

Package: causalSLSE (via r-universe)

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Title Semiparametric Least Squares Inference for Causal Effects

Description Several causal effects are measured using least squares regressions and basis function approximations. Backward and forward selection methods based on different criteria are used to select the basis functions.

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as.model	<i>Converter into Model Objects</i>
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Description

When the information about a model is available, it reconstructs it and returns a valid model object.

Usage

```
## S3 method for class 'slseFit'
```

```
as.model(x, ...)
```

```
## S3 method for class 'cslseFit'
```

```
as.model(x, ...)
```

```
## S3 method for class 'cslse'
```

```
as.model(x, ...)
```

Arguments

x	An object containing the model to extract.
...	Other arguments to pass to other methods. Currently not used.

Value

The method returns an object of class `slseModel` or `cslseModel`.

Examples

```

data(simDat3)
mod <- cslseModel(Y ~ Z | ~ X1 * X2, data = simDat3)
fit <- estSLSE(mod)

## Extract the model from a cslseModel object

as.model(fit)

## Extract the model from a cslse object

cs <- causalSLSE(mod)
as.model(cs)

```

causalSLSE

Causal Effect Based on Semiparametric Least Squares Models

Description

This is the main method to estimate the causal effects using the semiparametric least squares method. It returns an object of class `cslse` and is registered for objects of class `cslseModel` and `cslseFit`.

Usage

```

## S3 method for class 'cslseModel'
causalSLSE(object,
            selType=c("SLSE", "BLSE", "FLSE"),
            selCrit = c("AIC", "BIC", "PVT"),
            selVcov = c("HC0", "Classical", "HC1", "HC2", "HC3"),
            causal = c("ALL", "ACT", "ACE", "ACN"),
            pvalT = function(p) 1/log(p),
            vcov.=vcovHC, reSelect=FALSE, ...)

## S3 method for class 'cslseFit'
causalSLSE(object, causal = c("ALL", "ACT", "ACE", "ACN"),
            vcov.=vcovHC, ...)

## S3 method for class 'formula'
causalSLSE(object, data, nbasis=function(n) n^0.3,
            knots,
            selType=c("SLSE", "BLSE", "FLSE"),
            selCrit = c("AIC", "BIC", "PVT"),
            selVcov = c("HC0", "Classical", "HC1", "HC2", "HC3"),
            causal = c("ALL", "ACT", "ACE", "ACN"),
            pvalT = function(p) 1/log(p),
            vcov.=vcovHC, reSelect=FALSE, ...)

```

Arguments

object	An object of class <code>cslseModel</code> created by the <code>cslseModel</code> function, <code>cslseFit</code> created by <code>estSLSE</code> or formula.
data	A <code>data.frame</code> with all variables included in form.
nbasis	A function to determined the number of basis functions. It has to be a function of one argument, the sample size.
knots	A list of knots for the treated and nontreated groups. The list must be named using the group names. Each element of the list is also a list of length equal to the number of confounders. The choice for each confounders is <code>NULL</code> for no knots or numeric for specific values. If missing, the knots are automatically generated.
selType	The method for selecting the knots. By default (SLSE), all knots from the model are used.
selCrit	The criterion to select the knots.
causal	What causal effect should we compute.
pvalT	A function to determine the p-value threshold for the significance of the coefficients. It has to be a function of one parameter, which is the average number of knots in the model. This value may differ across treatment group.
selVcov	The type of least squares covariance matrix used to compute the p-values needed for the selection.
vcov.	An alternative function to compute the covariance matrix of the least squares estimators. The default is <code>vcovHC</code> . This function is used to compute the standard errors of the causal effect estimators and the SLSE coefficients.
reSelect	By default, the stored selections are used. If <code>reSelect</code> is set to <code>TRUE</code> , the selection is re-computed.
...	Additional arguments to pass to <code>vcov</code> .

Value

It returns an object of class `cslse`, which inherits from the class `cslseFit`. It is a list with the following elements:

treated, nontreated	They are objects of class <code>slseFit</code> obtained by <code>estSLSE</code> .
ACE, ACT, ACN	Estimates of the average causal effect, the causal effect on the treated and the causal effect on the nontreated. Each of them is a vector of two elements: the estimate and its estimated standard error. All three are included only if the argument <code>causal</code> is set to "ALL".

Also, the object contains the following additional attributes:

treatedVar	The name of the variable in the dataset that represents the treatment indicator.
groupInd	A named vector with the value of the treatment indicator corresponding do each treatment group.

See Also

[estSLSE](#) for the estimation of the model, [slseKnots](#) for the format of knots, and [selSLSE](#) and [update](#) for the knots selection and to understand how stored selections are used.

Examples

```
data(simDat3)

## A causal SLSE model with the outcome Y
## the treatment indicator Z and the confounders X1, X2 and X1:X2

mod1 <- cslseModel(Y ~ Z | ~ X1 * X2, data = simDat3)

## The causal effects are estimated using the backward method and AIC criterion
## The HC1 type is used for the least squares covariance matrix

fit1 <- causalSLSE(mod1, selType = "BLSE", type = "HC1")

## This is the same for formula objects

fit2 <- causalSLSE(Y ~ Z | ~ X1 * X2, data = simDat3, selType = "BLSE", type = "HC1")
```

cslseModel

Semiparametric Least Squares Estimator Model

Description

This function creates an object of class `cslseModel` or `slseModel`. The former is a list of `slseModel` objects, one for each treatment group. The model objects contain all specifications about the model being estimated including the starting knots used to construct the basis functions.

Usage

```
cslseModel(form, data, nbasis = function(n) n^0.3,
           knots, groupInd=c(treated=1, nontreated=0))
slseModel(form, data, nbasis = function(n) n^0.3,
          knots)
```

Arguments

<code>form</code>	A formula for the regression in the case of a simple spline regression, or two formulas separated by <code> </code> , one for the outcome versus the treatment indicator and one of the confounders. See the details and examples below.
<code>data</code>	A <code>data.frame</code> that contains all variables included in <code>form</code> .
<code>nbasis</code>	A function to determine the number of basis functions. It has to be a function of one argument, the sample size.

knots	An optional list of knots. For <code>cslseModel</code> , it must be a named list using the group names (treated and nontreated). Each element of the list is also a list of length equal to the number of covariates. The choice for each covariate is NULL for no knots or a numeric vector for specific knots. If missing, the knots are automatically generated.
groupInd	A named vector with the group names and values. By default, the treatment indicator defined in <code>form</code> is equal to 1 for the treated and 0 for the nontreated. For now, these are the only allowed names, but the values can differ. In particular, the treatment indicator may be a factor with character values. See the example below.

Details

An object of class `s1seModel` is a regression model with a dependent variable and a set of covariates. We assume that the model can be written as $Y = f(X) + e$, where $f(x)$ is an unknown function of the covariates that we approximate using linear combinations of basis functions. For now, we only consider local linear splines defined by sets of knots, one for each covariate generated by the `model.matrix` of `form`. The knots are automatically determined unless specified by the user.

An object of class `cslseModel` is a list of `s1seModel` objects, one for each treatment group. The assignment to a group is determined by a treatment indicator, which is the right-hand side variable in the formula to the left of `|` specified in `form`. The formula to the right of `|` is used by `model.matrix` to generate a set of confounders for each treatment group. See the example below and the vignette to more details.

Value

The function `s1seModel` creates a semiparametric least squares model or a class `s1seModel` object. It is a list with the following elements:

<code>s1seForm</code>	The formula for the semiparametric estimation. It links the outcome variable with the matrix of basis functions.
<code>form</code>	The <code>form</code> argument passed to the function <code>s1seModel</code> .
<code>nameY</code>	The name of the variable representing the outcome
<code>data</code>	The dataset passed to the function <code>s1seModel</code> with the missing values, if any, omitted.
<code>knots</code>	An object of class <code>s1seKnots</code> . It is the set of knots that define the local linear splines.
<code>nameX</code>	The names of all covariates included in the model. They are all variables implied by the formula <code>form</code> , which may include interactions and functions of variables.
<code>nameS</code>	The prefix added to the covariate names when the basis functions are generated. By default, it is equal to "U."
<code>xlevels</code>	When relevant, a list of levels of the factors included in the model.
<code>na</code>	A vector of integer representing the observations omitted because of missing values. It is NULL when there are no missing values.

Note that other elements may be present if a knots selection has been applied to the model.

The function `cslseModel` creates an object of class `cslseModel`. It is a list of `slseModel` objects, one for each treatment group. It also contains the following additional attributes:

<code>treatedVar</code>	The name of the variable in the dataset that represents the treatment indicator.
<code>groupInd</code>	A named vector with the value of the treatment indicator corresponding do each treatment group.

See Also

[selSLSE](#) for additional elements that can be included in the `slseModel` object, [estSLSE](#) to see how the model is estimated and [slseKnots](#) for the format of the knots

Examples

```
data(simDat3)

## Y is the outcome and Z the treatment indicator
## The confounders are X1, X2 and X1:X2:

mod1 <- cslseModel(Y ~ Z | ~ X1 * X2, data = simDat3)
mod1

## A simple SLSE for a regression model

mod2 <- slseModel(Y ~ X1 * X2, data = simDat3)
mod2

## If the treatment indicator differs from 0-1

data(simDat4)
mod3 <- cslseModel(Y ~ treat | ~ X1 * X2, data = simDat4,
                  groupInd = c(treated = "treat", nontreated = "notreat" ))
mod3
```

estSLSE

Least Squares Estimate of cslseModel or slseModel Objects

Description

This is the main function to estimate `cslseModel` or `slseModel` objects. It generates the basis functions based on the knots specified in the model and estimates it by least squares.

Usage

```
## S3 method for class 'cslseModel'
estSLSE(model, selKnots, ...)

## S3 method for class 'slseModel'
estSLSE(model, selKnots, ...)
```

Arguments

model	A model of class <code>cslseModel</code> or <code>slseModel</code> .
selKnots	An optional list of integers to select the knots from the list of knots specified by the model. If the model is a <code>cslseModel</code> object, it must be a named list with the names being either "treated", "nontreated" or both. By default, the knots from the model stored in the element <code>knots</code> are used.
...	Additional arguments to pass to other methods. Currently not used.

Details

The method for `slseModel` objects generates the matrix of basis functions implied by the set of knots included in the model and estimate the model by the least squares. Let Y be the outcome and U be the matrix of basis functions. Then, the function estimates the model using the code `lm(Y~U)`. For `cslseModel`, we could estimate the model using `lm(Y~Z+I(Z-1)+I(U0*(1-Z))+I(U1*Z))`, where Z is a binary variable equal to 1 for the treated and 0 for the nontreated, and U_0 and U_1 are the matrices of basis functions for the nontreated and treated, but the model is estimated separately for each group. Therefore, the function `estSLSE.cslseModel` calls the function `estSLSE.slseModel` for each `slseModel` objects included in the `cslseModel` object.

Value

It returns an object of class `slseFit` or `cslseFit` depending on which method is called. An object of class `slseFit` is a list with the following elements:

LSE	This is the least squares estimate of the semiparametric model. It is an object of class <code>lm</code> .
model	An object of class <code>slseModel</code> . It is the model that is being estimated. The model may have been created by <code>slseModel</code> or modified by <code>selSLSE</code> or <code>update</code> .

An object of class `cslseFit` is a list of `slseFit` objects, one for each treatment group. It also contains the following additional attributes:

treatedVar	The name of the variable in the dataset that represents the treatment indicator.
groupInd	A named vector with the value of the treatment indicator corresponding do each treatment group.

Examples

```
data(simDat3)

## Estimating a causal semiparametric model

mod1 <- cslseModel(Y ~ Z | ~ X1 * X2, data = simDat3)
fit1 <- estSLSE(mod1)

## Estimating a semiparametric model

mod2 <- slseModel(Y ~ X1 * X2, data = simDat3)
fit2 <- estSLSE(mod2)
```

extract,cslse-method [extract](#) Method for cslse Objects

Description

[extract](#) method for cslse objects created by the [causalSLSE](#) function.

Usage

```
## S4 method for signature 'cslse'
extract(
  model,
  include.nobs = TRUE,
  include.nknots = TRUE,
  include.numcov = TRUE,
  include.rsquared = TRUE,
  include.adjrs=TRUE,
  separated.rsquared = FALSE,
  which = c("ALL", "ACE", "ACT", "ACN", "ACE-ACT", "ACE-ACN", "ACT-ACN"),
  ...)
```

Arguments

<code>model</code>	A causal effect object created by causalSLSE .
<code>include.nobs</code>	Report the number of observations?
<code>include.nknots</code>	Report the total number of knots for each group?
<code>include.numcov</code>	Report the total number of covariates (including interactions is any) per group?
<code>include.rsquared</code>	Report the R-squared of the final regression estimation
<code>include.adjrs</code>	Report the adjusted R-squared of the final regression estimation
<code>separated.rsquared</code>	Should we print the R-squared separately for each group? This applies as well to the adjusted R-squared.
<code>which</code>	Which causal effect measures should be printed?
<code>...</code>	Custom parameters, which are handed over to subroutines. Currently not in use.

Value

It returns an object of class [texreg](#).

extract,slseFit-method

[extract](#) *Method for slseFit Objects*

Description

[extract](#) method for `slseFit` objects created by the `estSLSE` function.

Usage

```
## S4 method for signature 'slseFit'
extract(
  model,
  include.rsquared = TRUE,
  include.adjrs = TRUE,
  include.nobs = TRUE,
  include.fstatistic = FALSE,
  include.rmse = FALSE,
  ...)
```

Arguments

<code>model</code>	A model estimated by <code>estSLSE</code> .
<code>include.rsquared</code>	Report the R-squared of the final regression estimation
<code>include.adjrs</code>	Report the adjusted R-squared of the final regression estimation
<code>include.nobs</code>	Report the number of observations?
<code>include.fstatistic</code>	Report the F-statistics?
<code>include.rmse</code>	Report the RMSE?
<code>...</code>	Custom parameters, which are handed over to subroutines. Currently not in use.

Value

It returns an object of class `texreg`.

Description

It generates a matrix of basis functions using local linear splines. The number of basis functions and the breaking points are determined by the knot specifications of the `slseModel` or `cslseModel` model.

Usage

```
## S3 method for class 'slseModel'  
llSplines(object, ...)  
## S3 method for class 'cslseModel'  
llSplines(object, ...)
```

Arguments

`object` A model of class `slseModel` or `cslseModel`.
`...` Additional arguments to pass to other methods. Currently not used.

Value

The function returns a matrix of basis functions used to estimate the semiparametric model. It is used directly as regressor in `lm` as shown in the example below.

Examples

```
data(simDat3)  
  
## We manually estimate the semiparametric model  
  
mod1 <- slseModel(Y ~ X1 * X2, data = simDat3)  
U <- llSplines(mod1)  
fit1 <- lm(Y ~ U, data = simDat3)  
  
## We use estSLSE instead (results are identical)  
  
fit2 <- estSLSE(mod1)
```

nsw	<i>Lalonde Subsample of the National Supported Work Demonstration Data (NSW)</i>
-----	--

Description

This data was collected to evaluate the National Supported Work (NSW) Demonstration project in Lalonde (1986).

Usage

```
data(nsw)
```

Format

A data frame containing 9 variables.

treat Treatment assignment

age Age

ed Years of Education

black 1 if Black, 0 otherwise

hisp 1 if Hispanic 0 otherwise

married 1 if married 0 otherwise

nodeg 1 if no college degree 0 otherwise

re75 1975 earnings

re78 1978 earnings

Details

The dataset was obtained from the ATE package (see reference).

Source

"NSW Data Files" from Rajeev Dehejia's website. URL: <http://users.nber.org/~rdehejia/data/.nswdata2.html>

"National Supported Work Evaluation Study, 1975-1979: Public Use Files." from the Interuniversity Consortium for Political and Social Research. URL: <http://www.icpsr.umich.edu/icpsrweb/ICPSR/studies/7865>

References

- Lalonde, R. (1986). "Evaluating the Econometric Evaluations of Training Programs," *American Economic Review*, 76(4), 604-620.
- Dehejia R. and Wahba S. (1999). "Causal Effects in Non-Experimental Studies: Re-Evaluating the Evaluation of Training Programs," *JASA* 94 (448), 1053-1062.
- Asad Haris and Gary Chan (2015). ATE: Inference for Average Treatment Effects using Covariate Balancing. R package version 0.2.0. <https://CRAN.R-project.org/package=ATE>

plot

Plot of Predicted Outcome

Description

For objects of class `slseFit`, the method plots the predicted outcome with respect to a given covariate. It is the same for objects of class `cslseFit`, but the predicted outcome is plotted for each treatment group separately.

Usage

```
## S3 method for class 'cslseFit'
plot(x, y, which = y, interval = c("none", "confidence"),
     level = 0.95, fixedCov = list(),
     vcov. = vcovHC, add = FALSE, addToLegend = NULL,
     addPoints = FALSE, FUN = mean,
     plot=TRUE, graphPar=list(), ...)

## S3 method for class 'slseFit'
plot(x, y, which = y, interval = c("none", "confidence"),
     level = 0.95, fixedCov = NULL,
     vcov. = vcovHC, add = FALSE,
     addPoints = FALSE, FUN = mean,
     plot=TRUE, graphPar=list(), ...)
```

Arguments

- | | |
|-----------------------|---|
| <code>x</code> | Object of class <code>cslseFit</code> or <code>slseFit</code> created by estSLSE . |
| <code>y</code> | alias for which for compatibility with <code>plot</code> . |
| <code>which</code> | Which covariate to plot against the outcome variable. It could be an integer or a character. |
| <code>interval</code> | The type of confidence interval. The default is none. |
| <code>level</code> | The confidence interval level if included. |
| <code>fixedCov</code> | List of covariates to fix to specific values for the nontreated and treated groups. By default, covariates not selected by <code>which</code> are set to their group specific sample means. |

<code>vcov.</code>	An alternative function to compute the covariance matrix of the least squares estimates. The default is the <code>vcovHC</code> method for <code>lm</code> objects.
<code>add</code>	Should the plot be added to an existing one?
<code>addToLegend</code>	A character string to add to the legend next to treated and control.
<code>addPoints</code>	Should we add the scatterplot of the outcome and covariate on the graph?
<code>FUN</code>	The function to determine the fixed value for the covariates not fixed by <code>fixedCov</code> . The default is <code>mean</code> .
<code>plot</code>	If set to <code>FALSE</code> , a <code>data.frame</code> is returned for each group with the covariate selected by <code>which</code> and the prediction.
<code>graphPar</code>	A list of graphical parameters. See Details.
<code>...</code>	Additional argument to pass to the <code>vcov.</code> function.

Details

The default set of parameters can be obtained by running the command `causalSLSE:::initPar()`. It returns a list of four elements: `treated` for the parameter of the lines or points of the treated, `nontreated` for the parameters of the nontreated, `common` for the common parameters not specific to a group like the main title or the axis labels, and `legend` for the legend parameters. The elements `treated` and `nontreated` are lists of two elements: `points` and `lines`, which are lists of graphical parameters for the scatterplot (when `addPoints` is `TRUE`) and the lines. The argument `graphPar` can be used to modify existing parameters or to add new ones. It must respect the structure of the initial list. See the example below.

To fix covariates to the same values for both groups, `fixedCov` is just a named list with values for the covariates associated with the names. To fix the covariates to different values for the treated and nontreated, `fixedCov` is a named list of at least 1 element with names being either `treated`, `nontreated` or `both`. If only one group is specified, the covariates for the other groups are determined by `FUN`.

Value

It returns `NULL` invisibly if the argument `plot` is `TRUE`. Otherwise, it returns a `data.frame` with the necessary variables to manually create the plot. For `slseModel` objects, it is a list with the following elements:

<code>Outcome</code>	The outcome variable from the model. The name of this element is the actual name of the outcome variable in the model.
<code>which</code>	The covariate selected by the argument <code>which</code> . The name of this element is the name of the selected covariate.
<code>fit</code>	The predicted outcome from the model fit.
<code>lower, upper</code>	The lower and upper bounds of the confidence interval. It is only available when the argument <code>interval</code> is set to <code>"confidence"</code> .

Note that all returned variables are ordered with respect to the selected covariates. See the example below.

For `slseModel`, the above list of variables is returned separately for each treatment group.

Examples

```

data(simDat3)

## For cslse objects

mod <- cslseModel(Y ~ Z | ~ X1 + X2, data = simDat3)
fit <- causalSLSE(mod)
plot(fit, "X1")

## Let's change the type of points for the treated and lines for the
## nontreated

gpar <- list(treated = list(points = list(pch = 24, col = 5)),
             nontreated = list(lines = list(lty = 5, col =
             "darkgreen")), common = list(xlab = "New X", main =
             "Plot with changed parameters"))

plot(fit, "X1", addPoints = TRUE, graphPar = gpar)

## For slseModel objects:

mod2 <- slseModel(Y ~ X1 + X2, data = simDat3)
fit2 <- estSLSE(mod2)
plot(fit2, "X1", interval = "confidence", addPoints = TRUE)

## The same graph produced manually

p2 <- plot(fit2, "X1", interval = "confidence", plot = FALSE)
plot(p2$X1, p2$Y, pch = 21, main = "Y against X1", xlab = "X1", ylab =
     "Y")
lines(p2$X1, p2$fit, lwd = 2)
lines(p2$X1, p2$lower, lty = 3, lwd = 2)
lines(p2$X1, p2$upper, lty = 3, lwd = 2)

```

predict

Outcome Prediction

Description

The method computes the predicted outcome for each group with standard errors and confidence intervals.

Usage

```

## S3 method for class 'cslseFit'
predict(object, interval=c("none","confidence"),
        se.fit=FALSE, newdata=NULL, level=0.95,
        vcov.=vcovHC, ...)
## S3 method for class 'slseFit'

```

```
predict(object, interval=c("none","confidence"),
        se.fit=FALSE, newdata=NULL, level=0.95,
        vcov.=vcovHC, ...)
```

Arguments

<code>object</code>	Object of class <code>cslseFit</code> or <code>slseFit</code> created by <code>estSLSE</code> .
<code>interval</code>	If set to "confidence", it returns the predicted values along with the lower and upper bounds of the confidence interval.
<code>se.fit</code>	Should the function return the standard errors of the predicted values?
<code>level</code>	The confidence interval level if <code>interval</code> is set to "confidence".
<code>newdata</code>	A <code>data.frame</code> of new data. It must include values for all covariates, and for the treatment indicator in the case of <code>cslseFit</code> objects.
<code>vcov.</code>	An alternative function to compute the covariance matrix of the least squares estimates. The default is the <code>vcovHC</code> .
<code>...</code>	Additional argument to pass to the <code>vcov.</code> function.

Value

For `slseFit` objects, it returns the predicted outcome if `se.fit` is `FALSE` or a list of the following two elements otherwise:

<code>fit</code>	The predicted outcome.
<code>se.fit</code>	The standard errors of the predicted outcomes.

If the argument `confidence` is set to "interval", the predicted outcome is a matrix with the predicted outcome, and the lower and upper bounds of the confidence intervals.

For objects of class `'cslseFit'`, the same is returned for each treatment group in a list. The elements of the list are `treated` and `nontreated` (until the package allows for more than one treatment).

Examples

```
data(simDat3)
mod <- cslseModel(Y ~ Z | ~ X1 + X2, data = simDat3)
fit <- causalSLSE(mod)

## Predicting outcome for all observations
pr <- predict(fit, interval = "confidence")

## Predicting outcome with new data
ndat <- data.frame(X1 = c(-2, 1, 2, 3), X2 = c(-4, -2, 0, 1),
                  Z = c(1, 1, 0, 0))
predict(fit, newdata = ndat)
```

print

Print Methods

Description

Print methods for different objects from the package.

Usage

```
## S3 method for class 'cslseModel'
print(x, which=c("Model", "selKnots", "Pvalues"),
      digits = max(3L, getOption("digits") - 3L), ...)
## S3 method for class 'slseModel'
print(x, which=c("Model", "selKnots", "Pvalues"),
      digits = max(3L, getOption("digits") - 3L), ...)
## S3 method for class 'cslse'
print(x, digits = max(3L, getOption("digits") - 3L), ...)
## S3 method for class 'cslseFit'
print(x, digits = max(3L, getOption("digits") - 3L), ...)
## S3 method for class 'slseFit'
print(x, digits = max(3L, getOption("digits") - 3L), ...)
## S3 method for class 'summary.cslse'
print(x, digits = max(3L, getOption("digits") - 3L),
      signif.stars = getOption("show.signif.stars"),
      beta=FALSE, knots = FALSE,...)
## S3 method for class 'summary.cslseFit'
print(x, groups, digits = max(3L, getOption("digits") - 3L),
      signif.stars = getOption("show.signif.stars"),
      ...)
## S3 method for class 'summary.slseFit'
print(x, digits = max(3L, getOption("digits") - 3L),
      signif.stars = getOption("show.signif.stars"),
      ...)
## S3 method for class 'slseKnots'
print(x, header=c("None", "All", "Select"),
      digits = max(3L, getOption("digits") - 3L), ...)
```

Arguments

x	A model or a fit object.
digits	The number of digits to print.
signif.stars	Should we print the significant stars?
beta	Should we print the coefficient matrix of the basis functions?
knots	Should we print the set of knots?

which	What info we should print? It prints a summary of the model when set to "Model", the current knots when set to "selKnots" and the p-values and initial knots when set to "Pvalues". For the latter, it is only available when a knots selection method has been applied to the object.
header	What description of the object should be printed.
groups	The names of the group to be printed. By default, they are all printed.
...	Argument for other types of objects.

Value

All methods return NULL invisibly.

Examples

```
## For cslseModel objects

data(simDat3)
mod1 <- cslseModel(Y ~ Z | ~ X1 * X2, data = simDat3)
print(mod1)

## For cslse objects

causal <- causalSLSE(mod1)
print(causal)

## For summary.cslse objects

s <- summary(causal)
print(s)

## For cslseFit objects

fit <- estSLSE(mod1)
print(fit)

## For summary.cslseFit objects

summary(fit)
```

selSLSE

Knots Selection Method

Description

This is the main function to select the knots in cslseModel or slseModel objects. It returns a model with an optimal set of knots.

Usage

```
## S3 method for class 'cslseModel'
selSLSE(model, selType=c("BLSE", "FLSE"),
        selCrit = c("AIC", "BIC", "PVT"),
        pvalT = function(p) 1/log(p),
        vcovType = c("HC0", "Classical", "HC1", "HC2", "HC3"),
        reSelect=FALSE, ...)
## S3 method for class 'slseModel'
selSLSE(model, selType=c("BLSE", "FLSE"),
        selCrit = c("AIC", "BIC", "PVT"),
        pvalT = function(p) 1/log(p),
        vcovType = c("HC0", "Classical", "HC1", "HC2", "HC3"),
        reSelect=FALSE, ...)
```

Arguments

model	A model of class <code>cslseModel</code> or <code>slseModel</code> .
selType	The selection method: backward ("BLSE") or forward ("FLSE").
selCrit	The criterion to select the piecewise polynomial knots.
pvalT	A function to determine the p-value threshold for the significance of the coefficients. It has to be a function of one parameter, which is the average number of knots in the model.
vcovType	The type of least squares covariance matrix used to compute the p-values needed for the selection.
reSelect	By default, the stored selections are used. If <code>reSelect</code> is set to TRUE, the selection is re-computed.
...	Additional arguments to pass to other methods. Currently not used.

Details

It selects the knots using one of the two methods, FLSE or BLSE, with either the AIC, BIC or a p-value threshold (see the vignette for more details). Any of these selection methods requires several least squares estimations and it is performed only if the method has not been applied yet and `reSelect` is set to TRUE. This is possible because any new knots selection is saved into the returned model. A model may have more than one selection saved into it. The active knots (the ones used when we estimate the model) is stored into the element `knots` of the model and the saved selections are stored into the element `selections`. See below for what is included in this element.

Note that the selections for the three criteria AIC, BIC and PVT are computed and saved automatically in the returned model when `selCrit` is set to either "AIC" or "BIC", because it does not require many more operations to select them all once we do it for AIC or BIC. However, it is only computed for PVT when `selCrit` is set to "PVT".

The knots are selected jointly for all treatment groups in `cslseModel` objects. However, the active knots and all saved selections are stored separately for each treatment group. For example, the active knots for the treated in the `cslseModel` object `mod` are stored in `mod$treated$knots`. See the Vignette for more details.

Value

The method returns an object of class `slseModel` or `cslseModel` depending on which object it is applied to. When it does not already exist, the element selections is added to the `slseModel` object (or to each `slseModel` object in `cslseModel` objects). The element selections is a list with one or more of the following elements:

`originalKnots` The original knots as selected initially by `slseModel` or `cslseModel`. This element is always included.

`FLSE, BLSE` This is where selections based on the forward (FLSE) and the backward (BLSE) methods are stored.

Finally, BLSE and FLSE are lists that may contain the following elements:

`AIC, BIC` A list of integer vectors, one for each covariate in the `slseModel` model. The vectors of integers indicate which sets the original set of knots minimizes the AIC or the BIC.

`PVT` Same as the AIC and BIC, but the selection is based on a p-value threshold.

`JAIC, JBIC` This criteria is for `cslseModel` models. It is like the AIC and BIC, but it indicates that the AIC or BIC was computed by estimating the models of all treatment groups jointly. This selection is not the same as a selection group by group.

`Threshold` The p-value threshold used for the PVT criterion.

`pval` A list of p-values, one for each original knots. See vignette for a definition of the p-values.

See Also

`slseModel` and `cslseModel` for model objects description and `update` for ways of selecting stored selections

Examples

```
data(simDat3)
mod1 <- cslseModel(Y~Z|~X1*X2, data=simDat3)
mod1 <- selSLSE(mod1, selType="FLSE", selCrit="AIC")

## The following does not require additional computation
## because the selection is stored in mod1

mod1 <- selSLSE(mod1, selType="FLSE", selCrit="BIC")

## But the following does

mod1 <- selSLSE(mod1, selType="BLSE", selCrit="BIC")

## See one selection:

mod1$treated$selections$BLSE$JBIC
```

simDat1	<i>Simulated Data</i>
---------	-----------------------

Description

This dataset is used in several documentation files to illustrate the different functionalities of the package.

Usage

```
data("simDat1")
```

Format

A data frame with 300 observations on the following 9 variables.

X Continuous Covariate

Z Treatment indicator.

Y Observed outcome.

Y1 Outcome for the treated (not observed in practice).

Y0 Outcome for the nontreated (not observed in practice).

U01, U02 The true basis functions for the nontreated.

U11, U12 The true basis functions for the treated.

simDat2	<i>Simulated Data</i>
---------	-----------------------

Description

This dataset is used in several documentation files to illustrate the different functionalities of the package.

Usage

```
data("simDat2")
```

Format

A data frame with 300 observations on the following 11 variables.

X Continuous Covariate

Z Treatment indicator.

Y Observed outcome.

Y1 Outcome for the treated (not observed in practice).

Y0 Outcome for the nontreated (not observed in practice).

U01, U02, U03 The true basis functions for the nontreated.

U11, U12, U13 The true basis functions for the treated.

simDat3

Simulated Data

Description

This dataset is used in several documentation files to illustrate the different functionalities of the package.

Usage

```
data("simDat3")
```

Format

A data frame with 300 observations on the following 16 variables.

X1 Continuous Covariate

X2 Continuous Covariate

Z Treatment indicator.

Y Observed outcome.

Y1 Outcome for the treated (not observed in practice).

Y0 Outcome for the nontreated (not observed in practice).

U01, U02, U03, U04, U05 The true basis functions for the nontreated.

U11, U12, U13, U14, U15 The true basis functions for the treated.

simDat4

Simulated Data.

Description

This dataset is used in several documentation files to illustrate the different functionalities of the package.

Usage

```
data("simDat4")
```

Format

A data frame with 500 observations on the following 7 variables.

Z Treatment indicator.

Y Observed outcome.

X1 Continuous covariates with a large proportion of zeros.

X2 A categorical variable with 3 character levels (not expressed as factor).

X3 A categorical variable with 3 numerical levels (not expressed as factor).

X4 A binary variable.

treat A character variable.

simDat5

Simulated Data

Description

This dataset is used in several documentation files to illustrate the different functionalities of the package.

Usage

```
data("simDat5")
```

Format

A data frame with 300 observations on the following 16 variables.

X1 Continuous Covariate

X2 Continuous Covariate

Z Treatment indicator.

Y Observed outcome.

Y1 Outcome for the treated (not observed in practice).

Y0 Outcome for the nontreated (not observed in practice).

slseKnots

*Knots Creator for Basis Functions***Description**

The function creates an object of class `slseKnots` for semiparametric least squares models. It returns an object of class `slseKnots`.

Usage

```
slseKnots(form, data, X, nbasis = function(n) n^0.3,
          knots)
```

Arguments

<code>form</code>	A formula that determines the covariates to include in the regression. It is a regular formula and only the right hand side is considered.
<code>data</code>	A <code>data.frame</code> with all variables included in <code>form</code> . It is required when <code>X</code> is missing.
<code>nbasis</code>	A function to determine the number of basis functions. It has to be a function of one argument, the sample size.
<code>knots</code>	An optional list of knots. Each element of the list (one for each covariate) is either a vector of numeric knots or it is <code>NULL</code> . The latter implies that the number of knots is set to 0. If missing (the default), the knots are automatically generated (see the details below).
<code>X</code>	An optional matrix of covariates. When provided, <code>form</code> is not needed.

Details

The automatic selection is as follows. It is applied to each variable created by the `model.matrix` of `form` except for the intercept:

The number of knots is the ceiling of what the `nbasis` function returns minus 1. Let $p-1$ be the number of knots. Then, we compute the $p+1$ empirical quantiles of the variable for equally spaced probabilities going from 0 to 1 and drop the first and last ones. This is done using the function `quantile` with `type=1`. We then remove the duplicated values and the ones equal to either the `min` or the `max` of the variable. if the number of remaining knots is equal to 0, the set of knots for this variable is set to `NULL`.

For manual selection, see the vignette.

Value

It returns an object of class `slseKnots`. It is a list for which the length and names are respectively the number of columns and the column names of the `model.matrix` of `form` after the intercept has been removed. Each element is a numeric vector of knots, unless the number of knots is set to 0, in which case it is `NULL`.

Examples

```

data(simDat3)
k <- slseKnots(Y ~ X1 * X2, data = simDat3)
k

## We can extract the set for one variable

k$X1

```

summary

*Summary Method for Fitted Models***Description**

The method computes summary statistics for estimated semiparametric and causal models.

Usage

```

## S3 method for class 'cslse'
summary(object, ...)
## S3 method for class 'cslseFit'
summary(object, vcov.=vcovHC, ...)
## S3 method for class 'slseFit'
summary(object, vcov.=vcovHC, ...)

```

Arguments

object	A model estimated by estSLSE or causalSLSE .
vcov.	A function to compute the standard error of the least squares coefficients. The default is vcovHC .
...	Argument for other types of objects

Value

The function `summary.slseFit` returns the summary statistics of the least squares estimation of [slseModel](#) objects estimated by [estSLSE](#). It is an object of class `summary.slseFit`, which is a list with the following elements:

model	The slseModel model being estimated.
lseSum	An object of class summary.lm .

The function `summary.cslseFit` returns an object of class `summary.cslseFit`, which is a list of `summary.slseFit` objects, one for each treatment group.

The function `summary.cslse` returns the summary statistics of the causal effect estimates. It is an object of class `summary.cslse`, which is a list with the following elements:

causal	A matrix with the causal effect estimates in the first column, their standard errors in the second, their t-ratios in the third and p-values in the fourth.
lse	A list of <code>summary.slseFit</code> objects, one for each treatment group.

Examples

```
## A causal semiparametric model with causal effect estimates

data(simDat3)
mod1 <- cslseModel(Y ~ Z | ~ X1 * X2, data = simDat3)
CE <- causalSLSE(mod1, selType = "BLSE")
summary(CE)

## A causal semiparametric model

fit <- estSLSE(mod1)
summary(fit)

## A semiparametric model

mod2 <- slseModel(Y ~ X1 * X2, data = simDat3)
fit2 <- estSLSE(mod2)
summary(fit2)
```

update	<i>Update Methods</i>
--------	-----------------------

Description

The method updates an object by modifying its specification. Currently, it is used to change the set of knots by either selecting them manually or by specifying the selection methods.

Usage

```
## S3 method for class 'cslseModel'
update(object, selType, selCrit="AIC",
       selKnots, ...)

## S3 method for class 'slseModel'
update(object, selType, selCrit="AIC",
       selKnots, ...)

## S3 method for class 'slseKnots'
update(object, selKnots, ...)
```

Arguments

<code>object</code>	An object to be modified.
<code>selKnots</code>	An optional list of integers to select the knots from the original list of knots. If missing, the current knots are kept.
<code>selType</code>	The selection method: "BLSE" for the backward method, "FLSE" for the forward method or "none" to recover the original set of knots.
<code>selCrit</code>	The criterion to select the optimal set of knots.
<code>...</code>	Argument for other types of objects. Currently not used.

Details

The method for `slseKnots` is explained in the vignette, but it is mostly used internally. For the model objects, the method can be used to choose a set of knots already stored in the object. It avoids having to re-compute them which can be computationally intensive for large samples. It returns an error message if the type of selection requested does not exist. In that case, you need to run the `selSLSE` method. If the `selType` argument is set to `None`, the method returns the original model.

Value

It returns an object of the same class, but with a different set of knots.

See Also

[selSLSE](#) for more details on how stored knots can be selected from a model, [slseModel](#) for model description and [slseKnots](#) for the format of knots

Examples

```
data(simDat3)
mod1 <- cslseModel(Y ~ Z | ~ X1 * X2, data = simDat3)
mod2 <- selSLSE(mod1, "BLSE", "AIC")

## We changed the knots to the BLSE-BIC selection
## already stored in the model object

update(mod2, "BLSE", "BIC")

## We recover the original set of knots

update(mod2, "None")
```

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