`obj`

Examples

```
#using random sampling
sample <- sample_random()
tt <- train_test(sample, iris)

# distribution of train
table(tt$train$Species)

# preparing dataset into four folds
folds <- k_fold(sample, iris, 4)

# distribution of folds
tbl <- NULL
for (f in folds) {
  tbl <- rbind(tbl, table(f$Species))
}
head(tbl)
```

do_fit

do fit for time series

Description

The actual time series model fitting. This method should be override by descendants.

Usage

```
do_fit(obj, x, y = NULL)
```

Arguments

obj	object
x	input variable
y	output variable

Value

fitted object

do_predict	<i>do predict for time series</i>
------------	-----------------------------------

Description

The actual time series model prediction. This method should be override by descendants.

Usage

```
do_predict(obj, x)
```

Arguments

obj	object
x	input variable

Value

predicted values

dt_pca	<i>PCA</i>
--------	------------

Description

PCA (Principal Component Analysis) is an unsupervised dimensionality reduction technique used in data analysis and machine learning. It transforms a dataset of possibly correlated variables into a new set of uncorrelated variables called principal components.

Usage

```
dt_pca(attribute = NULL, components = NULL)
```

Arguments

attribute	target attribute to model building
components	number of components for PCA

Value

obj

Examples

```
mypca <- dt_pca("Species")
# Automatically fitting number of components
mypca <- fit(mypca, iris)
iris.pca <- transform(mypca, iris)
head(iris.pca)
head(mypca$pca.transf)
# Manual establishment of number of components
mypca <- dt_pca("Species", 3)
mypca <- fit(mypca, datasets::iris)
iris.pca <- transform(mypca, iris)
head(iris.pca)
head(mypca$pca.transf)
```

evaluate

evaluate

Description

evaluate learner performance. The actual evaluate varies according to the type of learner (clustering, classification, regression, time series regression)

Usage

```
evaluate(obj, ...)
```

Arguments

obj	object
...	optional arguments

Value

evaluation

Examples

```
data(iris)
slevels <- levels(iris$Species)
model <- cla_dtree("Species", slevels)
model <- fit(model, iris)
prediction <- predict(model, iris)
predictand <- adjust_class_label(iris[, "Species"])
test_eval <- evaluate(model, predictand, prediction)
test_eval$metrics
```

fit	<i>Fit</i>
-----	------------

Description

Fits a model.

Usage

```
fit(obj, ...)
```

Arguments

obj	object
...	optional arguments.

Value

obj

Examples

```
data(iris)
# an example is minmax normalization
trans <- minmax()
trans <- fit(trans, iris)
tiris <- action(trans, iris)
```

fit.cla_tune	<i>tune hyperparameters of ml model</i>
--------------	-----------------------------------------

Description

tune hyperparameters of ml model for classification

Usage

```
## S3 method for class 'cla_tune'
fit(obj, data, ranges, ...)
```

Arguments

obj	object
data	dataset
ranges	hyperparameters ranges
...	optional arguments

Value

fitted obj

 fit.cluster_dbscan *fit dbscan model*

Description

fit dbscan model

Usage

```
## S3 method for class 'cluster_dbscan'
fit(obj, data, ...)
```

Arguments

obj	object
data	dataset
...	optional arguments

Value

fitted obj

 fit_curvature_max *maximum curvature analysis*

Description

Fitting a curvature model in a sequence of observations. It extracts the the maximum curvature computed.

Usage

```
fit_curvature_max()
```

Value

Returns an object of class `fit_curvature_max`, which inherits from the `fit_curvature` and `dal_transform` classes. The object contains a list with the following elements:

- `x`: The position in which the maximum curvature is reached.
- `y`: The value where the the maximum curvature occurs.
- `yfit`: The value of the maximum curvature.

Examples

```
x <- seq(from=1,to=10,by=0.5)
dat <- data.frame(x = x, value = -log(x), variable = "log")
myfit <- fit_curvature_max()
res <- transform(myfit, dat$value)
head(res)
```

fit_curvature_min	<i>minimum curvature analysis</i>
-------------------	-----------------------------------

Description

Fitting a curvature model in a sequence of observations. It extracts the the minimum curvature computed.

Usage

```
fit_curvature_min()
```

Value

Returns an object of class `fit_curvature_max`, which inherits from the `fit_curvature` and `dal_transform` classes. The object contains a list with the following elements:

- `x`: The position in which the minimum curvature is reached.
- `y`: The value where the the minimum curvature occurs.
- `yfit`: The value of the minimum curvature.

Examples

```
x <- seq(from=1,to=10,by=0.5)
dat <- data.frame(x = x, value = log(x), variable = "log")
myfit <- fit_curvature_min()
res <- transform(myfit, dat$value)
head(res)
```

inverse_transform	<i>Inverse Transform</i>
-------------------	--------------------------

Description

Reverses the transformation applied to data.

Usage

```
inverse_transform(obj, ...)
```

Arguments

obj	a dal_transform object.
...	optional arguments.

Value

dataset inverse transformed.

Examples

```
#See ?minmax for an example of transformation
```

k_fold	<i>k-fold sampling</i>
--------	------------------------

Description

k-fold partition of a dataset using a sampling method

Usage

```
k_fold(obj, data, k)
```

Arguments

obj	object
data	dataset
k	number of folds

Value

k folds

Examples

```
#using random sampling
sample <- sample_random()

# preparing dataset into four folds
folds <- k_fold(sample, iris, 4)

# distribution of folds
tbl <- NULL
for (f in folds) {
  tbl <- rbind(tbl, table(f$Species))
}
head(tbl)
```

minmax

min-max normalization

Description

The minmax performs scales data between [0,1]. $\text{minmax} = (x - \min(x)) / (\max(x) - \min(x))$.

Usage

```
minmax()
```

Value

```
obj
```

Examples

```
data(iris)
head(iris)

trans <- minmax()
trans <- fit(trans, iris)
tiris <- transform(trans, iris)
head(tiris)

itiris <- inverse_transform(trans, tiris)
head(itiris)
```

MSE.ts

MSE

Description

Compute the mean squared error (MSE) between actual values and forecasts of a time series

Usage

```
MSE.ts(actual, prediction)
```

Arguments

actual	real observations
prediction	predicted observations

Value

A number, which is the calculated MSE

outliers

Outliers

Description

The outliers class uses box-plot definition for outliers. An outlier is a value that is below than $Q_1 - 1.5 \cdot IQR$ or higher than $Q_3 + 1.5 \cdot IQR$. The class remove outliers for numeric attributes. Users can set alpha to 3 to remove extreme values.

Usage

```
outliers(alpha = 1.5)
```

Arguments

alpha	boxplot outlier threshold (default 1.5, but can be 3.0 to remove extreme values)
-------	----------------------------------------------------------------------------------

Value

An outlier object

Examples

```
# code for outlier removal
out_obj <- outliers() # class for outlier analysis
out_obj <- fit(out_obj, iris) # computing boundaries
iris.clean <- transform(out_obj, iris) # returning cleaned dataset

#inspection of cleaned dataset
nrow(iris.clean)

idx <- attr(iris.clean, "idx")
table(idx)
iris.outliers <- iris[idx,]
iris.outliers
```

plot_bar

plot bar graph

Description

plot bar graph

Usage

```
plot_bar(data, label_x = "", label_y = "", colors = NULL, alpha = 1)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector
alpha	level of transparency

Value

ggplot graphic

Examples

```
#summarizing iris dataset
data <- iris |> dplyr::group_by(Species) |>
  dplyr::summarize(Sepal.Length=mean(Sepal.Length))
head(data)

#ploting data
grf <- plot_bar(data, colors="blue")
plot(grf)
```

plot_boxplot	<i>plot boxplot</i>
--------------	---------------------

Description

plot boxplot

Usage

```
plot_boxplot(data, label_x = "", label_y = "", colors = NULL, barwidth = 0.25)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector
barwidth	width of bar

Value

ggplot graphic

Examples

```
grf <- plot_boxplot(iris, colors="white")  
plot(grf)
```

plot_boxplot_class	<i>plot boxplot per class</i>
--------------------	-------------------------------

Description

plot boxplot per class

Usage

```
plot_boxplot_class(  
  data,  
  class_label,  
  label_x = "",  
  label_y = "",  
  colors = NULL  
)
```

Arguments

data	data.frame contain x, value, and variable
class_label	name of attribute for class label
label_x	x-axis label
label_y	y-axis label
colors	color vector

Value

ggplot graphic

Examples

```
grf <- plot_boxplot_class(iris |> dplyr::select(Sepal.Width, Species),
  class = "Species", colors=c("red", "green", "blue"))
plot(grf)
```

plot_density

plot density

Description

plot density

Usage

```
plot_density(
  data,
  label_x = "",
  label_y = "",
  colors = NULL,
  bin = NULL,
  alpha = 0.25
)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector
bin	bin width
alpha	level of transparency

Value

ggplot graphic

Examples

```
grf <- plot_density(iris |> dplyr::select(Sepal.Width), colors="blue")
plot(grf)
```

plot_density_class *plot density per class*

Description

plot density per class

Usage

```
plot_density_class(
  data,
  class_label,
  label_x = "",
  label_y = "",
  colors = NULL,
  bin = NULL,
  alpha = 0.5
)
```

Arguments

data	data.frame contain x, value, and variable
class_label	name of attribute for class label
label_x	x-axis label
label_y	y-axis label
colors	color vector
bin	bin width
alpha	level of transparency

Value

ggplot graphic

Examples

```
grf <- plot_density_class(iris |> dplyr::select(Sepal.Width, Species),
  class = "Species", colors=c("red", "green", "blue"))
plot(grf)
```

plot_groupedbar	<i>plot grouped bar</i>
-----------------	-------------------------

Description

plot grouped bar

Usage

```
plot_groupedbar(data, label_x = "", label_y = "", colors = NULL, alpha = 1)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector
alpha	level of transparency

Value

ggplot graphic

Examples

```
data <- iris |> dplyr::group_by(Species) |>
  dplyr::summarize(Sepal.Length=mean(Sepal.Length), Sepal.Width=mean(Sepal.Width))
grf <- plot_groupedbar(data, colors=c("blue", "red"))
plot(grf)
```

plot_hist	<i>plot histogram</i>
-----------	-----------------------

Description

plot histogram

Usage

```
plot_hist(data, label_x = "", label_y = "", color = "white", alpha = 0.25)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
color	color vector
alpha	transparency level

Value

ggplot graphic

Examples

```
grf <- plot_hist(iris |> dplyr::select(Sepal.Width), color=c("blue"))
plot(grf)
```

plot_lollipop	<i>plot lollipop</i>
---------------	----------------------

Description

plot lollipop

Usage

```
plot_lollipop(  
  data,  
  label_x = "",  
  label_y = "",  
  colors = NULL,  
  color_text = "black",  
  size_text = 3,  
  size_ball = 8,  
  alpha_ball = 0.2,  
  min_value = 0,  
  max_value_gap = 1  
)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector
color_text	color of text inside ball

size_text	size of text inside ball
size_ball	size of ball
alpha_ball	transparency of ball
min_value	minimum value
max_value_gap	maximum value gap

Value

ggplot graphic

Examples

```
#summarizing iris dataset
data <- iris |> dplyr::group_by(Species) |>
  dplyr::summarize(Sepal.Length=mean(Sepal.Length))
head(data)

#plotting data
grf <- plot_lollipop(data, colors="blue", max_value_gap=0.2)
plot(grf)
```

plot_pieplot	<i>plot pie</i>
--------------	-----------------

Description

plot pie

Usage

```
plot_pieplot(
  data,
  label_x = "",
  label_y = "",
  colors = NULL,
  textcolor = "white",
  bordercolor = "black"
)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector
textcolor	text color
bordercolor	border color

Value

ggplot graphic

Examples

```
#summarizing iris dataset
data <- iris |> dplyr::group_by(Species) |>
  dplyr::summarize(Sepal.Length=mean(Sepal.Length))
head(data)

#plotting data
grf <- plot_pieplot(data, colors=c("red", "green", "blue"))
plot(grf)
```

plot_points

plot points

Description

plot points

Usage

```
plot_points(data, label_x = "", label_y = "", colors = NULL)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector

Value

ggplot graphic

Examples

```
x <- seq(0, 10, 0.25)
data <- data.frame(x, sin=sin(x), cosine=cos(x)+5)
head(data)

grf <- plot_points(data, colors=c("red", "green"))
plot(grf)
```

plot_radar	<i>plot radar</i>
------------	-------------------

Description

plot radar

Usage

```
plot_radar(data, label_x = "", label_y = "", colors = NULL)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector

Value

ggplot graphic

Examples

```
data <- data.frame(name = "Petal.Length", value = mean(iris$Petal.Length))
data <- rbind(data, data.frame(name = "Petal.Width", value = mean(iris$Petal.Width)))
data <- rbind(data, data.frame(name = "Sepal.Length", value = mean(iris$Sepal.Length)))
data <- rbind(data, data.frame(name = "Sepal.Width", value = mean(iris$Sepal.Width)))

grf <- plot_radar(data, colors="red") + ggplot2::ylim(0, NA)
plot(grf)
```

plot_scatter	<i>scatter graph</i>
--------------	----------------------

Description

scatter graph

Usage

```
plot_scatter(data, label_x = "", label_y = "", colors = NULL)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector

Value

ggplot graphic

Examples

```
grf <- plot_scatter(iris |> dplyr::select(x = Sepal.Length,
  value = Sepal.Width, variable = Species),
  label_x = "Sepal.Length", label_y = "Sepal.Width",
  colors=c("red", "green", "blue"))
plot(grf)
```

plot_series

plot series

Description

plot series

Usage

```
plot_series(data, label_x = "", label_y = "", colors = NULL)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector

Value

plot

Examples

```
x <- seq(0, 10, 0.25)
data <- data.frame(x, sin=sin(x))
head(data)

grf <- plot_series(data, colors=c("red"))
plot(grf)
```

plot_stackedbar *plot stacked bar*

Description

plot stacked bar

Usage

```
plot_stackedbar(data, label_x = "", label_y = "", colors = NULL, alpha = 1)
```

Arguments

data	data.frame contain x, value, and variable
label_x	x-axis label
label_y	y-axis label
colors	color vector
alpha	level of transparency

Value

ggplot graphic

Examples

```
data <- iris |> dplyr::group_by(Species) |>
  dplyr::summarize(Sepal.Length=mean(Sepal.Length), Sepal.Width=mean(Sepal.Width))
grf <- plot_stackedbar(data, colors=c("blue", "red"))
plot(grf)
```

plot_ts *Plot a time series chart*

Description

The function receives six variables as a parameter, which are obj and y, yadj, main and xlabel. The graph is plotted with 3 lines: the original series (in black), the adjusted series (in blue) and the predicted series (in green)

Usage

```
plot_ts(x = NULL, y, label_x = "", label_y = "", color = "black")
```

Arguments

x	input variable
y	output variable
label_x	x-axis label
label_y	y-axis label
color	color for time series

Value

ggplot graphic

Examples

```
x <- seq(0, 10, 0.25)
data <- data.frame(x, sin=sin(x))
head(data)

grf <- plot_ts(x = data$x, y = data$sin, color=c("red"))
plot(grf)
```

plot_ts_pred *Plot a time series chart*

Description

The function receives six variables as a parameter, which are obj and y, yadj, main and xlabel. The graph is plotted with 3 lines: the original series (in black), the adjusted series (in blue) and the predicted series (in green)

Usage

```
plot_ts_pred(
  x = NULL,
  y,
  yadj,
  ypred = NULL,
  label_x = "",
  label_y = "",
  color = "black",
  color_adjust = "blue",
  color_prediction = "green"
)
```

Arguments

x	time index
y	time series
yadj	adjustment of time series
ypred	prediction of the time series
label_x	x-axis title
label_y	y-axis title
color	color for the time series
color_adjust	color for the adjusted values
color_prediction	color for the predictions

Value

ggplot graphic

Examples

```
data(sin_data)
ts <- ts_data(sin_data$y, 0)
ts_head(ts, 3)

samp <- ts_sample(ts, test_size= 5)
io_train <- ts_projection(samp$train)
io_test <- ts_projection(samp$test)

model <- ts_arima()
model <- fit(model, x=io_train$input, y=io_train$output)
adjust <- predict(model, io_train$input)

prediction <- predict(model, x=io_test$input, steps_ahead=5)
prediction <- as.vector(prediction)

yvalues <- c(io_train$output, io_test$output)
grf <- plot_ts_pred(y=yvalues, yadj=adjust, ypre=prediction)
plot(grf)
```

predictor

DAL Predict

Description

Ancestor class for regression and classification. It provides basis for fit and predict methods. Besides, action method proxies to predict.

An example of learner is a decision tree (cla_dtree)

Usage

```
predictor()
```

Value

a predictor object

Examples

```
#See ?cla_dtree for a classification example using a decision tree
```

R2.ts

R2

Description

Compute the R-squared (R2) between actual values and forecasts of a time series

Usage

```
R2.ts(actual, prediction)
```

Arguments

actual	real observations
prediction	predicted observations

Value

A number, which is the calculated R2

regression

Regression

Description

Ancestor class for regression problems

Usage

```
regression(attribute)
```

Arguments

attribute	attribute target to model building
-----------	------------------------------------

Value

regression object

Examples

```
#See ?reg_dtree for a regression example using a decision tree
```

reg_dtree

Decision Tree for regression

Description

Creates a regression object that uses the Decision Tree method for regression It wraps the tree library.

Usage

```
reg_dtree(attribute)
```

Arguments

attribute attribute target to model building.

Value

A decision tree regression object

Examples

```
data(Boston)
model <- reg_dtree("medv")

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, Boston)
train <- sr$train
test <- sr$test

model <- fit(model, train)

test_prediction <- predict(model, test)
test_predictand <- test[, "medv"]
test_eval <- evaluate(model, test_predictand, test_prediction)
test_eval$metrics
```

reg_knn	<i>knn regression</i>
---------	-----------------------

Description

Creates a regression object that uses the K-Nearest Neighbors (knn) method for regression

Usage

```
reg_knn(attribute, k)
```

Arguments

attribute	attribute target to model building
k	number of k neighbors

Value

A knn regression object

Examples

```
data(Boston)
model <- reg_knn("medv", k=3)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, Boston)
train <- sr$train
test <- sr$test

model <- fit(model, train)

test_prediction <- predict(model, test)
test_predictand <- test[, "medv"]
test_eval <- evaluate(model, test_predictand, test_prediction)
test_eval$metrics
```

reg_mlp	<i>MLP for regression</i>
---------	---------------------------

Description

Creates a regression object that uses the Multi-Layer Perceptron (MLP) method. It wraps the nnet library.

Usage

```
reg_mlp(attribute, size = NULL, decay = 0.05, maxit = 1000)
```

Arguments

attribute	attribute target to model building
size	number of neurons in hidden layers
decay	decay learning rate
maxit	number of maximum iterations for training

Value

obj

Examples

```
data(Boston)
model <- reg_mlp("medv", size=5, decay=0.54)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, Boston)
train <- sr$train
test <- sr$test

model <- fit(model, train)

test_prediction <- predict(model, test)
test_predictand <- test[, "medv"]
test_eval <- evaluate(model, test_predictand, test_prediction)
test_eval$metrics
```

reg_rf

Random Forest for regression

Description

Creates a regression object that uses the Random Forest method. It wraps the randomForest library.

Usage

```
reg_rf(attribute, nodesize = 1, ntree = 10, mtry = NULL)
```

Arguments

attribute	attribute target to model building
nodesize	node size
ntree	number of trees
mtry	number of attributes to build tree

Value

obj

Examples

```
data(Boston)
model <- reg_rf("medv", ntree=10)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, Boston)
train <- sr$train
test <- sr$test

model <- fit(model, train)

test_prediction <- predict(model, test)
test_predictand <- test[, "medv"]
test_eval <- evaluate(model, test_predictand, test_prediction)
test_eval$metrics
```

reg_svm

SVM for regression

Description

Creates a regression object that uses the Support Vector Machine (SVM) method for regression. It wraps the e1071 library.

Usage

```
reg_svm(attribute, epsilon = 0.1, cost = 10, kernel = "radial")
```

Arguments

attribute	attribute target to model building
epsilon	parameter that controls the width of the margin around the separating hyperplane
cost	parameter that controls the trade-off between having a wide margin and correctly classifying training data points
kernel	the type of kernel function to be used in the SVM algorithm (linear, radial, polynomial, sigmoid)

Value

A SVM regression object

Examples

```
data(Boston)
model <- reg_svm("medv", epsilon=0.2, cost=40.000)

# preparing dataset for random sampling
sr <- sample_random()
sr <- train_test(sr, Boston)
train <- sr$train
test <- sr$test

model <- fit(model, train)

test_prediction <- predict(model, test)
test_predictand <- test[, "medv"]
test_eval <- evaluate(model, test_predictand, test_prediction)
test_eval$metrics
```

reg_tune

Regression Tune

Description

Regression Tune

Usage

```
reg_tune(base_model, folds = 10)
```

Arguments

base_model	base model for tuning
folds	number of folds for cross-validation

Value

a reg_tune object.

Examples

```
# preparing dataset for random sampling
data(Boston)
sr <- sample_random()
sr <- train_test(sr, Boston)
train <- sr$train
test <- sr$test

# hyper parameter setup
tune <- reg_tune(reg_mlp("medv"))
ranges <- list(size=c(3), decay=c(0.1, 0.5))
```

```
# hyper parameter optimization
model <- fit(tune, train, ranges)

test_prediction <- predict(model, test)
test_predictand <- test[, "medv"]
test_eval <- evaluate(model, test_predictand, test_prediction)
test_eval$metrics
```

sample_random

Sample Random

Description

The `sample_random` function in R is used to generate a random sample of specified size from a given data set.

Usage

```
sample_random()
```

Value

obj

Examples

```
#using random sampling
sample <- sample_random()
tt <- train_test(sample, iris)

# distribution of train
table(tt$train$Species)

# preparing dataset into four folds
folds <- k_fold(sample, iris, 4)

# distribution of folds
tbl <- NULL
for (f in folds) {
  tbl <- rbind(tbl, table(f$Species))
}
head(tbl)
```

sample_stratified	<i>sample_stratified</i>
-------------------	--------------------------

Description

The `sample_stratified` function in R is used to generate a stratified random sample from a given dataset. Stratified sampling is a statistical method that is used when the population is divided into non-overlapping subgroups or strata, and a sample is selected from each stratum to represent the entire population. In stratified sampling, the sample is selected in such a way that it is representative of the entire population and the variability within each stratum is minimized.

Usage

```
sample_stratified(attribute)
```

Arguments

`attribute` attribute target to model building

Value

`obj`

Examples

```
#using stratified sampling
sample <- sample_stratified("Species")
tt <- train_test(sample, iris)

# distribution of train
table(tt$train$Species)

# preparing dataset into four folds
folds <- k_fold(sample, iris, 4)

# distribution of folds
tbl <- NULL
for (f in folds) {
  tbl <- rbind(tbl, table(f$Species))
}
head(tbl)
```

select_hyper	<i>Selection hyper parameters</i>
--------------	-----------------------------------

Description

Selection hyper parameters from a k-fold cross-validation execution

Usage

```
select_hyper(obj, hyperparameters)
```

Arguments

obj	object
hyperparameters	data set with hyper parameters and quality measure from execution

Value

index of selected hyper parameter

select_hyper.cla_tune	<i>selection of hyperparameters</i>
-----------------------	-------------------------------------

Description

selection of hyperparameters (maximizing classification metric)

Usage

```
## S3 method for class 'cla_tune'  
select_hyper(obj, hyperparameters)
```

Arguments

obj	object
hyperparameters	hyperparameters dataset

Value

optimized key number of hyperparameters

```
select_hyper.ts_tune  selection of hyperparameters (time series)
```

Description

selection of hyperparameters (minimizing error)

Usage

```
## S3 method for class 'ts_tune'
select_hyper(obj, hyperparameters)
```

Arguments

```
obj          object
hyperparameters  hyperparameters dataset
```

Value

optimized key number of hyperparameters

```
set_params          Assign parameters
```

Description

set_params function assigns all parameters to the attributes presented in the object. It returns the object with the parameters set.

Usage

```
set_params(obj, params)
```

Arguments

```
obj          object of class dal_base
params      parameters to set obj
```

Value

obj with parameters set

Examples

```
obj <- set_params(dal_base(), list(x = 0))
```

set_params.default	<i>Assign parameters</i>
--------------------	--------------------------

Description

This function receives the obj and params variables as parameters. It returns the obj as it is.

Usage

```
## Default S3 method:  
set_params(obj, params)
```

Arguments

obj	object
params	parameters

Value

obj

sin_data	<i>Time series example dataset</i>
----------	------------------------------------

Description

Synthetic dataset of sine function.

- x: correspond time from 0 to 10.
- y: dependent variable for time series modeling.

Usage

```
data(sin_data)
```

Format

data.frame.

Source

This dataset was generated for examples.

Examples

```
data(sin_data)  
head(sin_data)
```

`sMAPE.ts`*sMAPE*

Description

Compute the symmetric mean absolute percent error (sMAPE)

Usage

```
sMAPE.ts(actual, prediction)
```

Arguments

<code>actual</code>	real observations
<code>prediction</code>	predicted observations

Value

The sMAPE between the actual and prediction vectors

`smoothing`*Smoothing*

Description

Smoothing is a statistical technique used to reduce the noise in a signal or a dataset by removing the high-frequency components. The smoothing level is associated with the number of bins used. There are alternative methods to establish the smoothing: equal interval, equal frequency, and clustering.

Usage

```
smoothing(n)
```

Arguments

<code>n</code>	number of bins
----------------	----------------

Value

`obj`

Examples

```
data(iris)
obj <- smoothing_inter(n = 2)
obj <- fit(obj, iris$Sepal.Length)
sl.bi <- transform(obj, iris$Sepal.Length)
table(sl.bi)
obj$interval

entro <- evaluate(obj, as.factor(names(sl.bi)), iris$Species)
entro$entropy
```

smoothing_cluster	<i>Smoothing by cluster</i>
-------------------	-----------------------------

Description

Uses clustering method to perform data smoothing. The input vector is divided into clusters using the k-means algorithm. The mean of each cluster is then calculated and used as the smoothed value for all observations within that cluster.

Usage

```
smoothing_cluster(n)
```

Arguments

n	number of bins
---	----------------

Value

```
obj
```

Examples

```
data(iris)
obj <- smoothing_cluster(n = 2)
obj <- fit(obj, iris$Sepal.Length)
sl.bi <- transform(obj, iris$Sepal.Length)
table(sl.bi)
obj$interval

entro <- evaluate(obj, as.factor(names(sl.bi)), iris$Species)
entro$entropy
```

smoothing_freq *Smoothing by Freq*

Description

The 'smoothing_freq' function is used to smooth a given time series data by aggregating observations within a fixed frequency.

Usage

```
smoothing_freq(n)
```

Arguments

n number of bins

Value

obj

Examples

```
data(iris)
obj <- smoothing_freq(n = 2)
obj <- fit(obj, iris$Sepal.Length)
sl.bi <- transform(obj, iris$Sepal.Length)
table(sl.bi)
obj$interval

entro <- evaluate(obj, as.factor(names(sl.bi)), iris$Species)
entro$entropy
```

smoothing_inter *Smoothing by interval*

Description

The "smoothing by interval" function is used to apply a smoothing technique to a vector or time series data using a moving window approach.

Usage

```
smoothing_inter(n)
```

Arguments

n number of bins

Value

obj

Examples

```

data(iris)
obj <- smoothing_inter(n = 2)
obj <- fit(obj, iris$Sepal.Length)
sl.bi <- transform(obj, iris$Sepal.Length)
table(sl.bi)
obj$interval

entro <- evaluate(obj, as.factor(names(sl.bi)), iris$Species)
entro$entropy

```

train_test	<i>training and test</i>
------------	--------------------------

Description

training and test partition of a dataset using a sampling method

Usage

```
train_test(obj, data, perc = 0.8, ...)
```

Arguments

obj	object
data	dataset
perc	percentage for training
...	optional arguments.

Value

train and test sets

Examples

```

#using random sampling
sample <- sample_random()
tt <- train_test(sample, iris)

# distribution of train
table(tt$train$Species)

```

`train_test_from_folds` *k-fold training and test partition object*

Description

k-fold training and test partition object

Usage

`train_test_from_folds(folds, k)`

Arguments

`folds` data partitioned into folds
`k` k-fold for test set, all reminder for training set

Value

train and test folds

`transform` *Transform*

Description

Defines a transformation method.

Usage

`transform(obj, ...)`

Arguments

`obj` a `dal_transform` object.
`...` optional arguments.

Value

transformed data.

Examples

`#See ?minmax for an example of transformation`

ts_arma	<i>ARIMA</i>
---------	--------------

Description

Creates a time series prediction object that uses the AutoRegressive Integrated Moving Average (ARIMA). It wraps the forecast library.

Usage

```
ts_arma()
```

Value

a ts_arma object.

Examples

```
data(sin_data)
ts <- ts_data(sin_data$y, 0)
ts_head(ts, 3)

samp <- ts_sample(ts, test_size = 5)
io_train <- ts_projection(samp$train)
io_test <- ts_projection(samp$test)

model <- ts_arma()
model <- fit(model, x=io_train$input, y=io_train$output)

prediction <- predict(model, x=io_test$input[1,], steps_ahead=5)
prediction <- as.vector(prediction)
output <- as.vector(io_test$output)

ev_test <- evaluate(model, output, prediction)
ev_test
```

ts_conv1d	<i>Conv1D</i>
-----------	---------------

Description

Creates a time series prediction object that uses the Conv1D. It wraps the pytorch library.

Usage

```
ts_conv1d(preprocess = NA, input_size = NA, epochs = 10000L)
```

Arguments

preprocess	normalization
input_size	input size for machine learning model
epochs	maximum number of epochs

Value

a ts_conv1d object.

Examples

```
#Use the same example of ts_mlp changing the constructor to:
model <- ts_conv1d(ts_norm_gminmax(), input_size=4, epochs = 10000L)
```

ts_data	<i>ts_data</i>
---------	----------------

Description

Time series data structure used in DAL Toolbox. It receives a vector (representing a time series) or a matrix *y* (representing a sliding windows). Internal *ts_data* is matrix of sliding windows with size *sw*. If *sw* equals to zero, it store a time series as a single matrix column.

Usage

```
ts_data(y, sw = 1)
```

Arguments

<i>y</i>	output variable
<i>sw</i>	integer: sliding window size.

Value

a ts_data object.

Examples

```
data(sin_data)
head(sin_data)

data <- ts_data(sin_data$y)
ts_head(data)

data10 <- ts_data(sin_data$y, 10)
ts_head(data10)
```

`ts_elm`*ELM*

Description

Creates a time series prediction object that uses the Extreme Learning Machine (ELM). It wraps the `elmNNRcpp` library.

Usage

```
ts_elm(preprocess = NA, input_size = NA, nhid = NA, actfun = "purelin")
```

Arguments

<code>preprocess</code>	normalization
<code>input_size</code>	input size for machine learning model
<code>nhid</code>	ensemble size
<code>actfun</code>	defines the type to use, possible values: 'sig', 'radbas', 'tribas', 'relu', 'purelin' (default).

Value

a `ts_elm` object.

Examples

```
data(sin_data)
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)

samp <- ts_sample(ts, test_size = 5)
io_train <- ts_projection(samp$train)
io_test <- ts_projection(samp$test)

model <- ts_elm(ts_norm_gminmax(), input_size=4, nhid=3, actfun="purelin")
model <- fit(model, x=io_train$input, y=io_train$output)

prediction <- predict(model, x=io_test$input[1,], steps_ahead=5)
prediction <- as.vector(prediction)
output <- as.vector(io_test$output)

ev_test <- evaluate(model, output, prediction)
ev_test
```

ts_head	<i>ts_head</i>
---------	----------------

Description

Returns the first n observations from a ts_data

Usage

```
ts_head(x, n = 6L, ...)
```

Arguments

x	ts_data
n	number of rows to return
...	optional arguments

Value

The first n observations of a ts_data

Examples

```
data(sin_data)
data10 <- ts_data(sin_data$y, 10)
ts_head(data10)
```

ts_knn	<i>knn time series prediction</i>
--------	-----------------------------------

Description

Creates a prediction object that uses the K-Nearest Neighbors (knn) method for time series regression

Usage

```
ts_knn(preprocess = NA, input_size = NA, k = NA)
```

Arguments

preprocess	normalization
input_size	input size for machine learning model
k	number of k neighbors

Value

a ts_knn object.

Examples

```
data(sin_data)
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)

samp <- ts_sample(ts, test_size = 5)
io_train <- ts_projection(samp$train)
io_test <- ts_projection(samp$test)

model <- ts_knn(ts_norm_gminmax(), input_size=4, k=3)
model <- fit(model, x=io_train$input, y=io_train$output)

prediction <- predict(model, x=io_test$input[1,], steps_ahead=5)
prediction <- as.vector(prediction)
output <- as.vector(io_test$output)

ev_test <- evaluate(model, output, prediction)
ev_test
```

ts_lstm

LSTM

Description

Creates a time series prediction object that uses the LSTM. It wraps the pytorch library.

Usage

```
ts_lstm(preprocess = NA, input_size = NA, epochs = 10000L)
```

Arguments

preprocess	normalization
input_size	input size for machine learning model
epochs	maximum number of epochs

Value

a ts_lstm object.

Examples

```
#Use the same example of ts_mlp changing the constructor to:
model <- ts_lstm(ts_norm_gminmax(), input_size=4, epochs = 10000L)
```

`ts_mlp`*MLP*

Description

Creates a time series prediction object that uses the Multilayer Perceptron (MLP). It wraps the nnet library.

Usage

```
ts_mlp(preprocess = NA, input_size = NA, size = NA, decay = 0.01, maxit = 1000)
```

Arguments

<code>preprocess</code>	normalization
<code>input_size</code>	input size for machine learning model
<code>size</code>	number of neurons inside hidden layer
<code>decay</code>	decay parameter for MLP
<code>maxit</code>	maximum number of iterations

Value

a `ts_mlp` object.

Examples

```
data(sin_data)
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)

samp <- ts_sample(ts, test_size = 5)
io_train <- ts_projection(samp$train)
io_test <- ts_projection(samp$test)

model <- ts_mlp(ts_norm_gminmax(), input_size=4, size=4, decay=0)
model <- fit(model, x=io_train$input, y=io_train$output)

prediction <- predict(model, x=io_test$input[1,], steps_ahead=5)
prediction <- as.vector(prediction)
output <- as.vector(io_test$output)

ev_test <- evaluate(model, output, prediction)
ev_test
```

ts_norm_an	<i>Time Series Adaptive Normalization</i>
------------	-------------------------------------------

Description

Transform data to a common scale while taking into account the changes in the statistical properties of the data over time.

Usage

```
ts_norm_an(remove_outliers = TRUE, nw = 0)
```

Arguments

remove_outliers	logical: if TRUE (default) outliers will be removed.
nw	integer: window size.

Value

a ts_norm_an object.

Examples

```
# time series to normalize
data(sin_data)

# convert to sliding windows
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)
summary(ts[,10])

# normalization
preproc <- ts_norm_an()
preproc <- fit(preproc, ts)
tst <- transform(preproc, ts)
ts_head(tst, 3)
summary(tst[,10])
```

ts_norm_diff	<i>Time Series Diff</i>
--------------	-------------------------

Description

It receives as parameter the variable `remove_outliers`. This function calculates the difference between the values of a time series

Usage

```
ts_norm_diff(remove_outliers = TRUE)
```

Arguments

`remove_outliers`
logical: if TRUE (default) outliers will be removed.

Value

a `ts_norm_diff` object.

Examples

```
# time series to normalize
data(sin_data)

# convert to sliding windows
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)
summary(ts[,10])

# normalization
preproc <- ts_norm_diff()
preproc <- fit(preproc, ts)
tst <- transform(preproc, ts)
ts_head(tst, 3)
summary(tst[,9])
```

ts_norm_ean	<i>Time Series Adaptive Normalization (Exponential Moving Average - EMA)</i>
-------------	------------------------------------------------------------------------------

Description

It takes 2 parameters: `remove_outliers` and `nw`

Usage

```
ts_norm_ean(remove_outliers = TRUE, nw = 0)
```

Arguments

```
remove_outliers      logical: if TRUE (default) outliers will be removed.  
nw                   windows size
```

Value

a ts_norm_ean object.

Examples

```
# time series to normalize  
data(sin_data)  
  
# convert to sliding windows  
ts <- ts_data(sin_data$y, 10)  
ts_head(ts, 3)  
summary(ts[,10])  
  
# normalization  
preproc <- ts_norm_ean()  
preproc <- fit(preproc, ts)  
tst <- transform(preproc, ts)  
ts_head(tst, 3)  
summary(tst[,10])
```

ts_norm_gminmax	<i>Time Series Global Min-Max</i>
-----------------	-----------------------------------

Description

Rescales data, so the minimum value is mapped to 0 and the maximum value is mapped to 1.

Usage

```
ts_norm_gminmax(remove_outliers = TRUE)
```

Arguments

```
remove_outliers      logical: if TRUE (default) outliers will be removed.
```

Value

a ts_norm_gminmax object.

Examples

```
# time series to normalize
data(sin_data)

# convert to sliding windows
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)
summary(ts[,10])

# normalization
preproc <- ts_norm_gminmax()
preproc <- fit(preproc, ts)
tst <- transform(preproc, ts)
ts_head(tst, 3)
summary(tst[,10])
```

`ts_norm_swminmax`*Time Series Sliding Window Min-Max*

Description

It takes as parameter the variable `remove_outliers`. The `ts_norm_swminmax` function creates an object for normalizing a time series based on the "sliding window min-max scaling" method

Usage

```
ts_norm_swminmax(remove_outliers = TRUE)
```

Arguments

```
remove_outliers
logical: if TRUE (default) outliers will be removed.
```

Value

a `ts_norm_swminmax` object.

Examples

```
# time series to normalize
data(sin_data)

# convert to sliding windows
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)
summary(ts[,10])

# normalization
preproc <- ts_norm_swminmax()
```



```
preproc <- fit(preproc, ts)
tst <- transform(preproc, ts)
ts_head(tst, 3)
summary(tst[,10])
```

ts_projection	<i>Time Series Projection</i>
---------------	-------------------------------

Description

Separates the ts_data into input and output.

Usage

```
ts_projection(ts)
```

Arguments

ts matrix or data.frame containing the time series.

Value

a ts_projection object.

Examples

```
#setting up a ts_data
data(sin_data)
ts <- ts_data(sin_data$y, 10)

io <- ts_projection(ts)

#input data
ts_head(io$input)

#output data
ts_head(io$output)
```

ts_reg	<i>TSReg</i>
--------	--------------

Description

Time Series Regression directly from time series Ancestral class for non-sliding windows implementation.

Usage

```
ts_reg()
```

Value

A ts_reg object

Examples

```
#See ?ts_arima for an example using Auto-regressive Integrated Moving Average
```

ts_regsw	<i>TSRegSW</i>
----------	----------------

Description

Time Series Regression from Sliding Windows. Ancestral class for Machine Learning Implementation.

Usage

```
ts_regsw(preprocess = NA, input_size = NA)
```

Arguments

preprocess	normalization
input_size	input size for machine learning model

Value

A ts_regsw object

Examples

```
#See ?ts_elm for an example using Extreme Learning Machine
```

`ts_rf`*Random Forest*

Description

Creates a time series prediction object that uses the Random Forest. It wraps the randomForest library.

Usage

```
ts_rf(preprocess = NA, input_size = NA, nodesize = 1, ntree = 10, mtry = NULL)
```

Arguments

<code>preprocess</code>	normalization
<code>input_size</code>	input size for machine learning model
<code>nodesize</code>	node size
<code>ntree</code>	number of trees
<code>mtry</code>	number of attributes to build tree

Value

a `ts_rf` object.

Examples

```
data(sin_data)
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)

samp <- ts_sample(ts, test_size = 5)
io_train <- ts_projection(samp$train)
io_test <- ts_projection(samp$test)

model <- ts_rf(ts_norm_gminmax(), input_size=4, nodesize=3, ntree=50)
model <- fit(model, x=io_train$input, y=io_train$output)

prediction <- predict(model, x=io_test$input[1,], steps_ahead=5)
prediction <- as.vector(prediction)
output <- as.vector(io_test$output)

ev_test <- evaluate(model, output, prediction)
ev_test
```

ts_sample	<i>Time Series Sample</i>
-----------	---------------------------

Description

Separates the `ts_data` into training and test. It separates the test size from the last observations minus an offset. The offset is important to allow replication under different recent origins. The data for train uses the number of rows of a `ts_data` minus the test size and offset.

Usage

```
ts_sample(ts, test_size = 1, offset = 0)
```

Arguments

<code>ts</code>	time series.
<code>test_size</code>	integer: size of test data (default = 1).
<code>offset</code>	integer: starting point (default = 0).

Value

A list with the two samples

Examples

```
#setting up a ts_data
data(sin_data)
ts <- ts_data(sin_data$y, 10)

#separating into train and test
test_size <- 3
samp <- ts_sample(ts, test_size)

#first five rows from training data
ts_head(samp$train, 5)

#last five rows from training data
ts_head(samp$train[-c(1:(nrow(samp$train)-5)),])

#testing data
ts_head(samp$test)
```

ts_svm	SVM
--------	-----

Description

Creates a time series prediction object that uses the Support Vector Machine (SVM). It wraps the e1071 library.

Usage

```
ts_svm(
  preprocess = NA,
  input_size = NA,
  kernel = "radial",
  epsilon = 0,
  cost = 10
)
```

Arguments

preprocess	normalization
input_size	input size for machine learning model
kernel	SVM kernel (linear, radial, polynomial, sigmoid)
epsilon	error threshold
cost	cost

Value

a ts_svm object.

Examples

```
data(sin_data)
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)

samp <- ts_sample(ts, test_size = 5)
io_train <- ts_projection(samp$train)
io_test <- ts_projection(samp$test)

model <- ts_svm(ts_norm_gminmax(), input_size=4)
model <- fit(model, x=io_train$input, y=io_train$output)

prediction <- predict(model, x=io_test$input[1,], steps_ahead=5)
prediction <- as.vector(prediction)
output <- as.vector(io_test$output)

ev_test <- evaluate(model, output, prediction)
ev_test
```

`ts_tune`*Time Series Tune*

Description

Time Series Tune

Usage

```
ts_tune(input_size, base_model, folds = 10)
```

Arguments

<code>input_size</code>	input size for machine learning model
<code>base_model</code>	base model for tuning
<code>folds</code>	number of folds for cross-validation

Valuea `ts_tune` object.**Examples**

```
data(sin_data)
ts <- ts_data(sin_data$y, 10)
ts_head(ts, 3)

samp <- ts_sample(ts, test_size = 5)
io_train <- ts_projection(samp$train)
io_test <- ts_projection(samp$test)

tune <- ts_tune(input_size=c(3:5), base_model = ts_elm(ts_norm_gminmax()))
ranges <- list(nhid = 1:5, actfun=c('purelin'))

# Generic model tuning
model <- fit(tune, x=io_train$input, y=io_train$output, ranges)

prediction <- predict(model, x=io_test$input[1,], steps_ahead=5)
prediction <- as.vector(prediction)
output <- as.vector(io_test$output)

ev_test <- evaluate(model, output, prediction)
ev_test
```

zscore	<i>z-score normalization</i>
--------	------------------------------

Description

Scale data using z-score normalization. $zscore = (x - \text{mean}(x))/sd(x)$.

Usage

```
zscore(nmean = 0, nsd = 1)
```

Arguments

nmean	new mean for normalized data
nsd	new standard deviation for normalized data

Value

z-score transformation object

Examples

```
data(iris)
head(iris)

trans <- zscore()
trans <- fit(trans, iris)
tiris <- transform(trans, iris)
head(tiris)

itiris <- inverse_transform(trans, tiris)
head(itiris)
```

[.ts_data	<i>Extract a subset of a time series stored in an object</i>
-----------	--------------------------------------------------------------

Description

Receives as parameters the variables x, i, j ...

Usage

```
## S3 method for class 'ts_data'
x[i, j, ...]
```

Arguments

x	input variable
i	row i
j	column j
...	optional arguments

Value

A new ts_data object

Examples

```
data(sin_data)
data10 <- ts_data(sin_data$y, 10)
ts_head(data10)
#single line
data10[12,]

#range of lines
data10[12:13,]

#single column
data10[,1]

#range of columns
data10[,1:2]

#range of rows and columns
data10[12:13,1:2]

#single line and a range of columns
#'data10[12,1:2]

#range of lines and a single column
data10[12:13,1]

#single observation
data10[12,1]
```


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