Package ‘quint’

June 16, 2020

Type Package

Title Qualitative Interaction Trees

Version 2.1.0

Date 2020-06-12

Maintainer Elise Dusseldorp <elise.dusseldorp@fsw.leidenuniv.nl>

Description Grows a qualitative interaction tree. Quint is a tool for subgroup analysis, suitable for data from a two-arm randomized controlled trial.

URL https://doi.org/10.3758/s13428-015-0594-z

Depends R (>= 3.0.2), partykit, Formula, rpart, stats, grid, utils, graphics

Suggests knitr, rmarkdown

VignetteBuilder knitr

License GPL-2

LazyData TRUE

RoxygenNote 7.1.0

Author Elise Dusseldorp [aut, cre, cph], Lisa Doove [aut], Jeanne van de Put [aut], Cor Ninaber [ctb] (supported with the plot function), Iven Van Mechelen [aut, cph], Juan Claramunt Gonzalez [aut]

NeedsCompilation no

Repository CRAN

Date/Publication 2020-06-15 23:50:03 UTC

R topics documented:

quint-package ........................................................................ 2
bcrp .................................................................................. 3
plot.quint ............................................................................. 4
R package for Qualitative Treatment-Subgroup Interactions

Description

When two treatment alternatives (say A and B) are available for some problem, one may be interested in qualitative treatment-subgroup interactions. Such interactions imply the existence of subgroups of persons (patients) which are such that in one subgroup Treatment A outperforms Treatment B, whereas the reverse holds in another subgroup. Obviously, this type of interactions is crucial for optimal treatment assignment of future patients. Given baseline characteristics and outcome data from a two-arm Randomized Controlled Trial (RCT), QUalitative INteraction Trees (QUINT) is a tool to identify subgroups that are involved in meaningful qualitative treatment-subgroup interactions. The result of QUINT is a tree that partitions the total group of participants (patients) on the basis of their baseline characteristics into three subgroups (i.e., partition classes): Subgroup 1: Those for whom Treatment A is better than Treatment B (P1), Subgroup 2: Those for whom Treatment B is better than Treatment A (P2), and Subgroup 3: Those for whom it does not make any difference (P3).

Details

Package: quint
Type: Package
Version: 2.2.0
Date: 2020-02-03
License: GPL

This method is suitable for a continuous outcome variable. From version 1.2 onwards the baseline variables for growing a tree may have numerical or integer values (such as continuous, ordinal or dichotomous variables) or may be nominal (categorical variables with factors). Previously only numerical or dichotomous variables were supported. Another new feature of this version is that the output of a quint object can now also display results for either the raw difference in means or the effect size with corresponding standard error. This depends on the criterion specified. Furthermore a predict function predict.quint is newly included in this package. The final new feature is a validate function quint.validate for estimating the bias (i.e., optimism) of a grown QUINT tree. From version 2.0 onwards the qualitative treatment-subgroup interaction is checked during the
prune of the tree and not at the beginning of QUINT. Furthermore, it is possible to obtain outcomes from the summary and predict functions when the tree only contains the root node.

The core function of the package is `quint`.

**Author(s)**

Maintainer: Elise Dusseldorp <elise.dusseldorp@fsw.leidenuniv.nl>

**References**


**See Also**

`quint`, `summary.quint`, `quint.control`, `prune.quint`, `predict.quint`, `quint.validate`, `quint.bootstrapCI`

---

**bcrp**

*Breast Cancer Recovery Project*

**Description**

Data from a three-arm randomized controlled trial. Women with early-stage breast cancer were randomly assigned to a nutrition intervention (n = 85), an education intervention (n = 83) or standard care (n = 84). They were measured before and after treatment. These data contain the baseline measurement and the 9-month follow-up.

**Usage**

`bcrp`

**Format**

A data frame with 252 observations on the following 14 variables:

- `physt1` physical functioning (from SF-36) at baseline.
- `cesdt1` depression score (CESD) at baseline.
- `physt3` physical functioning (from SF-36) at 9 months follow-up.
- `cesdt3` depression score (CESD) at 9 months follow-up.
- `negsoct1` negative social interaction at baseline.
uncomt1 unmitigated communion at baseline.
disopt1 dispositional optimism at baseline.
comorbid number of comorbidities (e.g. diabetes, migraines, arthritis, or angina).
age age at baseline.
wcht1 weight change since diagnosis: yes [1] or no [0].
nationality Caucasian [1] or not [0].
marital married [1] or not [0].
text treatment extensiveness index: lumpectomy without or with one form of adjuvant therapy (radiation or chemo) [-1.77], lumpectomy with radiation and chemotherapy [0.26], mastectomy without or with lumpectomy, and without or with one form of adjuvant therapy [0.56], mastectomy without or with lumpectomy, and radiation and chemotherapy [2.59].
cond experimental condition: nutrition [1], education [2] or standard care [3].

Details

IMPORTANT: for questions about these data contact Elise Dusseldorp: elise.dusseldorp@fsw.leidenuniv.nl.

Source

The authors thank M.F. Scheier for making his data available.

References


---

**plot.quint**

*Visualisation of a Qualitative Interaction Tree*

**Description**

Plot function for a *quint* object. The plot shows the result of *quint*: a binary tree with (a) splitting variable(s) and split point(s). The colors of the leaves of the tree correspond to the final subgroups: Subgroup 1 (P1), those patients for whom the mean treatment outcome (Y) is higher for Treatment A than B, is GREEN; Subgroup 2 (P2), those patients for whom the mean treatment outcome (Y) is higher for Treatment B than A, is RED, and Subgroup 3 (P3), those for whom the mean treatment
outcome (Y) is about the same for both treatments, is GREY. Within the leaves the effect size \( d \) is displayed, with its 95 percent confidence interval. This effect size is the standardized mean difference between Treatment A and B. The plot function uses the plot method from the package \texttt{partykit} of Hothorn & Zeileis (2013).

**Usage**

```r
## S3 method for class 'quint'
plot(x, digits = 2, 
```

**Arguments**

- \( x \): fitted tree of class \texttt{quint}.
- \( digits \): specified number of decimal places of the splitpoints in the graph (default is 2).
- \( ... \): additional arguments to be passed.

**Details**

For categorical variables we recommend to use short names for levels to avoid overlapping labels at split points.

**Author(s)**

Cor Ninaber and Elise Dusseldorp

**References**


**See Also**

- \texttt{quint,quint.control,bcrp}

**Description**

Predicts for (new) subjects the treatment subgroups (P1, P2 or P3) based on a fitted \texttt{quint} object. The meaning of the subgroups are based on the two treatment categories used to fit the \texttt{quint} object.

```r
## S3 method for class 'quint'
predict(object, newdata, type = "class", 
```
Arguments

object an object of the class “quint”.

newdata a data frame with data on new subjects for whom predictions should be made. The data frame should contain at least the variables used in the splits of the fitted tree. It is not necessary to include the treatment variable.

type character string denoting the type of predicted object to be returned. The default is set to type="class": a vector with predicted treatment subgroup classes per subject is returned. If set to "matrix", a matrix is returned with the leaf and corresponding node of the tree to which a subject is assigned.

... optional additional arguments.

Value

One of the following objects is returned depending on output type specified in the function:

If type="class": vector of predicted treatment classes for every individual in the data set. Returns NA for subjects with missing values on one or more of the splitting variables.

If type="matrix": a matrix with predicted locations of subjects within the fitted tree. The leaf numbers are in the first column and the corresponding node numbers in the second column. Returns NA for subjects with missing values on one or more of the splitting variables.

See Also

quint, prune.quint

Examples

data(bcrp)
formula1<- I(cesdt1-cesdt3)~cond | nationality+marital+wcht1+age+trext+comorbid+disopt1+uncomt1+negsoct1

set.seed(10)
control1<-quint.control(maxl=5,B=2)
quint1<-quint(formula1, data= subset(bcrp,cond<3),control=control1) #Grow a QUINT tree

prquint1<-prune(quint1) #Prune QUINT tree to optimal size

#Predict for the same data set the treatment classes for patients individually:
predquint1<-predict(prquint1, newdata=subset(bcrp,cond<3), type='class')
predquint1
**Description**

Determines the optimally pruned size of the tree by applying the one standard error rule to the results from the bias-corrected bootstrap procedure. At the end of the pruning procedure, it is checked whether the pruned tree satisfies the qualitative interaction condition. If this condition is not met, it is concluded that there is no qualitative treatment-subgroup interaction and a tree containing only the root node is returned.

**Usage**

```r
## S3 method for class 'quint'
prune(tree, pp = 1, ...)
```

**Arguments**

- `tree`: fitted tree of the class `quint`.
- `pp`: pruning parameter, the constant \(c\) to be used in the \(c\) standard error rule. The default value is 1.
- `...`: optional additional arguments.

**Details**

The pruning algorithm of `quint` is explained in Dusseldorp & Van Mechelen (2014), Appendix B of the online supplementary material. It is based on the bias-corrected bootstrap pruning procedure (Le Blanc & Crowley, 1993) and the one standard error rule (Breiman, Friedman, Olshen, & Stone, 1984). The one standard error rule for `quint` uses the estimates of the bias-corrected criterion value \(C\) and its standard error for each value of \(L\) (= maximum number of leaves). The optimally pruned tree corresponds to the smallest tree with a bias-corrected \(C\) higher or equal to the maximum bias-corrected \(C\) minus its standard error.

**Value**

Returns an object of class `quint`. The number of leaves of this object is equal to the optimally pruned size of the tree.

**References**


See Also

quint.control, quint.quint.bootstrapCI

Examples

data(bcrp)
formula2 <- I(cesdt1-cesdt3)~cond |age+trext+uncomt1+disopt1+negsoct1
#Adjust the control parameters only to save computation time in the example;
#The default control parameters are preferred
control2 <- quint.control(maxl=5,B=2)
set.seed(2) #this enables you to repeat the results of the bootstrap procedure
quint2 <- quint(formula2, data= subset(bcrp,cond<3),control=control2)
quint2pr <- prune(quint2)
summary(quint2pr)

Description

This is the core function of the package. It performs a subgroup analysis by QUalitative INteraction
Trees (QUINT; Dusseldorp & Van Mechelen, 2014) and is suitable for data from a two-arm ran-
donized controlled trial. Ingredients of the analysis are: one continuous outcome variable \(Y\) (the
effect variable), one dichotomous treatment variable \(T\) (indicating two treatment conditions, e.g., A
and B), and several background characteristics \(X_1, \ldots, X_J\). These background characteristics are
measured at baseline and may have a numeric or ordinal measurement level (i.e., in R a numeric or
integer variable) or a nominal measurement level (i.e., in R a factor). They are used to identify the
following subgroups (i.e., partition classes): Subgroup 1: Those patients for whom Treatment A is
better than Treatment B (P1); Subgroup 2: Those for whom Treatment B is better than Treatment A
(P2), and Subgroup 3: Those for whom it does not make any difference (P3).

Usage

quint(formula, data, control = NULL)

Arguments

formula a description of the model to be fit. The format is \(Y \sim T \mid X_1 + \ldots + X_J\), where
the variable before the | represents the dichotomous treatment variable \(T\) and the
variables after the | are the baseline characteristics used for partitioning. If the
data are in the order \(Y, T, X_1, \ldots, X_J\), no formula is needed. The lay-out of this
formula is based on Zeileis & Croissant (2010).

data a dataframe containing the variables in the model. The treatment variable can
be a numeric or a factor variable with two values (or levels).
control a list with control parameters as returned by quint.control.
Details
The method QUINT uses a sequential partitioning algorithm. The algorithm starts with a tree consisting of a single node, that is, the root node containing all patients. Next, it follows a stepwise binary splitting procedure. This procedure implies that in each step a node, a baseline characteristic, a split of that characteristic, and an assignment of the leaves of the current tree to partition classes 1, 2, and 3 (P1 to P3) are chosen that maximize the partitioning criterion. Note that this means that after each split, all leaves of the tree are re-assigned afresh to the partition classes P1, P2, and P3.

Value
Returns an object of class quint with components:

- **call**: the call that created the object.
- **crit**: the partitioning criterion used to grow the tree. The default is the Effect size criterion. Use crit="dm" for the Difference in means criterion.
- **control**: the control parameters used in the analysis.
- **fi**: the fit information of the final tree.
- **si**: the split information of the final tree.
- **li**: the leaf information of the final tree. Treatment A is denoted with \( T=1 \), and treatment B is denoted with \( T=2 \). Can display either the output for Difference in Means (crit='dm') or Cohen’s \( d \) effect size (crit='es').
- **data**: the data used to grow the tree.
- **orig_data**: the original data used as input.
- **nind**: an \( N \times L \) matrix indicating leaf membership.
- **siboot**: an \( L \times 9 \times B \) array with split information for each bootstrap sample: \( C\text{\_boot} = \) value of \( C \); \( C\text{\_compdif} = \) value of Difference in treatment outcome component; checkdif = indicates if pooled Difference in treatment outcome component in test set (i.e., original sample) is positive, with values: 0 = yes, 1 = negative in P1, 2 = negative in P2, 3 = negative in P1 and P2; \( C\text{\_compcard} = \) value of Cardinality component; checkcard = indicates if value of pooled cardinality in test set is zero, with values: 0 = no, 1 = zero in P1, 2 = zero in P2, 3 = zero in P1 and P2; opt = value of optimism (\( C\text{\_boot}-C\text{\_orig} \)).
- **indexboot**: an \( N \times B \) matrix indicating bootstrap sample membership.
- **formula**: a description of the model to be fit.
- **pruned**: a boolean indicating whether the tree has been already pruned or not.

References


See Also

summary.quint, quint.control, prune.quint, bcpr, quint.bootstrapCI

Examples

#EXAMPLE with data from the Breast Cancer Recovery Project
data(bcpr)
#Start with expliciting the model for quint
#The outcome Y is a change score between timepoint 3 and timepoint 1
#A positive Y value indicates an improvement in depression (i.e., a decrease)
formula1<- I(cesdt1-cesdt3)~cond | nationality+marital+wcht1+age+
trext+comorbid+disopt1+uncomt1+negsoct1

#Perform a quint analysis
#The BCRP data contain 3 conditions. Quint only works now for 2 conditions.
#For the example, we disregard the control condition
#To save computation time, we also adjust the control parameters
set.seed(2)
control1<-quint.control(maxl=5,B=2) #The recommended number of bootstraps is 25.
quint1<-quint(formula1, data= subset(bcpr,cond<3),control=control1)
quint1pr<-prune(quint1)

#Inspect the main results of the analysis:
summary(quint1pr)

#plot the tree
plot(quint1pr)

quint.bootstrapCI

Bootstrap method to compute confidence intervals for Qualitative Interaction Trees (Quint)

Description

A bootstrap algorithm based on Loh et al. (2015) to estimate the confidence intervals of the difference in mean outcome between the two treatments in each leaf.

Usage

quint.bootstrapCI(tree, n_boot, boot_r = 1)
Arguments

- `tree` - a (pruned) quint object of class `quint`.
- `n_boot` - number of bootstrap samples.
- `boot_r` - bootstrap sample size expressed as proportion of total sample size. Default value is 1.

Details

The details of this validation procedure are described in "Instability of QUalitative INteraction Trees: Quantifying uncertainty in decision trees." (https://openaccess.leidenuniv.nl/handle/1887/83059)

Value

Returns two lists: A first one ($tree) containing an object of the class `quint`, and a list ($bootinfo) with estimates obtained from the bootstrap procedure containing the following components:

- `nleaves` - vector containing the number of leaves in each of the estimated trees in the bootstrap samples.
- `meanT_1` - a matrix containing for each bootstrap sample (= rows) the mean outcome for Treatment A (T=1) in each leaf of the input quint tree (= columns) using the subjects in the intersection.
- `meanT_2` - a matrix containing for each bootstrap sample (= rows) the mean outcome for Treatment B (T=2) in each leaf of the input quint tree (= columns) using the subjects in the intersection.
- `meandif` - a matrix containing the difference in means between Treatment A and Treatment B in each leaf for each bootstrap sample.
- `bias_est` - vector containing the bias in each leaf of the quint tree.
- `meanboot` - vector containing the bootstrap estimates of the difference of means between treatments in each leaf.
- `CIs` - vector containing the confidence intervals of the estimate of the difference of means between treatments in each leaf.
- `se_est` - vector containing the new estimates of the standard error of the difference of means between treatments in each leaf.

References


See Also

- `quint.prune.quint`
- `quint.control`
Examples

```r
## Not run: data(bcrp)
formula1<- I(cesdt1-cesdt3)~cond | nationality+marital+wcht1+age+
       trext+comorbid+disopt1+uncomt1+negsoct1

set.seed(10)
control1<-quint.control(maxl=5, B=2, crit="dm")
quint1<-quint(formula1, data= subset(bcrp,bcrp$cond<3),control=control1) #Grow a QUINT tree

prquint1<-prune(quint1) #Prune tree to optimal size

bootquint1<-quint.bootstrapCI(prquint1, n_boot = 5) #apply the bootstrap procedure
#
summary of the tree with the new standard errors obtained from the bootstrap procedure
summary(bootquint1$tree)

#all results of the bootstrap procedure
bootquint1$bootinfo

#plot wiht 95% confidence intervals using the new standard errors
plot(bootquint1$tree)
## End(Not run)
```

quint.control

Control Parameters for QUINT Algorithm

Description

Various parameters that control aspects of the “quint” algorithm. Appendix A of Dusseldorp & Van Mechelen (2013) gives a detailed overview of the choices that can be made.

Usage

```r
quint.control(
  crit = "es",
  maxl = 10,
  a1 = NULL,
  a2 = NULL,
  w = NULL,
  Bootstrap = TRUE,
  B = 25,
  dmin = 0.3
)
```

Arguments

- **crit**: the type of difference in treatment outcome used in the partitioning criterion: "es" (Treatment effect sizes) or "dm" (Difference in treatment means).
**quint.control**

maxl maximum number of leaves ($L$) of the tree. Default value is 10.

a1 the minimal sample size of Treatment A ($T=1$) in a leaf. If NULL, a1 is set to 1/10 of the sample size of the Treatment A group (assignment is done in the function quint). The minimum value is 2.

a2 the minimal sample size of Treatment B ($T=2$) in a leaf. If NULL, a2 is set to 1/10 of the sample size of the Treatment B group (assignment is done in the function quint). The minimum value is 2.

w a vector with $w_1$ and $w_2$ representing the weights of, respectively, the Difference in treatment outcome component and the Cardinality component of the partitioning criterion. If crit = "dm", the default value of $w_1$ is $1/\log(1+\text{IQR}(Y))$. If crit = "es", the default value of $w_1$ is $1/\log(1+3)$. The default of $w_2$ is $1/\log(0.50N)$.

Bootstrap whether the bias-corrected bootstrap procedure should be performed. The default is TRUE.

B the number of bootstrap samples to be drawn. The default is 25. We recommend a number of bootstraps of at least 25.

dmin the minimum absolute standardized mean difference in treatment outcome in one of the leaves assigned to P1 and one of the leaves assigned to P2 of the pruned tree. This value is used to check whether a qualitative interaction is present in the data (the qualitative interaction condition); dmin controls the balance between Type I error and Type II error. The default value of dmin is 0.30.

**Value**

A list containing the options.

**References**


**See Also**

quint

**Examples**

```r
# Load data
data(bcrp)
formula1<- I(cesdt1-cesdt3)~cond | nationality+marital+wcht1+age+tret+comorbid+disopt1+uncomt1+negsoct1
#Specify the Difference in treatment outcome as Difference in means
#and skip the bias-corrected bootstrap procedure
#and change the maximum number of leaves
control3<-quint.control(crit="dm",Bootstrap=FALSE,maxl=3)
quint3<-quint(formula1, data= subset(bcrp,cond<3),control=control3)
summary(quint3)
```
#Set number of bootstrap samples at 30
control4<-quint.control(B=30)

#Set minimal sample size in each treatment group at 5
control5<-quint.control(a1=5,a2=5)

---

## quint.validate

Validation of a Qualitative Interaction Tree

### Description

A bootstrap-based validation procedure to estimate the optimism in the effect sizes of a QUINT tree which gives insight in the generalizability of the results.

### Usage

```r
quint.validate(object, B = 10, allresults = FALSE)
```

### Arguments

- **object**: a (pruned) QUINT tree object of class `quint`.
- **B**: number of bootstrap samples. Default number is 10; for better accuracy $B=1000$ is recommended.
- **allresults**: option to return an extended list of output. Default is set to FALSE. See Value section for details.

### Details

In this procedure bootstrap trees are grown of the same leaf size as the (pruned) QUINT tree. The bootstrap samples are drawn from the data used to grow the original tree. For every bootstrap tree the largest and smallest (i.e., largest negative) treatment mean differences (or treatment effect sizes) of two leaves are saved. Treatment mean differences in the leaves are then predicted using the original data set as input for each bootstrapped tree. From these predictions, the largest and smallest treatment mean differences are saved. For each bootstrap tree, the largest predicted treatment effect is subtracted from the largest treatment effect in the bootstrap sample. The average of these values is the bias (i.e., the optimism) for the largest treatment effects. This is done likewise for the smallest treatment effects. Subsequently, the bias is computed as the difference between the bias for the largest effects minus the bias for the smallest effects.

The details of this validation procedure are described in Appendix C of Dusseldorp & Van Mechelen (2014).
quint.validate

Value

Returns a list with the following components:

- **estopt**: the estimated optimism for either the treatment effect size (biasd) or the raw treatment mean difference (biasdif).
- **li**: a data frame with leaf information output similar to the leaf information output of the (pruned) QUINT tree object. An extra column is added for the bias-corrected differences in treatment outcomes (d or diff). The bias-corrected values are only computed for the leaves with the most extreme values, i.e. the largest and smallest treatment effects. Hence, the other leaves get the value NA in this column.
- **optd**: a matrix with computed estimated optimism of the treatment effect size per bootstrapped tree. The first column contains the difference between the largest and smallest effect size of the bootstrapped tree. The second column contains the difference between the largest and smallest predicted effect size. Returned when allresults is set to TRUE and crit='es' is specified in the QUINT object.
- **optdif**: a matrix with computed estimated optimism of the raw mean difference bootstrapped tree. The first column contains the difference between the largest and smallest raw mean difference of the bootstrapped tree. The second column contains the difference between the largest and smallest predicted raw mean difference. Returned when allresults is set to TRUE and crit='es' is specified in the QUINT object.
- **resultd**: a vector with the estimated overall mean optimism, the mean bias for the smallest and for the largest effect size. Returned when allresults is set to TRUE and crit="es".
- **resultdif**: a vector with the estimated overall mean optimism, the mean bias for the smallest and largest raw mean difference. Returned when allresults is set to TRUE and crit="dm".

References


See Also

- **quint**, **prune.quint**, **quint.control**, **quint.bootstrapCI**

Examples

```r
## Not run: data(bcrp)
formula1<- I(cesdt1-cesdt3)~cond | nationality+marital+wcht1+age+
text+comorbid+disopt1+uncomt1+negsoct1

set.seed(10)
control1<-quint.control(maxl=5,B=2)
quint1<-quint(formula1, data= subset(bcrp,cond<3),control=control1) #Grow a QUINT tree
```
prquant1<-prune(quint1) #Prune tree to optimal size

set.seed(3)
valquant1<-quant.validate(prquant1, B = 5) #estimate the optimism by bootstrapping 5 times
valquant1

## End(Not run)

Description

Data generated sampling from a multivariate normal distribution using as parameters samp = 150, es = 1.0, J = 20, rho = 0 where samp is the sample size, es is the effect size, J is the number of covariates, and rho is the correlation between pairs of covariates. The treatment condition (A) is sampled from a binomial distribution with \( p = 0.5 \) and the treatment outcome (Y) is obtained using the following regression

\[
Y = 1 + (0.25 \times X_1) + (0.25 \times X_2) - (0.25 \times X_5) - (es \times ((A - 1)^2)) + \text{error}
\]

where the error is sampled from a standard normal distribution.

Usage

SimData_1

Format

A data frame with 150 observations on the following variables:

Y Outcome variable. It is obtained by regression according to the previously explained formula.
A This is the treatment condition variable. It is sampled from a binomial distribution.
X1 Simulated covariate. It is sampled from a multivariate normal distribution.
X2 Simulated covariate. It is sampled from a multivariate normal distribution.
X3 Simulated covariate. It is sampled from a multivariate normal distribution.
X4 Simulated covariate. It is sampled from a multivariate normal distribution.
X5 Simulated covariate. It is sampled from a multivariate normal distribution.
X6 Simulated covariate. It is sampled from a multivariate normal distribution.
X7 Simulated covariate. It is sampled from a multivariate normal distribution.
X8 Simulated covariate. It is sampled from a multivariate normal distribution.
X9 Simulated covariate. It is sampled from a multivariate normal distribution.
X10 Simulated covariate. It is sampled from a multivariate normal distribution.
X11 Simulated covariate. It is sampled from a multivariate normal distribution.
X12 Simulated covariate. It is sampled from a multivariate normal distribution.
X13 Simulated covariate. It is sampled from a multivariate normal distribution.
**X14** Simulated covariate. It is sampled from a multivariate normal distribution.

**X15** Simulated covariate. It is sampled from a multivariate normal distribution.

**X16** Simulated covariate. It is sampled from a multivariate normal distribution.

**X17** Simulated covariate. It is sampled from a multivariate normal distribution.

**X18** Simulated covariate. It is sampled from a multivariate normal distribution.

**X19** Simulated covariate. It is sampled from a multivariate normal distribution.

**X20** Simulated covariate. It is sampled from a multivariate normal distribution.

**gopt** Variable indicating whether the data is simulated such that there is qualitative interaction (0) or there is no qualitative interaction (1)

**Details**

IMPORTANT: for questions about these data contact Elise Dusseldorp: elise.dusseldorp@fsw.leidenuniv.nl.

---

**summary.quint**  
**Summarizing Qualitative Interaction Tree Information**

---

**Description**

Summary method for an object of class `quint`.

**Usage**

```r
## S3 method for class 'quint'
summery(object, digits = 2, ...)  
```

**Arguments**

- `object` a quint object. This can be the output of `quint`.
- `digits` specified number of decimal places (default is 2).
- `...` optional additional arguments.

**Details**

This function is a method for the generic function `summary` for class `quint`. It extracts the following essential components from a quint object: 1) Specification of the partitioning criterion; 2) Fit information; 3) Split information, and 4) Leaf information.

**Value**

prints a summarized version of the quint output.
Examples

data(bcrp)
formula1 <- I(cesdt1 - cesdt3) ~ cond | nationality + marital + wcht1 + age + trext + comorbid + disopt1 + uncomt1 + negsoct1
control1 <- quint.control(maxl = 5, Bootstrap = FALSE)
quint1 <- quint(formula1, data = subset(bcrp, cond < 3), control = control1)
summary(quint1)

# Example with only root node tree as outcome

data(SimData_1)
formula <- Y ~ A | X1 + X2 + X3 + X4 + X5
# Adjust the control parameters only to save computation time in the example;
# The default control parameters are preferred
control <- quint.control(maxl = 5, B = 2)
set.seed(2) # this enables you to repeat the results of the bootstrap procedure
quint_1 <- quint(formula, data = SimData_1, control = control)
quint_1pr <- prune(quint_1)
summary(quint_1pr)
Index

*Topic as.party
  plot.quint, 4
*Topic cluster
  quint, 8
*Topic datasets
  bcrp, 3
  SimData_1, 16
*Topic package
  quint-package, 2
*Topic plot
  plot.quint, 4
*Topic summary
  summary.quint, 17
*Topic tree
  prune.quint, 7
  quint, 8

bcrp, 3, 5, 10
plot.quint, 4
predict.quint, 3, 5
prune.quint, 3, 6, 7, 10, 11, 15
quint, 3, 5, 6, 8, 8, 11, 13, 15, 17
quint-package, 2
quint.bootstrapCI, 3, 8, 10, 10, 15
quint.control, 3, 5, 8, 10, 11, 12, 15
quint.validate, 3, 14

SimData_1, 16
summary.quint, 3, 10, 17