

Package ‘metaforest’

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Type Package

Title Exploring Heterogeneity in Meta-Analysis using Random Forests

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Description Conduct random forests-based meta-analysis, obtain partial dependence plots for metaforest and classic meta-analyses, and cross-validate and tune metaforest- and classic meta-analyses in conjunction with the caret package. A requirement of classic meta-analysis is that the studies being aggregated are conceptually similar, and ideally, close replications. However, in many fields, there is substantial heterogeneity between studies on the same topic. Classic meta-analysis lacks the power to assess more than a handful of univariate moderators. MetaForest, by contrast, has substantial power to explore heterogeneity in meta-analysis. It can identify important moderators from a larger set of potential candidates (Van Lissa, 2020). This is an appealing quality, because many meta-analyses have small sample sizes. Moreover, MetaForest yields a measure of variable importance which can be used to identify important moderators, and offers partial prediction plots to explore the shape of the marginal relationship between moderators and effect size.

Depends R (>= 3.5.0), ggplot2, metafor, ranger, data.table, methods

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coef_test	<i>Test coefficients of a model</i>
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Description

Conduct a t-test or z-test for coefficients of a model.

Usage

```
coef_test(x, par1, par2, distribution = "pt")
```

Arguments

x	A model.
par1	Numeric or character. Name or position of the first parameter.
par2	Numeric or character. Name or position of the second parameter.
distribution	Character. Which distribution to use. Currently, can be one of c("pt", "pnorm"), for a t-test or z-test, respectively. Defaults to "pt".

Value

Named vector.

Examples

```
dat <- escalc(measure="RR", ai=tpos, bi=tneg, ci=cpos, di=cneg, data=dat.bcg)
res <- rma(yi, vi, mods = ~alloc-1, data=dat, method="REML")
coef_test(res, 1, 2)
```

curry

*Happy to Help?***Description**

A systematic review and meta-analysis of the effects of performing acts of kindness on the well-being of the actor.

Usage

```
data(curry)
```

Format

A data.frame with 56 rows and 18 columns.

Details

study_id	factor	Unique identifier of the study
effect_id	integer	Unique identifier of the effect size
d	numeric	Standardized mean difference between the control group and intervention group
vi	numeric	Variance of the effect size
n1i	numeric	Number of participants in the intervention group
n1c	numeric	Number of participants in the control group
sex	numeric	Percentage of male participants
age	numeric	Mean age of participants
location	character	Geographical location of the study
donor	character	From what population did the donors (helpers) originate?
donorcode	factor	From what population did the donors (helpers) originate? Dichotomized to Anxious or Typ
interventioniv	character	Description of the intervention / independent variable
interventioncode	factor	Description of the intervention / independent variable, categorized to Acts of Kindness, Pro
control	character	Description of the control condition
controlcode	factor	Description of the control condition, categorized to Neutral Activity, Nothing, or Self Help
recipients	character	Who were the recipients of the act of kindness?
outcomedv	character	What was the outcome, or dependent variable, of the study?
outcomecode	factor	What was the outcome, or dependent variable, of the study? Categorized into Happiness, L

Source

[doi:10.1016/j.jesp.2018.02.014](https://doi.org/10.1016/j.jesp.2018.02.014)

References

Curry, O. S., Rowland, L. A., Van Lissa, C. J., Zlotowitz, S., McAlaney, J., & Whitehouse, H. (2018). Happy to help? A systematic review and meta-analysis of the effects of performing acts of

kindness on the well-being of the actor. *Journal of Experimental Social Psychology*, 76, 320-329.
[doi:10.1016/j.jesp.2018.02.014](https://doi.org/10.1016/j.jesp.2018.02.014)

extract_proximity *Extract proximity matrix for a MetaForest object.*

Description

Extract proximity matrix for a MetaForest object.

Usage

```
extract_proximity(fit, newdata)
```

Arguments

fit object of class `'MetaForest'`.
newdata new data with the same columns as the data used for fit

Value

an $n \times n$ matrix where position i, j gives the proportion of times observation i and j are in the same terminal node across all trees.

Examples

fukkink_lont *Does training matter? A meta-analysis of caregiver training studies*

Description

A review of 17 experimental studies published between 1980 and 2005 on the effect of specialized training on the competency of caregivers in childcare.

Usage

```
data(fukkink_lont)
```

Format

A data.frame with 78 rows and 30 columns.

Details

id_exp	integer	Unique identifier of the study
yi	numeric	Standardized mean difference between the control group and
vi	numeric	Variance of the effect size
Journal	factor	Publication type (scientific journal or other publications)
Setting	factor	Setting (center-based care or family daycare)
Integrated	factor	Whether the training was integrated into childcare practice
Supervision	factor	Whether supervision was part of the training
Scope	factor	Scope of the training (narrow or broad)
Location	factor	Location of the training (one-site or multi-site)
Curriculum	factor	Fixed curriculum
Control	factor	Alternative treatment for control group
Assignment	factor	Random assignment or matching (at the level of the individual caregiver or childcare center)
Train_Knowledge	factor	Explicit focus on knowledge
Train_Skills	factor	Explicit focus on skills
Train_Attitude	factor	Explicit focus on attitude
Video	factor	Use of video feedback
Design	factor	Single group, or two-group experimental design
Pre_Post	factor	Pretest/posttest design (yes/no)
Blind	factor	Was a blinding procedure used?
Attrition	numeric	Attrition from the experimental condition (percentage)
Pretest_es	numeric	Pre-test effect size
Self_report	factor	Self-report measures of caregiver competencies versus ‘objective’ test or observation by in
DV_Knowledge	factor	Test focused on knowledge
DV_Skills	factor	Test focused skills
DV_Attitude	factor	Test focused on attitudes
DV_Aligned	factor	Test aligned with the content of the training (yes/no)
Two_group_design	factor	Single group, or two-group experimental design
Trainee_Age	numeric	Trainees’ age
Trainee_Experience	numeric	Trainees’ working experience
n_total	integer	Total n at post-test

Source

[doi:10.1016/j.ecresq.2007.04.005](https://doi.org/10.1016/j.ecresq.2007.04.005)

References

Fukkink, R. G., & Lont, A. (2007). Does training matter? A meta-analysis and review of caregiver training studies. *Early childhood research quarterly*, 22(3), 294-311. [doi:10.1016/j.ecresq.2007.04.005](https://doi.org/10.1016/j.ecresq.2007.04.005)

Description

MetaForest uses a weighted random forest to explore heterogeneity in meta-analytic data. MetaForest is a wrapper for [ranger](#) (Wright & Ziegler, 2015). As input, MetaForest takes the study effect sizes and their variances (these can be computed, for example, using the [metafor](#) package), as well as the moderators that are to be included in the model. By default, MetaForest uses random-effects weights, and estimates the between-studies variance using a restricted maximum-likelihood estimator. However, it may be beneficial to first conduct an unweighted MetaForest, and then use the estimated residual heterogeneity from this model as the estimate of τ^2 for a random-effects weighted MetaForest.

Usage

```
MetaForest(
  formula,
  data,
  vi = "vi",
  study = NULL,
  whichweights = "random",
  num.trees = 500,
  mtry = NULL,
  method = "REML",
  tau2 = NULL,
  ...
)
```

Arguments

formula	Formula. Specify a formula for the MetaForest model, for example, $y_i \sim .$ to predict the outcome y_i from all moderators in the data. Only additive formulas are allowed (i.e., $x_1+x_2+x_3$). Interaction terms and non-linear terms are not required, as the random forests algorithm inherently captures these associations.
data	A data.frame containing the effect size, moderators, and the variance of the effect size.
vi	Character. Specify the name of the column in the data that contains the variances of the effect sizes. This column will be removed from the data prior to analysis. Defaults to "vi".
study	Character. Optionally, specify the name of the column in the data that contains the study id. Use this when the data includes multiple effect sizes per study. This column can be a vector of integers, or a factor. This column will be removed from the data prior to analysis. See Details for more information about analyzing dependent data.
whichweights	Character. Indicate what type of weights are required. A random-effects MetaForest is grown by specifying <code>whichweights = "random"</code> . A fixed-effects MetaForest is grown by specifying <code>whichweights = "fixed"</code> . An unweighted MetaForest is grown by specifying <code>whichweights = "unif"</code> . Defaults to "random".
num.trees	Atomic integer. Specify the number of trees in the forest. Defaults to 500.

mtry	Atomic integer. Number of candidate moderators available for each split. Defaults to the square root of the number moderators (rounded down).
method	Character. Specify the method by which to estimate the residual variance. Can be set to one of the following: "DL", "HE", "SJ", "ML", "REML", "EB", "HS", or "GENQ". Default is "REML". See the metafor package for more information about these estimators.
tau2	Numeric. Specify a predetermined value for the residual heterogeneity. Entering a value here supersedes the estimated tau2 value. Defaults to NULL.
...	Additional arguments are passed directly to ranger . It is recommended not to use additional arguments.

Details

For dependent data, a clustered MetaForest analysis is more appropriate. This is because the predictive performance of a MetaForest analysis is evaluated on out-of-bootstrap cases, and when cases out of the bootstrap sample originate from the same study, the model will be overly confident in its ability to predict their value. When the MetaForest is clustered by the study variable, the dataset is first split into two cross-validation samples by study. All dependent effect sizes from each study are thus included in the same cross-validation sample. Then, two random forests are grown on these cross-validation samples, and for each random forest, the other sample is used to calculate prediction error and variable importance, see [doi:10.1007/s1163401602764](https://doi.org/10.1007/s1163401602764).

Value

List of length 3. The "forest" element of this list is an object of class "ranger", containing the results of the random forests analysis. The "rma_before" element is an object of class "rma.uni", containing the results of a random-effects meta-analysis on the raw data, without moderators. The "rma_after" element is an object of class "rma.uni", containing the results of a random-effects meta-analysis on the residual heterogeneity, or the difference between the effect sizes predicted by MetaForest and the observed effect sizes.

Examples

```
#Example 1:
#Simulate data with a univariate linear model
set.seed(42)
data <- SimulateSMD()
#Conduct unweighted MetaForest analysis
mf.unif <- MetaForest(formula = yi ~ ., data = data$training,
  whichweights = "unif", method = "DL")
#Print model
mf.unif
#Conduct random-effects weighted MetaForest analysis
mf.random <- MetaForest(formula = yi ~ ., data = data$training,
  whichweights = "random", method = "DL",
  tau2 = 0.0116)
#Print summary
summary(mf.random)

#Example 2: Real data from metafor
```

```

#Load and clean data
data <- dat.bangertdrowns2004
data[, c(4:12)] <- apply(data[, c(4:12)], 2, function(x){
  x[is.na(x)] <- median(x, na.rm = TRUE)
  x})
data$subject <- factor(data$subject)
data$yi <- as.numeric(data$yi)
#Conduct MetaForest analysis
mf.bd2004 <- MetaForest(formula = yi~ grade + length + minutes + wic+
                        meta, data, whichweights = "unif")

#Print MetaForest object
mf.bd2004
#Check convergence plot
plot(mf.bd2004)
#Check summary
summary(mf.bd2004, digits = 4)
#Examine variable importance plot
VarImpPlot(mf.bd2004)

```

ModelInfo_mf

Returns a MetaForest ModelInfo list for use with caret

Description

This function allows users to rely on the powerful `caret` package for cross-validating and tuning a MetaForest analysis. Methods for MetaForest are not included in the `caret` package, because the interface of `caret` is not entirely compatible with MetaForest's model call. Specifically, MetaForest is not compatible with the `train` methods for classes 'formula' or 'recipe', because the variance of the effect size must be a column of the training data `x`. The name of this column is specified using the argument 'vi'.

Usage

```
ModelInfo_mf()
```

Details

To train a clustered MetaForest, for nested data structures, simply provide the optional argument 'study' to the `train` function, to specify the study ID. This should again refer to a column of `x`.

When training a clustered MetaForest, make sure to use 'index = groupKfold(your_study_id_variable, k = 10))' in `traincontrol`, to sample by study ID when creating cross-validation partitions; otherwise the testing error will be positively biased.

Value

ModelInfo list of length 17.

Examples

```

## Not run:
# Prepare data
data <- dat.bangertdrowns2004
data[, c(4:12)] <- apply(data[, c(4:12)], 2, function(x){
  x[is.na(x)] <- median(x, na.rm = TRUE)
  x})
data$subject <- factor(data$subject)
data$yi <- as.numeric(data$yi)
# Load caret
library(caret)
set.seed(999)
# Specify the resampling method as 10-fold CV
fit_control <- trainControl(method = "cv", number = 10)
cv_mf_fit <- train(y = data$yi, x = data[,c(3:13, 16)],
  method = ModelInfo_mf(), trControl = fit_control)

# Cross-validated clustered MetaForest
data <- get(data(dat.bourassa1996))
data <- escalc(measure = "OR", ai = lh.le, bi = lh.re, ci = rh.le, di = rh.re,
  data = data, add = 1/2, to = "all")
data$mage[is.na(data$mage)] <- median(data$mage, na.rm = TRUE)
data[c(5:8)] <- lapply(data[c(5:8)], factor)
data$yi <- as.numeric(data$yi)
# Set up 10-fold grouped CV
fit_control <- trainControl(method = "cv", index = groupKFold(data$sample,
  k = 10))
# Set up a custom tuning grid for the three tuning parameters of MetaForest
rf_grid <- expand.grid(whichweights = c("random", "fixed", "unif"),
  mtry = c(2, 4, 6),
  min.node.size = c(2, 4, 6))
# Train the model
cv.mf.cluster <- train(y = data$yi, x = data[, c("selection", "investigator",
  "hand_assess", "eye_assess",
  "mage", "sex", "vi",
  "sample")],
  study = "sample", method = ModelInfo_mf(),
  trControl = fit_control,
  tuneGrid = rf_grid)

## End(Not run)

```

ModelInfo_rma

Returns an rma ModelInfo list for use with caret

Description

This function allows users to rely on the powerful caret package for cross-validating and tuning a rma analysis. Methods for rma are not included in the caret package, because the interface of

caret is not entirely compatible with rma's model call. Specifically, rma is not compatible with the train methods for classes 'formula' or 'recipe'. The variance of the effect sizes can be passed to the 'weights' parameter of train.

Usage

```
ModelInfo_rma()
```

Details

When using clustered data (effect sizes within studies), make sure to use 'index = groupKFold(your_study_id_variable, k = 10))' in traincontrol, to sample by study ID when creating cross-validation partitions; otherwise the testing error will be positively biased.

Value

ModelInfo list of length 13.

Examples

```
## Not run:
# Prepare data
dat <- escalc(measure="RR", ai=tpos, bi=tneg, ci=cpos, di=cneg, data=dat.bcg)
dat$yi <- as.numeric(dat$yi)
dat$alloc <- factor(dat$alloc)
# Run rma
rma.model <- rma(y = dat$yi, mods = dat[, c("ablat", "year")], vi = dat$vi)
# R^2 is estimated to be .64
rma.model$R2
# Now, use cross-validation to see how well this model generalizes
# Leave-one-out cross-validation is more appropriate than 10-fold cv because
# the sample size is very small
fit_control <- trainControl(method = "LOOCV")
# Train the model without tuning, because rma has no tuning parameters
cv.mf.cluster <- train(y = dat$yi, x = dat[, c("ablat", "year")],
  weights = dat$vi,
  method = ModelInfo_rma(),
  trControl = fit_control)
# Cross-validated R^2 is .08, suggesting substantial overfitting of the
# original rma model
cv.mf.cluster$results$Rsquared

## End(Not run)
```

Description

Partial dependence plots

Usage

```

PartialDependence(
  x,
  vars = NULL,
  pi = NULL,
  rawdata = FALSE,
  bw = FALSE,
  resolution = NULL,
  moderator = NULL,
  mod_levels = NULL,
  output = "plot",
  ...
)

```

Arguments

<code>x</code>	Model object.
<code>vars</code>	Character vector containing the moderator names for which to plot partial dependence plots. If empty, all moderators are plotted.
<code>pi</code>	Numeric (0-1). What percentile interval should be plotted for the partial dependence predictions? Defaults to NULL. To obtain a 95% interval, set to .95.
<code>rawdata</code>	Logical, indicating whether to plot weighted raw data. Defaults to FALSE. Uses the same weights as the model object passed to the <code>x</code> argument.
<code>bw</code>	Logical, indicating whether the plot should be black and white, or color.
<code>resolution</code>	Integer vector of length two, giving the resolution of the partial predictions. The first element indicates the resolution of the partial predictions; for Monte-Carlo integration, the second element gives the number of rows of the data to be sampled without replacement when averaging over values of the other predictors.
<code>moderator</code>	Atomic character vector, referencing the name of one variable in the model. Results in partial prediction plots, conditional on the moderator. If <code>moderator</code> references a factor variable, separate lines/boxplots are plotted for each factor level. If <code>moderator</code> references a numeric variable, heatmaps are plotted - unless the moderator is categorized using the <code>mod_levels</code> argument.
<code>mod_levels</code>	Vector. If <code>moderator</code> is continuous, specify thresholds for the <code>cut</code> function. The continuous moderator is categorized, and predictions are based on the median moderator value within each category. You can call <code>quantile</code> to cut the moderator at specific quantiles. If <code>moderator</code> is a factor variable, you can use <code>mod_levels</code> to specify a character vector with the factor levels to retain in the plot (i.e., dropping the other factor levels).
<code>output</code>	Character. What type of output should be returned? Defaults to "plot", which returns and plots a gtable object. To obtain a list of ggplot objects instead, provide the argument "list".
<code>...</code>	Additional arguments to be passed to and from functions.

Details

Plots partial dependence plots (predicted effect size as a function of the value of each predictor variable) for a MetaForest- or rma model object. For rma models, it is advisable to mean-center numeric predictors, and to not include plot_int effects, except when the rma model is bivariate, and the plot_int argument is set to TRUE.

Value

A gtable object.

Examples

```
## Not run:
#' # Partial dependence plot for MetaForest() model:
set.seed(42)
data <- SimulateSMD(k_train = 200, model = es * x[, 1] + es * x[, 2] + es *
                    x[, 1] * x[, 2])$training
data$X2 <- cut(data$X2, breaks = 2, labels = c("Low", "High"))
mf.random <- MetaForest(formula = yi ~ ., data = data,
                       whichweights = "random", method = "DL",
                       tau2 = 0.2450)
# Examine univariate partial dependence plot for all variables in the model:
PartialDependence(mf.random, pi = .8)
# Examine bivariate partial dependence plot the plot_int between X1 and X2:
pd.plot <- PartialDependence(mf.random, vars = c("X1", "X2"), plot_int = TRUE)
# Save to pdf file
pdf("pd_plot.pdf")
grid.draw(pd.plot)
dev.off()
# Partial dependence plot for metafor rma() model:
dat <- escalc(measure="RR", ai=tpos, bi=tneg, ci=cpos, di=cneg, data=dat.bcg)
dat$yi <- as.numeric(dat$yi)
dat$alloc <- factor(dat$alloc)
dat$ablat_d <- cut(dat$ablat, breaks = 2, labels = c("low", "high"))
# Demonstrate partial dependence plot for a bivariate plot_int
rma.model.int <- rma(yi, vi, mods=cbind(ablat, tpos),
                    data=dat, method="REML")
PartialDependence(rma.model.int, rawdata = TRUE, pi = .95,
                  plot_int = TRUE)

# Compare partial dependence for metaforest and rma
dat2 <- dat
dat2[3:7] <- lapply(dat2[3:7],
                  function(x){as.numeric(scale(x, scale = FALSE))})
mf.model.all <- MetaForest(yi ~ ., dat2[, c(3:11)])
rma.model.all <- rma(dat$yi, dat2$vi,
                    mods = model.matrix(yi~., dat2[, c(3:10)]), [-1],
                    method="REML")
PartialDependence(mf.model.all, rawdata = TRUE, pi = .95)
PartialDependence(rma.model.all, rawdata = TRUE, pi = .95)

## End(Not run)
```

plot.MetaForest	<i>Plots cumulative MSE for a MetaForest object.</i>
-----------------	--

Description

Plots cumulative MSE for a MetaForest object.

Usage

```
## S3 method for class 'MetaForest'
plot(x, y, ...)
```

Arguments

x	MetaForest object.
y	not used for plot.MetaForest
...	Arguments to be passed to methods, not used for plot.MetaForest

Value

A ggplot object, visualizing the number of trees on the x-axis, and the cumulative mean of the MSE of that number of trees on the y-axis. As a visual aid to assess convergence, a dashed gray line is plotted at the median cumulative MSE value.

Examples

predict.MetaForest	<i>MetaForest prediction</i>
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Description

MetaForest prediction

Usage

```
## S3 method for class 'MetaForest'
predict(object, data = NULL, type = "response", ...)
```

Arguments

object	MetaForest object.
data	New test data of class data.frame.
type	Type of prediction. One of 'response', 'se', 'terminalNodes' with default 'response'. See below for details.
...	further arguments passed to or from other methods.

Value

Object of class `MetaForest.prediction` with elements

<code>predictions</code>	Predicted classes/values (only for classification and regression)
<code>num.trees</code>	Number of trees.
<code>num.independent.variables</code>	Number of independent variables.
<code>treetype</code>	Type of forest/tree. Classification, regression or survival.
<code>num.samples</code>	Number of samples.

See Also

[ranger](#)

Examples

```
set.seed(56)
data <- SimulateSMD(k_train = 100, model = es * x[,1] * x[,2])
#Conduct fixed-effects MetaForest analysis
mf.fixed <- MetaForest(formula = yi ~ ., data = data$training,
                      whichweights = "fixed", method = "DL")
predicted <- predict(mf.fixed, data = data$testing)$predictions
r2_cv <- sum((predicted - mean(data$training$yi)) ^ 2) /
        sum((data$testing$yi - mean(data$training$yi)) ^ 2)
```

```
preselect
```

Preselect variables for MetaForest analysis

Description

Takes a [MetaForest](#) object, and applies different algorithms for variable selection.

Usage

```
preselect(x, replications = 100L, algorithm = "replicate", ...)
```

Arguments

<code>x</code>	Model to perform variable selection for. Accepts <code>MetaForest</code> objects.
<code>replications</code>	Integer. Number of replications to run for variable preselection. Default: 100.
<code>algorithm</code>	Character. Preselection method to apply. Currently, 'replicate', 'recursive', and 'bootstrap' are available.
<code>...</code>	Other arguments to be passed to and from functions.

Details

Currently, available methods under `algorithm` are:

replicate This simply replicates the analysis, which means the forest has access to the full data set, but the trees are grown on different bootstrap samples across replications (thereby varying monte carlo error).

bootstrap This replicates the analysis on bootstrapped samples, which means each replication has access to a different sub-sample of the full data set. When selecting this algorithm, cases are either bootstrap-sampled by study, or a new study column is generated, and a clustered MetaForest is grown (because some of the rows in the data will be duplicated), and this would lead to an under-estimation of the OOB error.

recursive Starting with all moderators, the variable with the most negative variable importance is dropped from the model, and the analysis re-run. This is repeated until only variables with a positive variable importance are left, or no variables are left. The proportion of final models containing each variable reflects its importance.

Value

An object of class 'mf_preselect'

Examples

```
## Not run:
data <- get(data(dat.bourassa1996))
data <- escalc(measure = "OR", ai = lh.le, bi = lh.re, ci = rh.le, di = rh.re,
              data = data, add = 1/2, to = "all")
data$mage[is.na(data$mage)] <- median(data$mage, na.rm = TRUE)
data[c(5:8)] <- lapply(data[c(5:8)], factor)
data$yi <- as.numeric(data$yi)
mf.model <- MetaForest(formula = yi ~ selection + investigator + hand_assess + eye_assess +
                       mage + sex,
                       data, study = "sample",
                       whichweights = "unif", num.trees = 300)
preselect(mf.model,
          replications = 10,
          algorithm = "bootstrap")

## End(Not run)
```

```
preselect_vars
```

Extract variable names from mf_preselect object

Description

Returns a vector of variable names from an `mf_preselect` object, based on a cutoff criterion provided.

Usage

```
preselect_vars(x, cutoff = NULL, criterion = NULL)
```

Arguments

x	Object of class mf_preselect.
cutoff	Numeric. Must be a value between 0 and 1. By default, uses .95 for bootstrapped preselection, and .1 for recursive preselection.
criterion	Character. Which criterion to use. See Details for more information. By default, uses 'ci' (confidence interval) for bootstrapped preselection, and 'p' (proportion) for recursive preselection.

Details

For criterion = 'p', the function evaluates the proportion of replications in which a variable achieved a positive (>0) variable importance. For criterion = 'ci', the function evaluates whether the lower bound of a confidence interval of a variable's importance across replications exceeds zero. The width of the confidence interval is determined by cutoff.

For recursive preselection, any variable not included in a final model is assigned zero importance.

Value

Character vector.

Examples

```
## Not run:
data <- get(data(dat.bourassa1996))
data <- escalc(measure = "OR", ai = lh.le, bi = lh.re, ci = rh.le, di = rh.re,
              data = data, add = 1/2, to = "all")
data$mage[is.na(data$mage)] <- median(data$mage, na.rm = TRUE)
data[c(5:8)] <- lapply(data[c(5:8)], factor)
data$yi <- as.numeric(data$yi)
preselected <- preselect(formula = yi~ selection + investigator + hand_assess + eye_assess +
                        mage +sex,
                        data, study = "sample",
                        whichweights = "unif", num.trees = 300,
                        replications = 10,
                        algorithm = "bootstrap")
preselect_vars(preselected)

## End(Not run)
```

```
print.summary.MetaForest
```

Prints summary.MetaForest object.

Description

Prints summary.MetaForest object.

Usage

```
## S3 method for class 'summary.MetaForest'
print(x, digits, ...)
```

Arguments

`x` an object used to select a method.

`digits` minimal number of significant digits, see `print.default`.

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

Examples

SimulateSMD	<i>Simulates a meta-analytic dataset</i>
-------------	--

Description

This function simulates a meta-analytic dataset based on the random-effects model. The simulated effect size is Hedges' G , an estimator of the Standardized Mean Difference. The functional form of the model can be specified, and moderators can be either normally distributed or Bernoulli-distributed. See Van Lissa, 2018, for a detailed explanation of the simulation procedure.

Usage

```
SimulateSMD(
  k_train = 20,
  k_test = 100,
  mean_n = 40,
  es = 0.5,
  tau2 = 0.04,
  moderators = 5,
  distribution = "normal",
  model = es * x[, 1]
)
```

Arguments

`k_train` Atomic integer. The number of studies in the training dataset. Defaults to 20.

`k_test` Atomic integer. The number of studies in the testing dataset. Defaults to 100.

`mean_n` Atomic integer. The mean sample size of each simulated study in the meta-analytic dataset. Defaults to 40. For each simulated study, the sample size n is randomly drawn from a normal distribution with mean `mean_n`, and sd `mean_n/3`.

es	Atomic numeric vector. The effect size, also known as beta, used in the model statement. Defaults to .5.
tau2	Atomic numeric vector. The residual heterogeneity. Defaults to 0.04.
moderators	Atomic integer. The number of moderators to simulate for each study. Make sure that the number of moderators to be simulated is at least as large as the number of moderators referred to in the model parameter. Internally, the matrix of moderators is referred to as "x". Defaults to 5.
distribution	Atomic character. The distribution of the moderators. Can be set to either "normal" or "bernoulli". Defaults to "normal".
model	Expression. An expression to specify the model from which to simulate the mean true effect size, mu. This formula may use the terms "es" (referring to the es parameter of the call to SimulateSMD), and "x[, 1]" (referring to the matrix of moderators, x). Thus, to specify that the mean effect size, mu, is a function of the effect size and the first moderator, one would pass the value model = es * x[, 1]. Defaults to es * x[, 1].

Value

List of length 4. The "training" element of this list is a data.frame with k_{train} rows. The columns are the variance of the effect size, v_i ; the effect size, y_i , and the moderators, X. The "testing" element of this list is a data.frame with k_{test} rows. The columns are the effect size, y_i , and the moderators, X. The "housekeeping" element of this list is a data.frame with $k_{\text{train}} + k_{\text{test}}$ rows. The columns are n, the sample size n for each simulated study; μ_i , the mean true effect size for each simulated study; and θ_i , the true effect size for each simulated study.

References

Van Lissa, C. J. (2020). Small sample meta-analyses: exploring heterogeneity using metaForest. In R. Van De Schoot & M. Miočević (Eds.), Small sample size solutions (open access): A guide for applied researchers and practitioners. CRC Press (pp.186–202). doi:10.4324/9780429273872-16
 Van Lissa, C. J. (2018). MetaForest: Exploring heterogeneity in meta-analysis using random forests. PsyArxiv. doi:10.31234/osf.io/myg6s

Examples

```
set.seed(8)
SimulateSMD()
SimulateSMD(k_train = 50, distribution = "bernoulli")
SimulateSMD(distribution = "bernoulli", model = es * x[,1] * x[,2])
```

 VarImpPlot

Plots variable importance for a MetaForest object.

Description

Plots variable importance for a MetaForest object.

Usage

```
VarImpPlot(mf, n.var = 30, sort = TRUE, ...)
```

Arguments

mf	MetaForest object.
n.var	Number of moderators to plot.
sort	Should the moderators be sorted from most to least important?
...	Parameters passed to and from other functions.

Value

A ggplot object.

Examples

```
set.seed(42)
data <- SimulateSMD()
mf.random <- MetaForest(formula = yi ~ ., data = data$training,
                        whichweights = "random", method = "DL",
                        tau2 = 0.0116)
VarImpPlot(mf.random)
VarImpPlot(mf.random, n.var = 2)
VarImpPlot(mf.random, sort = FALSE)
```

WeightedScatter	<i>Plots weighted scatterplots for meta-analytic data. Can plot effect size as a function of either continuous (numeric, integer) or categorical (factor, character) predictors.</i>
-----------------	--

Description

Plots weighted scatterplots for meta-analytic data. Can plot effect size as a function of either continuous (numeric, integer) or categorical (factor, character) predictors.

Usage

```
WeightedScatter(
  data,
  yi = "yi",
  vi = "vi",
  vars = NULL,
  tau2 = NULL,
  summarize = TRUE
)
```

Arguments

data	A data.frame.
yi	Character. The name of the column in data that contains the meta-analysis effect sizes. Defaults to "yi".
vi	Character. The name of the column in the data that contains the variances of the effect sizes. Defaults to "vi". By default, vi is used to calculate fixed-effects weights, because fixed effects weights summarize the data set at hand, rather than generalizing to the population.
vars	Character vector containing the names of specific moderator variables to plot. When set to NULL, the default, all moderators are plotted.
tau2	Numeric. Provide an optional value for tau2. If this value is provided, random-effects weights will be used instead of fixed-effects weights.
summarize	Logical. Should summary stats be displayed? Defaults to FALSE. If TRUE, a smooth trend line is displayed for continuous variables, using [stats::loess()] for less than 1000 observations, and [mgcv::gam()] for larger datasets. For categorical variables, box-and-whiskers plots are displayed. Outliers are omitted, because the raw data fulfill this function.

Value

A gtable object.

Examples

```
## Not run:
set.seed(42)
data <- SimulateSMD(k_train = 100, model = es * x[, 1] + es * x[, 2] + es *
  x[, 1] * x[, 2])$training
data$X2 <- cut(data$X2, breaks = 2, labels = c("Low", "High"))
data$X3 <- cut(data$X3, breaks = 2, labels = c("Small", "Big"))
WeightedScatter(data, summarize = FALSE)
WeightedScatter(data, vars = c("X3"))
WeightedScatter(data, vars = c("X1", "X3"))

## End(Not run)
```

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