Package: C443 (via r-universe)

October 13, 2024

Type Package				
Title See a Forest for the Trees				
Version 3.4.0				
Imports MASS, partykit, rpart, RColorBrewer, grDevices, gridExtra, ggplot2, cluster, parallel, foreach, igraph, stats, graphics, plyr, ranger, randomForest, methods, doParallel				
LazyData true				
Encoding UTF-8				
Date 2023-06-21				
Description Get insight into a forest of classification trees, by calculating similarities between the trees, and subsequently clustering them. Each cluster is represented by it's most central cluster member. The package implements the methodology described in Sies & Van Mechelen (2020) <doi:10.1007 s00357-019-09350-4="">.</doi:10.1007>				
<pre>URL https://github.com/KULeuven-PPW-OKPIV/C443</pre>				
<pre>BugReports https://github.com/KULeuven-PPW-OKPIV/C443/issues</pre>				
License GPL (>= 2)				
RoxygenNote 7.2.3				
Repository https://kuleuven-ppw-okpiv.r-universe.dev				
RemoteUrl https://github.com/kuleuven-ppw-okpiv/c443				
RemoteRef HEAD				
RemoteSha acb9b60d42c560675110c9c6689e636db45e20f1				
Contents				
clusterforest				

2 clusterforest

	drugs	7
	medoidtrees	8
	medoidtrees.clusterforest	9
	medoidtrees.default	9
	plot.clusterforest	10
	print.clusterforest	11
	summary.clusterforest	12
	treesimilarities	12
	treesimilarities.clusterforest	13
	treesimilarities.default	13
	treesource	14
	treesource.clusterforest	15
	treesource.default	16
Index		17

clusterforest

Clustering the classification trees in a forest based on similarities

Description

A function to get insight into a forest of classification trees by clustering the trees in a forest using Partitioning Around Medoids (PAM, Kaufman & Rousseeuw, 2009), based on user provided similarities, or based on similarities calculated by the package using a similarity measure chosen by the user (see Sies & Van Mechelen, 2020).

Usage

```
clusterforest(
  observeddata,
  treedata = NULL,
  trees,
  simmatrix = NULL,
  m = NULL,
  tol = NULL,
  weight = NULL,
  fromclus = 1,
  toclus = 1,
  treecov = NULL,
  sameobs = FALSE,
  seed = NULL,
  no_cores = detectCores(logical = FALSE)
)
```

clusterforest 3

Arguments

observeddata The entire observed dataset

treedata A list of dataframes on which the trees are based. Not necessary if the data set

is included in the tree object already.

trees A list of trees of class party, classes inheriting from party (e.g., glmtree), classes

that can be coerced to party (i.e., rpart, Weka_tree, XMLnode), or a randomFor-

est or ranger object.

simmatrix A similarity matrix with the similarities between all trees. Should be square,

symmetric and have ones on the diagonal. Default=NULL

m Similarity measure that should be used to calculate similarities, in the case that

no similarity matrix was provided by the user. Default=NULL. m=1 is based on counting common predictors; m=2 is based on counting common predictor-split point combinations; m=3 is based on common ordered sets of predictor-range part combinations (see Shannon & Banks (1999)); m=4 is based on the agreement of partitions implied by leaf membership (Chipman, 1998); m=5 is based on the agreement of partitions implied by class labels (Chipman, 1998); m=6 is based on the number of predictor occurrences in definitions of leaves with same class label; m=7 is based on the number of predictor-split point combinations in definitions of leaves with same class label m=8 measures closeness to logical

equivalence (applicable in case of binary predictors only)

tol A vector with for each predictor a number that defines the tolerance zone within

which two split points of the predictor in question are assumed equal. For example, if the tolerance for predictor X is 1, then a split on that predictor in tree A will be assumed equal to a split in tree B as long as the splitpoint in tree B is within the splitpoint in tree A + or - 1. Only applicable for m=1 and m=6.

Default=NULL

weight If 1, the number of dissimilar paths in the Shannon and Banks measure (m=2),

should be weighted by 1/their length (Otherwise they are weighted equally).

Only applicable for m=2. Default=NULL

from Clus The lowest number of clusters for which the PAM algorithm should be run.

Default=1.

toclus The highest number of clusters for which the PAM algorithm should be run.

Default=1.

treecov A vector/dataframe with the covariate value(s) for each tree in the forest (1 col-

umn per covariate) in the case of known sources of variation underlying the

forest, that should be linked to the clustering solution.

sameobs Are the same observations included in every tree data set? For example, in the

case of subsamples or bootstrap samples, the answer is no. Default=FALSE

seed A seed number that should be used for the multi start procedure (based on which

initial medoids are assigned). Default=NULL.

no_cores Number of CPU cores used for computations. Default=detectCores(logical=FALSE)

Details

The user should provide the number of clusters that the solution should contain, or a range of numbers that should be explored. In the latter case, the resulting clusterforest object will contain

4 clusterforest

clustering results for each solution. On this clusterforest object, several methods, such as plot, print and summary, can be used.

Value

The function returns an object of class clusterforest, with attributes:

medoids the position of the medoid trees in the forest (i.e., which element of the list of

partytrees)

medoidtrees the medoid trees

clusters The cluster to which each tree in the forest is assigned

avgsilwidth The average silhouette width for each solution (see Kaufman and Rousseeuw,

2009)

accuracy For each solution, the accuracy of the predicted class labels based on the medoids.

agreement For each solution, the agreement between the predicted class label for each ob-

servation based on the forest as a whole, and those based on the medoids only

(see Sies & Van Mechelen, 2020)

withinsim Within cluster similarity for each solution (see Sies & Van Mechelen, 2020)

treesimilarities

Similarity matrix on which clustering was based

treecov covariate value(s) for each tree in the forest

seed seed number that was used for the multi start procedure (based on which initial

medoids were assigned)

References

Kaufman, L., & Rousseeuw, P. J. (2009). Finding groups in data: an introduction to cluster analysis (Vol. 344). John Wiley & Sons.

Sies, A. & Van Mechelen I. (2020). C443: An R-package to see a forest for the trees. Journal of Classification.

Shannon, W. D., & Banks, D. (1999). Combining classification trees using MLE. Statistics in medicine, 18(6), 727-740.

Chipman, H. A., George, E. I., & McCulloh, R. E. (1998). Making sense of a forest of trees. Computing Science and Statistics, 84-92.

Examples

```
require(MASS)
require(ranger)
require(rpart)
#Function to draw a bootstrap sample from a dataset
DrawBoots <- function(dataset, i){
set.seed(2394 + i)
Boot <- dataset[sample(1:nrow(dataset), size = nrow(dataset), replace = TRUE),]
return(Boot)
}</pre>
```

clusters 5

```
#Function to grow a tree using rpart on a dataset
GrowTree <- function(x,y,BootsSample, minsplit = 40, minbucket = 20, maxdepth =3){
controlrpart <- rpart.control(minsplit = minsplit, minbucket = minbucket, maxdepth = maxdepth,</pre>
 maxsurrogate = 0, maxcompete = 0)
tree <- rpart(as.formula(paste(noquote(paste(y, "~")), noquote(paste(x, collapse="+")))),</pre>
data = BootsSample, control = controlrpart)
return(tree)
#Use functions to draw 10 boostrapsamples and grow a tree on each sample
Boots<- lapply(1:10, function(k) DrawBoots(Pima.tr ,k))</pre>
Trees <- lapply(1:10, function (i) GrowTree(x=c("npreg", "glu", "bp", "skin",</pre>
"bmi", "ped", "age"), y="type", Boots[[i]] ))
#Clustering the trees in this forest
ClusterForest<- clusterforest(observeddata=Pima.tr,treedata=Boots,trees=Trees,m=1,
fromclus=1, toclus=2, sameobs=FALSE, no_cores=2)
#Example RandomForest
Pima.tr.ranger <- ranger(type ~ ., data = Pima.tr, keep.inbag = TRUE, num.trees=20,
max.depth=3)
ClusterForest<- clusterforest(observeddata=Pima.tr,trees=Pima.tr.ranger,m=5,
                           fromclus=1, toclus=2, sameobs=FALSE, no_cores=2)
```

clusters

Get the cluster assignments for a solution of a clusterforest object

Description

A function to get the cluster assignments for a given solution of a clusterforest object.

Usage

```
clusters(clusterforest, solution)
```

Arguments

clusterforest A clusterforest object

solution The solution for which cluster assignments should be returned. Default = 1

6 clusters.default

clusters.clusterforest

Get the cluster assignments for a solution of a clusterforest object

Description

A function to get the cluster assignments for a given solution of a clusterforest object.

Usage

```
## S3 method for class 'clusterforest'
clusters(clusterforest, solution = 1)
```

Arguments

clusterforest The clusterforest object

solution The solution

clusters.default

Get the cluster assignments for a solution of a clusterforest object

Description

A function to get the cluster assignments for a given solution of a clusterforest object.

Usage

```
## Default S3 method:
clusters(clusterforest, solution)
```

Arguments

clusterforest The clusterforest object

solution The solution

drugs 7

drugs

Drug consumption data set

Description

A dataset collected by Fehrman et al. (2017), freely available on the UCI Machine Learning Repository (Lichman, 2013) containing records of 1885 respondents regarding their use of 18 types of drugs, and their measurements on 12 predictors. #' All predictors were originally categorical and were quantified by Fehrman et al. (2017). The meaning of the values can be found on https://archive.ics.uci.edu/dataset/373/drug+consumption+quantified. The original response categories for each drug were: never used the drug, used it over a decade ago, or in the last decade, year, month, week, or day. We transformed these into binary response categories, where 0 (non-user) consists of the categories never used the drug and used it over a decade ago and 1 (user) consists of all other categories.

Usage

drugs

Format

A data frame with 1185 rows and 32 variables:

ID Respondent ID

Age Age of respondent

Gender Gender of respondent, where 0.48 denotes female and -0.48 denotes male

Edu Level of education of participant

Country Country of current residence of participant

Ethn Ethnicity of participant

Neuro NEO-FFI-R Neuroticism score

Extr NEO-FFI-R Extraversion score

Open NEO-FFI-R Openness to experience score

Agree NEO-FFI-R Agreeableness score

Consc NEO-FFI-R Conscientiousness score

Impul Impulsiveness score measured by BIS-11

Sensat Sensation seeking score measured by ImpSS

Alc Alcohol user (1) or non-user (0)

Amphet Amphetamine user (1) or non-user (0)

Amyl Amyl nitrite user (1) or non-user (0)

Benzos Benzodiazepine user (1) or non-user (0)

Caff Caffeine user (1) or non-user (0)

Can Cannabis user (1) or non-user (0)

8 medoidtrees

Choco Chocolate user (1) or non-user (0)

Coke Coke user (1) or non-user (0)

Crack Crack user (1) or non-user (0)

Ecst Ecstacy user (1) or non-user (0)

Her Heroin user (1) or non-user (0)

Ket Ketamine user (1) or non-user (0)

Leghighs Legal Highs user (1) or non-user (0)

LSD LSD user (1) or non-user (0)

Meth Methadone user (1) or non-user (0)

Mush Magical Mushroom user (1) or non-user (0)

Nico Nicotine user (1) or non-user (0)

Semeron Semeron user (1) or non-user (0), fictitious drug to identify over-claimers

VSA volatile substance abuse user(1) or non-user (0)

Source

https://archive.ics.uci.edu/dataset/373/drug+consumption+quantified

References

Fehrman, E., Muhammad, A. K., Mirkes, E. M., Egan, V., & Gorban, A. N. (2017). The Five Factor Model of personality and evaluation of drug consumption risk. In Data Science (pp. 231-242). Springer, Cham. Lichman, M. (2013). UCI machine learning repository.

medoidtrees

Get the medoid trees for a solution of a clusterforest object

Description

A function to get the medoid trees for a given solution of a clusterforest object.

Usage

```
medoidtrees(clusterforest, solution)
```

Arguments

clusterforest A clusterforest object

solution The solution for which medoid trees should be returned. Default = 1

medoidtrees.clusterforest 9

```
medoidtrees.clusterforest
```

Get the medoid trees for a solution of a clusterforest object

Description

A function to get the medoid trees for a given solution of a clusterforest object.

Usage

```
## S3 method for class 'clusterforest'
medoidtrees(clusterforest, solution = 1)
```

Arguments

clusterforest A clusterforest object

solution The solution for which medoid trees should be returned. Default = 1

medoidtrees.default

Get the medoid trees for a solution of a clusterforest object

Description

A function to get the medoid trees for a given solution of a clusterforest object.

Usage

```
## Default S3 method:
medoidtrees(clusterforest, solution)
```

Arguments

clusterforest A clusterforest object

solution The solution for which medoid trees should be returned. Default = 1

10 plot.clusterforest

plot.clusterforest

Plot a clusterforest object

Description

A function that can be used to plot a clusterforest object, either by returning plots with information such as average silhouette width and within cluster silliarity on the cluster solutions, or plots of the medoid trees of each solution.

Usage

```
## S3 method for class 'clusterforest'
plot(x, solution = NULL, predictive_plots = FALSE, ...)
```

Arguments

x A clusterforest object

solution

The solution to plot the medoid trees from. If NULL, plots with the average silhouette width, within cluster similarty (and predictive accuracy) per solution are returned. Default = NULL

predictive_plots

Indicating whether predictive plots should be returned: A plot showing the predictive accuracy when making predictions based on the medoid trees, and a plot of the agreement between the class label for each object predicted on the basis of the random forest as a whole versus based on the medoid trees. Default = FALSE.

Additional arguments that can be used in generic plot function, or in plot.party.

Details

This function can be used to plot a clusterforest object in two ways. If it's used without specifying a solution, then the average silhouette width, and within cluster similarity measures are plotted for each solution. If additionally, predictive_plots=TRUE, two more plots are returned, namely a plot showing for each solution the predictive accuracy when making predictions based on the medoid trees, and a plot showing for each solution the agreement between the class label for each object predicted on the basis of the random forest as a whole versus based on the medoid trees. These plots may be helpful in deciding how many clusters are needed to summarize the forest (see Sies & Van Mechelen, 2020).

If the function is used with the clusterforest object and the number of the solution, then the medoid tree(s) of that solution are plotted.

References

Sies, A. & Van Mechelen I. (2020). C443: An R-package to see a forest for the trees. Journal of Classification.

print.clusterforest 11

Examples

```
require(MASS)
require(rpart)
#Function to draw a bootstrap sample from a dataset
DrawBoots <- function(dataset, i){</pre>
set.seed(2394 + i)
Boot <- dataset[sample(1:nrow(dataset), size = nrow(dataset), replace = TRUE),]</pre>
return(Boot)
#Function to grow a tree using rpart on a dataset
GrowTree <- function(x,y,BootsSample, minsplit = 40, minbucket = 20, maxdepth =3){
 controlrpart <- rpart.control(minsplit = minsplit, minbucket = minbucket,</pre>
 maxdepth = maxdepth, maxsurrogate = 0, maxcompete = 0)
 tree <- rpart(as.formula(paste(noquote(paste(y, "~")),</pre>
 noquote(paste(x, collapse="+")))), data = BootsSample,
 control = controlrpart)
 return(tree)
#Use functions to draw 20 boostrapsamples and grow a tree on each sample
Boots<- lapply(1:10, function(k) DrawBoots(Pima.tr ,k))</pre>
Trees <- lapply(1:10, function (i) GrowTree(x=c("npreg", "glu", "bp",</pre>
 "skin", "bmi", "ped", "age"), y="type",
Boots[[i]] ))
ClusterForest<- clusterforest(observeddata=Pima.tr,treedata=Boots,trees=Trees,m=1,
fromclus=1, toclus=5, sameobs=FALSE, no_cores=2)
plot(ClusterForest)
plot(ClusterForest,2)
```

print.clusterforest

Print a clusterforest object

Description

A function that can be used to print a clusterforest object.

Usage

```
## S3 method for class 'clusterforest'
print(x, solution = 1, ...)
```

Arguments

x A clusterforest object
 solution The solution to print the medoid trees from. Default = NULL
 ... Additional arguments that can be used in the generic print function.

12 treesimilarities

```
summary.clusterforest Summarize a clusterforest object
```

Description

A function to summarize a clusterforest object.

Usage

```
## S3 method for class 'clusterforest'
summary(object, ...)
```

Arguments

object A clusterforest object

... Additional arguments that can be used in the generic summary function.

treesimilarities

Get the similarity matrix that wast used to create a clusterforest object

Description

A function to get the similarity matrix used to obtain a clusterforest object.

Usage

```
treesimilarities(clusterforest)
```

Arguments

clusterforest A clusterforest object

treesimilarities.clusterforest 13

```
treesimilarities.clusterforest
```

Get the similarity matrix that wast used to create a clusterforest object

Description

A function to get the similarity matrix used to obtain a clusterforest object.

Usage

```
## S3 method for class 'clusterforest'
treesimilarities(clusterforest)
```

Arguments

clusterforest A clusterforest object

treesimilarities.default

Get the similarity matrix that wast used to create a clusterforest object

Description

A function to get the similarity matrix used to obtain a clusterforest object.

Usage

```
## Default S3 method:
treesimilarities(clusterforest)
```

Arguments

clusterforest A clusterforest object

14 treesource

treesource	Mapping the tree clustering solution to a known source of variation underlying the forest

Description

A function that can be used to get insight into a clusterforest solution, in the case that there are known sources of variation underlying the forest. These known sources of variation must be included in the clusterforest object (and thus must be defined when running the clusterforest function) In case of a categorical covariate, it visualizes the number of trees from each value of the covariate that belong to each cluster. In case of a continuous covariate, it returns the mean and standard deviation of the covariate in each cluster.

Usage

```
treesource(clusterforest, solution)
```

Arguments

clusterforest The clusterforest object, indluding the treecov attribute.

solution The solution

Value

multiplot In case of categorical covariate, for each value of the covariate, a bar plot with the number of trees that belong to each cluster

heatmap In case of a categorical covariate, a heatmap with for each value of the covariate, the number of trees that belong to each cluster

clustermeans In case of a continuous covariate, the mean of the covariate in each cluster

clusterstds In case of a continuous covariate, the standard deviation of the covariate in each cluster

Examples

treesource.clusterforest 15

```
#Function to grow a tree using rpart on a dataset
GrowTree <- function(x,y,BootsSample, minsplit = 40, minbucket = 20, maxdepth =3){
controlrpart <- rpart.control(minsplit = minsplit, minbucket = minbucket, maxdepth = maxdepth,</pre>
 maxsurrogate = 0, maxcompete = 0)
tree <- rpart(as.formula(paste(noquote(paste(y, "~")), noquote(paste(x, collapse="+")))),</pre>
  data = BootsSample, control = controlrpart)
 return(tree)
#Draw bootstrap samples and grow trees
BootsA<- lapply(1:5, function(k) DrawBoots(data_Amphet,k))</pre>
BootsC<- lapply(1:5, function(k) DrawBoots(data_cocaine,k))</pre>
Boots = c(BootsA,BootsC)
TreesA <- lapply(1:5, function (i) GrowTree(x=c ("Age", "Gender", "Edu", "Neuro",</pre>
"Extr", "Open", "Agree", "Consc", "Impul", "Sensat"), y="Amphet", BootsA[[i]] ))
TreesC <- lapply(1:5, function (i) GrowTree(x=c ( "Age", "Gender", "Edu", "Neuro",
"Extr", "Open", "Agree", "Consc", "Impul", "Sensat"), y="Coke", BootsC[[i]] ))
Trees=c(TreesA,TreesC)
#Cluster the trees
ClusterForest<- clusterforest(observeddata=drugs,treedata=Boots,trees=Trees,m=1,
fromclus=2, toclus=2, treecov=rep(c("Amphet","Coke"),each=5), sameobs=FALSE, no_cores=2)
#Link cluster result to known source of variation
treesource(ClusterForest, 2)
```

treesource.clusterforest

Mapping the tree clustering solution to a known source of variation underlying the forest

Description

A function that can be used to get insight into a clusterforest solution, in the case that there is a known source of variation underlying the forest. It visualizes the number of trees from each source that belong to each cluster.

Usage

```
## S3 method for class 'clusterforest'
treesource(clusterforest, solution)
```

Arguments

clusterforest The clusterforest object solution The solution

16 treesource.default

treesource.default Mapping the tree underlying the for	clustering solution to a known source of variation est
--	---

Description

A function that can be used to get insight into a clusterforest solution, in the case that there is a known source of variation underlying the forest. It visualizes the number of trees from each source that belong to each cluster.

Usage

```
## Default S3 method:
treesource(clusterforest, solution)
```

Arguments

clusterforest The clusterforest object

solution The solution

Index

```
\ast datasets
    drugs, 7
clusterforest, 2
clusters, 5
clusters.clusterforest, 6
clusters.default, 6
drugs, 7
medoidtrees, 8
medoidtrees.clusterforest, 9
{\tt medoidtrees.default,9}
\verb|plot.clusterforest|, 10
print.clusterforest, 11
summary.clusterforest, 12
treesimilarities, 12
tree similarities. cluster for est, \\ 13
treesimilarities.default, 13
treesource, 14
treesource.clusterforest, 15
treesource.default, 16
```