Package ‘wsrf’

March 13, 2022

Type Package
Title Weighted Subspace Random Forest for Classification
Version 1.7.27
Date 2022-03-13
Description A parallel implementation of Weighted Subspace Random Forest. The Weighted Subspace Random Forest algorithm was proposed in the International Journal of Data Warehousing and Mining by Baoxun Xu, Joshua Zhexue Huang, Graham Williams, Qiang Wang, and Yunming Ye (2012) <DOI:10.4018/jdwm.2012040103>. The algorithm can classify very high-dimensional data with random forests built using small subspaces. A novel variable weighting method is used for variable subspace selection in place of the traditional random variable sampling. This new approach is particularly useful in building models from high-dimensional data.
License GPL (>= 2)
BugReports https://github.com/SimonYansenZhao/wsrf/issues
Depends parallel, R (>= 3.3.0), Rcpp (>= 0.10.2), stats
LinkingTo Rcpp
Suggests knitr (>= 1.5), randomForest (>= 4.6.7), stringr (>= 0.6.2), rmarkdown (>= 1.6)
VignetteBuilder knitr
NeedsCompilation yes
SystemRequirements C++11

Classification/ACM-2012 Computing methodologies ~ Classification and regression trees, Computing methodologies ~ Supervised learning by classification, Computing methodologies ~ Massively parallel and high-performance simulations, Computing methodologies ~ Distributed simulation
combine.wsrf

**Description**

Combine two or more ensembles of trees into one.

**Usage**

```
combine(...)  
```

**Arguments**

```
...  
```

Two or more objects of class `randomForest`, to be combined into one.

**Value**

An object of class `wsrf`.

**See Also**

`subset`
Examples

library("wsrf")

# Prepare parameters.
ds <- iris
target <- "Species"
vars <- names(ds)
if (sum(is.na(ds[vars]))) ds[vars] <- randomForest::na.roughfix(ds[vars])
ds[target] <- as.factor(ds[[target]])
form <- as.formula(paste(target, "~ "))
set.seed(42)
train.1 <- sample(nrow(ds), 0.7*nrow(ds))
test.1 <- setdiff(seq_len(nrow(ds)), train.1)

set.seed(49)
train.2 <- sample(nrow(ds), 0.7*nrow(ds))
test.2 <- setdiff(seq_len(nrow(ds)), train.2)

# Build model. We disable parallelism here, since CRAN Repository
# Policy (https://cran.r-project.org/web/packages/policies.html)
# limits the usage of multiple cores to save the limited resource of
# the check farm.
model.wsrf.1 <- wsrf(form, data=ds[train.1, vars], parallel=FALSE)
model.wsrf.2 <- wsrf(form, data=ds[train.2, vars], parallel=FALSE)

# Merge two models.
model.wsrf.big <- combine.wsrf(model.wsrf.1, model.wsrf.2)
print(model.wsrf.big)
cl <- predict(model.wsrf.big, newdata=ds[test.1, vars], type="response")$response
actual <- ds[test.1, target]
(accuracy.wsrf <- mean(cl==actual))

correlation.wsrf

<table>
<thead>
<tr>
<th>Correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Give the measure for the diversity of the trees in the forest model built from wsrf.</td>
</tr>
</tbody>
</table>

Usage

## S3 method for class 'wsrf'
correlation(object, ...)
Details
The measure was introduced in Breiman (2001).

Value
A numeric value.

Author(s)
He Zhao and Graham Williams (SIAT, CAS)

References

See Also
wsrf

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### importance.wsrf

Extract Variable Importance Measure

#### Description
This is the extractor function for variable importance measures as produced by `wsrf`.

#### Usage
```r
# S3 method for class 'wsrf'
importance(x, type=NULL, class=NULL, scale=TRUE, ...)
```

#### Arguments
- `x` an object of class `wsrf`.
- `type` either 1 or 2, specifying the type of importance measure (1=mean decrease in accuracy, 2=mean decrease in node impurity).
- `class` for classification problem, which class-specific measure to return.
- `scale` for permutation based measures, should the measures be divided their “standard errors”?
- `...` not used.
Details
Here are the definitions of the variable importance measures. The first measure is computed from
permuting OOB data: For each tree, the prediction error on the out-of-bag portion of the data is
recorded. Then the same is done after permuting each predictor variable. The difference between
the two are then averaged over all trees, and normalized by the standard deviation of the differences.
The second measure is the total decrease in node impurities from splitting on the variable, averaged
over all trees. The node impurity is measured by the Information Gain Ratio index.

Value
A matrix of importance measure, one row for each predictor variable. The column(s) are different
importance measures.

See Also
randomForest

Description
Return out-of-bag error rate for "wsrf" model.

Usage
## S3 method for class 'wsrf'
oob.error.rate(object, tree, ...)

Arguments
object object of class wsrf.
tree logical or an integer vector for the index of a specific tree in the forest model. If
provided as an integer vector, oobErrorRate.wsrf will give the corresponding
out-of-bag error rates of the exact trees specified by tree. If TRUE, all error
rates will be presented. If FALSE or missing, the gross error rate for the forest
will be given.

Value
return a vector of error rates.

Author(s)
He Zhao and Graham Williams (SIAT, CAS)
predict.wsrf

See Also

wsrf

Description

Give the predictions for the new data by the forest model built from wsrf.

Usage

## S3 method for class 'wsrf'
predict(object, newdata, type=c("response", "class", "vote", "prob", "aprob", "waprob"), ...)

Arguments

object       object of class wsrf.
newdata      the data that needs to be predicted. Its format should be the same as that for wsrf.
type         the type of prediction required, a character vector indicating the types of output, and can be one of the values below:
              vote  matrix of vote counts
              response predicted values.
              class  the same as response.
              prob   matrix of class probabilities. The probability is the proportion of trees in the forest voting for the particular outcome (prob = votes / ntree)
              aprob  the average score from the decision trees for each class rather than the proportion of decision trees for each class (aprob = scores / ntree)
              waprob the weighted average, weighted by the accuracy of the tree (waprob = scores * accuracy / sum(accuracy))
              ...
              optional additional arguments. At present no additional arguments are used.

Value

a list of predictions for the new data with corresponding components for each type of predictions. For type=class or type=class, a vector of length nrow(newdata), otherwise, a matrix of nrow(newdata) * (number of class label). For example, if given type=c("class", "prob") and the return value is res, then res$class is a vector of predicted class labels of length nrow(newdata), and res$prob is a matrix of class probabilities.

Author(s)

He Zhao and Graham Williams (SIAT, CAS)
print.wsrf

See Also

wsrf

---

**Description**

Print a summary of the forest model or one specific tree in the forest model built from `wsrf`.

**Usage**

```r
## S3 method for class 'wsrf'
print(x, trees, ...)  
```

**Arguments**

- `x`: object of class `wsrf`.
- `trees`: the index of a specific tree. If missing, `print` will print a summary of the model.
- `...`: optional additional arguments. At present no additional arguments are used.

**Author(s)**

He Zhao and Graham Williams (SIAT, CAS)

**See Also**

wsrf

---

strength.wsrf  

**Description**

Give the measure for the collective performance of individual trees in the forest model built from `wsrf`.

**Usage**

```r
## S3 method for class 'wsrf'
strength(object, ...)  
```

**Arguments**

- `object`: object of class `wsrf`.
- `...`: optional additional arguments. At present no additional arguments are used.
Details
The measure was introduced in Breiman (2001).

Value
A numeric value.

Author(s)
He Zhao and Graham Williams (SIAT, CAS)

References

See Also
wsrf

description
Obtain a subset of a forest.

Usage
## S3 method for class 'wsrf'
subset(x, trees, ...)

Arguments
x an object of class wsr.
trees which trees should be included in the sub-forest. An integer vector, which indi-
cates the index of the trees.
... not used.

Value
An object of class wsr.

See Also
combine
Examples

```r
library("wsrf")

# Prepare parameters.
ds <- iris
target <- "Species"
vars <- names(ds)
if (sum(is.na(ds[vars]))) ds[vars] <- randomForest::na.roughfix(ds[vars])
ds[target] <- as.factor(ds[[target]])
form <- as.formula(paste(target, "~ "))
set.seed(42)
train <- sample(nrow(ds), 0.7*nrow(ds))
test <- setdiff(seq_len(nrow(ds)), train)

# Build model. We disable parallelism here, since CRAN Repository
# Policy (https://cran.r-project.org/web/packages/policies.html)
# limits the usage of multiple cores to save the limited resource of
# the check farm.

model.wsrf <- wsrf(form, data=ds[train, vars], parallel=FALSE)
print(model.wsrf)

# Subset.
submodel.wsrf <- subset.wsrf(model.wsrf, 1:200)
print(submodel.wsrf)
cl <- predict(submodel.wsrf, newdata=ds[test, vars], type="response")$response
actual <- ds[test, target]
(accuracy.wsrf <- mean(cl==actual))
```

<table>
<thead>
<tr>
<th>varCounts.wsrf</th>
<th>Number of Times of Variables Selected as Split Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

Return the times of each variable being selected as split condition. For evaluating the bias of wsrf towards attribute types (categorical and numerical) and the number of values each attribute has.

Usage

```r
## S3 method for class 'wsrf'
varCounts(object)
```

Arguments

- `object` object of class `wsrf`.

Value

A vector of integer. The length is the same as the training data for building that `wsrf` model.
build a forest of weighted subspace decision trees

Build weighted subspace C4.5-based decision trees to construct a forest.

Usage

```r
## S3 method for class 'formula'
wsrf(formula, data, ...)
## Default S3 method:
wsrf(x, y, mtry=floor(log2(length(x))+1), ntree=500,
     weights=TRUE, parallel=TRUE, na.action=na.fail,
     importance=FALSE, nodesize=2, clusterlogfile, ...)
```

Arguments

- `x`, `formula`: a data frame or a matrix of predictors, or a formula with a response but no interaction terms.
- `y`: a response vector.
- `data`: a data frame in which to interpret the variables named in the formula.
- `ntree`: number of trees to grow. By default, 500.
- `mtry`: number of variables to choose as candidates at each node split, by default, floor(log2(length(x))+1).
- `weights`: logical. TRUE for weighted subspace selection, which is the default; FALSE for random selection, and the trees are based on C4.5.
- `na.action`: a function indicate the behaviour when encountering NA values in data. By default, na.fail. If NULL, do nothing.
- `parallel`: whether to run multiple cores (TRUE), nodes, or sequentially (FALSE).
- `importance`: should importance of predictors be assessed?
- `nodesize`: minimum size of leaf node, i.e., minimum number of observations a leaf node represents. By default, 2.
- `clusterlogfile`: character. The pathname of the log file when building model in a cluster. For debug.
- `...`: optional parameters to be passed to the low level function `wsrf.default`. 
**Details**

See Xu, Huang, Williams, Wang, and Ye (2012) for more details of the algorithm, and Zhao, Williams, Huang (2017) for more details of the package.

Currently, `wsrf` can only be used for classification. When `weights=FALSE`, C4.5-based trees (Quinlan (1993)) are grown by `wsrf`, where binary split is used for continuous predictors (variables) and \( k \)-way split for categorical ones. For continuous predictors, each of the values themselves is used as split points, no discretization used. The only stopping condition for split is the minimum node size must not less than `nodesize`.

**Value**

An object of class `wsrf`, which is a list with the following components:

- `confusion`: the confusion matrix of the prediction (based on OOB data).
- `oob.times`: number of times cases are ‘out-of-bag’ (and thus used in computing OOB error estimate)
- `predicted`: the predicted values of the input data based on out-of-bag samples.
- `useweights`: logical. Whether weighted subspace selection is used? NULL if the model is obtained by combining multiple `wsrf` model and one of them has different value of ‘useweights’.
- `mtry`: integer. The number of variables to be chosen when splitting a node.

**Author(s)**

He Zhao and Graham Williams (SIAT, CAS)

**References**

Xu, B. and Huang, J. Z. and Williams, G. J. and Wang, Q. and Ye, Y. 2012 "Classifying very high-dimensional data with random forests built from small subspaces". *International Journal of Data Warehousing and Mining (IJDWM)*, 8(2), 44–63.


**Examples**

```
library("wsrf")

# Prepare parameters.
ds <- iris
dim(ds)
names(ds)
target <- "Species"
vars <- names(ds)
if (sum(is.na(ds[vars]))) ds[vars] <- randomForest::na.roughfix(ds[vars])
ds[target] <- as.factor(ds[[target]])
```
(tt <- table(ds[target]))
form <- as.formula(paste(target, "- "))
set.seed(42)
train <- sample(nrow(ds), 0.7*nrow(ds))
test <- setdiff(seq_len(nrow(ds)), train)

# Build model. We disable parallelism here, since CRAN Repository
# Policy (https://cran.r-project.org/web/packages/policies.html)
# limits the usage of multiple cores to save the limited resource of
# the check farm.
model.wsrf <- wsrf(form, data=ds[train, vars], parallel=FALSE)

# View model.
print(model.wsrf)
print(model.wsrf, tree=1)

# Evaluate.
strength(model.wsrf)
correlation(model.wsrf)
res <- predict(model.wsrf, newdata=ds[test, vars], type=c("response", "waprob"))
actual <- ds[test, target]
(accuracy.wsrf <- mean(res$response==actual))

# Different type of prediction.
c1 <- apply(res$waprob, 1, which.max)
c1 <- factor(c1, levels=1:ncol(res$waprob), labels=levels(actual))
(accuracy2.wsrf <- mean(c1==actual))
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