Package 'PRIM'

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Description

Function for determining on one box out of a "peel"-object.

Usage

```
define_fixbox(prim, step)
```

Arguments

prim object of class "peel" from which the box should be defined.

step number of the step (i.e. row in the argument box of the "peel"-object.) of the

requested box.

Value

define_fixbox returns a object of class "fixbox", which is the basically same as a "peel"-object, but only contains one box. It is a list containing at least the following elements:

f target function of the selected box.

beta support of the selected box.

box a data. frame with one row defining the borders of the box. The columns with

"min." and "max." describe the lower and upper boundaries of the at least ordinal covariates. Therefore the value taken is the last one that is **not** included in

the box.

For the nominal variables there are columns for every category they can take. If the category is removed from the box the value FALSE is taken. The names of

these columns are structured like: <variable name>.<category>

For each variable with missing values (only if use_NAs = TRUE) there is also a column taking the value FALSE if the NAs of this variable are removed from the box. The names of these columns are structured like: <variable name>.NA

box_metric, box_nom, box_na

easier to handle definitions of the box for other functions

subset logical vector indicating the subset (i.e. the observations that lie in the box).

data_orig original dataset that was used for the peeling.

inbox 3

| inbox | Defining Subset lying in a Box | |
|-------|--------------------------------|--|
| | | |

Description

Help function that returns the subset of the whole data set which is contained in one box (fixbox). This function is needed in other functions like PRIM_peel_bs.

Usage

```
inbox(X, fixbox_metric = NULL, fixbox_nom = NULL, fixbox_na = NULL)
```

Arguments

```
X data.frame from which the subset should be taken.

fixbox_metric one row of a box_metric from a "peel"-object.

fixbox_nom one element of a box_nom list from a "peel"-object.

fixbox_na one element of a box_na list from a "peel"-object.
```

Details

The arguments fixbox_metric, fixbox_nom, fixbox_na can also come from a "fixbox"-object.

Value

inbox returns a logical vector indicating the observations that lie in the box.

| inter_diss |
|------------|
|------------|

Description

This Function calculates the interbox dissimilarity between two boxes defined on one data set.

Usage

```
inter_diss(fixbox1, fixbox2, data)
```

Arguments

| fixbox1 | first object of class "fixbox". |
|---------|---|
| fixbox2 | second object of class "fixbox". |
| data | original data used to calculate the supports needed for the interbox dissimilarity. If this argument is missing data_orig from fixbox1 is used as data. |

PRIM PRIM

Details

The interbox dissimilarity is a diagnostic tool of PRIM measuring the dissimilarity between two boxes B_k and B_l . It is defined as the difference between the smallest box B_{kl} that covers both boxes and the support of their union.

The interbox dissimilarity can assume values between 0 and 1. While nested boxes have a dissimilarity of 0, it gets bigger the more different the two boxes are.

plot.peel

Plotting a Trajectory

Description

This function plots the trajectory of a "peel"-object. In the trajectory the target functions evaluated on the data in the box are plotted against the supports beta.

Usage

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

Arguments

x object of class "pee1".... further arguments of the plot-function.

PRIM

Combined Function for the Patient Rule Induction Method (PRIM)

Description

This function is a automated implementation of PRIM as suggested by Friedman and Fisher (1999). It includes multiple peeling (PRIM_peel_bs), pasting (PRIM_paste) and the covering stretegy to find more than one box.

```
PRIM(formula, data, f_min, beta_min = 0.2, max_boxes = Inf,
   peel_alpha = seq(0.01, 0.4, 0.03), B = 0, target = mean,
   alter_crit = TRUE, use_NAs = TRUE, seed, print_position = FALSE,
   paste_alpha = 0.01, max_steps = 50, stop_by_dec = TRUE)
```

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Arguments

| formula | an object of class "formula" with a response but no interaction terms. It indicates the response over which the target function should be maximized and the covariates that are used for the later box definitions. |
|----------------|---|
| data | an object of class data. frame containing the variables named in the formula. |
| f_min | minimum target the final box must have. From all boxes, that fulfill this criterion, the one with the biggest support is taken after the peeling. If this argument is missing the box with the biggest target having at least a support of beta_min is taken. |
| beta_min | minimum support that one box must have. This proportion always refers to the whole data set. |
| max_boxes | maximum number of boxes to be found. |
| peel_alpha | vector of a sequence of different alpha-fractions used for the peelings. |
| В | number of bootstrap samples on which the peeling is applied to for each alpha. For B $=$ 0 no bootstraps are created. |
| target | target-function to be maximized. In most cases the mean is a useful target, although other functions like e.g. the median are also possible here. |
| alter_crit | logical. If TRUE the alternative criterion is used for peeling. I.e. "target/beta" is maximized during peeling instead of "target", so that large subboxes are not prefered to be peeled off. This is important especially in case of nominal covariates. |
| use_NAs | logical. If TRUE observations with missing values are included in the analysis. |
| seed | seed to be set before the first iteration. Only useful for $B > \emptyset$. |
| print_position | logical. If TRUE the current position of the algorithm is printed out. |
| paste_alpha | alpha-fraction that is pasted to the box at each pasting step |
| max_steps | maximum number of pasting steps the function should make. |
| stop_by_dec | logical. If TRUE the pasting stops if the target at one step is lower than the target of the last step. |

Details

This function repeats the peeling and pasting algorithm for the same settings of the metaparameters until a stop ctiterion is reached. After each iteration the observations already included in a box are removed from the data, on which the next box is built. This strategy is called covering. This iteration stops if either max_boxes is reached or if the target function of the "best" box is lower than the overall target.

In each iteration step this function does a multiple peeling characterized by the sequenz alpha_peel and B. From the peeling output the box defined by beta_min and f_min is chosen. After that the pasting function seeks for boxes with bigger supports and bigger targets and takes the one with the highest target function within the box.

The function can also cope with survival outcomes (Surv-object). Therefore the hazard rate is used as target function as suggested in Ott and Hapfelmeier (2017). The value of the input parameter target is ignored in this case.

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Value

PRIM returns an object of class "prim", which is a list containing the following components:

f vector of the target functions evaluated on each box. The last element is the

target of all observations not lying in a box.

beta vector of the supports of each box. The last element is the fraction of observa-

tions not lying in a box.

box a data. frame defining the borders of the boxes. Each row belongs to one box.

The columns with "min." and "max." describe the lower and upper boundaries of the at least ordinal covariates. Therefore the value taken is the last one that is

not included in the current box.

For the nominal variables there are columns for every category they can take. If the category is removed from the box the value FALSE is taken. The names of

these columns are structured like: <variable name>.<category>

For each variable with missing values (only if use_NAs = TRUE) there is also a column taking the value FALSE if the NAs of this variable are removed from the current box. The names of these columns are structured like: <variable name>.NA

box_metric, box_nom, box_na

easier to handle definitions of the boxes for other functions

subsets list of logical vectors indicating the subsets (i.e. the observations that lie in

each box)

fixboxes list of all fixbox'es defining the final boxes.

data_orig original dataset that is used.

References

Friedman, J. H. and Fisher, N. I., 'Bump hunting in high-dimensional data', Statistics and Computing 9 (2) (1999), 123-143

Ott, A. and Hapfelmeier, A., 'Nonparametric Subgroup Identification by PRIM and CART: A Simulation and Application Study', Computational and Mathematical Methods in Medicine, vol. 2017 (2017), 17 pages, Article ID 5271091

See Also

```
PRIM_peel_bs, PRIM_paste, define_fixbox
```

Examples

```
# generating random data:
set.seed(123)
n <- 500
x1 <- runif(n = n, min = -1)
x2 <- runif(n = n, min = -1)
x3 <- runif(n = n, min = -1)
cat <- as.factor(sample(c("a","b","c", "d"), size = n, replace = TRUE))
wsk <- (1-sqrt(x1^2+x2^2)/sqrt(2))
y <- as.logical(rbinom(n = n, prob = wsk, size = 1))
dat <- cbind.data.frame(y, x1, x2, x3, cat)
#plot(dat$x1, dat$x2, col=dat$y+1, pch=16)
remove(x1, x2, x3, y, wsk, cat, n)
# apply the PRIM function to find the best boxes with a support of at least 0.1:</pre>
```

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```
p <- PRIM(y^{-}., data=dat, beta_min = 0.1, max_boxes = 3) p
```

| Function | Pasting-Function | PRIM_paste |
|----------|------------------|------------|
|----------|------------------|------------|

Description

This function is an implementation of the Pasting-Algorithm as suggested by Friedman and Fisher (1999). In each iteration the fraction alpha is pasted to one edge of the current box.

Usage

```
PRIM_paste(fixbox, paste_alpha = 0.01, max_steps = 50, stop_by_dec = TRUE)
```

Arguments

fixbox an object of class fixbox, which was defined after the peeling function and now should be used for pasting.

paste_alpha alpha-fraction that is pasted to the box at each iteration.

max_steps maximum number of pasting steps the function should make.

stop_by_dec logical. If TRUE the pasting stops if the target at one step is lower than the target

of the last step.

Details

The outcome of this function is also a "peel"-object, because it has basically the same structure as the outcome of the peeling functions. The only difference is, that pasting goes from small supports to bigger ones, while by peeling its the other way round.

Value

PRIM_paste returns an object of class "peel", which is a list containing at least the following components:

f vector of the target functions evaluated on the box at each pasting step.

beta vector of the supports beta of the boxes at each pasting step.

box a data. frame defining the borders of the boxes. Each row belongs to one past-

ing step. The columns with "min." and "max." describe the lower and upper boundaries of the at least ordinal covariates. Therefore the value taken is the last

one that is **not** included in the current box.

For the nominal variables there are columns for every category they can take. If the category is removed from the box the value FALSE is taken. The names of

these columns are structured like: <variable name>.<category>

For each variable with missing values (only if use_NAs = TRUE) there is also a column taking the value FALSE if the NAs of this variable are removed from the current box. The names of these columns are structured like: <variable name>.NA

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```
box_metric, box_nom, box_na
```

easier to handle definitions of the boxes for other functions

subsets list of logical vectors indicating the subsets at each pasting step (i.e. the ob-

servations that lie in the box)

data_orig original dataset that is used (extracted from fixbox).

References

Friedman, J. H. and Fisher, N. I., 'Bump hunting in high-dimensional data', Statistics and Computing 9 (2) (1999), 123-143

Ott, A. and Hapfelmeier, A., 'Nonparametric Subgroup Identification by PRIM and CART: A Simulation and Application Study', Computational and Mathematical Methods in Medicine, vol. 2017 (2017), 17 pages, Article ID 5271091

Examples

```
# generating random data:
set.seed(123)
n <- 500
x1 \leftarrow runif(n = n, min = -1)
x2 \leftarrow runif(n = n, min = -1)
x3 \leftarrow runif(n = n, min = -1)
cat <- as.factor(sample(c("a","b","c", "d"), size = n, replace = TRUE))</pre>
wsk <- (1-sqrt(x1^2+x2^2)/sqrt(2))
y <- as.logical(rbinom(n = n, prob = wsk, size = 1))
dat <- cbind.data.frame(y, x1, x2, x3, cat)</pre>
#plot(dat$x1, dat$x2, col=dat$y+1, pch=16)
remove(x1, x2, x3, y, wsk, cat, n)
# apply the PRIM_peel function:
prim <- PRIM_peel(y ~ ., data = dat, beta_min = .01, peel_alpha = .1)</pre>
abline(h=prim$f[17], v=prim$beta[17]) # box decided to paste
fix <- define_fixbox(prim, 17) # define fixbox</pre>
# apply the PRIM_paste function:
paste <- PRIM_paste(fix, stop_by_dec = FALSE)</pre>
head(cbind(paste$box, paste$f, paste$beta))
```

PRIM_peel

Peeling-Function

Description

This function is an implementation of the (singular) Peeling-Algorithm as suggested by Friedman and Fisher (1999). In each iteration the fraction alpha is peeled from one edge of the current box.

```
PRIM_peel(formula, data, peel_alpha = 0.05, beta_min = 0.01,
  target = mean, alter_crit = TRUE, use_NAs = TRUE)
```

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Arguments

formula an object of class "formula" with a response but no interaction terms. It indicates the response over which the target function should be maximized and the covariates that are used for the later box definitions. If this argument is missing,

the argument data is used as the model.frame.

data an object of class data. frame containing the variables named in the formula.

peel_alpha alpha-fraction used for the peeling.

beta_min minimum support that one Box should have (stop-criterion).

target target-function to be maximized. In most cases the mean is a useful target,

although other functions like e.g. the median are also possible here.

alter_crit logical. If TRUE the alternative criterion is used for peeling. I.e. "target/beta"

is maximized during peeling instead of "target", so that large subboxes are not prefered to be peeled off. This is important especially in case of nominal covari-

ates.

use_NAs logical. If TRUE observations with missing values are included in the analysis.

Details

The outcome of the formula can either be numeric, logical or a survival object (see Surv). If it is a survival object the target is set to the number of events per amount of time.

This function is the main part of the multiple Version PRIM_peel_bs.

Value

PRIM_peel returns an object of class "peel", which is a list containing at least the following components:

f vector of the target functions evaluated on the box at each peeling step.

beta vector of the supports beta of the boxes at each peeling step.

box a data.frame defining the borders of the boxes. Each row belongs to one peeling step. The columns with "min." and "max." describe the lower and upper

boundaries of the at least ordinal covariates. Therefore the value taken is the last

one that is **not** included in the current box.

For the nominal variables there are columns for every category they can take. If the category is removed from the box the value FALSE is taken. The names of

these columns are structured like: <variable name>.<category>

For each variable with missing values (only if use_NAs = TRUE) there is also a column taking the value FALSE if the NAs of this variable are removed from the current box. The names of these columns are structured like: <variable name>.NA

box_metric, box_nom, box_na

easier to handle definitions of the boxes for other functions

subsets list of logical vectors indicating the subsets at each peeling step (i.e. the ob-

servations that lie in the box)

data_orig original dataset that is used for the peeling.

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References

Friedman, J. H. and Fisher, N. I., 'Bump hunting in high-dimensional data', Statistics and Computing 9 (2) (1999), 123-143

Ott, A. and Hapfelmeier, A., 'Nonparametric Subgroup Identification by PRIM and CART: A Simulation and Application Study', Computational and Mathematical Methods in Medicine, vol. 2017 (2017), 17 pages, Article ID 5271091

Examples

```
# generating random data:
set.seed(123)
n <- 500
x1 \leftarrow runif(n = n, min = -1)
x2 \leftarrow runif(n = n, min = -1)
x3 \leftarrow runif(n = n, min = -1)
cat <- as.factor(sample(c("a","b","c", "d"), size = n, replace = TRUE))</pre>
wsk \leftarrow (1-sqrt(x1^2+x2^2)/sqrt(2))
y <- as.logical(rbinom(n = n, prob = wsk, size = 1))
dat <- cbind.data.frame(y, x1, x2, x3, cat)</pre>
#plot(dat$x1, dat$x2, col=dat$y+1, pch=16)
remove(x1, x2, x3, y, wsk, cat, n)
# apply the PRIM_peel function:
prim <- PRIM_peel(formula=y ~ ., data=dat, beta_min = .01, peel_alpha = .1)</pre>
plot(prim) # trajectory
prim$box # box definitions
```

PRIM_peel_bs

Multiple Peeling-Function

Description

This function is an implementation of the multiple Peeling-Algorithm as suggested by Friedman and Fisher (1999). The singular peeling function PRIM_peel is repeated for different alpha's and bootstrap samples out of the original data.

Usage

```
PRIM_peel_bs(formula, data, peel_alpha = seq(0.01, 0.4, 0.03), B = 0,
  beta_min = 0.01, target = mean, alter_crit = TRUE, use_NAs = TRUE,
  seed, print_position = TRUE)
```

Arguments

| formula | an object of class "formula" with a response but no interaction terms. It indi- |
|------------|---|
| | cates the response over which the target function should be maximized and the |
| | covariates that are used for the later box definitions. |
| data | an object of class data. frame containing the variables named in the formula. |
| peel_alpha | vector of a sequence of different alpha-fractions used for the peelings. |

PRIM_peel_bs

B number of bootstrap samples on which the peeling is applied to for each alpha.

For $B = \emptyset$ no bootstraps are created.

beta_min minimum support that one Box should have (stop-criterion).

target target-function to be maximized. In most cases the mean is a useful target,

although other functions like e.g. the median are also possible here.

alter_crit logical. If TRUE the alternative criterion is used for peeling. I.e. "target/beta"

is maximized during peeling instead of "target", so that large subboxes are not prefered to be peeled off. This is important especially in case of nominal covari-

ates.

use_NAs logical. If TRUE observations with missing values are included in the analysis.

seed seed to be set before the first iteration. Only useful for B > 0. print_position logical. If TRUE the current position of the algorithm is printed out.

Details

The outcome of the formula can either be numeric, logical or a survival object (see Surv). If it is a survival object the target is set to the number of events per amount of time.

The output of this function can become very large because all outputs of the singular peel function PRIM_peel are put together in one output. Therefore it is usefull to remove all the dominated boxes (see remove_dominated).

Value

PRIM_peel_bs returns an object of class "peel", which is a list containing at least the following components:

f vector of the target functions evaluated on the box at each peeling step.

beta vector of the supports beta of the boxes at each peeling step.

box a data. frame defining the borders of the boxes. Each row belongs to one peel-

ing step. The columns with "min." and "max." describe the lower and upper boundaries of the at least ordinal covariates. Therefore the value taken is the last

one that is **not** included in the current box.

For the nominal variables there are columns for every category they can take. If the category is removed from the box the value FALSE is taken. The names of

these columns are structured like: <variable name>.<category>

For each variable with missing values (only if use_NAs = TRUE) there is also a column taking the value FALSE if the NAs of this variable are removed from the

current box. The names of these columns are structured like: <variable name>.NA

box_metric, box_nom, box_na

easier to handle definitions of the boxes for other functions

subsets list of logical vectors indicating the subsets at each peeling step (i.e. the ob-

servations that lie in the box)

data_orig original dataset that is used for the peeling.

References

Friedman, J. H. and Fisher, N. I., 'Bump hunting in high-dimensional data', Statistics and Computing 9 (2) (1999), 123-143

Ott, A. and Hapfelmeier, A., 'Nonparametric Subgroup Identification by PRIM and CART: A Simulation and Application Study', Computational and Mathematical Methods in Medicine, vol. 2017 (2017), 17 pages, Article ID 5271091

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See Also

```
remove_dominated, PRIM_peel, PRIM_paste, PRIM
```

Examples

```
# generating random data:
set.seed(123)
n <- 500
x1 \leftarrow runif(n = n, min = -1)
x2 \leftarrow runif(n = n, min = -1)
x3 \leftarrow runif(n = n, min = -1)
cat <- as.factor(sample(c("a","b","c", "d"), size = n, replace = TRUE))</pre>
wsk <- (1-sqrt(x1^2+x2^2)/sqrt(2))
y <- as.logical(rbinom(n = n, prob = wsk, size = 1))
dat <- cbind.data.frame(y, x1, x2, x3, cat)</pre>
#plot(dat$x1, dat$x2, col=dat$y+1, pch=16)
remove(x1, x2, x3, y, wsk, cat, n)
# apply the PRIM_peel_bs function:
prim <- PRIM_peel_bs(formula=y ~ ., data=dat, beta_min = .01)</pre>
plot(prim) # multiple trajectory
head(prim$box) # box definitions
```

print.PRIM

Printing a "PRIM"-Object

Description

This function prints a PRIM-object. It is a method for the generic function print of class "PRIM".

Usage

```
## S3 method for class 'PRIM'
print(x, ...)
```

Arguments

x object of class "PRIM".... further arguments of the print-function.

relfreq

Relative Frequency Plots

Description

This Function creates the relative frequency plots of a box defined by define_fixbox.

```
relfreq(fixbox, plot_together = TRUE, ...)
```

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Arguments

```
fixbox object of class "fixbox" to be used to create the relative frequency plots.

plot_together logical. If TRUE all plots are drawn together in one figure by par(mfrow = ...).

further arguments of the functions plot and barplot.
```

Details

Relative frequency plots are a diagnostic tool for a result of PRIM (fixbox). These are histograms or barplots illustrating ratios of distributions r_{jk} for each variable x_j . This ratio is defined by the distribution of x_j in one box B_k divided by the distribution of this variable over the whole data set.

See Also

```
define_fixbox, PRIM
```

Examples

```
# generating random data:
set.seed(123)
n <- 500
x1 \leftarrow runif(n = n, min = -1)
x2 \leftarrow runif(n = n, min = -1)
x3 \leftarrow runif(n = n, min = -1)
cat <- as.factor(sample(c("a","b","c", "d"), size = n, replace = TRUE))
wsk <- (1-sqrt(x1^2+x2^2)/sqrt(2))
y <- as.logical(rbinom(n = n, prob = wsk, size = 1))
dat <- cbind.data.frame(y, x1, x2, x3, cat)</pre>
plot(dat$x1, dat$x2, col=dat$y+1, pch=16)
remove(x1, x2, x3, y, wsk, cat, n)
# apply the PRIM function to find the best box with a support of at least 0.1:
p <- PRIM(y~., data=dat, beta_min = 0.1, max_boxes = 1, print_position = FALSE)
# relative frequency plots:
relfreq(p$fixboxes[[1]])
```

remove_dominated

Remove Dominated Boxes from a Peeling Output

Description

This function removes all boxes from one "peel"-object that are dominated by another box. I.e. there is at least one other box with a better output and a bigger support beta.

```
remove_dominated(prim, sort = TRUE, dup.rm = TRUE)
```

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Arguments

| prim | object of class "pee1". |
|--------|---|
| sort | logical. If TRUE the boxes in the output are sorted decreasing by their box supports. |
| dup.rm | logical. Tf TRUE duplicated boxes in the output are removed. Boxes are dupli- |

cated if another box has exactly the same target and support.

Details

Dominated boxes mainly occur in outputs of the multiple peeling function (PRIM_peel_bs). So this function is practically only useful for "peel"-objects beeing the results of such a function. Without the dominated boxes the object gets much smaller by keeping all the relevant boxes. Also the trajectory (see plot.peel) without the not useful dominated boxes looks much clearer.

Value

This function returns another "peel"-object having the same structure as the input object, just without the dominated boxes.

See Also

```
PRIM_peel_bs, plot.peel
```

var_select

Sequential Relevance of Box-defining Variables

Description

This function creates a table of the sequential relevance of the variables used in the definition of a box defined by define_fixbox.

Usage

```
var_select(fixbox)
```

Arguments

fixbox object of class "fixbox" to be used to create the table of the sequential variable

relevances.

Details

This function does a sequential removal off all variables defining one box. In each iteration every variable left in the box definition is tried to be completley left out of the definition. From all variables this one is removed, which causes the least decrease of the target function evaluated on all observations lying in the box.

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Value

var_select returns a list with the following components:

table showing the sequential relevances of the variables used in the definition of

the fixbox.

fixboxes list of the boxes defined at every iteration step (by removing of one variable).

See Also

```
define_fixbox, PRIM
```

Examples

```
# generating random data:
set.seed(123)
n <- 500
x1 \leftarrow runif(n = n, min = -1)
x2 \leftarrow runif(n = n, min = -1)
x3 \leftarrow runif(n = n, min = -1)
cat <- as.factor(sample(c("a","b","c", "d"), size = n, replace = TRUE))</pre>
wsk <- (1-sqrt(x1^2+x2^2)/sqrt(2))
y <- as.logical(rbinom(n = n, prob = wsk, size = 1))
dat <- cbind.data.frame(y, x1, x2, x3, cat)</pre>
plot(dat$x1, dat$x2, col=dat$y+1, pch=16)
remove(x1, x2, x3, y, wsk, cat, n)
# apply the PRIM function to find the best box with a support of at least 0.1:
p <- PRIM(y~., data=dat, beta_min = 0.1, max_boxes = 1, print_position = FALSE)</pre>
# sequential variable relevances:
var_select(p$fixboxes[[1]])$tab
```

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