Package 'diversityForest'

June 9, 2022

Type Package

Title Innovative Complex Split Procedures in Random Forests Through Candidate Split Sampling

Version 0.3.4

Date 2022-06-08

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Description Implements interaction forests [1], which are specific diversity forests and the basic form of diversity forests that uses univariable, binary splitting [2]. Interaction forests (IFs) are ensembles of decision trees that model quantitative and qualitative interaction effects using bivariable splitting. IFs come with the Effect Importance Measure (EIM), which can be used to identify variable pairs that feature quantitative and qualitative interaction effects with high predictive relevance. IFs and EIM focus on well interpretable forms of interactions. The package also offers plot functions for visualising the estimated forms of interaction effects.

Categorical, metric, and survival outcomes are supported.

This is a fork of the R package 'ranger' (main author: Marvin N. Wright) that implements random forests using an efficient C++ implementation. References:

[1] Hornung, R. & Boulesteix, A.-L. (2022) Interaction Forests: Identifying and exploiting interpretable quantitative and qualitative interaction effects.

Computational Statistics & Data Analysis 171:107460, <doi:10.1016/j.csda.2022.107460>.

[2] Hornung, R. (2022) Diversity forests: Using split sampling to enable innovative complex split procedures in random forests.

SN Computer Science 3(2):1, <doi:10.1007/s42979-021-00920-1>.

SystemRequirements C++11

Encoding UTF-8

License GPL-3

Imports Rcpp (>= 0.11.2), Matrix, ggplot2, ggpubr, scales, nnet, sgeostat, rms, MapGAM, gam, rlang, grDevices, RColorBrewer, RcppEigen, survival

LinkingTo Rcpp, RcppEigen

Depends R (>= 3.5)

Suggests testthat, BOLTSSIRR

Additional_repositories https://romanhornung.github.io/drat

RoxygenNote 7.1.1 NeedsCompilation yes Repository CRAN

Date/Publication 2022-06-09 09:10:02 UTC

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diversityForest-package

Diversity Forests

Description

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The diversity forest algorithm is not a specific algorithm, but an alternative candidate split sampling scheme that makes complex split procedures in random forests possible computationally by drastically reducing the numbers of candidate splits that need to be evaluated for each split. It also avoids the well-known variable selection bias in conventional random forests that has the effect that variables with many possible splits are selected too frequently for splitting (Strobl et al., 2007). For details, see Hornung (2022).

Details

This package currently features two types of diversity forests:

• the basic form of diversity forests that uses univariable, binary splitting, which is also used in conventional random forests

• interaction forests (IFs) (Hornung & Boulesteix, 2022), which use bivariable splitting to model quantitative and qualitative interaction effects. IFs feature the Effect Importance Measure (EIM), which ranks the variable pairs with respect to the predictive importance of their quantitative and qualitative interaction effects. The individual variables can be ranked as well using EIM. For details, see Hornung & Boulesteix (2022).

Diversity forests with univariable splitting can be constructed using the function divfor and interaction forests using the function interactionfor. Both functions support categorical, metric, and survival outcomes.

This package is a fork of the R package 'ranger' that implements random forests using an efficient C++ implementation. The documentation is in large parts taken from 'ranger', where some parts of the documentation may not apply to (the current version of) the 'diversityForest' package.

Details on further functionalities of the code that are not presented in the help pages of 'diversity-Forest' are found in the help pages of 'ranger', version 0.11.0, because 'diversity-Forest' is based on the latter version of 'ranger'. The code in the example sections can be used as a template for all basic application scenarios with respect to classification, regression and survival prediction.

References

- Hornung, R. (2022). Diversity forests: Using split sampling to enable innovative complex split procedures in random forests. SN Computer Science 3(2):1, <doi: 10.1007/s42979021-009201>.
- Hornung, R., Boulesteix, A.-L. (2022). Interaction forests: Identifying and exploiting interpretable quantitative and qualitative interaction effects. Computational Statistics & Data Analysis 171:107460, <doi: 10.1016/j.csda.2022.107460>.
- Strobl, C., Boulesteix, A.-L., Zeileis, A., Hothorn, T. (2007). Bias in random forest variable importance measures: Illustrations, sources and a solution. BMC Bioinformatics 8:25, <doi: 10.1186/14712105825>.
- Wright, M. N., Ziegler, A. (2017). ranger: A fast Implementation of Random Forests for High Dimensional Data in C++ and R. Journal of Statistical Software 77:1-17, <doi: 10.18637/jss.v077.i01>.

divfor

Construct a basic diversity forest prediction rule that uses univariable, binary splitting.

Description

Implements the most basic form of diversity forests that uses univariable, binary splitting. Currently, categorical, metric, and survival outcomes are supported.

Usage

```
divfor(
  formula = NULL,
  data = NULL,
  num.trees = 500,
 mtry = NULL,
  importance = "none",
 write.forest = TRUE,
 probability = FALSE,
 min.node.size = NULL,
 max.depth = NULL,
  replace = TRUE,
  sample.fraction = ifelse(replace, 1, 0.632),
  case.weights = NULL,
  class.weights = NULL,
  splitrule = NULL,
  num.random.splits = 1,
  alpha = 0.5,
 minprop = 0.1,
  split.select.weights = NULL,
  always.split.variables = NULL,
  respect.unordered.factors = NULL,
  scale.permutation.importance = FALSE,
  keep.inbag = FALSE,
  inbag = NULL,
  holdout = FALSE,
  quantreg = FALSE,
  oob.error = TRUE,
  num.threads = NULL,
  save.memory = FALSE,
  verbose = TRUE,
  seed = NULL,
  dependent.variable.name = NULL,
  status.variable.name = NULL,
  classification = NULL,
  nsplits = 30,
 proptry = 1
)
```

Arguments

formula	Object of class formula or character describing the model to fit. Interactio terms supported only for numerical variables.	
data	Training data of class data.frame, matrix, dgCMatrix (Matrix) or gwaa.data (GenABEL).	
num.trees	s Number of trees. Default is 500.	
mtry	Artefact from 'ranger'. NOT needed for diversity forests.	

importance Variable importance mode, one of 'none', 'impurity', 'impurity_corrected', 'per-

mutation'. The 'impurity' measure is the Gini index for classification, the variance of the responses for regression and the sum of test statistics (see splitrule) for survival. NOTE: Currently, only "permutation" (and "none") work for diver-

sity forests.

write.forest Save divfor.forest object, required for prediction. Set to FALSE to reduce

memory usage if no prediction intended.

probability Grow a probability forest as in Malley et al. (2012). NOTE: Not yet imple-

mented for diversity forests!

min.node.size Minimal node size. Default 1 for classification, 5 for regression, 3 for survival,

and 5 for probability.

max.depth Maximal tree depth. A value of NULL or 0 (the default) corresponds to unlim-

ited depth, 1 to tree stumps (1 split per tree).

replace Sample with replacement.

sample.fraction

Fraction of observations to sample. Default is 1 for sampling with replacement and 0.632 for sampling without replacement. For classification, this can be a

vector of class-specific values.

case.weights Weights for sampling of training observations. Observations with larger weights

will be selected with higher probability in the bootstrap (or subsampled) samples

for the trees.

class.weights Weights for the outcome classes (in order of the factor levels) in the splitting

rule (cost sensitive learning). Classification and probability prediction only. For classification the weights are also applied in the majority vote in terminal nodes.

splittrule Splitting rule. For classification and probability estimation "gini" or "extratrees"

with default "gini". For regression "variance", "extratrees" or "maxstat" with default "variance". For survival "logrank", "extratrees", "C" or "maxstat" with default "logrank". NOTE: For diversity forests currently only the default splitting

rules are supported.

num.random.splits

Artefact from 'ranger'. NOT needed for diversity forests.

alpha For "maxstat" splitrule: Significance threshold to allow splitting. NOT needed

for diversity forests.

minprop For "maxstat" splitrule: Lower quantile of covariate distribution to be considered

for splitting. NOT needed for diversity forests.

split.select.weights

Numeric vector with weights between 0 and 1, representing the probability to select variables for splitting. Alternatively, a list of size num.trees, containing

split select weight vectors for each tree can be used.

always.split.variables

Currently not useable. Character vector with variable names to be always se-

lected.

respect.unordered.factors

Handling of unordered factor covariates. One of 'ignore' and 'order' (the option 'partition' possible in 'ranger' is not (yet) possible with diversity forests). Default is 'ignore'. Alternatively TRUE (='order') or FALSE (='ignore') can be

used.

scale.permutation.importance

Scale permutation importance by standard error as in (Breiman 2001). Only

applicable if permutation variable importance mode selected.

keep. inbag Save how often observations are in-bag in each tree.

inbag Manually set observations per tree. List of size num.trees, containing inbag

counts for each observation. Can be used for stratified sampling.

holdout Hold-out mode. Hold-out all samples with case weight 0 and use these for

variable importance and prediction error.

quantreg Prepare quantile prediction as in quantile regression forests (Meinshausen 2006).

Regression only. Set keep. inbag = TRUE to prepare out-of-bag quantile predic-

tion.

oob.error Compute OOB prediction error. Set to FALSE to save computation time, e.g. for

large survival forests.

num. threads Number of threads. Default is number of CPUs available.

save.memory Use memory saving (but slower) splitting mode. No effect for survival and

GWAS data. Warning: This option slows down the tree growing, use only if you

encounter memory problems. NOT needed for diversity forests.

verbose Show computation status and estimated runtime.

seed Random seed. Default is NULL, which generates the seed from R. Set to 0 to

ignore the R seed.

dependent.variable.name

Name of outcome variable, needed if no formula given. For survival forests this

is the time variable.

status.variable.name

Name of status variable, only applicable to survival data and needed if no for-

mula given. Use 1 for event and 0 for censoring.

classification Only needed if data is a matrix. Set to TRUE to grow a classification forest.

nsplits Number of candidate splits to sample for each split. Default is 30.

proptry Parameter that restricts the number of candidate splits considered for small

nodes. If nsplits is larger than proptry times the number of all possible splits, the number of candidate splits to draw is reduced to the largest integer smaller than proptry times the number of all possible splits. Default is 1, which corre-

sponds to always using nsplits candidate splits.

Value

Object of class divfor with elements

forest Saved forest (If write forest set to TRUE). Note that the variable IDs in the

split.varIDs object do not necessarily represent the column number in R.

predictions Predicted classes/values, based on out-of-bag samples (classification and regres-

sion only).

variable.importance

Variable importance for each independent variable.

prediction.error

Overall out-of-bag prediction error. For classification this is the fraction of missclassified samples, for probability estimation the Brier score, for regression the mean squared error and for survival one minus Harrell's C-index.

r.squared

R squared. Also called explained variance or coefficient of determination (regression only). Computed on out-of-bag data.

confusion.matrix

Contingency table for classes and predictions based on out-of-bag samples (classification only).

unique.death.times

Unique death times (survival only).

chf Estimated cumulative hazard function for each sample (survival only).

survival Estimated survival function for each sample (survival only).

call Function call.
num.trees Number of trees.
num.independent.variables

Number of independent variables.

min.node.size Value of minimal node size used.

treetype Type of forest/tree. classification, regression or survival.

importance.mode

Importance mode used.

 ${\color{blue} \textbf{num.samples}} \qquad \qquad Number\ of\ samples.$

splitrule Splitting rule.

replace Sample with replacement.

nsplits Value of nsplits used.

proptry Value of proptry used.

Author(s)

Roman Hornung, Marvin N. Wright

References

- Hornung, R. (2022). Diversity forests: Using split sampling to enable innovative complex split procedures in random forests. SN Computer Science 3(2):1, <doi: 10.1007/s42979021-009201>.
- Wright, M. N., Ziegler, A. (2017). ranger: A fast implementation of random forests for high dimensional data in C++ and R. Journal of Statistical Software 77:1-17, <doi: 10.18637/jss.v077.i01>.
- Breiman, L. (2001). Random forests. Machine Learning 45:5-32, <doi: 10.1023/A:1010933404324>.
- Malley, J. D., Kruppa, J., Dasgupta, A., Malley, K. G., & Ziegler, A. (2012). Probability machines: consistent probability estimation using nonparametric learning machines. Methods of Information in Medicine 51:74-81, <doi: 10.3414/ME00010052>.
- Meinshausen (2006). Quantile Regression Forests. Journal of Machine Learning Research 7:983-999.

See Also

```
predict.divfor
```

Examples

```
## Not run:
## Load package:
library("diversityForest")
## Set seed to obtain reproducible results:
set.seed(1234)
## Diversity forest with default settings (NOT recommended)
# Classification:
divfor(Species ~ ., data = iris, num.trees = 20)
# Regression:
iris2 <- iris; iris2$Species <- NULL; iris2$Y <- rnorm(nrow(iris2))</pre>
divfor(Y ~ ., data = iris2, num.trees = 20)
# Survival:
library("survival")
divfor(Surv(time, status) ~ ., data = veteran, num.trees = 20, respect.unordered.factors = "order")
# NOTE: num.trees = 20 is specified too small for practical
# purposes - the prediction performance of the resulting
# forest will be suboptimal!!
# In practice, num.trees = 500 (default value) or a
# larger number should be used.
## Diversity forest with specified values for nsplits and proptry (NOT recommended)
divfor(Species ~ ., data = iris, nsplits = 10, proptry = 0.4, num.trees = 20)
# NOTE again: num.trees = 20 is specified too small for practical purposes.
## Applying diversity forest after optimizing the values of nsplits and proptry (recommended)
tuneres <- tunedivfor(formula = Species ~ ., data = iris, num.trees.pre = 20)
# NOTE: num.trees.pre = 20 is specified too small for practical
# purposes - the out-of-bag error estimates of the forests
# constructed during optimization will be much too variable!!
# In practice, num.trees.pre = 500 (default value) or a
# larger number should be used.
divfor(Species ~ ., data = iris, nsplits = tuneres$nsplitsopt,
 proptry = tuneres$proptryopt, num.trees = 20)
# NOTE again: num.trees = 20 is specified too small for practical purposes.
## Prediction
train.idx <- sample(nrow(iris), 2/3 * nrow(iris))</pre>
iris.train <- iris[train.idx, ]</pre>
iris.test <- iris[-train.idx, ]</pre>
tuneres <- tunedivfor(formula = Species ~ ., data = iris.train, num.trees.pre = 20)
# NOTE again: num.trees.pre = 20 is specified too small for practical purposes.
rg.iris <- divfor(Species ~ ., data = iris.train, nsplits = tuneres$nsplitsopt,</pre>
 proptry = tuneres$proptryopt, num.trees = 20)
# NOTE again: num.trees = 20 is specified too small for practical purposes.
```

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```
pred.iris <- predict(rg.iris, data = iris.test)
table(iris.test$Species, pred.iris$predictions)

## Variable importance
rg.iris <- divfor(Species ~ ., data = iris, importance = "permutation", num.trees = 20)
# NOTE again: num.trees = 20 is specified too small for practical purposes.
rg.iris$variable.importance

## End(Not run)</pre>
```

importance.divfor

Diversity Forest variable importance

Description

Extract variable importance of divfor object.

Usage

```
## S3 method for class 'divfor'
importance(x, ...)
```

Arguments

x divfor object.

... Further arguments passed to or from other methods.

Value

Variable importance measures.

Author(s)

Marvin N. Wright

See Also

divfor

interactionfor

Construct an interaction forest prediction rule and calculate EIM values as described in Hornung & Boulesteix (2022).

Description

Implements interaction forests as described in Hornung & Boulesteix (2022). Currently, categorical, metric, and survival outcomes are supported. Interaction forests feature the effect importance measure (EIM), which can be used to rank the covariate variable pairs with respect to the impact of their interaction effects on prediction. This allows to identify relevant interaction effects. Interaction forests focus on well interpretable interaction effects. See the 'Details' section below for more details.

Usage

```
interactionfor(
  formula = NULL,
  data = NULL,
  importance = "both",
  num.trees = NULL,
  simplify.large.n = TRUE,
  num.trees.eim.large.n = NULL,
 write.forest = TRUE,
  probability = FALSE,
 min.node.size = NULL,
 max.depth = NULL,
  replace = FALSE,
  sample.fraction = ifelse(replace, 1, 0.7),
  case.weights = NULL,
  class.weights = NULL,
  splitrule = NULL,
  always.split.variables = NULL,
  keep.inbag = FALSE,
  inbag = NULL,
  holdout = FALSE,
  quantreg = FALSE,
  oob.error = TRUE,
  num.threads = NULL,
  verbose = TRUE,
  seed = NULL.
  dependent.variable.name = NULL,
  status.variable.name = NULL,
  npairs = NULL,
  classification = NULL
)
```

Arguments

formula Object of class formula or character describing the model to fit.

data Training data of class data.frame, matrix, dgCMatrix (Matrix) or gwaa.data

(GenABEL).

importance Effect importance mode. One of the following: "both" (the default), "qualita-

tive", "quantitative", "mainonly", "none". See the 'Details' section below for

explanation.

num. trees Number of trees. The default number is 20000, if EIM values should be com-

puted and 2000 otherwise. Note that if simplify.large.n = TRUE (default), the number of observations is larger than 1000, and EIM values should be calculated two forests are constructed, one for calculating the EIM values and one for prediction (cf. 'Details' section). In such cases, the default number of trees used for the forest for EIM value calculation is 20000 and the default number of

trees used for the forest for prediction is 2000.

simplify.large.n

Should restricted tree depths be used, when calculating EIM values for large data sets? See the 'Details' section below for more information. Default is TRUE.

num.trees.eim.large.n

Number of trees in the forest used for calculating the EIM values for large data sets. If num.trees is provided, but not num.trees.eim.large.n, the value given by num.trees will be used. The default number is 20000. Only used

when simplify.large.n = TRUE.

write.forest Save interaction.forest object, required for prediction. Set to FALSE to re-

duce memory usage if no prediction intended.

probability Grow a probability forest as in Malley et al. (2012).

min.node.size Minimal node size. Default 1 for classification, 5 for regression, 3 for survival,

and 5 for probability.

max.depth Maximal tree depth. A value of NULL or 0 (the default) corresponds to unlim-

ited depth, 1 to tree stumps (1 split per tree).

replace Sample with replacement. Default is FALSE.

sample.fraction

Fraction of observations to sample. Default is 1 for sampling with replacement and 0.7 for sampling without replacement. For classification, this can be a vector

of class-specific values.

case .weights Weights for sampling of training observations. Observations with larger weights

will be selected with higher probability in the bootstrap (or subsampled) samples

for the trees.

class.weights Weights for the outcome classes (in order of the factor levels) in the splitting

rule (cost sensitive learning). Classification and probability prediction only. For classification the weights are also applied in the majority vote in terminal nodes.

splitrule Splitting rule. For classification and probability estimation "gini" or "extratrees"

with default "gini". For regression "variance", "extratrees" or "maxstat" with default "variance". For survival "logrank", "extratrees", "C" or "maxstat" with default "logrank". NOTE: For interaction forests currently only the default split-

ting rules are supported.

always.split.variables

Currently not useable. Character vector with variable names to be always se-

lected.

keep. inbag Save how often observations are in-bag in each tree.

inbag Manually set observations per tree. List of size num.trees, containing inbag

counts for each observation. Can be used for stratified sampling.

holdout Hold-out mode. Hold-out all samples with case weight 0 and use these for vari-

able importance and prediction error. NOTE: Currently, not useable for interac-

tion forests.

quantreg Prepare quantile prediction as in quantile regression forests (Meinshausen 2006).

Regression only. Set keep. inbag = TRUE to prepare out-of-bag quantile predic-

tion. NOTE: Currently, not useable for interaction forests.

oob.error Compute OOB prediction error. Set to FALSE to save computation time, e.g. for

large survival forests.

num. threads Number of threads. Default is number of CPUs available.

verbose Show computation status and estimated runtime.

seed Random seed. Default is NULL, which generates the seed from R. Set to 0 to

ignore the R seed.

dependent.variable.name

Name of outcome variable, needed if no formula given. For survival forests this

is the time variable.

status.variable.name

Name of status variable, only applicable to survival data and needed if no for-

mula given. Use 1 for event and 0 for censoring.

npairs Number of variable pairs to sample for each split. Default is the square root of

the number of independent variables divided by 2 (this number is rounded up).

classification Only needed if data is a matrix. Set to TRUE to grow a classification forest.

Details

The effect importance measure (EIM) of interaction forests distinguishes quantitative and qualitative interaction effects (Peto, 1982). This is a common distinction as these two types of interaction effects are interpreted in different ways (see below). For both of these types, EIM values for each variable pair are obtained: the quantitative and qualitative EIM values.

Interaction forests target easily interpretable types of interaction effects. These can be communicated clearly using statements of the following kind: "The strength of the positive (negative) effect of variable A on the outcome depends on the level of variable B" for quantitative interactions, and "for observations with small values of variable B, the effect of variable A is positive (negative), but for observations with large values of B, the effect of A is negative (positive)" for qualitative interactions.

In addition to calculating EIM values for variable pairs, importance values for the individual variables are calculated as well, the univariable EIM values. These measure the variable importance as in the case of classical variable importance measures of random forests.

The effect importance mode can be set via the importance argument: "qualitative": Calculate only qualitative EIM values; "quantitative": Calculate only quantitative EIM values; "both"

(the default): Calculate qualitative and quantitative EIM values; "mainonly": Calculate only univariable EIM values.

The top variable pairs with largest quantitative and qualitative EIM values likely have quantitative and qualitative interactions, respectively, which have a considerable impact on prediction. The top variables with largest univariable EIM values likely have a considerable impact on prediction.

If the number of variables is larger than 100, not all possible variable pairs are considered, but, using a screening procedure, the 5000 variable pairs with the strongest indications of interaction effects are pre-selected.

NOTE: To make interpretations, it is crucial to investigate (visually) the forms the interaction effects of variable pairs with large quantitative and qualitative EIM values take. This can be done using the plot function plot.interactionfor (first overview) and plotEffects.

NOTE ALSO: As described in Hornung & Boulesteix (2022), in the case of data with larger numbers of variables (larger than 100, but more seriously for high-dimensional data), the univariable EIM values can be biased. Therefore, it is strongly recommended to interpret the univariable EIM values with caution, if the data are high-dimensional. If it is of interest to measure the univariable importance of the variables for high-dimensional data, an additional conventional random forest (e.g., using the ranger package) should be constructed and the variable importance measure values of this random forest be used for ranking the univariable effects.

For large data sets with many observations the calculation of the EIM values can become very costly - when using fully grown trees. Therefore, when calculating EIM values for data sets with more than 1000 observations we use the following maximum tree depths by default (argument: simplify.large.n = TRUE):

- if n < 1000: Use fully grown trees.
- if $1000 < n \le 2000$: Use tree depth 10.
- if $2000 < n \le 5000$: Use tree depth 7.
- if n > 5000: Use tree depth 5.

Extensive analyses in Hornung & Boulesteix (2022) suggest that by restricting the tree depth in this way, the EIM values that would result when using fully grown trees are approximated well. However, the prediction performance suffers, when using restricted trees. Therefore, we restrict the tree depth only when calculating the EIM values (if n>1000), but construct a second interaction forest with unrestricted tree depth, which is then used for prediction purposes.

Value

Object of class interactionfor with elements

predictions Predicted classes/values, based on out-of-bag samples (classification and regres-

sion only).

Number of independent variables.

unique.death.times

Unique death times (survival only).

min.node.size Value of minimal node size used.

npairs Number of variable pairs sampled for each split.

eim.univ.sorted

Univariable EIM values sorted in decreasing order.

eim.univ

Univariable EIM values.

eim.qual.sorted

Qualitative EIM values sorted in decreasing order.

eim.qual

Qualitative EIM values.

eim.quant.sorted

Quantitative EIM values sorted in decreasing order.

The labeling of these values provides the information on the type of quantitative interactions the respective variable pairs feature. For example, consider a variable pair A and B and say the label reads "A large AND B small". This would mean that if the value of A is large and, at the same time, the value of B is small, the expected value of the outcome variable is (considerably) different from all other cases. For this type of quantitative interaction, the effect of B is weak for small values of A and strong for large values of B. See Hornung & Boulesteix (2022) for more information on the types of quantitative interaction effects targeted by interaction forest.

eim.quant

Quantitative EIM values. These values are labeled analoguously as those in eim.quant.sorted.

prediction.error

Overall out-of-bag prediction error. For classification this is the fraction of misclassified samples, for probability estimation the Brier score, for regression the mean squared error and for survival one minus Harrell's C-index. This is 'NA' for data sets with more than 100 covariate variables, because for such data sets we pre-select the 5000 variable pairs with strongest indications of interaction effects. This pre-selection cannot be taken into account in the out-of-bag error estimation, which is why the out-of-bag error estimates would be (much) too optimistic for data sets with more than 100 covariate variables.

forest

Saved forest (If write.forest set to TRUE). Note that the variable IDs in the split.multvarIDs object do not necessarily represent the column number in R.

confusion.matrix

Contingency table for classes and predictions based on out-of-bag samples (classification only).

chf Estimated cumulative hazard function for each sample (survival only).

survival Estimated survival function for each sample (survival only).

splitrule Splitting rule.

treetype Type of forest/tree. classification, regression or survival.

r.squared R squared. Also called explained variance or coefficient of determination (re-

gression only). Computed on out-of-bag data.

call Function call.

importance.mode

Importance mode used.

num.samples Number of samples.

replace Sample with replacement.

eim.quant.rawlists

List containing the four vectors of un-adjusted 'raw' quantitative EIM values and the four vectors of adjusted EIM values. These are usually not required by the user.

For each of the four types of quantitative splits there exists a separate vector of raw quantitative EIM values. For example, eim.quant.large.small.raw contains the raw quantitative EIM values of the quantitative split type associated with quantitative interaction effects for which the expected values of the outcome variable are different, if the value of variable A is large and, at the same time, the value of variable B is small. The list entries of the un-adjusted 'raw' quantitative EIM values are labeled with the suffix .raw, while the list entries of the adjusted quantitative EIM values miss this suffix. See Hornung & Boulesteix (2022) for details on the raw and adjusted EIM values.

promispairs

List giving the indices of the variables in the pre-selected variable pairs. If the number of variables is at most 100, all variable pairs are considered.

plotres

List ob objects needed by the plot functions: eim.univ.order contains the sorting of the univariable EIM values in descending order, where the first element gives the index of the variable with largest EIM value, the second element the index of the variable with second-largest EIM value and so on; eim.qual.order / eim.quant.order contains the sorting in descending order of the qualitative / quantitative EIM values for the (pre-selected) variable pairs given by the object promispairs above. The first element gives the index of the (pre-selected) variable pair with largest qualitative / quantitative EIM value, the second element the index of the variable pair with second-largest qualitative / quantitative EIM value; data contains the data; yvarname is the name of the outcome variable (survival time for survival); statusvarname is the name of the status variable.

Author(s)

Roman Hornung, Marvin N. Wright

References

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See Also

```
predict.divfor, plot.interactionfor, plotEffects
```

Examples

```
## Not run:
## Load package:
library("diversityForest")
## Set seed to make results reproducible:
set.seed(1234)
## Construct interaction forests and calculate EIM values:
# Binary outcome:
data(zoo)
modelcat <- interactionfor(dependent.variable.name = "type", data = zoo,</pre>
  num.trees = 20)
# Metric outcome:
data(stock)
modelcont <- interactionfor(dependent.variable.name = "company10", data = stock,</pre>
  num.trees = 20)
# Survival outcome:
library("survival")
mgus2$id <- NULL # 'mgus2' data set is contained in the 'survival' package
# categorical variables need to be of factor format - important!!
mgus2$sex <- factor(mgus2$sex)</pre>
mgus2$pstat <- factor(mgus2$pstat)</pre>
# Remove the second time variable 'ptime':
mgus2$ptime <- NULL
# Remove missing values:
mgus2 <- mgus2[complete.cases(mgus2),]</pre>
# Take subset to make the calculations less computationally
```

```
# expensive for the example (in actual applications, we would of course
# use the whole data set):
mgus2sub <- mgus2[sample(1:nrow(mgus2), size=500),]</pre>
# Apply 'interactionfor':
modelsurv <- interactionfor(formula = Surv(futime, death) ~ ., data=mgus2sub, num.trees=20)</pre>
# NOTE: num.trees = 20 (in the above) would be much too small for practical
# purposes. This small number of trees was simply used to keep the
# runtime of the example short.
# The default number of trees is num.trees = 20000 if EIM values are calculated
# and num.trees = 2000 otherwise.
## Inspect the rankings of the variables and variable pairs with respect to
## the univariable, quantitative, and qualitative EIM values:
# Univariable EIM values:
modelcat$eim.univ.sorted
# Pairs with top quantitative EIM values:
modelcat$eim.quant.sorted[1:5]
# Pairs with top qualitative EIM values:
modelcat$eim.qual.sorted[1:5]
## Investigate visually the forms of the interaction effects of the variable pairs with
## largest quantitative and qualitative EIM values:
plot(modelcat)
plotEffects(modelcat, type="quant") # type="quant" is default.
plotEffects(modelcat, type="qual")
## Prediction:
# Separate 'zoo' data set randomly in training
# and test data:
data(zoo)
train.idx <- sample(nrow(zoo), 2/3 * nrow(zoo))</pre>
zoo.train <- zoo[train.idx, ]</pre>
zoo.test <- zoo[-train.idx, ]</pre>
# Construct interaction forest on training data:
# NOTE again: num.trees = 20 is specified too small for practical purposes.
modelcattrain <- interactionfor(dependent.variable.name = "type", data = zoo,</pre>
                                importance = "none", num.trees = 20)
# NOTE: Because we are only interested in prediction here, we do not
```

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```
# calculate EIM values (by setting importance = "none"), because this
# speeds up calculations.

# Predict class values of the test data:
pred.zoo <- predict(modelcattrain, data = zoo.test)

# Compare predicted and true class values of the test data:
table(zoo.test$type, pred.zoo$predictions)

## End(Not run)</pre>
```

plot.interactionfor

Plot method for interaction for objects

Description

Plot function for interaction objects that allows to obtain a first overview of the result of the interaction forest analysis. This function visualises the distributions of the EIM values and the estimated forms of the bivariable influences of the variable pairs with largest quantitative and qualitative EIM values. Further visual exploration of the result of the interaction forest analysis should be conducted using plotEffects.

Usage

```
## S3 method for class 'interactionfor'
plot(x, numpairsquant = 2, numpairsqual = 2, ...)
```

Arguments

x Object of class interactionfor.

numpairsquant The number of pairs with largest quantitative EIM values to plot. Default is 2.

The number of pairs with largest qualitative EIM values to plot. Default is 2.

... Further arguments passed to or from other methods.

Details

For details on the plots of the estimated forms of the bivariable influences of the variable pairs see plotEffects.

NOTE: The p-values shown in the plots are generally much too optimistic and **MUST NOT** be reported as the result of a statistical test for significance of interaction. To obtain adjusted p-values that would correspond to valid tests, it would be possible to multiply these p-values by the number of possible variable pairs, which would correspond to Bonferroni-adjusted p-values. See the 'Details' section of plotEffects for further explanation and guidance.

NOTE ALSO: As described in Hornung & Boulesteix (2022), in the case of data with larger numbers of variables (larger than 100, but more seriously for high-dimensional data), the univariable EIM values can be biased. Therefore, it is strongly recommended to interpret the univariable EIM

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values with caution, if the data are high-dimensional. If it is of interest to measure the univariable importance of the variables for high-dimensional data, an additional conventional random forest (e.g., using the ranger package) should be constructed and the variable importance measure values of this random forest be used for ranking the univariable effects.

Value

A ggplot2 plot.

Author(s)

Roman Hornung

References

- Hornung, R., Boulesteix, A.-L. (2022). Interaction forests: Identifying and exploiting interpretable quantitative and qualitative interaction effects. Computational Statistics & Data Analysis 171:107460, <doi: 10.1016/j.csda.2022.107460>.
- Hornung, R. (2022). Diversity forests: Using split sampling to enable innovative complex split procedures in random forests. SN Computer Science 3(2):1, <doi: 10.1007/s42979021-009201>.

See Also

```
plotEffects
```

Examples

```
## Not run:
## Load package:
library("diversityForest")

## Set seed to make results reproducible:
set.seed(1234)

## Construct interaction forest and calculate EIM values:
data(stock)
model <- interactionfor(dependent.variable.name = "company10", data = stock, num.trees = 20)

# NOTE: num.trees = 20 (in the above) would be much too small for practical # purposes. This small number of trees was simply used to keep the # runtime of the example short.</pre>
```

```
# The default number of trees is num.trees = 20000 if EIM values are calculated
# and num.trees = 2000 otherwise.

## When using the plot() function without further specifications,
## by default the estimated bivariable influences of the two pairs with largest quantitative
## and qualitative EIM values are shown:

plot(model)

# It is, however, also possible to change the numbers of
# pairs with largest quantitative and qualitative EIM values
# to be shown:

plot(model, numpairsquant = 4, numpairsqual = 3)

## End(Not run)
```

plotEffects

Interaction forest plots: exploring interaction forest results through visualisation

Description

This function allows to visualise the (estimated) bivariable influences of pairs of variables (with large quantitative and qualitative EIM values) on the outcome. This step is crucial, because to interpret interaction effects between variable pairs with large quantitative and qualitative EIM values, it is necessary to learn about the forms these interaction effects take.

Usage

```
plotEffects(
  intobj,
  type = "quant",
  numpairs = 5,
  indpairs = NULL,
  pairs = NULL,
  allwith = NULL,
  pvalues = TRUE,
  twoplots = TRUE,
  addtitles = TRUE,
  plotit = TRUE
```

Arguments

intobj Object of class interactionfor.

type This can be either "quant" or "qual" and determines whether the plotted pairs

are sorted according to either the quantitative or qualitative EIM values in de-

creasing order. Default is "quant".

numpairs The number of pairs to plot (default: 5). This is overwritten by indpairs.

indpairs Optional. The indices of the pairs in the sorted lists of quantitative (type="quant")

or qualitative EIM values to plot (type="qual"). This overwrites the numpairs

argument.

pairs This can be used to specify the pairs to plot. It is an optional list of charac-

ter string vectors, where each of these vectors has length two. Each list element corresponds to one pair, where the first character string gives the name of the first member of the respective pair to plot and the second character string gives the name of the second member. This argument overwrites numpairs and

indpairs.

allwith This is an optional character string that can be set to the name of one of the

variables. If provided, only variable pairs will be considered that feature the variable specified by this argument allwith. These pairs are again sorted in decreasing order according to the quantitative (type="quant") or qualitative (type="qual") EIM values and their number is restricted to the value given by numpairs. This argument allwith can be used, if it is of interest to learn whether a specific variable (e.g., sex or age) interacts with other variables in the

data set and if so, which forms these interactions take.

pvalues Set to TRUE (default) to add to the plots p-values from tests for interaction effect

obtained using classical parametric regression approaches. For categorical outcomes logistic regression is used, for metric outcomes linear regression and for survival outcomes Cox regression. NOTE: These p-values are generally much too optimistic and **MUST NOT** be reported as the result of a statistical test for

significance of interaction. See the 'Details' section below for further details.

twoplots Set to TRUE / FALSE if for each plot page the results of two / one pair(s) of

variables should be shown. Default is TRUE.

addtitles Set to TRUE (default) to add headings providing the names of the variables in

each pair. If type="quant", these headings also give information on the type of quantitative interaction effect. Setting addtitles to FALSE is, for example, useful, when the produced plots are intended for use in a publication, where

these headings might not be desirable.

plotit This states whether the plots are actually plotted or merely returned as ggplot

objects. Default is TRUE.

Details

For each considered pair the bivariable influence of both pair members on the outcome estimated using a two-dimensional flexible function is shown. Such visualisations make it possible to learn about the forms of the interaction effects between variable pairs with large EIM values. Moreover, these visualisations reveal (pathological) cases in which variable pairs do not show indications of

interaction effects despite featuring large EIM values.

For binary outcomes the estimated probabilities for the second class are predicted, for categorical outcomes with more than two classes the estimated probabilities for the largest class are predicted, for metric outcomes the means of the outcome are predicted, and for survival outcomes the log hazards ratio values compared to the median effect are predicted.

The kinds of estimates shown differ also according to whether both pair members are metric or only one of the two members is metric and the other one categorical or both pair members are categorical:

- If both pair members are metric and the outcome is categorical or metric we use two-dimensional LOESS regression, where in the case of categorical outcomes, to obtain probability estimates for the first class (or largest class for multi-class outcomes), we use the value '1' for the first class (largest class for multi-class outcomes) and the value '0' for the second class (all other classes for multi-class outcomes).
- If both pair members are metric and the outcome is survival we use a Cox proportional hazard additive model with a two-dimensional LOESS smooth (gamcox function from the 'MapGAM' package (version 1.2-5)) and in the rare cases for which the latter fails, we use classical Cox regression with an interaction term between the two covariates.
- If one pair member is metric and the other one categorical and the outcome is categorical or metric, we use LOESS regression between the outcome (coded as '0' and '1' in the case of categorical outcomes as described above) and the values of the metric variable separately for each category of the categorical variable. In the rare cases in which the LOESS regression fails we use classical linear regression.
- If one pair member is metric and the other one categorical and the outcome is survival, we use Cox regression with a linear tail-restricted cubic spline with four knots (univariable LOESS regression for survival outcomes does not seem to be available yet in R) separately for each category of the categorical variable. In cases in which the fitting of this spline regression fails we use classical Cox regression.
- If both pair members are categorical and the outcome is categorical or metric, we simply calculate the mean of the outcome (coded as '0' and '1' in the case of categorical outcomes as described above) for each possible combination of the categories of the two variables.
- If both pair members are categorical and the outcome is survival, we use classical Cox regression with an interaction term between the two variables (there is no need for any flexible modelling in this setting, because the Cox model with two categorical variables plus interaction term is saturated).

As described above (function argument: pvalues), there is an option to add p-values from tests for interaction effect to the plots. If at least one of the variables in the considered variable pair is categorical and features more than two categories, there are more than one interaction terms in the regression approaches used for testing, because the categorical variables are dummy-coded. Therefore, in these cases we obtain a p-value for each interaction term. to obtain a single p-value for the test for interaction we adjust these multiple p-values using the Holm-Bonferroni procedure and take the minimum of the adjusted p-values.

NOTE: These p-values are generally much too optimistic, in particular for small data sets and large numbers of variables. The reason for this overoptimism is that these p-values are not adjusted for the fact that we already used the data to find the variable pairs with strongest indications of interaction effects. This is similar to a multiple testing problem. Therefore, these p-values should only be seen as a rough guide to be interpreted very cautiously and **MUST NOT** be reported as the results of a

statistical test for significance of interaction. To obtain adjusted p-values that would correspond to valid tests, it would be possible to multiply these p-values by the number of possible pairs, which would correspond to Bonferroni-adjusted p-values. For example, assume that we have 30 covariate variables. In that case the number of possible pairs would be 'choose(30, 2) = 435', which is why we would need to multiply each p-value by 435 to obtain an adjusted p-value (or keep the original p-values and divide the significance level 0.05 by 435). Note, however, that Bonferroni-adjusted p-values deliver quite conservative results.

Value

A list of ggplot2 plots returned invisibly.

Author(s)

Roman Hornung

References

- Hornung, R., Boulesteix, A.-L. (2022). Interaction forests: Identifying and exploiting interpretable quantitative and qualitative interaction effects. Computational Statistics & Data Analysis 171:107460, <doi: 10.1016/j.csda.2022.107460>.
- Hornung, R. (2022). Diversity forests: Using split sampling to enable innovative complex split procedures in random forests. SN Computer Science 3(2):1, <doi: 10.1007/s42979021-009201>.

See Also

```
plot.interactionfor, plotPair
```

Examples

```
## Not run:
## Load package:
library("diversityForest")

## Set seed to make results reproducible:
set.seed(1234)

## Construct interaction forest and calculate EIM values:
data(stock)
model <- interactionfor(dependent.variable.name = "company10", data = stock, num.trees = 20)</pre>
```

```
# NOTE: num.trees = 20 (in the above) would be much too small for practical
# purposes. This small number of trees was simply used to keep the
# runtime of the example short.
# The default number of trees is num.trees = 20000 if EIM values are calculated
# and num.trees = 2000 otherwise.
## Obtain a first overview by applying the plot() function for
## interactionfor obects:
plot(model)
## Several possible application cases of the plotEffects() function:
# Visualise the estimated bivariable influences of the five variable pairs with the
# largest quantitative EIM values:
plotEffects(model) # type="quant" is default.
# Visualise the estimated bivariable influences of the five pairs with the
# largest qualitative EIM values:
plotEffects(model, type="qual")
# Visualise the estimated bivariable influences of all (eight) pairs that involve
# the variable "company7" sorted in decreasing order according to the
# qualitative EIM values:
plotEffects(model, allwith="company7", type="qual", numpairs=8)
# Visualise the estimated bivariable influences of the pairs with third and fifth
# largest qualitative EIM values:
plotEffects(model, type="qual", indpairs=c(3,5))
# Visualise the estimated bivariable influences of the pairs ("company3", "company5") and
# ("company1", "company9"):
plotEffects(model, pairs=list(c("company3", "company5"), c("company1", "company9")))
## Saving of plots generated with the plotEffects() function (e.g., for use in publications):
# Apply plotEffects() to obtain plots for the five variable pairs
# with the largest qualitative EIM values and store these plots in
```

```
# an object 'ps':
ps <- plotEffects(model, type="qual", pvalues=FALSE, twoplots=FALSE, addtitles=FALSE, plotit=FALSE)
# pvalues = FALSE states that no p-values should be shown in the plots,
# because these might not be desired in plots meant for publication.
# twoplots = FALSE ensures that we get one plot for each page instead of two plots per page.
# addtitles = FALSE removes the automatically generated titles, because these are likely
# not desired in publications.
# plotit = FALSE ensures that the plots are not displayed, but only returned (invisibly)
# by plotEffects().
# Save the plot with second largest qualitative EIM value:
p1 <- ps[[2]]
# Add title:
library("ggpubr")
p1 <- annotate_figure(p1, top = text_grob("My descriptive plot title 1", face = "bold", size = 14))
# Save as PDF:
# library("ggplot2")
# ggsave(file="mypathtofolder/FigureXY1.pdf", width=14, height=6)
# Save the plot with fifth largest qualitative EIM value:
p2 <- ps[[5]]
# Add title:
p2 <- annotate_figure(p2, top = text_grob("My descriptive plot title 2", face = "bold", size = 14))
p2
# Save as PDF:
# ggsave(file="mypathtofolder/FigureXY1.pdf", width=14, height=6)
# Combine both of the above plots:
p \leftarrow ggarrange(p1, p2, nrow = 2)
р
# Save the combined plot:
# ggsave(file="mypathtofolder/FigureXYcombined.pdf", width=14, height=11)
# NOTE: Using plotEffects() it is not possible to change the plots
# themselves (by e.g., increasing the label sizes or changing the
# axes ranges). However, the function plotPair() can be used to change
# the plots themselves.
## End(Not run)
```

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plotPair

Plot of the (estimated) simultaneous influence of two variables

Description

This function allows to visualise the (estimated) bivariable influence of a single specific pair of variables on the outcome. The estimation and plotting is performed in the same way as in plotEffects. However, plotPair does not require an interaction of object and can thus be used also without a constructed interaction forest.

Usage

```
plotPair(
  pair,
  yvarname,
  statusvarname = NULL,
  data,
  levelsorder1 = NULL,
  levelsorder2 = NULL,
  pvalue = TRUE,
  returnseparate = FALSE,
  intobj = NULL
)
```

Arguments

pair Character string vector of length two, where the first character string gives the

name of the first member of the respective pair to plot and the second character

string gives the name of the second member.

yvarname Name of outcome variable.

statusvarname Name of status variable, only applicable to survival data.

data Data frame containing the variables.

levelsorder1 Optional. Order the categories of the first variable should have in the plot (if it

is categorical). Character string vector, where the i-th entry contains the name of the category that should take the i-th place in the ordering of the categories of

the first variable.

levelsorder2 Optional. Order the categories of the second variable should have in the plot

(if it is categorical). Character string vector specified in an analogous way as

levels order 1.

pvalue Set to TRUE (default) to add to the plot a p-value from a test for interaction

effect obtained using a classical parametric regression approach. For categorical outcomes logistic regression is used, for metric outcomes linear regression and for survival outcomes Cox regression. See the 'Details' section of plotEffects

for further details.

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returnseparate Set to TRUE to return invisibly the two generated ggplot plots separately in the

form of a list. The latter option is useful, because it allows to manipulate the resulting plots (label size etc.) and makes it possible to consider only one of the two plots. The default is FALSE, which results in the two plots being returned

together in the form of a ggarrange object.

intobj Optional. Object of class interactionfor. If this is provided, the ordering of

the categories obtained when constructing the interaction forest will be used for categorical variables. See Hornung & Boulesteix (2021) for details.

Details

See the 'Details' section of plotEffects.

Value

A ggplot2 plot.

Author(s)

Roman Hornung

References

- Hornung, R., Boulesteix, A.-L. (2022). Interaction forests: Identifying and exploiting interpretable quantitative and qualitative interaction effects. Computational Statistics & Data Analysis 171:107460, <doi: 10.1016/j.csda.2022.107460>.
- Hornung, R. (2022). Diversity forests: Using split sampling to enable innovative complex split procedures in random forests. SN Computer Science 3(2):1, <doi: 10.1007/s42979021-009201>.

See Also

plotEffects, plot.interactionfor

Examples

```
## Not run:
## Load package:
library("diversityForest")

## Visualise the estimated bivariable influence of 'toothed' and 'feathers' on
## the probability of type="mammal":

data(zoo)
plotPair(pair = c("toothed", "feathers"), yvarname="type", data = zoo)
```

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```
## Visualise the estimated bivariable influence of 'creat' and 'hgb' on
## survival (more precisely, on the log hazards ratio compared to the
## median effect):
library("survival")
mgus2compl <- mgus2[complete.cases(mgus2),]</pre>
plotPair(pair=c("creat", "hgb"), yvarname="futime", statusvarname = "death", data=mgus2compl)
# Problem: The outliers in the left plot make it difficult to see what is going
# on in the region with creat values smaller than about two even though the
# majority of values lie there.
# --> Solution: We re-run the above line setting returnseparate = TRUE, because
# this allows to get the two ggplot plots separately, which can then be manipulated
# to change the x-axis range in order to remove the outliers:
ps <- plotPair(pair=c("creat", "hgb"), yvarname="futime", statusvarname = "death",</pre>
               data=mgus2compl, returnseparate = TRUE)
# Change the x-axis range:
library("ggplot2")
ps[[1]] + xlim(c(0.5,2))
# Save the plot:
# ggsave(file="mypathtofolder/FigureXY1.pdf", width=7, height=6)
# We can, for example, also change the label sizes of the second plot:
# With original label sizes:
ps[[2]]
# With larger label sizes:
ps[[2]] + theme(axis.title=element_text(size=15))
# Save the plot:
# library("ggplot2")
# ggsave(file="mypathtofolder/FigureXY2.pdf", width=7, height=6)
## End(Not run)
```

predict.divfor

Diversity Forest prediction

Description

Prediction with new data and a saved forest from divfor.

Usage

```
## S3 method for class 'divfor'
```

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```
predict(
  object,
  data = NULL,
  predict.all = FALSE,
  num.trees = object$num.trees,
  type = "response",
  se.method = "infjack",
  quantiles = c(0.1, 0.5, 0.9),
  seed = NULL,
  num.threads = NULL,
  verbose = TRUE,
  ...
)
```

Arguments

object urviol object	object	divfor object
----------------------	--------	---------------

data New test data of class data.frame or gwaa.data (GenABEL).

predict.all Return individual predictions for each tree instead of aggregated predictions for

all trees. Return a matrix (sample x tree) for classification and regression, a 3d array for probability estimation (sample x class x tree) and survival (sample x

time x tree).

num.trees Number of trees used for prediction. The first num.trees in the forest are used.

type Type of prediction. One of 'response', 'se', 'terminalNodes', 'quantiles' with

default 'response'. See below for details.

se.method Method to compute standard errors. One of 'jack', 'infjack' with default 'inf-

jack'. Only applicable if type = 'se'. See below for details.

quantiles Vector of quantiles for quantile prediction. Set type = 'quantiles' to use.

seed Random seed. Default is NULL, which generates the seed from R. Set to 0 to

ignore the R seed. The seed is used in case of ties in classification mode.

num. threads Number of threads. Default is number of CPUs available.

verbose Verbose output on or off.

... further arguments passed to or from other methods.

Details

This package is a fork of the R package 'ranger' that implements random forests using an efficient C++ implementation. More precisely, 'diversityForest' was written by modifying the code of 'ranger', version 0.11.0. Therefore, details on further functionalities of the code that are not presented in the help pages of 'diversityForest' are found in the help pages of 'ranger' (version 0.11.0). The code in the example sections of divfor and tunedivfor can be used as a template for all common application scenarios with respect to classification, regression and survival prediction using univariable, binary splitting. Some function arguments adopted from the 'ranger' package may not be useable with diversity forests (for the current package version).

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Value

Object of class divfor.prediction with elements

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predictions Predicted classes/values (only for classification and regression)

unique.death.times Unique death times (only for survival).

chf Estimated cumulative hazard function for each sample (only for survival).

survival Estimated survival function for each sample (only for survival).

num. trees Number of trees.

num.independent.variables Number of independent variables.

treetype Type of forest/tree. Classification, regression or survival.

num.samples Number of samples.

Author(s)

Marvin N. Wright

References

- Hornung, R. (2022). Diversity forests: Using split sampling to enable innovative complex split procedures in random forests. SN Computer Science 3(2):1, <doi: 10.1007/s42979021-009201>.
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- Wager, S., Hastie T., & Efron, B. (2014). Confidence Intervals for Random Forests: The Jackknife and the Infinitesimal Jackknife. Journal of Machine Learning Research 15:1625-1651.
- Meinshausen (2006). Quantile Regression Forests. Journal of Machine Learning Research 7:983-999.

See Also

divfor

predict.interactionfor

Interaction Forest prediction

Description

Prediction with new data and a saved interaction forest from interactionfor.

Usage

```
## S3 method for class 'interactionfor'
predict(
  object,
  data = NULL,
  predict.all = FALSE,
```

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```
num.trees = object$num.trees,
  type = "response",
  se.method = "infjack",
  quantiles = c(0.1, 0.5, 0.9),
  seed = NULL,
  num.threads = NULL,
  verbose = TRUE,
  ...
)
```

Arguments

object interactionfor object.

data New test data of class data. frame or gwaa. data (GenABEL).

predict.all Return individual predictions for each tree instead of aggregated predictions for

all trees. Return a matrix (sample x tree) for classification and regression, a 3d array for probability estimation (sample x class x tree) and survival (sample x

time x tree).

type Type of prediction. One of 'response', 'se', 'terminalNodes', 'quantiles' with

default 'response'. See below for details.

se.method Method to compute standard errors. One of 'jack', 'infjack' with default 'inf-

jack'. Only applicable if type = 'se'. See below for details.

quantiles Vector of quantiles for quantile prediction. Set type = 'quantiles' to use.

seed Random seed. Default is NULL, which generates the seed from R. Set to 0 to

ignore the R seed. The seed is used in case of ties in classification mode.

num. threads Number of threads. Default is number of CPUs available.

verbose Verbose output on or off.

... further arguments passed to or from other methods.

Details

Note that his package is a fork of the R package 'ranger' that implements random forests using an efficient C++ implementation. The documentation is in large parts taken from 'ranger', where some parts of the documentation may not apply to (the current version of) the 'diversityForest' package. Details on further functionalities of the code that are not presented in the help pages of 'diversityForest' are found in the help pages of 'ranger' (version 0.11.0).

Value

Object of class interaction.prediction with elements

predictions Predicted classes/values (only for classification and regression)

unique.death.times Unique death times (only for survival).

chf Estimated cumulative hazard function for each sample (only for survival).

survival Estimated survival function for each sample (only for survival).

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num. trees Number of trees.

num.independent.variables Number of independent variables.

treetype Type of forest/tree. Classification, regression or survival.

num.samples Number of samples.

Author(s)

Marvin N. Wright, Roman Hornung

References

- Hornung, R., Boulesteix, A.-L. (2022). Interaction forests: Identifying and exploiting interpretable quantitative and qualitative interaction effects. Computational Statistics & Data Analysis 171:107460, <doi: 10.1016/j.csda.2022.107460>.
- Hornung, R. (2022). Diversity forests: Using split sampling to enable innovative complex split procedures in random forests. SN Computer Science 3(2):1, <doi: 10.1007/s42979021-009201>.
- Wright, M. N., Ziegler, A. (2017). ranger: A fast Implementation of Random Forests for High Dimensional Data in C++ and R. Journal of Statistical Software 77:1-17, <doi: 10.18637/jss.v077.i01>.
- Wager, S., Hastie T., & Efron, B. (2014). Confidence Intervals for Random Forests: The Jackknife and the Infinitesimal Jackknife. Journal of Machine Learning Research 15:1625-1651.
- Meinshausen (2006). Quantile Regression Forests. Journal of Machine Learning Research 7:983-999.

See Also

interactionfor

stock

Data on stock prices of aerospace companies

Description

This data set contains 950 daily stock prices from January 1988 through October 1991, for ten aerospace companies. The names of the companies are anonymised and the stock prices for one of these companies (company10) were flagged as the outcome variable. Thus, for this data set, both the outcome and the covariates were metric.

Format

A data frame with 950 observations, nine covariates and one metric outcome variable

Details

The variables are as follows: covariates: company1, ..., company9, outcome variable: company10.

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Source

OpenML: data.name: stock, data.id: 223, link: https://www.openml.org/d/223/

References

 Vanschoren, J., van Rijn, J. N., Bischl, B., Torgo, L. (2013). OpenML: networked science in machine learning. SIGKDD Explorations 15(2):49-60, <doi: 10.1145/2641190.2641198>.

Examples

```
## Load data:
data(stock)

## Dimension of data:
dim(stock)

## First rows of data:
head(stock)
```

tunedivfor

Optimization of the values of the tuning parameters nsplits and proptry

Description

First, both for nsplits and proptry a grid of possible values may be provided, where default grids are used if no grids are provided. Second, for each pairwise combination of values from these two grids a forest is constructed. Third, that pair of nsplits and proptry values is used as the optimized set of parameter values that is associated with the smallest out-of-bag prediction error. If several pairs of parameter values are associated with the same smallest out-of-bag prediction error, the pair with the smallest (parameter) values is used.

Usage

```
tunedivfor(
  formula = NULL,
  data = NULL,
  nsplitsgrid = c(2, 5, 10, 30, 50, 100, 200),
  proptrygrid = c(0.05, 1),
  num.trees.pre = 500
)
```

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Arguments

formula Object of class formula or character describing the model to fit. Interaction

terms supported only for numerical variables.

data Training data of class data.frame, matrix, dgCMatrix (Matrix) or gwaa.data

(GenABEL).

nsplitsgrid Grid of values to consider for nsplits. Default grid: 2, 5, 10, 30, 50, 100, 200.

proptrygrid Grid of values to consider for proptry. Default grid: 0.05, 1.

num.trees.pre Number of trees used for each forest constructed during tuning parameter opti-

mization. Default is 500.

Value

List with elements

nsplitsopt Optimized value of nsplits.
proptryopt Optimized value of proptry.

tunegrid Two-dimensional data. frame, where each row contains one pair of values con-

sidered for nsplits (first entry) and proptry (second entry).

ooberrs The out-of-bag prediction errors obtained for each pair of values considered for

nsplits and proptry, where the ordering of pairs of values is the same as in

tunegrid (see above).

Author(s)

Roman Hornung

References

- Hornung, R. (2022). Diversity forests: Using split sampling to enable innovative complex split procedures in random forests. SN Computer Science 3(2):1, <doi: 10.1007/s42979021-009201>.
- Wright, M. N., Ziegler, A. (2017). ranger: A fast Implementation of Random Forests for High Dimensional Data in C++ and R. Journal of Statistical Software 77:1-17, <doi: 10.18637/jss.v077.i01>.

See Also

divfor

Examples

Load package:

library("diversityForest")

36 zoo

```
## Set seed to obtain reproducible results:
set.seed(1234)

## Tuning parameter optimization for the iris data set:

tuneres <- tunedivfor(formula = Species ~ ., data = iris, num.trees.pre = 20)
# NOTE: num.trees.pre = 20 is specified too small for practical
# purposes - the out-of-bag error estimates of the forests
# constructed during optimization will be much too variable!!
# In practice, num.trees.pre = 500 (default value) or a
# larger number should be used.

tuneres

tuneres$proptryopt
tuneres$proptryopt
tuneres$tunegrid
tuneres$ooberrs</pre>
```

Z00

Data on biological species

Description

This data set describes 101 different biological species using 16 simple attributes, where 15 of these are binary and one is metric (the number of legs). The outcome "mammal vs. other" (type) is binary.

Format

A data frame with 101 observations, 16 covariates and one binary outcome variable

Details

The variables are as follows:

- hair. factor. Presence of hairs (true = yes; false = no)
- feathers. factor. Presence of feathers (true = yes; false = no)
- eggs. factor. Does the species lay eggs? (true = yes; false = no)
- milk. factor. Does the species give milk? (true = yes; false = no)
- airborne. factor. Does the species fly? (true = yes; false = no)
- aquatic. factor. Does the species live in the water? (true = yes; false = no)
- predator. factor. Is the species a predator? (true = yes; false = no)
- toothed. factor. Presence of teeth (true = yes; false = no)

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- backbone. factor. Presence of backbone (true = yes; false = no)
- breathes. factor. Does the species breathe with lungs? (true = yes; false = no)
- venomous. factor. Is the species venomous? (true = yes; false = no)
- fins. factor. Presence of fins (true = yes; false = no)
- legs. metric. Number of legs
- tail. factor. Presence of tail (true = yes; false = no)
- domestic. factor. Is the species domestic? (true = yes; false = no)
- catsize. factor. Is the species large? (true = yes; false = no)
- type. factor. Binary outcome variable type of species ('mammal' vs. 'other')

The original openML dataset contains an additional variable animal, which is removed in this version of the data set. This variable provided the names of all species.

Source

OpenML: data.name: zoo, data.id: 965, link: https://www.openml.org/d/965/

References

- Vanschoren, J., van Rijn, J. N., Bischl, B., Torgo, L. (2013). OpenML: networked science in machine learning. SIGKDD Explorations 15(2):49-60, <doi: 10.1145/2641190.2641198>.
- Dua, D., Graff, C. (2019) UCI Machine Learning Repository. Irvine, CA: University of California, School of Information and Computer Science. https://archive.ics.uci.edu/ml/.

Examples

```
##' Load data:
data(zoo)

##' Numbers of observations in the two classes:
table(zoo$type)

##' Dimension of data:
dim(zoo)

##' First rows of data:
head(zoo)
```

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