

# Package ‘SSLR’

April 13, 2020

**Type** Package

**Title** Semi-Supervised Classification and Regression Methods

**Version** 0.9.1

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**URL** <https://dicits.ugr.es/software/SSLR/>

**Description** Providing a collection of techniques for semi-supervised classification and regression. In semi-supervised problem, both labeled and unlabeled data are used to train a classifier. The package includes a collection of semi-supervised learning techniques: self-training, co-training, democratic, decision tree, random forest, 'S3VM' ... etc, with a fairly intuitive interface that is easy to use.

**License** GPL-3

**ByteCompile** true

**Depends** R (>= 2.10)

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 7.1.0

**Imports** stats, parsnip, plyr, dplyr (>= 0.8.0.1), magrittr, purrr, rlang (>= 0.3.1), proxy, methods, generics, utils, RANN, foreach, RSSL

**LinkingTo** Rcpp, RcppArmadillo

**Suggests** caret, tidymodels, e1071, C50, kernlab, testthat, doParallel, tidyverse, survival, xgboost, covr, kknn, randomForest, ranger, MASS, nlme, knitr, rmarkdown

**VignetteBuilder** knitr

**NeedsCompilation** yes

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Repository CRAN

Date/Publication 2020-04-13 15:00:02 UTC

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abalone	<i>Abalone</i>
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### Description

Abalone

### Usage

`data(abalone)`

### Format

Predict the age of abalone from physical measurements

### Source

<https://archive.ics.uci.edu/ml/datasets/Abalone>

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best_split	<i>An S4 method to best split</i>
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---

### Description

An S4 method to best split

### Usage

`best_split(object, ...)`

### Arguments

object	DecisionTree object
...	This parameter is included for compatibility reasons.

---

best\_split, DecisionTreeClassifier-method  
*Best Split function*

---

### Description

Function to get best split in Decision Tree. Find the best split for node. "Beast" means that the mean of impurity is the least possible. To find the best division. Let's iterate through all the features. All threshold / feature pairs will be computed in the numerical features. In the features that are not numerical, We get the best group of possible values will be obtained based on an algorithm with the function get\_levels\_categoric

### Usage

```
## S4 method for signature 'DecisionTreeClassifier'
best_split(object, X, y, parms)
```

### Arguments

object	DecisionTree object
X	is data
y	is class values
parms	parms in function

### Value

A list with: best\_idx name of the feature with the best split or Null if it not be found best\_thr: threshold found in the best split, or Null if it not be found

---

breast	<i>Breast</i>
--------	---------------

---

### Description

Breast

### Usage

```
data(breast)
```

### Format

: Diagnostic Wisconsin Breast Cancer Database

### Source

[https://archive.ics.uci.edu/ml/datasets/Breast+Cancer+Wisconsin+\(Diagnostic\)](https://archive.ics.uci.edu/ml/datasets/Breast+Cancer+Wisconsin+(Diagnostic))

---

calculate_gini	<i>Function calculate gini</i>
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---

**Description**

Function to calculate gini index. Formula is:  $1 - n:\text{num\_classes} \sum \text{probabilitie\_class}^2$

**Usage**

```
calculate_gini(column_factor)
```

**Arguments**

column\_factor    class values

---

check_value	<i>Check value in leaf</i>
-------------	----------------------------

---

**Description**

Function to check value in leaf from numeric until character

**Usage**

```
check_value(value, threshold)
```

**Arguments**

value            is the value in leaf node  
threshold        in leaf node

**Value**

TRUE if  $\leq$  in numeric or `%in%` in factor

---

check_xy_interface	<i>Check interface x y</i>
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---

**Description**

Check interface

**Usage**

```
check_xy_interface(x, y)
```

**Arguments**

x	data without class labels
y	values class

---

coBC	<i>General Interface for CoBC model</i>
------	---

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**Description**

Co-Training by Committee (CoBC) is a semi-supervised learning algorithm with a co-training style. This algorithm trains N classifiers with the learning scheme defined in the learner argument using a reduced set of labeled examples. For each iteration, an unlabeled example is labeled for a classifier if the most confident classifications assigned by the other N-1 classifiers agree on the labeling proposed. The unlabeled examples candidates are selected randomly from a pool of size u. The final prediction is the average of the estimates of the N regressors.

**Usage**

```
coBC(learner, N = 3, perc.full = 0.7, u = 100, max.iter = 50)
```

**Arguments**

learner	model from parsnip package for training a supervised base classifier using a set of instances. This model need to have probability predictions in classification mode
N	The number of classifiers used as committee members. All these classifiers are trained using the gen.learner function. Default is 3.
perc.full	A number between 0 and 1. If the percentage of new labeled examples reaches this value the self-labeling process is stopped. Default is 0.7.
u	Number of unlabeled instances in the pool. Default is 100.
max.iter	Maximum number of iterations to execute in the self-labeling process. Default is 50.

## Details

For regression tasks, labeling data is very expensive computationally. Its so slow. This method trains an ensemble of diverse classifiers. To promote the initial diversity the classifiers are trained from the reduced set of labeled examples by Bagging. The stopping criterion is defined through the fulfillment of one of the following criteria: the algorithm reaches the number of iterations defined in the `max.iter` parameter or the portion of unlabeled set, defined in the `perc.full` parameter, is moved to the enlarged labeled set of the classifiers.

## Value

(When model fit) A list object of class "coBC" containing:

**model** The final N base classifiers trained using the enlarged labeled set.

**model.index** List of N vectors of indexes related to the training instances used per each classifier. These indexes are relative to the `y` argument.

**instances.index** The indexes of all training instances used to train the N models. These indexes include the initial labeled instances and the newly labeled instances. These indexes are relative to the `y` argument.

**model.index.map** List of three vectors with the same information in `model.index` but the indexes are relative to `instances.index` vector.

**classes** The levels of `y` factor in classification.

**pred** The function provided in the `pred` argument.

**pred.pars** The list provided in the `pred.pars` argument.

## References

Avrim Blum and Tom Mitchell.

*Combining labeled and unlabeled data with co-training.*

In Eleventh Annual Conference on Computational Learning Theory, COLT' 98, pages 92-100, New York, NY, USA, 1998. ACM. ISBN 1-58113-057-0. doi: 10.1145/279943.279962.

Mohamed Farouk Abdel-Hady, Mohamed Farouk Abdel-Hady and Günther Palm.

*Semi-supervised Learning for Regression with Cotraining by Committee*

Institute of Neural Information Processing University of Ulm D-89069 Ulm, Germany

## Examples

```
library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

data(wine)

set.seed(1)
train.index <- createDataPartition(wine$Wine, p = .7, list = FALSE)
train <- wine[ train.index, ]
test <- wine[ -train.index, ]
```



```
cls <- which(colnames(wine) == "Wine")

#% LABELED
labeled.index <- createDataPartition(wine$Wine, p = .2, list = FALSE)
train[~labeled.index,cls] <- NA

#We need a model with probability predictions from parsnip
#https://tidymodels.github.io/parsnip/articles/articles/Models.html
#It should be with mode = classification

#For example, with Random Forest
rf <- rand_forest(trees = 100, mode = "classification") %>%
  set_engine("randomForest")

m <- coBC(learner = rf, N = 3,
          perc.full = 0.7,
          u = 100,
          max.iter = 3) %>% fit(Wine ~ ., data = train)

#Accuracy
predict(m, test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)
```

---

coBCCombine

*Combining the hypothesis*

---

## Description

This function combines the probabilities predicted by the committee of classifiers.

## Usage

```
coBCCombine(h.prob, classes)
```

## Arguments

h.prob	A list of probability matrices.
classes	The classes in the same order that appear in the columns of each matrix in h.prob.

## Value

A probability matrix

coBCG

*CoBC generic method***Description**

CoBC is a semi-supervised learning algorithm with a co-training style. This algorithm trains  $N$  classifiers with the learning scheme defined in `gen.learner` using a reduced set of labeled examples. For each iteration, an unlabeled example is labeled for a classifier if the most confident classifications assigned by the other  $N-1$  classifiers agree on the labeling proposed. The unlabeled examples candidates are selected randomly from a pool of size  $u$ .

**Usage**

```
coBCG(y, gen.learner, gen.pred, N = 3, perc.full = 0.7, u = 100, max.iter = 50)
```

**Arguments**

<code>y</code>	A vector with the labels of training instances. In this vector the unlabeled instances are specified with the value NA.
<code>gen.learner</code>	A function for training $N$ supervised base classifiers. This function needs two parameters, <code>indexes</code> and <code>cls</code> , where <code>indexes</code> indicates the instances to use and <code>cls</code> specifies the classes of those instances.
<code>gen.pred</code>	A function for predicting the probabilities per classes. This function must be two parameters, <code>model</code> and <code>indexes</code> , where the <code>model</code> is a classifier trained with <code>gen.learner</code> function and <code>indexes</code> indicates the instances to predict.
<code>N</code>	The number of classifiers used as committee members. All these classifiers are trained using the <code>gen.learner</code> function. Default is 3.
<code>perc.full</code>	A number between 0 and 1. If the percentage of new labeled examples reaches this value the self-labeling process is stopped. Default is 0.7.
<code>u</code>	Number of unlabeled instances in the pool. Default is 100.
<code>max.iter</code>	Maximum number of iterations to execute in the self-labeling process. Default is 50.

**Details**

coBCG can be helpful in those cases where the method selected as base classifier needs a `learner` and `pred` functions with other specifications. For more information about the general coBC method, please see `coBC` function. Essentially, `coBC` function is a wrapper of `coBCG` function.

**Value**

A list object of class "coBCG" containing:

**model** The final  $N$  base classifiers trained using the enlarged labeled set.

**model.index** List of  $N$  vectors of indexes related to the training instances used per each classifier. These indexes are relative to the `y` argument.

**instances.index** The indexes of all training instances used to train the N models. These indexes include the initial labeled instances and the newly labeled instances. These indexes are relative to the y argument.

**model.index.map** List of three vectors with the same information in `model.index` but the indexes are relative to `instances.index` vector.

**classes** The levels of y factor.

## Examples

```
library(SSLR)
library(caret)
## Load Wine data set
data(wine)

cls <- which(colnames(wine) == "Wine")
x <- wine[, - cls] # instances without classes
y <- wine[, cls] # the classes
x <- scale(x) # scale the attributes

## Prepare data
set.seed(20)
# Use 50% of instances for training
tra.idx <- sample(x = length(y), size = ceiling(length(y) * 0.5))
xtrain <- x[tra.idx,] # training instances
ytrain <- y[tra.idx] # classes of training instances
# Use 70% of train instances as unlabeled set
tra.na.idx <- sample(x = length(tra.idx), size = ceiling(length(tra.idx) * 0.7))
ytrain[tra.na.idx] <- NA # remove class information of unlabeled instances

# Use the other 50% of instances for inductive testing
tst.idx <- setdiff(1:length(y), tra.idx)
xitest <- x[tst.idx,] # testing instances
yitest <- y[tst.idx] # classes of testing instances

## Example: Training from a set of instances with 1-NN (knn3) as base classifier.
gen.learner1 <- function(indexes, cls)
  caret::knn3(x = xtrain[indexes,], y = cls, k = 1)
gen.pred1 <- function(model, indexes)
  predict(model, xtrain[indexes,])

set.seed(1)

trControl_coBCG <- list(gen.learner = gen.learner1, gen.pred = gen.pred1)
md1 <- train_generic(ytrain, method = "coBCG", trControl = trControl_coBCG)

# Predict probabilities per instances using each model
h.prob <- lapply(
  X = md1$model,
  FUN = function(m) predict(m, xitest)
)
# Combine the predictions
```

```

cls1 <- coBCCombine(h.prob, md1$classes)
table(cls1, yitest)

confusionMatrix(cls1, yitest)$overall[1]

## Example: Training from a distance matrix with 1-NN (oneNN) as base classifier.
dtrain <- as.matrix(proxy::dist(x = xtrain, method = "euclidean", by_rows = TRUE))
gen.learner2 <- function(indexes, cls) {
  m <- SSLR::oneNN(y = cls)
  attr(m, "tra.idx") <- indexes
  m
}

gen.pred2 <- function(model, indexes) {
  tra.idx <- attr(model, "tra.idx")
  d <- dtrain[indexes, tra.idx]
  prob <- predict(model, d, distance.weighting = "none")
  prob
}

set.seed(1)

trControl_coBCG2 <- list(gen.learner = gen.learner2, gen.pred = gen.pred2)
md2 <- train_generic(ytrain, method = "coBCG", trControl = trControl_coBCG2)

# Predict probabilities per instances using each model
ditest <- proxy::dist(x = xitest, y = xtrain[md2$instances.index,],
  method = "euclidean", by_rows = TRUE)

h.prob <- list()
ninstances <- nrow(dtrain)
for (i in 1:length(md2$model)) {
  m <- md2$model[[i]]
  D <- ditest[, md2$model.index.map[[i]]]
  h.prob[[i]] <- predict(m, D)
}
# Combine the predictions
cls2 <- coBCCombine(h.prob, md2$classes)
table(cls2, yitest)

confusionMatrix(cls2, yitest)$overall[1]

```

**Description**

coBCReg is based on an ensemble of  $N$  diverse regressors. At each iteration and for each regressor, the companion committee labels the unlabeled examples then the regressor select the most informative newly-labeled examples for itself, where the selection confidence is based on estimating the validation error. The final prediction is the average of the estimates of the  $N$  regressors.

**Usage**

```
coBCReg(learner, N = 3, perc.full = 0.7, u = 100, max.iter = 50)
```

**Arguments**

learner	model from parsnip package for training a supervised base classifier using a set of instances. This model need to have probability predictions
N	The number of classifiers used as committee members. All these classifiers are trained using the gen. learner function. Default is 3.
perc.full	A number between 0 and 1. If the percentage of new labeled examples reaches this value the self-labeling process is stopped. Default is 0.7.
u	Number of unlabeled instances in the pool. Default is 100.
max.iter	Maximum number of iterations to execute in the self-labeling process. Default is 50.

**Details**

For regression tasks, labeling data is very expensive computationally. Its so slow.

**References**

Mohamed Farouk Abdel-Hady, Mohamed Farouk Abdel-Hady and Günther Palm.  
*Semi-supervised Learning for Regression with Cotraining by Committee*  
 Institute of Neural Information Processing University of Ulm D-89069 Ulm, Germany

---

 coBCRegG

*Generic Interface coBCReg model*


---

**Description**

coBCReg is based on an ensemble of  $N$  diverse regressors. At each iteration and for each regressor, the companion committee labels the unlabeled examples then the regressor select the most informative newly-labeled examples for itself, where the selection confidence is based on estimating the validation error. The final prediction is the average of the estimates of the  $N$  regressors.

**Usage**

```
coBCRegG(
  y,
  gen.learner,
  gen.pred,
  N = 3,
  perc.full = 0.7,
  u = 100,
  max.iter = 50,
  gr = 1
)
```

**Arguments**

<code>y</code>	A vector with the labels of training instances. In this vector the unlabeled instances are specified with the value NA.
<code>gen.learner</code>	A function for training N supervised base classifiers. This function needs two parameters, indexes and cls, where indexes indicates the instances to use and cls specifies the classes of those instances.
<code>gen.pred</code>	A function for predicting the probabilities per classes. This function must be two parameters, model and indexes, where the model is a classifier trained with <code>gen.learner</code> function and indexes indicates the instances to predict.
<code>N</code>	The number of classifiers used as committee members. All these classifiers are trained using the <code>gen.learner</code> function. Default is 3.
<code>perc.full</code>	A number between 0 and 1. If the percentage of new labeled examples reaches this value the self-labeling process is stopped. Default is 0.7.
<code>u</code>	Number of unlabeled instances in the pool. Default is 100.
<code>max.iter</code>	Maximum number of iterations to execute in the self-labeling process. Default is 50.
<code>gr</code>	growing rate

**Details**

For regression tasks, labeling data is very expensive computationally. Its so slow.

**References**

Mohamed Farouk Abdel-Hady, Mohamed Farouk Abdel-Hady and Günther Palm.  
*Semi-supervised Learning for Regression with Cotraining by Committee*  
 Institute of Neural Information Processing University of Ulm D-89069 Ulm, Germany

---

coffee	<i>Time series data set</i>
--------	-----------------------------

---

**Description**

A dataset containing 56 times series z-normalized. Time series length is 286.

**Usage**

```
data(coffee)
```

**Format**

A data frame with 56 rows and 287 variables including the class.

**Source**

[https://www.cs.ucr.edu/~eamonn/time\\_series\\_data\\_2018/](https://www.cs.ucr.edu/~eamonn/time_series_data_2018/)

---

COREG	<i>General Interface for COREG model</i>
-------	--

---

**Description**

COREG is a semi-supervised learning for regression with a co-training style. This technique uses two kNN regressors with different distance metrics. For each iteration, each regressor labels the unlabeled example which can be most confidently labeled for the other learner, where the labeling confidence is estimated through considering the consistency of the regressor with the labeled example set. The final prediction is made by averaging the predictions of both the refined kNN regressors

**Usage**

```
COREG(max.iter = 50, k1 = 3, k2 = 5, p1 = 3, p2 = 5, u = 100)
```

**Arguments**

<code>max.iter</code>	maximum number of iterations to execute the self-labeling process. Default is 50.
<code>k1</code>	parameter in first KNN
<code>k2</code>	parameter in second KNN
<code>p1</code>	distance order 1. Default is 3
<code>p2</code>	distance order 1. Default is 5
<code>u</code>	Number of unlabeled instances in the pool. Default is 100.

**Details**

labeling data is very expensive computationally. Its so slow. For executing this model, we need RANN installed.

**References**

Zhi-Hua Zhou and Ming Li.  
*Semi-Supervised Regression with Co-Training.*  
 National Laboratory for Novel Software Technology Nanjing University, Nanjing 210093, China

**Examples**

```
library(SSLR)

m <- COREG(max.iter = 1)
```

---

DecisionTreeClassifier-class  
*Class DecisionTreeClassifier*

---

**Description**

Class DecisionTreeClassifier Slots: max\_depth, n\_classes\_, n\_features\_, tree\_, classes, min\_samples\_split, min\_samples\_leaf

---

democratic                      *General Interface for Democratic model*

---

**Description**

Democratic Co-Learning is a semi-supervised learning algorithm with a co-training style. This algorithm trains N classifiers with different learning schemes defined in list gen. learners. During the iterative process, the multiple classifiers with different inductive biases label data for each other.

**Usage**

```
democratic(learners, schemes = NULL)
```

**Arguments**

learners	List of models from parsnip package for training a supervised base classifier using a set of instances. This model need to have probability predictions
schemes	List of schemes (col x names in each learner). Default is null, it means that learner uses all x columns



## Details

This method trains an ensemble of diverse classifiers. To promote the initial diversity the classifiers must represent different learning schemes. When `x.inst` is `FALSE` all learners defined must be able to learn a classifier from the precomputed matrix in `x`. The iteration process of the algorithm ends when no changes occurs in any model during a complete iteration. The generation of the final hypothesis is produced via a weighed majority voting.

## Value

(When model fit) A list object of class "democratic" containing:

**W** A vector with the confidence-weighted vote assigned to each classifier.

**model** A list with the final `N` base classifiers trained using the enlarged labeled set.

**model.index** List of `N` vectors of indexes related to the training instances used per each classifier. These indexes are relative to the `y` argument.

**instances.index** The indexes of all training instances used to train the `N` models. These indexes include the initial labeled instances and the newly labeled instances. These indexes are relative to the `y` argument.

**model.index.map** List of three vectors with the same information in `model.index` but the indexes are relative to `instances.index` vector.

**classes** The levels of `y` factor.

**preds** The functions provided in the `preds` argument.

**preds.pars** The set of lists provided in the `preds.pars` argument.

**x.inst** The value provided in the `x.inst` argument.

## Examples

```
library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

data(wine)

set.seed(1)
train.index <- createDataPartition(wine$Wine, p = .7, list = FALSE)
train <- wine[ train.index,]
test <- wine[-train.index,]

cls <- which(colnames(wine) == "Wine")

#% LABELED
labeled.index <- createDataPartition(wine$Wine, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

#We need a model with probability predictions from parsnip
#https://tidymodels.github.io/parsnip/articles/articles/Models.html
#It should be with mode = classification
```

```

rf <- rand_forest(trees = 100, mode = "classification") %>%
  set_engine("randomForest")

bt <- boost_tree(trees = 100, mode = "classification") %>%
  set_engine("xgboost")

m <- democratic(learners = list(rf,bt)) %>% fit(Wine ~ ., data = train)

#' \donttest{
#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)

#With schemes
set.seed(1)
m <- democratic(learners = list(rf,bt),
               schemes = list(c("Malic.Acid","Ash"), c("Magnesium","Proline"))) %>%
  fit(Wine ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)

#'}

```

---

democraticCombine      *Combining the hypothesis of the classifiers*

---

## Description

This function combines the probabilities predicted by the set of classifiers.

## Usage

```
democraticCombine(pred, W, classes)
```

## Arguments

pred	A list with the prediction for each classifier.
W	A vector with the confidence-weighted vote assigned to each classifier during the training process.
classes	the classes.

**Value**

The classification proposed.

---

 democraticG

*Democratic generic method*


---

**Description**

Democratic is a semi-supervised learning algorithm with a co-training style. This algorithm trains N classifiers with different learning schemes defined in list `gen.learners`. During the iterative process, the multiple classifiers with different inductive biases label data for each other.

**Usage**

```
democraticG(y, gen.learners, gen.preds)
```

**Arguments**

<code>y</code>	A vector with the labels of training instances. In this vector the unlabeled instances are specified with the value NA.
<code>gen.learners</code>	A list of functions for training N different supervised base classifiers. Each function needs two parameters, <code>indexes</code> and <code>cls</code> , where <code>indexes</code> indicates the instances to use and <code>cls</code> specifies the classes of those instances.
<code>gen.preds</code>	A list of functions for predicting the probabilities per classes. Each function must be two parameters, <code>model</code> and <code>indexes</code> , where the <code>model</code> is a classifier trained with <code>gen.learner</code> function and <code>indexes</code> indicates the instances to predict.

**Details**

`democraticG` can be helpful in those cases where the method selected as base classifier needs a learner and pred functions with other specifications. For more information about the general democratic method, please see [democratic](#) function. Essentially, `democratic` function is a wrapper of `democraticG` function.

**Value**

A list object of class "democraticG" containing:

**W** A vector with the confidence-weighted vote assigned to each classifier.

**model** A list with the final N base classifiers trained using the enlarged labeled set.

**model.index** List of N vectors of indexes related to the training instances used per each classifier. These indexes are relative to the `y` argument.

**instances.index** The indexes of all training instances used to train the N models. These indexes include the initial labeled instances and the newly labeled instances. These indexes are relative to the `y` argument.

**model.index.map** List of three vectors with the same information in `model.index` but the indexes are relative to `instances.index` vector.

**classes** The levels of `y` factor.

## References

Yan Zhou and Sally Goldman.

*Democratic co-learning.*

In IEEE 16th International Conference on Tools with Artificial Intelligence (ICTAI), pages 594-602. IEEE, Nov 2004. doi: 10.1109/ICTAI.2004.48.

---

EMLeastSquaresClassifierSSLR

*General Interface for EMLeastSquaresClassifier model*

---

## Description

model from RSSL package

An Expectation Maximization like approach to Semi-Supervised Least Squares Classification

As studied in Krijthe & Loog (2016), minimizes the total loss of the labeled and unlabeled objects by finding the weight vector and labels that minimize the total loss. The algorithm proceeds similar to EM, by subsequently applying a weight update and a soft labeling of the unlabeled objects. This is repeated until convergence.

By default (`method="block"`) the weights of the classifier are updated, after which the unknown labels are updated. `method="simple"` uses LBFGS to do this update simultaneously. `Objective="responsibility"` corresponds to the responsibility based, instead of the label based, objective function in Krijthe & Loog (2016), which is equivalent to hard-label self-learning.

## Usage

```
EMLeastSquaresClassifierSSLR(
  x_center = FALSE,
  scale = FALSE,
  verbose = FALSE,
  intercept = TRUE,
  lambda = 0,
  eps = 1e-09,
  y_scale = FALSE,
  alpha = 1,
  beta = 1,
  init = "supervised",
  method = "block",
  objective = "label",
  save_all = FALSE,
  max_iter = 1000
)
```

**Arguments**

x_center	logical; Should the features be centered?
scale	Should the features be normalized? (default: FALSE)
verbose	logical; Controls the verbosity of the output
intercept	logical; Whether an intercept should be included
lambda	numeric; L2 regularization parameter
eps	Stopping criterion for the minimization
y_scale	logical; whether the target vector should be centered
alpha	numeric; the mixture of the new responsibilities and the old in each iteration of the algorithm (default: 1)
beta	numeric; value between 0 and 1 that determines how much to move to the new solution from the old solution at each step of the block gradient descent
init	objective character; "random" for random initialization of labels, "supervised" to use supervised solution as initialization or a numeric vector with a coefficient vector to use to calculate the initialization
method	character; one of "block", for block gradient descent or "simple" for LBFGS optimization (default="block")
objective	character; "responsibility" for hard label self-learning or "label" for soft-label self-learning
save_all	logical; saves all classifiers trained during block gradient descent
max_iter	integer; maximum number of iterations

**References**

Krijthe, J.H. & Loog, M., 2016. Optimistic Semi-supervised Least Squares Classification. In International Conference on Pattern Recognition (To Appear).

**Examples**

```
library(tidyverse)
#' \donttest{
library(tidymodels)
library(caret)
library(SSLR)

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$Class, p = .7, list = FALSE)
train <- breast[ train.index,]
test <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

#% LABELED
labeled.index <- createDataPartition(breast$Class, p = .2, list = FALSE)
```

```

train[~labeled.index,cls] <- NA

m <- EMLeastSquaresClassifierSSLR() %>% fit(Class ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)
#' }

```

---

EMNearestMeanClassifierSSLR

*General Interface for EMNearestMeanClassifier model*


---

### Description

model from RSSL package Semi-Supervised Nearest Mean Classifier using Expectation Maximization

Expectation Maximization applied to the nearest mean classifier assuming Gaussian classes with a spherical covariance matrix.

Starting from the supervised solution, uses the Expectation Maximization algorithm (see Dempster et al. (1977)) to iteratively update the means and shared covariance of the classes (Maximization step) and updates the responsibilities for the unlabeled objects (Expectation step).

### Usage

```
EMNearestMeanClassifierSSLR(method = "EM", scale = FALSE, eps = 1e-04)
```

### Arguments

method	character; Currently only "EM"
scale	Should the features be normalized? (default: FALSE)
eps	Stopping criterion for the maximimization

### References

Dempster, A., Laird, N. & Rubin, D., 1977. Maximum likelihood from incomplete data via the EM algorithm. *Journal of the Royal Statistical Society. Series B*, 39(1), pp.1-38.

### Examples

```

library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

```

```

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$class, p = .7, list = FALSE)
train <- breast[ train.index,]
test <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

#% LABELED
labeled.index <- createDataPartition(breast$class, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- EMNearestMeanClassifierSSLR() %>% fit(Class ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)

```

---

EntropyRegularizedLogisticRegressionSSLR

*General Interface for EntropyRegularizedLogisticRegression model*


---

## Description

model from RSSL package R Implementation of entropy regularized logistic regression implementation as proposed by Grandvalet & Bengio (2005). An extra term is added to the objective function of logistic regression that penalizes the entropy of the posterior measured on the unlabeled examples.

## Usage

```

EntropyRegularizedLogisticRegressionSSLR(
  lambda = 0,
  lambda_entropy = 1,
  intercept = TRUE,
  init = NA,
  scale = FALSE,
  x_center = FALSE
)

```

## Arguments

lambda	l2 Regularization
lambda_entropy	Weight of the labeled observations compared to the unlabeled observations
intercept	logical; Whether an intercept should be included

init	Initial parameters for the gradient descent
scale	logical; Should the features be normalized? (default: FALSE)
x_center	logical; Should the features be centered?

## References

Grandvalet, Y. & Bengio, Y., 2005. Semi-supervised learning by entropy minimization. In L. K. Saul, Y. Weiss, & L. Bottou, eds. *Advances in Neural Information Processing Systems 17*. Cambridge, MA: MIT Press, pp. 529-536.

## Examples

```
library(tidyverse)
library(caret)
library(tidymodels)
library(SSLR)

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$Class, p = .7, list = FALSE)
train <- breast[ train.index,]
test <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

#% LABELED
labeled.index <- createDataPartition(breast$Class, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- EntropyRegularizedLogisticRegressionSSLR() %>% fit(Class ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)
```

---

fit.model_sslr	<i>Fit with formula and data</i>
----------------	----------------------------------

---

## Description

Funtion to fit through the formula

## Usage

```
## S3 method for class 'model_sslr'
fit(object, formula = NULL, data = NULL, ...)
```



**Arguments**

object	is the model
formula	is the formula
data	is the total data train
...	unused in this case

---

fit\_decision\_tree      *An S4 method to fit decision tree.*

---

**Description**

An S4 method to fit decision tree.

**Usage**

```
fit_decision_tree(object, ...)
```

**Arguments**

object	DecisionTree object
...	This parameter is included for compatibility reasons.

---

fit\_decision\_tree, DecisionTreeClassifier-method  
*Fit decision tree*

---

**Description**

method in class DecisionTreeClassifier used to build a Decision Tree

**Usage**

```
## S4 method for signature 'DecisionTreeClassifier'
fit_decision_tree(
  object,
  X,
  y,
  min_samples_split = 20,
  min_samples_leaf = ceiling(min_samples_split/3),
  w = 0.5
)
```

**Arguments**

object	A DecisionTreeClassifier object
X	A object that can be coerced as data.frame. Training instances
y	A vector with the labels of the training instances. In this vector the unlabeled instances are specified with the value NA.
min_samples_split	the minimum number of observations to do split
min_samples_leaf	the minimum number of any terminal leaf node
w	weight parameter ranging from 0 to 1

---

fit\_random\_forest, RandomForestSemisupervised-method  
*Fit Random Forest*

---

**Description**

method in classRandomForestSemisupervised used to build a Decision Tree

**Usage**

```
## S4 method for signature 'RandomForestSemisupervised'
fit_random_forest(
  object,
  X,
  y,
  mtry = 2,
  trees = 500,
  min_n = 2,
  w = 0.5,
  replace = TRUE,
  tree_max_depth = Inf,
  sampsize = if (replace) nrow(X) else ceiling(0.632 * nrow(X)),
  min_samples_leaf = if (!is.null(y) && !is.factor(y)) 5 else 1,
  allowParallel = TRUE
)
```

**Arguments**

object	A RandomForestSemisupervised object
X	A object that can be coerced as data.frame. Training instances
y	A vector with the labels of the training instances. In this vector the unlabeled instances are specified with the value NA.
mtry	number of features in each decision tree

trees	number of trees. Default is 5
min_n	number of minimum samples in each tree
w	weight parameter ranging from 0 to 1
replace	replacing type in sampling
tree_max_depth	maximum tree depth. Default is Inf
sampsize	Size of sample. Default if (replace) nrow(x) else ceiling(.632*nrow(x))
min_samples_leaf	the minimum number of any terminal leaf node
allowParallel	Execute Random Forest in parallel if doParallel is loaded. Default is TRUE

**Value**

list of decision trees

---

fit_xy.model_sslr	<i>Fit with x and y</i>
-------------------	-------------------------

---

**Description**

Funtion to fit with x and y

**Usage**

```
## S3 method for class 'model_sslr'
fit_xy(object, x = NULL, y = NULL, ...)
```

**Arguments**

object	is the model
x	is a data frame or matrix with train dataset without objective feature. X have labeled and unlabeled data
y	is objective feature with labeled values and NA values in unlabeled data
...	unused in this case

---

fit_x_u	<i>fit_x_u object</i>
---------	-----------------------

---

**Description**

fit\_x\_u

**Usage**

fit\_x\_u(object, ...)

**Arguments**

object	object
...	other parameters to be passed

---

fit_x_u.model_sslr	<i>Fit with x , y (labeled data) and unlabeled data (x_U)</i>
--------------------	---

---

**Description**

Funtion to fit with x and y and x\_U. Function calcule y with NA values and append in y param

**Usage**

```
## S3 method for class 'model_sslr'
fit_x_u(object, x = NULL, y = NULL, x_U = NULL, ...)
```

**Arguments**

object	is the model
x	is a data frame or matrix with train dataset without objective feature. X only have labeled data
y	is objective feature with labeled values
x_U	train unlabeled data without objective feature
...	This parameter is included for compatibility reasons.

---

get\_class\_max\_prob      *Get most frequented*

---

**Description**

Get value most frequented in vector Used in predictions. It calls a predict with type = "prob" in Decision Tree

**Usage**

```
get_class_max_prob(trees, input)
```

**Arguments**

trees	trees list
input	is input to be predicted

---

get\_function      *FUNCTION TO GET FUNCTION METHOD*

---

**Description**

FUNCTION TO GET FUNCTION METHOD SPECIFIC

**Usage**

```
get_function(met)
```

**Arguments**

met	character
-----	-----------

**Value**

method\_train (function)

---

get\_function\_generic *FUNCTION TO GET FUNCTION METHOD*

---

**Description**

FUNCTION TO GET FUNCTION METHOD GENERIC

**Usage**

```
get_function_generic(met)
```

**Arguments**

met                    character

**Value**

method\_train (function)

---

get\_levels\_categoric *Function to get group from gini index*

---

**Description**

Function to get group from gini index. Used in categorical variable From: <https://freakonometrics.hypotheses.org/20736>

**Usage**

```
get_levels_categoric(column, Y)
```

**Arguments**

column                is the column  
Y                      values

---

`get_most_frequented`    *Get most frequented*

---

**Description**

Get value most frequented in vector Used in predictions

**Usage**

`get_most_frequented(elements)`

**Arguments**

`elements`            vector with values

---

`get_value_mean`            *Get value mean*

---

**Description**

Get value most frequented in vector Used in predictions. It calls a predict with type = "numeric" in Decision Tree

**Usage**

`get_value_mean(trees, input)`

**Arguments**

`trees`                trees list  
`input`                is input to be predicted

---

`get_x_y`*FUNCTION TO GET REAL X AND Y WITH FORMULA AND DATA*

---

**Description**

FUNCTION TO GET REAL X AND Y WITH FORMULA AND DATA

**Usage**

```
get_x_y(form, data)
```

**Arguments**

form	formula
data	data values, matrix, dataframe..

**Value**

x (matrix,dataframe...) and y(factor)

---

`gini_or_variance`*Gini or Variance by column*

---

**Description**

function used to calculate the gini coefficient or variance according to the type of the column. This function is called for the creation of the decision tree

**Usage**

```
gini_or_variance(X)
```

**Arguments**

X	column to calculate variance or gini
---	--------------------------------------



---

gini_prob	<i>Function to compute Gini index</i>
-----------	---------------------------------------

---

**Description**

Function to compute Gini index From: <https://freakonometrics.hypotheses.org/20736>

**Usage**

```
gini_prob(y, classe)
```

**Arguments**

y	values
classe	classes

---

grow_tree	<i>An S4 method to grow tree.</i>
-----------	-----------------------------------

---

**Description**

An S4 method to grow tree.

**Usage**

```
grow_tree(object, ...)
```

**Arguments**

object	DecisionTree object
...	This parameter is included for compatibility reasons.

---

grow\_tree, DecisionTreeClassifier-method  
*Function grow tree*

---

**Description**

Function to grow tree in Decision Tree

**Usage**

```
## S4 method for signature 'DecisionTreeClassifier'  
grow_tree(object, X, y, parms, depth = 0)
```

**Arguments**

object	DecisionTree instance
X	data values
y	classes
parms	parameters for grow tree
depth	depth in tree

---

knn\_regression      *knn\_regression*

---

**Description**

create model knn

**Usage**

```
knn_regression(k, x, y, p)
```

**Arguments**

k	parameter in KNN model
x	data
y	vector labeled data
p	distance order

**Description**

model from RSSL package Manifold regularization applied to the support vector machine as proposed in Belkin et al. (2006). As an adjacency matrix, we use the k nearest neighbour graph based on a chosen distance (default: euclidean).

**Usage**

```
LaplacianSVMSSLR(
  lambda = 1,
  gamma = 1,
  scale = TRUE,
  kernel = kernlab::vanilladot(),
  adjacency_distance = "euclidean",
  adjacency_k = 6,
  normalized_laplacian = FALSE,
  eps = 1e-09
)
```

**Arguments**

lambda	numeric; L2 regularization parameter
gamma	numeric; Weight of the unlabeled data
scale	logical; Should the features be normalized? (default: FALSE)
kernel	kernlab::kernel to use
adjacency_distance	character; distance metric used to construct adjacency graph from the dist function. Default: "euclidean"
adjacency_k	integer; Number of of neighbours used to construct adjacency graph.
normalized_laplacian	logical; If TRUE use the normalized Laplacian, otherwise, the Laplacian is used
eps	numeric; Small value to ensure positive definiteness of the matrix in the QP formulation

**References**

Belkin, M., Niyogi, P. & Sindhwani, V., 2006. Manifold regularization: A geometric framework for learning from labeled and unlabeled examples. *Journal of Machine Learning Research*, 7, pp.2399-2434.

**Examples**

```

library(tidyverse)
library(caret)
library(tidymodels)
library(SSLR)

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$Class, p = .7, list = FALSE)
train <- breast[ train.index,]
test <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

#% LABELED
labeled.index <- createDataPartition(breast$Class, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

library(kernlab)
m <- LaplacianSVMSSLR(kernel=kernlab::vanilladot()) %>%
  fit(Class ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)

```

---

 LinearTSVMSSLR

*General Interface for LinearTSVM model*


---

**Description**

model from RSSL package Implementation of the Linear Support Vector Classifier. Can be solved in the Dual formulation, which is equivalent to [SVM](#) or the Primal formulation.

**Usage**

```

LinearTSVMSSLR(
  C = 1,
  Cstar = 0.1,
  s = 0,
  x_center = FALSE,
  scale = FALSE,
  eps = 1e-06,
  verbose = FALSE,
  init = NULL
)

```

**Arguments**

C	Cost variable
Cstar	numeric; Cost parameter of the unlabeled objects
s	numeric; parameter controlling the loss function of the unlabeled objects
x_center	logical; Should the features be centered?
scale	Whether a z-transform should be applied (default: TRUE)
eps	Small value to ensure positive definiteness of the matrix in QP formulation
verbose	logical; Controls the verbosity of the output
init	numeric; Initial classifier parameters to start the convex concave procedure

**Examples**

```
library(tidyverse)
library(caret)
library(tidymodels)
library(SSLR)

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$Class, p = .7, list = FALSE)
train <- breast[ train.index,]
test <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

#% LABELED
labeled.index <- createDataPartition(breast$Class, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- LinearTSMVSSLR() %>% fit(Class ~ ., data = train)

#Accuracy
predict(m, test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)
```

---

load\_parsnip

*Load parsnip*


---

**Description**

function to load parsnip package

**Usage**

```
load_parsnip()
```

---

load_RANN	<i>Load parsnip</i>
-----------	---------------------

---

**Description**

function to load parsnip package

**Usage**

```
load_RANN()
```

---

load_RSSL	<i>Load RSSL</i>
-----------	------------------

---

**Description**

function to load RSSL package

**Usage**

```
load_RSSL()
```

---

MCNearestMeanClassifierSSLR	<i>General Interface for MCNearestMeanClassifier (Moment Constrained Semi-supervised Nearest Mean Classifier) model</i>
-----------------------------	---

---

**Description**

model from RSSL package Update the means based on the moment constraints as defined in Loog (2010). The means estimated using the labeled data are updated by making sure their weighted mean corresponds to the overall mean on all (labeled and unlabeled) data. Optionally, the estimated variance of the classes can be re-estimated after this update is applied by setting `update_sigma` to TRUE. To get the true nearest mean classifier, rather than estimate the class priors, set them to equal priors using, for instance `prior=matrix(0.5, 2)`.

**Usage**

```
MCNearestMeanClassifierSSLR(
  update_sigma = FALSE,
  prior = NULL,
  x_center = FALSE,
  scale = FALSE
)
```

**Arguments**

update_sigma	logical; Whether the estimate of the variance should be updated after the means have been updated using the unlabeled data
prior	matrix; Class priors for the classes
x_center	logical; Should the features be centered?
scale	logical; Should the features be normalized? (default: FALSE)

**References**

Loog, M., 2010. Constrained Parameter Estimation for Semi-Supervised Learning: The Case of the Nearest Mean Classifier. In Proceedings of the 2010 European Conference on Machine learning and Knowledge Discovery in Databases. pp. 291-304.

**Examples**

```
library(tidyverse)
library(caret)
library(tidymodels)
library(SSLR)

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$Class, p = .7, list = FALSE)
train <- breast[ train.index,]
test <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

#% LABELED
labeled.index <- createDataPartition(breast$Class, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- MCNearestMeanClassifierSSLR() %>% fit(Class ~ ., data = train)

#Accuracy
predict(m, test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)
```

---

newDecisionTree	<i>Function to create DecisionTree</i>
-----------------	--

---

**Description**

Function to create DecisionTree

**Usage**

```
newDecisionTree(max_depth)
```

**Arguments**

max_depth	max depth in tree
-----------	-------------------

---

Node-class	<i>Class Node for Decision Tree</i>
------------	-------------------------------------

---

**Description**

Class Node for Decision Tree Slots: gini, num\_samples, num\_samples\_per\_class, predicted\_class\_value, feature\_index threshold, left, right, probabilities

---

nullOrNumericOrCharacter-class	<i>An S4 class to represent a class with more types values: null, numeric or character</i>
--------------------------------	--

---

**Description**

An S4 class to represent a class with more types values: null, numeric or character



---

oneNN	<i>1-NN supervised classifier builder</i>
-------	---

---

**Description**

Build a model using the given data to be able to predict the label or the probabilities of other instances, according to 1-NN algorithm.

**Usage**

```
oneNN(x = NULL, y)
```

**Arguments**

x	This argument is not used, the reason why he gets is to fulfill an agreement
y	a vector with the labels of training instances

**Value**

A model with the data needed to use 1-NN

---

predict,DecisionTreeClassifier-method	<i>Function to predict inputs in Decision Tree</i>
---------------------------------------	--

---

**Description**

Function to predict inputs in Decision Tree

**Usage**

```
## S4 method for signature 'DecisionTreeClassifier'
predict(object, inputs, type = "class")
```

**Arguments**

object	The Decision Tree object
inputs	data to be predicted
type	Is param to define the type of predict. It can be "class", to get class labels Or "prob" to get probabilties for class in each input. Default is "class"

---

predict, RandomForestSemisupervised-method

*Function to predict inputs in Decision Tree*

---

### Description

Function to predict inputs in Decision Tree

### Usage

```
## S4 method for signature 'RandomForestSemisupervised'
predict(
  object,
  inputs,
  type = "class",
  confident = "max_prob",
  allowParallel = TRUE
)
```

### Arguments

object	The Decision Tree object
inputs	data to be predicted
type	class raw
confident	Is param to define the type of predict. It can be "max_prob", to get class with sum of probability is the maximum Or "vote" to get the most frequented class in all trees. Default is "max_prob"
allowParallel	Execute Random Forest in parallel if doParallel is loaded.

---

predict.coBC

*Predictions of the coBC method*

---

### Description

Predicts the label of instances according to the coBC model.

### Usage

```
## S3 method for class 'coBC'
predict(object, x, ...)
```

**Arguments**

object	coBC model built with the <a href="#">coBC</a> function.
x	An object that can be coerced to a matrix. Depending on how the model was built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
...	This parameter is included for compatibility reasons.

**Details**

For additional help see [coBC](#) examples.

**Value**

Vector with the labels assigned.

---

predict.COREG	<i>Predictions of the COREG method</i>
---------------	--

---

**Description**

Predicts the label of instances according to the COREG model.

**Usage**

```
## S3 method for class 'COREG'
predict(object, x, type = "numeric", ...)
```

**Arguments**

object	Self-training model built with the <a href="#">COREG</a> function.
x	A object that is data
type	of predict in principal model (numeric)
...	This parameter is included for compatibility reasons.

**Details**

For additional help see [COREG](#) examples.

**Value**

Vector with the labels assigned (numeric).

---

predict.democratic      *Predictions of the Democratic method*

---

### Description

Predicts the label of instances according to the democratic model.

### Usage

```
## S3 method for class 'democratic'
predict(object, x, ...)
```

### Arguments

object	Democratic model built with the <a href="#">democratic</a> function.
x	A object that can be coerced as matrix. Depending on how was the model built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
...	This parameter is included for compatibility reasons.

### Details

For additional help see [democratic](#) examples.

### Value

Vector with the labels assigned.

---

predict.EMLeastSquaresClassifierSSLR  
*Predict EMLeastSquaresClassifierSSLR*

---

### Description

Predict EMLeastSquaresClassifierSSLR

### Usage

```
## S3 method for class 'EMLeastSquaresClassifierSSLR'
predict(object, x, ...)
```

### Arguments

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

```
predict.EMNearestMeanClassifierSSLR  
Predict EMNearestMeanClassifierSSLR
```

---

### **Description**

Predict EMNearestMeanClassifierSSLR

### **Usage**

```
## S3 method for class 'EMNearestMeanClassifierSSLR'  
predict(object, x, ...)
```

### **Arguments**

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

```
predict.EntropyRegularizedLogisticRegressionSSLR  
Predict EntropyRegularizedLogisticRegressionSSLR
```

---

### **Description**

Predict EntropyRegularizedLogisticRegressionSSLR

### **Usage**

```
## S3 method for class 'EntropyRegularizedLogisticRegressionSSLR'  
predict(object, x, ...)
```

### **Arguments**

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

predict.LaplacianSVMSSLR

*Predict LaplacianSVMSSLR*

---

### Description

Predict LaplacianSVMSSLR

### Usage

```
## S3 method for class 'LaplacianSVMSSLR'  
predict(object, x, ...)
```

### Arguments

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

predict.LinearTSVMSSLR

*Predict LinearTSVMSSLR*

---

### Description

Predict LinearTSVMSSLR

### Usage

```
## S3 method for class 'LinearTSVMSSLR'  
predict(object, x, ...)
```

### Arguments

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

```
predict.MCNearestMeanClassifierSSLR
  Predict MCNearestMeanClassifierSSLR
```

---

**Description**

Predict MCNearestMeanClassifierSSLR

**Usage**

```
## S3 method for class 'MCNearestMeanClassifierSSLR'
predict(object, x, ...)
```

**Arguments**

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

```
predict.model_sslr_fitted
  Predictions of model_sslr_fitted class
```

---

**Description**

Predicts from model. There are different types: class, prob, raw class returns tibble with one column  
 prob returns tibble with probabilities class columns raw returns factor or numeric values

**Usage**

```
## S3 method for class 'model_sslr_fitted'
predict(object, x, type = NULL, ...)
```

**Arguments**

object	model_sslr_fitted model built.
x	A object that can be coerced as matrix. Depending on how was the model built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
type	of predict in principal model: class, raw, prob, vote, max_prob, numeric
...	This parameter is included for compatibility reasons.

**Value**

tibble or vector.

---

`predict.OneNN`                      *Model Predictions*

---

**Description**

This function predicts the class label of instances or its probability of pertaining to each class based on the distance matrix.

**Usage**

```
## S3 method for class 'OneNN'
predict(object, dists, type = "prob", ...)
```

**Arguments**

<code>object</code>	A model of class <code>OneNN</code> built with <a href="#">oneNN</a>
<code>dists</code>	A matrix of distances between the instances to classify (by rows) and the instances used to train the model (by column)
<code>type</code>	A string that can take two values: "class" for computing the class of the instances or "prob" for computing the probabilities of belonging to each class.
<code>...</code>	Currently not used.

**Value**

If `type` is equal to "class" a vector of length equal to the rows number of matrix `dists`, containing the predicted labels. If `type` is equal to "prob" it returns a matrix which has `nrow(dists)` rows and a column for every class, where each cell represents the probability that the instance belongs to the class, according to 1NN.

---

`predict.RandomForestSemisupervised_fitted`  
*Predictions of the `SSLRDecisionTree_fitted` method*

---

**Description**

Predicts the label of instances according to the `RandomForestSemisupervised_fitted` model.

**Usage**

```
## S3 method for class 'RandomForestSemisupervised_fitted'
predict(object, x, type = "class", confident = "max_prob", ...)
```



**Arguments**

object	RandomForestSemisupervised_fitted.
x	A object that can be coerced as matrix. Depending on how was the model built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
type	of predict in principal model
confident	Is param to define the type of predict. It can be "max_prob", to get class with sum of probability is the maximum Or "vote" to get the most frequented class in all trees. Default is "max_prob"
...	This parameter is included for compatibility reasons.

**Value**

Vector with the labels assigned.

---

predict.selfTraining *Predictions of the Self-training method*

---

**Description**

Predicts the label of instances according to the selfTraining model.

**Usage**

```
## S3 method for class 'selfTraining'
predict(object, x, type = "class", ...)
```

**Arguments**

object	Self-training model built with the <a href="#">selfTraining</a> function.
x	A object that can be coerced as matrix. Depending on how was the model built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
type	of predict in principal model
...	This parameter is included for compatibility reasons.

**Details**

For additional help see [selfTraining](#) examples.

**Value**

Vector with the labels assigned.

---

predict.setred      *Predictions of the SETRED method*

---

### Description

Predicts the label of instances according to the setred model.

### Usage

```
## S3 method for class 'setred'  
predict(object, x, col_name = ".pred_class", ...)
```

### Arguments

object	SETRED model built with the <a href="#">setred</a> function.
x	A object that can be coerced as matrix. Depending on how was the model built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
col_name	is the colname from returned tibble in class type. The same from parsnip and tidymodels Default is .pred_clas
...	This parameter is included for compatibility reasons.

### Details

For additional help see [setred](#) examples.

### Value

Vector with the labels assigned.

---

predict.snnrce      *Predictions of the SNNRCE method*

---

### Description

Predicts the label of instances according to the snnrce model.

### Usage

```
## S3 method for class 'snnrce'  
predict(object, x, ...)
```

**Arguments**

object	SNNRCE model built with the <a href="#">snnrce</a> function.
x	A object that can be coerced as matrix. Depending on how was the model built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
...	This parameter is included for compatibility reasons.

**Details**

For additional help see [snnrce](#) examples.

**Value**

Vector with the labels assigned.

---

predict.snnrceG

*Predictions of the SNNRCE method*

---

**Description**

Predicts the label of instances according to the snnrceG model.

**Usage**

```
## S3 method for class 'snnrceG'  
predict(object, D, ...)
```

**Arguments**

object	model instance
D	distance matrix
...	This parameter is included for compatibility reasons.

---

```
predict.SSLRDecisionTree_fitted
```

*Predictions of the SSLRDecisionTree\_fitted method*

---

### Description

Predicts the label of instances SSLRDecisionTree\_fitted model.

### Usage

```
## S3 method for class 'SSLRDecisionTree_fitted'
predict(object, x, type = "class", ...)
```

### Arguments

object	model SSLRDecisionTree_fitted.
x	A object that can be coerced as matrix. Depending on how was the model built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
type	of predict in principal model
...	This parameter is included for compatibility reasons.

### Value

Vector with the labels assigned.

---

```
predict.triTraining
```

*Predictions of the Tri-training method*

---

### Description

Predicts the label of instances according to the triTraining model.

### Usage

```
## S3 method for class 'triTraining'
predict(object, x, ...)
```

### Arguments

object	Tri-training model built with the <a href="#">triTraining</a> function.
x	A object that can be coerced as matrix. Depending on how was the model built, x is interpreted as a matrix with the distances between the unseen instances and the selected training instances, or a matrix of instances.
...	This parameter is included for compatibility reasons.

**Details**

For additional help see [triTraining](#) examples.

**Value**

Vector with the labels assigned.

---

predict.TSVMSSLR	<i>Predict TSVMSSLR</i>
------------------	-------------------------

---

**Description**

Predict TSVMSSLR

**Usage**

```
## S3 method for class 'TSVMSSLR'
predict(object, x, ...)
```

**Arguments**

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

predict.USMLeastSquaresClassifierSSLR	<i>Predict USMLeastSquaresClassifierSSLR</i>
---------------------------------------	--

---

**Description**

Predict USMLeastSquaresClassifierSSLR

**Usage**

```
## S3 method for class 'USMLeastSquaresClassifierSSLR'
predict(object, x, ...)
```

**Arguments**

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

predict.WellSVMSSLR    *Predict WellSVMSSLR*

---

### Description

Predict WellSVMSSLR

### Usage

```
## S3 method for class 'WellSVMSSLR'
predict(object, x, ...)
```

### Arguments

object	is the object
x	is the dataset
...	This parameter is included for compatibility reasons.

---

predict\_inputs    *An S4 method to predict inputs.*

---

### Description

An S4 method to predict inputs.

### Usage

```
predict_inputs(object, ...)
```

### Arguments

object	DecisionTree object
...	This parameter is included for compatibility reasons.

---

*predict\_inputs,DecisionTreeClassifier-method*  
*Predict inputs Decision Tree*

---

**Description**

Function to predict one input in Decision Tree

**Usage**

```
## S4 method for signature 'DecisionTreeClassifier'  
predict_inputs(object, inputs, type = "class")
```

**Arguments**

object	DecisionTree object
inputs	inputs to be predicted
type	type prediction, class or prob

---

*print.model\_sslr*      *Print model SSLR*

---

**Description**

Print model SSLR

**Usage**

```
## S3 method for class 'model_sslr'  
print(object)
```

**Arguments**

object	model_sslr object to print
--------	----------------------------

---

*RandomForestSemisupervised-class*  
*Class Random Forest*

---

**Description**

Class Random Forest Slots: mtry, trees, min\_n, w, classes, mode

---

`selfTraining`*General Interface for Self-training model*

---

### Description

Self-training is a simple and effective semi-supervised learning classification method. The self-training classifier is initially trained with a reduced set of labeled examples. Then it is iteratively retrained with its own most confident predictions over the unlabeled examples. Self-training follows a wrapper methodology using a base supervised classifier to establish the possible class of unlabeled instances.

### Usage

```
selfTraining(learner, max.iter = 50, perc.full = 0.7, thr.conf = 0.5)
```

### Arguments

<code>learner</code>	model from parsnip package for training a supervised base classifier using a set of instances. This model need to have probability predictions (or optionally a distance matrix) and it's corresponding classes.
<code>max.iter</code>	maximum number of iterations to execute the self-labeling process. Default is 50.
<code>perc.full</code>	A number between 0 and 1. If the percentage of new labeled examples reaches this value the self-training process is stopped. Default is 0.7.
<code>thr.conf</code>	A number between 0 and 1 that indicates the confidence threshold. At each iteration, only the newly labelled examples with a confidence greater than this value ( <code>thr.conf</code> ) are added to the training set.

### Details

For predicting the most accurate instances per iteration, `selfTraining` uses the predictions obtained with the learner specified. To train a model using the learner function, it is required a set of instances (or a precomputed matrix between the instances if `x.inst` parameter is `FALSE`) in conjunction with the corresponding classes. Additional parameters are provided to the learner function via the `learner.pars` argument. The model obtained is a supervised classifier ready to predict new instances through the `pred` function. Using a similar idea, the additional parameters to the `pred` function are provided using the `pred.pars` argument. The `pred` function returns the probabilities per class for each new instance. The value of the `thr.conf` argument controls the confidence of instances selected to enlarge the labeled set for the next iteration.

The stopping criterion is defined through the fulfillment of one of the following criteria: the algorithm reaches the number of iterations defined in the `max.iter` parameter or the portion of the unlabeled set, defined in the `perc.full` parameter, is moved to the labeled set. In some cases, the process stops and no instances are added to the original labeled set. In this case, the user must assign a more flexible value to the `thr.conf` parameter.



**Value**

(When model fit) A list object of class "selfTraining" containing:

**model** The final base classifier trained using the enlarged labeled set.

**instances.index** The indexes of the training instances used to train the model. These indexes include the initial labeled instances and the newly labeled instances. Those indexes are relative to x argument.

**classes** The levels of y factor.

**pred** The function provided in the pred argument.

**pred.pars** The list provided in the pred.pars argument.

**References**

David Yarowsky.

*Unsupervised word sense disambiguation rivaling supervised methods.*

In Proceedings of the 33rd annual meeting on Association for Computational Linguistics, pages 189-196. Association for Computational Linguistics, 1995.

**Examples**

```
library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

data(wine)

set.seed(1)
train.index <- createDataPartition(wine$Wine, p = .7, list = FALSE)
train <- wine[ train.index,]
test <- wine[-train.index,]

cls <- which(colnames(wine) == "Wine")

#% LABELED
labeled.index <- createDataPartition(train$Wine, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

#We need a model with probability predictions from parsnip
#https://tidymodels.github.io/parsnip/articles/articles/Models.html
#It should be with mode = classification

#For example, with Random Forest
rf <- rand_forest(trees = 100, mode = "classification") %>%
  set_engine("randomForest")

m <- selfTraining(learner = rf,
                  perc.full = 0.7,
                  thr.conf = 0.5, max.iter = 10) %>% fit(Wine ~ ., data = train)
```

```
#Accuracy
predict(m, test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)
```

---

selfTrainingG

*Self-training generic method*


---

### Description

Self-training is a simple and effective semi-supervised learning classification method. The self-training classifier is initially trained with a reduced set of labeled examples. Then it is iteratively retrained with its own most confident predictions over the unlabeled examples. Self-training follows a wrapper methodology using one base supervised classifier to establish the possible class of unlabeled instances.

### Usage

```
selfTrainingG(
  y,
  gen.learner,
  gen.pred,
  max.iter = 50,
  perc.full = 0.7,
  thr.conf = 0.5
)
```

### Arguments

y	A vector with the labels of training instances. In this vector the unlabeled instances are specified with the value NA.
gen.learner	A function for training a supervised base classifier. This function needs two parameters, indexes and cls, where indexes indicates the instances to use and cls specifies the classes of those instances.
gen.pred	A function for predicting the probabilities per classes. This function must be two parameters, model and indexes, where the model is a classifier trained with gen.learner function and indexes indicates the instances to predict.
max.iter	Maximum number of iterations to execute the self-labeling process. Default is 50.
perc.full	A number between 0 and 1. If the percentage of new labeled examples reaches this value the self-training process is stopped. Default is 0.7.
thr.conf	A number between 0 and 1 that indicates the confidence threshold. At each iteration, only the newly labelled examples with a confidence greater than this value (thr.conf) are added to the training set.

## Details

SelfTrainingG can be helpful in those cases where the method selected as base classifier needs learner and pred functions with other specifications. For more information about the general self-training method, please see the [selfTraining](#) function. Essentially, the selfTraining function is a wrapper of the selfTrainingG function.

## Value

A list object of class "selfTrainingG" containing:

**model** The final base classifier trained using the enlarged labeled set.

**instances.index** The indexes of the training instances used to train the model. These indexes include the initial labeled instances and the newly labeled instances. Those indexes are relative to the y argument.

## Examples

```
library(SSLR)

## Load Wine data set
data(wine)
cls <- which(colnames(wine) == "Wine")
x <- wine[, - cls] # instances without classes
y <- wine[, cls] # the classes
x <- scale(x)

set.seed(20)

# Use 50% of instances for training
tra.idx <- sample(x = length(y), size = ceiling(length(y) * 0.5))
xtrain <- x[tra.idx,]
ytrain <- y[tra.idx]

# Use 70% of train instances as unlabeled set
tra.na.idx <- sample(x = length(tra.idx), size = ceiling(length(tra.idx) * 0.7))
ytrain[tra.na.idx] <- NA

# Use the other 50% of instances for inductive testing
tst.idx <- setdiff(1:length(y), tra.idx)
xitest <- x[tst.idx,] # testing instances
yitest <- y[tst.idx] # classes of instances in xitest
# Use the unlabeled examples for transductive testing
xttest <- x[tra.idx[tra.na.idx],] # transductive testing instances
yttest <- y[tra.idx[tra.na.idx]] # classes of instances in xttest

library(caret)

#PREPARE DATA
data <- cbind(xtrain, Class = ytrain)
```

```

dtrain <- as.matrix(proxy::dist(x = xtrain, method = "euclidean", by_rows = TRUE))
ditest <- as.matrix(proxy::dist(x = xitest, y = xtrain, method = "euclidean", by_rows = TRUE))

ddata <- cbind(dtrain, Class = ytrain)
ddata <- as.data.frame(ddata)

ktrain <- as.matrix(exp(-0.048 * dtrain ^ 2))
kdata <- cbind(ktrain, Class = ytrain)
kdata <- as.data.frame(kdata)

ktrain <- as.matrix(exp(-0.048 * dtrain ^ 2))
kitest <- as.matrix(exp(-0.048 * ditest ^ 2))

## Example: Training from a set of instances with 1-NN (knn3) as base classifier.
gen.learner <- function(indexes, cls)
  caret::knn3(x = xtrain[indexes,], y = cls, k = 1)
gen.pred <- function(model, indexes)
  predict(model, xtrain[indexes,])

trControl_selfTrainingG1 <- list(gen.learner = gen.learner, gen.pred = gen.pred)
md1 <- train_generic(ytrain, method = "selfTrainingG", trControl = trControl_selfTrainingG1)

p1 <- predict(md1$model, xitest, type = "class")
table(p1, yitest)

confusionMatrix(p1, yitest)$overall[1]

## Example: Training from a distance matrix with 1-NN (oneNN) as base classifier.
dtrain <- as.matrix(proxy::dist(x = xtrain, method = "euclidean", by_rows = TRUE))
gen.learner <- function(indexes, cls) {
  m <- SSLR::oneNN(y = cls)
  attr(m, "tra.idx") <- indexes
  m
}

gen.pred <- function(model, indexes) {
  tra.idx <- attr(model, "tra.idx")
  d <- dtrain[indexes, tra.idx]
  prob <- predict(model, d, distance.weighting = "none")
  prob
}

trControl_selfTrainingG2 <- list(gen.learner = gen.learner, gen.pred = gen.pred)
md2 <- train_generic(ytrain, method = "selfTrainingG", trControl = trControl_selfTrainingG2)

ditest <- proxy::dist(x = xitest, y = xtrain[md2$instances.index,],

```

```

                                method = "euclidean", by_rows = TRUE)
p2 <- predict(md2$model, ditest, type = "class")
table(p2, yitest)

confusionMatrix(p2, yitest)$overall[1]

```

---

setred

*General Interface for SETRED model*


---

## Description

SETRED (SElf-TRaining with EDiting) is a variant of the self-training classification method (as implemented in the function `selfTraining`) with a different addition mechanism. The SETRED classifier is initially trained with a reduced set of labeled examples. Then, it is iteratively retrained with its own most confident predictions over the unlabeled examples. SETRED uses an amending scheme to avoid the introduction of noisy examples into the enlarged labeled set. For each iteration, the mislabeled examples are identified using the local information provided by the neighborhood graph.

## Usage

```

setred(
  dist = "Euclidean",
  learner,
  theta = 0.1,
  max.iter = 50,
  perc.full = 0.7,
  D = NULL
)

```

## Arguments

<code>dist</code>	A distance function or the name of a distance available in the proxy package to compute. Default is "Euclidean" the distance matrix in the case that <code>D</code> is <code>NULL</code> .
<code>learner</code>	model from <code>parsnip</code> package for training a supervised base classifier using a set of instances. This model need to have probability predictions (or optionally a distance matrix) and it's corresponding classes.
<code>theta</code>	Rejection threshold to test the critical region. Default is 0.1.
<code>max.iter</code>	maximum number of iterations to execute the self-labeling process. Default is 50.
<code>perc.full</code>	A number between 0 and 1. If the percentage of new labeled examples reaches this value the self-training process is stopped. Default is 0.7.
<code>D</code>	A distance matrix between all the training instances. This matrix is used to construct the neighborhood graph. Default is <code>NULL</code> , this means the method create a matrix with <code>dist</code> param

## Details

SETRED initiates the self-labeling process by training a model from the original labeled set. In each iteration, the `learner` function detects unlabeled examples for which it makes the most confident prediction and labels those examples according to the `pred` function. The identification of mislabeled examples is performed using a neighborhood graph created from the distance matrix. Most examples possess the same label in a neighborhood. So if an example locates in a neighborhood with too many neighbors from different classes, this example should be considered problematic. The value of the `theta` argument controls the confidence of the candidates selected to enlarge the labeled set. The lower this value is, the more restrictive is the selection of the examples that are considered good. For more information about the self-labeled process and the rest of the parameters, please see [selfTraining](#).

## Value

(When model fit) A list object of class "setred" containing:

**model** The final base classifier trained using the enlarged labeled set.

**instances.index** The indexes of the training instances used to train the model. These indexes include the initial labeled instances and the newly labeled instances. Those indexes are relative to `x` argument.

**classes** The levels of `y` factor.

**pred** The function provided in the `pred` argument.

**pred.pars** The list provided in the `pred.pars` argument.

## References

Ming Li and ZhiHua Zhou.

*Setred: Self-training with editing.*

In *Advances in Knowledge Discovery and Data Mining*, volume 3518 of *Lecture Notes in Computer Science*, pages 611-621. Springer Berlin Heidelberg, 2005. ISBN 978-3-540-26076-9. doi: 10.1007/11430919\_71.

## Examples

```
library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

data(wine)

set.seed(1)
train.index <- createDataPartition(wine$Wine, p = .7, list = FALSE)
train <- wine[ train.index,]
test <- wine[-train.index,]

cls <- which(colnames(wine) == "Wine")

#% LABELED
```

```

labeled.index <- createDataPartition(wine$Wine, p = .2, list = FALSE)
train[~labeled.index,cls] <- NA

#We need a model with probability predictions from parsnip
#https://tidymodels.github.io/parsnip/articles/articles/Models.html
#It should be with mode = classification

#For example, with Random Forest
rf <- rand_forest(trees = 100, mode = "classification") %>%
  set_engine("randomForest")

m <- setred(learner = rf,
            theta = 0.1,
            max.iter = 2,
            perc.full = 0.7) %>% fit(Wine ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)

#Another example, with dist matrix

distance <- as.matrix(proxy::dist(train[,-cls], method = "Euclidean",
                                  by_rows = TRUE, diag = TRUE, upper = TRUE))

m <- setred(learner = rf,
            theta = 0.1,
            max.iter = 2,
            perc.full = 0.7,
            D = distance) %>% fit(Wine ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)

```

---

setredG

*SETRED generic method*


---

## Description

SETRED is a variant of the self-training classification method ([selfTraining](#)) with a different addition mechanism. The SETRED classifier is initially trained with a reduced set of labeled examples. Then it is iteratively retrained with its own most confident predictions over the unlabeled examples. SETRED uses an amending scheme to avoid the introduction of noisy examples into

the enlarged labeled set. For each iteration, the mislabeled examples are identified using the local information provided by the neighborhood graph.

### Usage

```
setredG(
  y,
  D,
  gen.learner,
  gen.pred,
  theta = 0.1,
  max.iter = 50,
  perc.full = 0.7
)
```

### Arguments

y	A vector with the labels of training instances. In this vector the unlabeled instances are specified with the value NA.
D	A distance matrix between all the training instances. This matrix is used to construct the neighborhood graph.
gen.learner	A function for training a supervised base classifier. This function needs two parameters, indexes and cls, where indexes indicates the instances to use and cls specifies the classes of those instances.
gen.pred	A function for predicting the probabilities per classes. This function must be two parameters, model and indexes, where the model is a classifier trained with gen.learner function and indexes indicates the instances to predict.
theta	Rejection threshold to test the critical region. Default is 0.1.
max.iter	Maximum number of iterations to execute the self-labeling process. Default is 50.
perc.full	A number between 0 and 1. If the percentage of new labeled examples reaches this value the self-training process is stopped. Default is 0.7.

### Details

SetredG can be helpful in those cases where the method selected as base classifier needs a learner and pred functions with other specifications. For more information about the general setred method, please see [setred](#) function. Essentially, setred function is a wrapper of setredG function.

### Value

A list object of class "setredG" containing:

**model** The final base classifier trained using the enlarged labeled set.

**instances.index** The indexes of the training instances used to train the model. These indexes include the initial labeled instances and the newly labeled instances. Those indexes are relative to the y argument.



**Examples**

```

library(SSLR)
library(caret)

## Load Wine data set
data(wine)

cls <- which(colnames(wine) == "Wine")
x <- wine[, - cls] # instances without classes
y <- wine[, cls] # the classes
x <- scale(x) # scale the attributes

## Prepare data
set.seed(20)
# Use 50% of instances for training
tra.idx <- sample(x = length(y), size = ceiling(length(y) * 0.5))
xtrain <- x[tra.idx,] # training instances
ytrain <- y[tra.idx] # classes of training instances
# Use 70% of train instances as unlabeled set
tra.na.idx <- sample(x = length(tra.idx), size = ceiling(length(tra.idx) * 0.7))
ytrain[tra.na.idx] <- NA # remove class information of unlabeled instances

# Use the other 50% of instances for inductive testing
tst.idx <- setdiff(1:length(y), tra.idx)
xitest <- x[tst.idx,] # testing instances
yitest <- y[tst.idx] # classes of testing instances

# Compute distances between training instances
D <- as.matrix(proxy::dist(x = xtrain, method = "euclidean", by_rows = TRUE))

## Example: Training from a set of instances with 1-NN (knn3) as base classifier.
# Compute distances between training instances
D <- as.matrix(proxy::dist(x = xtrain, method = "euclidean", by_rows = TRUE))

## Example: Training from a set of instances with 1-NN (knn3) as base classifier.
gen.learner <- function(indexes, cls)
  caret::knn3(x = xtrain[indexes,], y = cls, k = 1)
gen.pred <- function(model, indexes)
  predict(model, xtrain[indexes,])

trControl_SETRED1 <- list(D = D, gen.learner = gen.learner,
                        gen.pred = gen.pred)
md1 <- train_generic(ytrain, method = "setredG", trControl = trControl_SETRED1)

'md1 <- setredG(y = ytrain, D, gen.learner, gen.pred)'

cls1 <- predict(md1$model, xitest, type = "class")
table(cls1, yitest)

confusionMatrix(cls1, yitest)$overall[1]

```

```

## Example: Training from a distance matrix with 1-NN (oneNN) as base classifier
gen.learner <- function(indexes, cls) {
  m <- SSLR::oneNN(y = cls)
  attr(m, "tra.idx") <- indexes
  m
}

gen.pred <- function(model, indexes) {
  tra.idx <- attr(model, "tra.idx")
  d <- D[indexes, tra.idx]
  prob <- predict(model, d, distance.weighting = "none")
  prob
}

trControl_SETRED2 <- list(D = D, gen.learner = gen.learner,
  gen.pred = gen.pred)
md2 <- train_generic(ytrain, method = "setredG", trControl = trControl_SETRED2)

ditest <- proxy::dist(x = xitest, y = xtrain[md2$instances.index,],
  method = "euclidean", by_rows = TRUE)

cls2 <- predict(md2$model, ditest, type = "class")
table(cls2, yitest)

confusionMatrix(cls2, yitest)$overall[1]

```

---

snnrce

*General Interface for SNNRCE model*


---

## Description

SNNRCE (Self-training Nearest Neighbor Rule using Cut Edges) is a variant of the self-training classification method ([selfTraining](#)) with a different addition mechanism and a fixed learning scheme (1-NN). SNNRCE uses an amending scheme to avoid the introduction of noisy examples into the enlarged labeled set. The mislabeled examples are identified using the local information provided by the neighborhood graph. A statistical test using cut edge weight is used to modify the labels of the missclassified examples.

## Usage

```
snnrce(x.inst = TRUE, dist = "Euclidean", alpha = 0.1)
```

**Arguments**

<code>x.inst</code>	A boolean value that indicates if <code>x</code> is or not an instance matrix. Default is <code>TRUE</code> .
<code>dist</code>	A distance function available in the <code>proxy</code> package to compute the distance matrix in the case that <code>x.inst</code> is <code>TRUE</code> .
<code>alpha</code>	Rejection threshold to test the critical region. Default is 0.1.

**Details**

SNNRCE initiates the self-labeling process by training a 1-NN from the original labeled set. This method attempts to reduce the noise in examples by labeling those instances with no cut edges in the initial stages of self-labeling learning. These highly confident examples are added into the training set. The remaining examples follow the standard self-training process until a minimum number of examples will be labeled for each class. A statistical test using cut edge weight is used to modify the labels of the missclassified examples. The value of the `alpha` argument defines the critical region where the candidates examples are tested. The higher this value is, the more relaxed it is the selection of the examples that are considered mislabeled.

**Value**

(When model fit) A list object of class "snnrce" containing:

**model** The final base classifier trained using the enlarged labeled set.

**instances.index** The indexes of the training instances used to train the model. These indexes include the initial labeled instances and the newly labeled instances. Those indexes are relative to `x` argument.

**classes** The levels of `y` factor.

**x.inst** The value provided in the `x.inst` argument.

**dist** The value provided in the `dist` argument when `x.inst` is `TRUE`.

**xtrain** A matrix with the subset of training instances referenced by the indexes `instances.index` when `x.inst` is `TRUE`.

**References**

Yu Wang, Xiaoyan Xu, Haifeng Zhao, and Zhongsheng Hua.  
*Semisupervised learning based on nearest neighbor rule and cut edges.*  
 Knowledge-Based Systems, 23(6):547-554, 2010. ISSN 0950-7051. doi: <http://dx.doi.org/10.1016/j.knosys.2010.03.012>.

**Examples**

```
library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

data(wine)
set.seed(1)
train.index <- createDataPartition(wine$Wine, p = .7, list = FALSE)
train <- wine[ train.index,]
```

```

test <- wine[-train.index,]

cls <- which(colnames(wine) == "Wine")

#% LABELED
labeled.index <- createDataPartition(wine$Wine, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- snnrce(x.inst = TRUE,
            dist = "Euclidean",
            alpha = 0.1) %>% fit(Wine ~ ., data = train)

predict(m, test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)

```

---

SSLRDecisionTree

*General Interface Decision Tree model*


---

### Description

Decision Tree is a simple and effective semi-supervised learning method. Based on the article "Semi-supervised classification trees". It also offers many parameters to modify the behavior of this method. It is the same as the traditional Decision Tree algorithm, but the difference is how the gini coefficient is calculated (classification). In regression we use SSE metric (different from the original investigation) It can be used in classification or regression. If Y is numeric is for regression, classification in another case

### Usage

```

SSLRDecisionTree(
  max_depth = 30,
  w = 0.5,
  min_samples_split = 20,
  min_samples_leaf = ceiling(min_samples_split/3)
)

```

### Arguments

max_depth	A number from 1 to Inf. Is the maximum number of depth in Decision Tree Default is 30
w	weight parameter ranging from 0 to 1. Default is 0.5
min_samples_split	the minimum number of observations to do split. Default is 20
min_samples_leaf	the minimum number of any terminal leaf node. Default is ceiling(min_samples_split/3)

**Details**

In this model we can make predictions with prob type

**References**

Jurica Levati, Michelangelo Ceci, Dragi Kocev, Saso Dzeroski.  
*Semi-supervised classification trees*.  
 Published online: 25 March 2017 © Springer Science Business Media New York 2017

**Examples**

```
library(tidyverse)
library(caret)
library(SSLR)
library(tidymodels)

data(wine)

set.seed(1)
train.index <- createDataPartition(wine$Wine, p = .7, list = FALSE)
train <- wine[ train.index,]
test <- wine[-train.index,]

cls <- which(colnames(wine) == "Wine")

#% LABELED
labeled.index <- createDataPartition(wine$Wine, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- SSLRDecisionTree(min_samples_split = round(length(labeled.index) * 0.25),
                      w = 0.3,
                      ) %>% fit(Wine ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)

#For probabilities
predict(m,test, type = "prob")
```

**Description**

Random Forest is a simple and effective semi-supervised learning method. It is the same as the traditional Random Forest algorithm, but the difference is that it use Semi supervised Decision Trees It can be used in classification or regression. If Y is numeric is for regression, classification in another case

**Usage**

```
SSLRRandomForest(
  mtry = NULL,
  trees = 500,
  min_n = NULL,
  w = 0.5,
  replace = TRUE,
  tree_max_depth = Inf,
  sampsize = NULL,
  min_samples_leaf = NULL,
  allowParallel = TRUE
)
```

**Arguments**

mtry	number of features in each decision tree. Default is null. This means that $mtry = \log(n\_features) + 1$
trees	number of trees. Default is 500
min_n	number of minimum samples in each tree Default is null. This means that uses all training data
w	weight parameter ranging from 0 to 1. Default is 0.5
replace	replacing type in sampling. Default is true
tree_max_depth	maximum tree depth. Default is Inf
sampsize	Size of sample. Default if (replace) $nrow(x)$ else $\text{ceiling}(.632 * nrow(x))$
min_samples_leaf	the minimum number of any terminal leaf node. Default is 1
allowParallel	Execute Random Forest in parallel if doParallel is loaded. Default is TRUE

**Details**

We can use paralleling processing with doParallel package and allowParallel = TRUE.

**References**

Jurica Levati, Michelangelo Ceci, Dragi Koccev, Saso Dzeroski.  
*Semi-supervised classification trees*.  
 Published online: 25 March 2017 © Springer Science Business Media New York 2017

**Examples**

```
library(tidyverse)
library(caret)
library(SSLR)
library(tidymodels)

data(wine)

set.seed(1)
train.index <- createDataPartition(wine$Wine, p = .7, list = FALSE)
train <- wine[ train.index,]
test <- wine[-train.index,]

cls <- which(colnames(wine) == "Wine")

#% LABELED
labeled.index <- createDataPartition(wine$Wine, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- SSLRRandomForest(trees = 5, w = 0.3) %>% fit(Wine ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)
```

---

train\_generic

*FUNCTION TO TRAIN GENERIC MODEL*

---

**Description**

FUNCTION TO TRAIN GENERIC MODEL

**Usage**

```
train_generic(y, ...)
```

**Arguments**

y (optional) factor (classes)  
... list parms trControl (method...)

**Value**

model trained

---

 triTraining

 General Interface for Tri-training model
 

---

### Description

Tri-training is a semi-supervised learning algorithm with a co-training style. This algorithm trains three classifiers with the same learning scheme from a reduced set of labeled examples. For each iteration, an unlabeled example is labeled for a classifier if the other two classifiers agree on the labeling proposed.

### Usage

```
triTraining(learner)
```

### Arguments

**learner** model from parsnip package for training a supervised base classifier using a set of instances. This model need to have probability predictions (or optionally a distance matrix) and it's corresponding classes.

### Details

Tri-training initiates the self-labeling process by training three models from the original labeled set, using the learner function specified. In each iteration, the algorithm detects unlabeled examples on which two classifiers agree with the classification and includes these instances in the enlarged set of the third classifier under certain conditions. The generation of the final hypothesis is produced via the majority voting. The iteration process ends when no changes occur in any model during a complete iteration.

### Value

A list object of class "triTraining" containing:

**model** The final three base classifiers trained using the enlarged labeled set.

**model.index** List of three vectors of indexes related to the training instances used per each classifier. These indexes are relative to the y argument.

**instances.index** The indexes of all training instances used to train the three models. These indexes include the initial labeled instances and the newly labeled instances. These indexes are relative to the y argument.

**model.index.map** List of three vectors with the same information in model.index but the indexes are relative to instances.index vector.

**classes** The levels of y factor.

**pred** The function provided in the pred argument.

**pred.pars** The list provided in the pred.pars argument.

**x.inst** The value provided in the x.inst argument.



## References

ZhiHua Zhou and Ming Li.  
*Tri-training: exploiting unlabeled data using three classifiers.*  
 IEEE Transactions on Knowledge and Data Engineering, 17(11):1529-1541, Nov 2005. ISSN 1041-4347. doi: 10.1109/TKDE.2005. 186.

## Examples

```
library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

data(wine)

set.seed(1)
train.index <- createDataPartition(wine$Wine, p = .7, list = FALSE)
train <- wine[ train.index,]
test <- wine[-train.index,]

cls <- which(colnames(wine) == "Wine")

#% LABELED
labeled.index <- createDataPartition(wine$Wine, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

#We need a model with probability predictions from parsnip
#https://tidymodels.github.io/parsnip/articles/articles/Models.html
#It should be with mode = classification

#For example, with Random Forest
rf <- rand_forest(trees = 100, mode = "classification") %>%
  set_engine("randomForest")

m <- triTraining(learner = rf) %>% fit(Wine ~ ., data = train)

#Accuracy
predict(m, test) %>%
  bind_cols(test) %>%
  metrics(truth = "Wine", estimate = .pred_class)
```

---

triTrainingCombine      *Combining the hypothesis*

---

## Description

This function combines the predictions obtained by the set of classifiers.

**Usage**

```
triTrainingCombine(pred)
```

**Arguments**

pred                    A list with the predictions of each classifiers

**Value**

A vector of classes

---

triTrainingG	<i>Tri-training generic method</i>
--------------	------------------------------------

---

**Description**

Tri-training is a semi-supervised learning algorithm with a co-training style. This algorithm trains three classifiers with the same learning scheme from a reduced set of labeled examples. For each iteration, an unlabeled example is labeled for a classifier if the other two classifiers agree on the labeling proposed.

**Usage**

```
triTrainingG(y, gen.learner, gen.pred)
```

**Arguments**

y                      A vector with the labels of training instances. In this vector the unlabeled instances are specified with the value NA.

gen.learner            A function for training three supervised base classifiers. This function needs two parameters, indexes and cls, where indexes indicates the instances to use and cls specifies the classes of those instances.

gen.pred              A function for predicting the probabilities per classes. This function must be two parameters, model and indexes, where the model is a classifier trained with gen.learner function and indexes indicates the instances to predict.

**Details**

TriTrainingG can be helpful in those cases where the method selected as base classifier needs a learner and pred functions with other specifications. For more information about the general triTraining method, please see the [triTraining](#) function. Essentially, the triTraining function is a wrapper of the triTrainingG function.

**Value**

A list object of class "triTrainingG" containing:

**model** The final three base classifiers trained using the enlarged labeled set.

**model.index** List of three vectors of indexes related to the training instances used per each classifier. These indexes are relative to the y argument.

**instances.index** The indexes of all training instances used to train the three models. These indexes include the initial labeled instances and the newly labeled instances. These indexes are relative to the y argument.

**model.index.map** List of three vectors with the same information in model.index but the indexes are relative to instances.index vector.

**Examples**

```
library(SSLR)
library(caret)

## Load Wine data set
data(wine)

cls <- which(colnames(wine) == "Wine")
x <- wine[, - cls] # instances without classes
y <- wine[, cls] # the classes
x <- scale(x) # scale the attributes

## Prepare data
set.seed(20)
# Use 50% of instances for training
tra.idx <- sample(x = length(y), size = ceiling(length(y) * 0.5))
xtrain <- x[tra.idx,] # training instances
ytrain <- y[tra.idx] # classes of training instances
# Use 70% of train instances as unlabeled set
tra.na.idx <- sample(x = length(tra.idx), size = ceiling(length(tra.idx) * 0.7))
ytrain[tra.na.idx] <- NA # remove class information of unlabeled instances

# Use the other 50% of instances for inductive testing
tst.idx <- setdiff(1:length(y), tra.idx)
xitest <- x[tst.idx,] # testing instances
yitest <- y[tst.idx] # classes of testing instances

## Example: Training from a set of instances with 1-NN (knn3) as base classifier.
gen.learner <- function(indexes, cls)
  caret::knn3(x = xtrain[indexes,], y = cls, k = 1)
gen.pred <- function(model, indexes)
  predict(model, xtrain[indexes,])

# Train
set.seed(1)

trControl_triTraining1 <- list(gen.learner = gen.learner,
                              gen.pred = gen.pred)
```

```

md1 <- train_generic(ytrain, method = "triTrainingG", trControl = trControl_triTraining1)

# Predict testing instances using the three classifiers
pred <- lapply(
  X = md1$model,
  FUN = function(m) predict(m, xitest, type = "class")
)
# Combine the predictions
cls1 <- triTrainingCombine(pred)
table(cls1, yitest)

confusionMatrix(cls1, yitest)$overall[1]

## Example: Training from a distance matrix with 1-NN (oneNN) as base classifier.
dtrain <- as.matrix(proxy::dist(x = xtrain, method = "euclidean", by_rows = TRUE))
gen.learner <- function(indexes, cls) {
  m <- SSLR::oneNN(y = cls)
  attr(m, "tra.idx") <- indexes
  m
}

gen.pred <- function(model, indexes) {
  tra.idx <- attr(model, "tra.idx")
  d <- dtrain[indexes, tra.idx]
  prob <- predict(model, d, distance.weighting = "none")
  prob
}

# Train
set.seed(1)

trControl_triTraining2 <- list(gen.learner = gen.learner,
                              gen.pred = gen.pred)
md2 <- train_generic(ytrain, method = "triTrainingG", trControl = trControl_triTraining2)

# Predict
ditest <- proxy::dist(x = xitest, y = xtrain[md2$instances.index,],
                     method = "euclidean", by_rows = TRUE)

# Predict testing instances using the three classifiers
pred <- mapply(
  FUN = function(m, indexes) {
    D <- ditest[, indexes]
    predict(m, D, type = "class")
  },
  m = md2$model,
  indexes = md2$model.index.map,
  SIMPLIFY = FALSE
)
# Combine the predictions

```

```

cls2 <- triTrainingCombine(pred)
table(cls2, yitest)

confusionMatrix(cls2, yitest)$overall[1]

```

TSVMSSLR

*General Interface for TSVM (Transductive SVM classifier using the convex concave procedure) model*

## Description

model from RSSL package Transductive SVM using the CCCP algorithm as proposed by Collobert et al. (2006) implemented in R using the quadprog package. The implementation does not handle large datasets very well, but can be useful for smaller datasets and visualization purposes.  $C$  is the cost associated with labeled objects, while  $C_{star}$  is the cost for the unlabeled objects.  $s$  control the loss function used for the unlabeled objects: it controls the size of the plateau for the symmetric ramp loss function. The balancing constraint makes sure the label assignments of the unlabeled objects are similar to the prior on the classes that was observed on the labeled data.

## Usage

```

TSVMSSLR(
  C = 1,
  Cstar = 0.1,
  kernel = kernlab::vanilladot(),
  balancing_constraint = TRUE,
  s = 0,
  x_center = TRUE,
  scale = FALSE,
  eps = 1e-09,
  max_iter = 20,
  verbose = FALSE
)

```

## Arguments

$C$	numeric; Cost parameter of the SVM
$C_{star}$	numeric; Cost parameter of the unlabeled objects
kernel	kernlab::kernel to use
balancing_constraint	logical; Whether a balancing constraint should be enforced that causes the fraction of objects assigned to each label in the unlabeled data to be similar to the label fraction in the labeled data.
$s$	numeric; parameter controlling the loss function of the unlabeled objects (generally values between -1 and 0)
$x_{center}$	logical; Should the features be centered?

scale	If TRUE, apply a z-transform to all observations in X and X_u before running the regression
eps	numeric; Stopping criterion for the maximimization
max_iter	integer; Maximum number of iterations
verbose	logical; print debugging messages, only works for vanilladot() kernel (default: FALSE)

## References

Collobert, R. et al., 2006. Large scale transductive SVMs. *Journal of Machine Learning Research*, 7, pp.1687-1712.

## Examples

```
library(tidyverse)
library(caret)
library(tidymodels)
library(SSLR)

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$Class, p = .7, list = FALSE)
train <- breast[ train.index,]
test <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

#% LABELED
labeled.index <- createDataPartition(breast$Class, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

library(kernlab)
m <- TSVMSSLR(kernel = kernlab::vanilladot()) %>% fit(Class ~ ., data = train)

#Accuracy
predict(m, test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)
```

---

USMLeastSquaresClassifierSSLR

*General Interface for USMLeastSquaresClassifier (Updated Second Moment Least Squares Classifier) model*

---

**Description**

model from RSSL package This methods uses the closed form solution of the supervised least squares problem, except that the second moment matrix ( $X'X$ ) is exchanged with a second moment matrix that is estimated based on all data. See for instance *Shaffer1991*, where in this implementation we use all data to estimate  $E(X'X)$ , instead of just the labeled data. This method seems to work best when the data is first centered `x_center=TRUE` and the outputs are scaled using `y_scale=TRUE`.

**Usage**

```
USMLEastSquaresClassifierSSLR(
  lambda = 0,
  intercept = TRUE,
  x_center = FALSE,
  scale = FALSE,
  y_scale = FALSE,
  ...,
  use_Xu_for_scaling = TRUE
)
```

**Arguments**

<code>lambda</code>	numeric; L2 regularization parameter
<code>intercept</code>	logical; Whether an intercept should be included
<code>x_center</code>	logical; Should the features be centered?
<code>scale</code>	logical; Should the features be normalized? (default: FALSE)
<code>y_scale</code>	logical; whether the target vector should be centered
<code>...</code>	Not used
<code>use_Xu_for_scaling</code>	logical; whether the unlabeled objects should be used to determine the mean and scaling for the normalization

**References**

Shaffer, J.P., 1991. The Gauss-Markov Theorem and Random Regressors. *The American Statistician*, 45(4), pp.269-273.

**Examples**

```
library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$class, p = .7, list = FALSE)
train <- breast[ train.index,]
```

```

test <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

## LABELED
labeled.index <- createDataPartition(breast$Class, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- USMLeastSquaresClassifierSSLR() %>% fit(Class ~ ., data = train)

#Accuracy
predict(m, test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)

```

---

WellSVMSSLR

*General Interface for WellSVM model*


---

## Description

model from RSSL package WellSVM is a minimax relaxation of the mixed integer programming problem of finding the optimal labels for the unlabeled data in the SVM objective function. This implementation is a translation of the Matlab implementation of Li (2013) into R.

## Usage

```

WellSVMSSLR(
  C1 = 1,
  C2 = 0.1,
  gamma = 1,
  x_center = TRUE,
  scale = FALSE,
  use_Xu_for_scaling = FALSE,
  max_iter = 20
)

```

## Arguments

C1	double; A regularization parameter for labeled data, default 1;
C2	double; A regularization parameter for unlabeled data, default 0.1;
gamma	double; Gaussian kernel parameter, i.e., $k(x,y) = \exp(-\text{gamma}^2 \ x-y\ ^2 / \text{avg})$ where avg is the average distance among instances; when gamma = 0, linear kernel is used. default gamma = 1;
x_center	logical; Should the features be centered?
scale	logical; Should the features be normalized? (default: FALSE)



```

use_Xu_for_scaling      logical; whether the unlabeled objects should be used to determine the mean and
                        scaling for the normalization
max_iter                integer; Maximum number of iterations

```

## References

Y.-F. Li, I. W. Tsang, J. T. Kwok, and Z.-H. Zhou. Scalable and Convex Weakly Labeled SVMs. *Journal of Machine Learning Research*, 2013.

R.-E. Fan, P.-H. Chen, and C.-J. Lin. Working set selection using second order information for training SVM. *Journal of Machine Learning Research* 6, 1889-1918, 2005.

## Examples

```

library(tidyverse)
library(tidymodels)
library(caret)
library(SSLR)

data(breast)

set.seed(1)
train.index <- createDataPartition(breast$Class, p = .7, list = FALSE)
train <- breast[ train.index,]
test  <- breast[-train.index,]

cls <- which(colnames(breast) == "Class")

#% LABELED
labeled.index <- createDataPartition(breast$Class, p = .2, list = FALSE)
train[-labeled.index,cls] <- NA

m <- WellSVMSSLR() %>% fit(Class ~ ., data = train)

#Accuracy
predict(m,test) %>%
  bind_cols(test) %>%
  metrics(truth = "Class", estimate = .pred_class)

```

---

wine

*Wine recognition data*

---

## Description

This dataset is the result of a chemical analysis of wine grown in the same region in Italy but derived from three different cultivars. The analysis determined the quantities of 13 constituents found in each of the three types of wines.

**Usage**

```
data(wine)
```

**Format**

A data frame with 178 rows and 14 variables including the class.

**Details**

The dataset is taken from the UCI data repository, to which it was donated by Riccardo Leardi, University of Genova. The attributes are as follows:

- Alcohol
- Malic acid
- Ash
- Alcalinity of ash
- Magnesium
- Total phenols
- Flavanoids
- Nonflavanoid phenols
- Proanthocyanins
- Color intensity
- Hue
- OD280/OD315 of diluted wines
- Proline
- Wine (class)

**Source**

<https://archive.ics.uci.edu/ml/datasets/Wine>

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