Semiparametric Thresholding Least Squares Inference for Causal Effects with R

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Abstract

The vignette explains how to use the causalTLSE package to estimate different causal effects using a semiparametric thresholding least squares method.

Introduction

This document presents the causalTLSE package explaining in details all functions. It is intended for users interested in all the details about the procedure presented in the paper and how it is implemented.

The main model is

$$Y = \beta_0(1-Z) + \beta_1 Z + f_0(X) + f_1(X) + \varepsilon$$

and it is approximated by the regression

$$Y = \beta_0(1 - Z) + \beta_1 Z + \psi_0' U_0(X) + \psi_1' U_1(X) + u,$$

where $U_0(X)$ and $U_1(X)$ are spline matrices satisfying $U_0(X_i) = 0$ if $Z_i = 1$ and $U_1(X_i) = 0$ if $Z_i = 0$. If X is a $k \times 1$ matrix of covariates. We can separate $U_j(X)$, for j = 0, 1, into a block matrix $\{U_{1j}(X), U_{2j}(X), ..., U_{kj}(X)\}$, where $U_{ij}(X)$ is matrix of basis functions to approximate the function $f_j(X_i)$. The paper proposes a data-driven method for selecting the matrices $U_0(X)$ and $U_1(X)$.

To understand the package, it is important to know how the $U_{ij}(X)$'s are defined. To simplify the notation, we remove the subscripts i from X and i and j from $U_{ij}(X)$. We just need to keep in mind that U(X) is different for the treated and control groups. We want to approximate f(X) by a linear spline basis function. Let $\{\kappa_1, ..., \kappa_{p-1}\}$ be a set of p-1 knots strictly inside the sample range of the X satisfying $\kappa_1 < \kappa_2 <, ..., < \kappa_{p-1}$. For a realization x and $p \ge 3$, we have the following bases.

$$\begin{array}{lcl} U_1(x) & = & xI(x \leq \kappa_1) + \kappa_1 I(x > \kappa_1) \\ U_2(x) & = & (x - \kappa_{p-1})I(x > \kappa_{p-1}) \\ U_k(x) & = & (x - \kappa_{k-1})I(\kappa_{k-1} \leq x \leq \kappa_k) + (\kappa_k - \kappa_{k-1})I(x > \kappa_k) \,, \end{array}$$

where the last $U_k(x)$ is defined for 2 < k < p. Therefore, if the number of knots is equal to 1, we only have the first two bases. Since knots must be strictly inside the sample range of X, any categorical variable with two levels, which includes as a special case binary variables, the number of knots must be equal to zero. When this is the case, U(X) = X. For general categorical variables, the number fo knots cannot exceed the number of levels minus two.

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The causalTLSE package

Setting up the Model

The first step to estimate the causal effect is to define a model. A model contains the information about the outcome, the treatment indicator, the covariates and their knots. This is the starting point before applying any basis selection method. To illustrate how to use the package, we are using the dataset from Lalonde (1986). It contains some continuous and categorical variables, so we can illustrate how knots are selected initially. The dataset is available from the package.

```
library(causalTLSE)
data(nsw)
```

The outcome is the real income in 1978 (re78) and the purpose is to measure the impact of a training program (treat) on the outcome. The dataset includes also covariates such as age (age), education (ed) past real income (re75) and some categorical variables (black, hisp, married and nodeg). We start by considering the covariates age, re75, ed and 'married. We can create a model simply by running the following command.

```
model1 <- setModel(re78~treat | ~age+re75+ed+married, data=nsw)</pre>
```

The left of | is for the formula linking the outcome and the treatment indicator only. The covariates are entered after | as a formula without a dependent variable. It works like for formulas in lm. For example, we can add interactions, functions of the variables, etc. The following is an example:

```
modEx <- setModel(re78~treat | ~age+I(age^2)+re75+ed*married, data=nsw)</pre>
```

This will create the vector of covariates $\{age, age^2, re75, ed, married, ed \times married\}$. The function returns an object of class tlseModel with its own print method. We will present it later. The following sub-sections explain all arguments of the function.

The starting knots

By default, the function automatically generates knots for each variable based on the following procedure. This procedure is applied separately for the treated and control groups. Therefore, the term sample size means the number of observations in the treated or control group.

- 1. The starting number of knots is a function of the sample size and is determined by the argument nknots, a function of one argument, the sample size. The starting number of knots is equal to the floor of what the function returns minus 1 (or 0 if this operation results in a negative number). The default function is function(n) n^0.3. For example, if the total sample size is 500, with 200 treated and 300 control, the starting number of knots in the treated and control groups are respectively equal to 3 (floor(200^0.3)-1) and 4 (floor(300^0.3)-1). It is possible to have a number of knots that does not depend on the sample size. All we need is to set the argument nknots to a function that returns an integer.
- 2. Let (p-1) be the number of knots determined by the previous step. The knots are obtained by computing p+1 quantiles of X for equally spaced probabilities from 0 to 1, and by dropping the first and last ones. For example, if the number of knots is equal to 3, we compute the quantiles for the probabilities $\{0.25, 0.5, 0.75\}$.
- 3. We drop any duplicated knots and any knots equal to either the max or the min of X. If the resulting number of knots is equal to 0, the vector of knots is set to NULL. When the knots is NULL for a variable X, it means that U(X) = X.

The last step implies that the number of knots for all categorical variables with two levels, which includes as a special case binary variables, is equal to 0. For other categorical variables with a small number of levels, the number of knots may be smaller than the ones defined by nknots. For example, when the number of levels is three, the number of knots cannot exceed 1.

The starting knots can be extracted from the object. The elements knots0 and knots1 are the list of knots for the control and treated groups. For example, the knots for the treated are:

model1\$knots1

```
## $age
## 20% 40% 60% 80%
    19 22 25 28
##
##
## $re75
##
         40%
                   60%
                             80%
##
    357.9499 1961.8640 5588.6640
##
## $ed
## 20% 40% 60% 80%
##
     9 10 11 12
##
## $married
## NULL
```

We see that it is set to NULL for married, because it is a binary variable. The number of treated workers is 297. Given the default nknots, it implies a number of starting knots equal to 4. This is the number of knots we have for ed and age, but not for re75. The reason is that re75 contains many zeros. Since the 20% quantile is equal to 0 and 0 in also the minimum value of ed75, it is dropped (the type argument is to replicate what is implemented in the package).

```
quantile(nsw[nsw$treat==1,'re75'], c(.2,.4,.6,.8), type=1)
## 20% 40% 60% 80%
## 0.0000 357.9499 1961.8640 5588.6640
```

By printing the object, we see a summary of the model. It includes the list of variable with a positive number of knots and the ones with no knots.

model1

```
## Semiparametric Thresholding LSE Model
## ****************
##
## Number of treated: 297
## Number of control: 425
## Number of missing values: 0
## Selection Method: SLSE
## Covariates being approximated by a piecewise function:
## age, re75, ed
## Covariates not being approximated by a piecewise function:
## ## Tovariates not being approximated by a piecewise function:
## ## Tovariates not being approximated by a piecewise function:
## married
```

SLSE: We see that the selection method is set to SLSE, which stands for Semiparametric Least Squares Estimator. We refer to this when the knots are automatically selected by the method described above. Later in the document, we will present methods for selecting a subset of this SLSE selection.

As another example, the simulated dataset simDat4 contains special types of covariates. It help illustrate better how the knots are determined. The dataset contains a continuous variable X1 with a large proportion of zeros, categorical variables X2 and X3 with 2 and 3 levels, respectively, and a binary variable X4.

```
data(simDat4)
model2 <- setModel(Y~Z |~X1+X2+X3+X4, data=simDat4)</pre>
```

model2\$knots0

```
## $X1
##
          40%
                      60%
                                  80%
    0.2531388 2.9118507 12.1110772
##
## $X2
## NULL
##
## $X3
## 40%
##
##
## $X4
## NULL
```

We see that the number of knots for the two categorical variables with 2 levels is set to 0 and it is equal to 1 for the one with two levels.

Setting the number of knots to 0 for specific variables

To avoid having a positive number of knots for a variable, we can enter its name in the argument userRem. For example, if we want the number of knots to be zero for ed and age, we can create the model as follows:

```
## Semiparametric Thresholding LSE Model
## *****************
##
## Number of treated: 297
## Number of control: 425
## Number of missing values: 0
## Selection Method: SLSE
## Covariates being approximated by a piecewise function:
## re75
## Covariates not being approximated by a piecewise function:
## age, ed, married
```

We see that only re75 has a positive number of knots.

Setting the knots manually

We have the control over the knots through the arguments knots0 and knots1. When the arguments are missing (the default), all knots are set automatically. One way to set the number of knots to 0 for all variables in a given group is to set the argument to NULL. For example, the number of knots is equal to 0 for all variables of the treated group in the following:

```
## Selection Method: User Based
## Covariates being approximated by a piecewise function:
## Treated: None
## Control: age, re75, ed
## Covariates not being approximated by a piecewise function:
## Treated: age, re75, ed, married
## Control: married
```

Notice that the selection method is defined as "User Based" whenever knots are provided manually by the user. The other option is to provide a list of knots. The list must have the same length as the number of covariates. For each element, we have three options:

- NA: The knots are set automatically for this variable only.
- NULL: The number of knots is set to 0 for this variable only.
- A numeric vector: The vector cannot contain missing or duplicated values and must be strictly inside the range of the variable for the group.

Suppose you want to set for the control group an automatic selection for age, no knots for ed and the knots {1000,5000,10000} for re75, and let the knots be automatically selected for the treated group, we proceed this way. Note that setting the value to NA or NULL has the same effect for the binary variable married. In the following, the argument knots=TRUE is added to the print method to only print the knots.

```
model <- setModel(re78~treat | ~age+re75+ed+married, data=nsw,
                 knots0=list(NA, c(1000,5000,10000), NULL, NA))
print(model, knots=TRUE)
## Lists of knots for the treated group
## ***********
## age:
## 20%
       40%
            60%
                 80%
##
   19
        22
             25
                  28
## re75:
                              80%
##
        40%
                   60%
   357.9499
             1961.8640
##
                        5588.6640
## ed:
##
  20%
       40%
            60%
                 80%
##
    9
        10
                  12
             11
## married:
## None
##
## Lists of knots for the Control group
## **********
## age:
## 16.66667%
             33.33333%
                                   66.66667%
                              50%
##
         18
                    20
                               23
                                         26
                                                    30
## re75:
##
     k1
            k2
                   k3
##
   1000
          5000 10000
## ed:
## None
## married:
## None
```

Estimating the model

Given the set of knots from the model object, the estimation is just a least squares method. We want to estimate the model

$$Y = \beta_0(1-Z) + \beta_1 Z + \psi_0' U_0(X) + \psi_1' U_1(X) + u,$$

where $U_0(X)$ and $U_1(X)$ are the bases defined above and depends on the model knots. The function that estimate the model is estModel. The function has three arguments, but two of them are mostly used internally by other functions. We present it in case it is needed. The arguments are:

- model: A model created by the function setModel.
- w0: A list of integers to select knots for the control group from the model. By default, all the knots are used.
- w1: A list of integers to select knots for the treated group from the model. By default, all the knots are used

We illustrate with a simple model containing only two covariates and one knot per eligible variables.

```
model <- setModel(re78~treat | ~age+married, data=nsw,</pre>
                   nknots=function(n) 2)
fit <- estModel(model)</pre>
## Semiparametric Thresholding LSE Estimate
##
##
  factor(treat)0
                    factor(treat)1
                                                             Xf0age_2
                                                                             Xf0married
                                            Xf0age_1
       4558.28061
                         3754.98326
                                            27.79868
                                                             -12.51415
                                                                             -115.81593
##
##
                           Xf1age_2
                                          Xf1married
         Xf1age_1
                           22.22331
##
         89.25358
                                          1435,28205
```

The object has its own print method to returns the coefficient estimates. A more detrailed presentation of the results can be obtained using the summary method. The following is an example with just a one knot per eligible variable.

```
summary(fit)
```

```
## Semiparametric Thresholding LSE Estimate
##
##
                   Estimate Std. Error t value Pr(>|t|)
## factor(treat)0 4558.28
                               3380.43
                                          1.348
                                                   0.178
## factor(treat)1
                   3754.98
                               4043.48
                                          0.929
                                                    0.353
## Xf0age_1
                      27.80
                                164.59
                                          0.169
                                                   0.866
## Xf0age 2
                     -12.51
                                         -0.186
                                                   0.852
                                 67.11
                    -115.82
## XfOmarried
                                         -0.135
                                859.66
                                                   0.893
## Xf1age_1
                      89.25
                                 194.19
                                          0.460
                                                   0.646
## Xf1age_2
                      22.22
                                 76.46
                                          0.291
                                                   0.771
## Xf1married
                    1435.28
                               1014.69
                                          1.415
                                                   0.157
## Multiple R-squared: 0.009618,
                                      Adjusted R-squared:
                                                           -9.119e-05
Note that the R^2 and adjusted R^2 are different from what we obtain using the summary of the 1m object:
summary(fit$lm.out)[c("r.squared","adj.r.squared")]
```

```
## $r.squared
## [1] 0.4379272
```

```
## ## $adj.r.squared
## [1] 0.4316295
```

This is because R thinks that our model does not contain an intercept and the R^2 is computed differently for models without an intercept. The definition of the R^2 used by R is the following (RSS means residual sum of squares):

$$R^2 = 1 - \frac{\text{RSS for the model with the regressors}}{\text{RSS for the model without the regressors}}$$

In a model with an intercept, the residual of the model without the regressors is $Y_i - \bar{Y}$, but it is equal to Y_i when the model does not have an intercept. As a result, the R^2 with and without an intercept are

$$R_{with}^{2} = 1 - \frac{\sum_{i=1}^{n} \hat{e}_{i}^{2}}{\sum_{i=1}^{n} (Y_{i} - \bar{Y})^{2}}$$
$$R_{without}^{2} = 1 - \frac{\sum_{i=1}^{n} \hat{e}_{i}^{2}}{\sum_{i=1}^{n} Y_{i}^{2}}$$

However, our model does contain an intercept since we include a binary variable for both the control and treated groups. To illustrate the issue, the following two regression models are identical in terms of goodness of fit, because the sets of regressors span the same vector space:

$$re78 = \beta_1 + \beta_2 married + u$$
$$re78 = \alpha_1 married + \alpha_2 (1 - married) + u$$

But R computes very different R^2 :

```
summary(lm(re78~married, nsw))$r.squared

## [1] 0.001505512
summary(lm(re78~factor(married)-1, nsw))$r.squared
```

```
## [1] 0.4333229
```

The second R^2 overestimates the goodness of fit of our model and should not be used. The one returned by estModel is the right one.

The predict and plot method

The predict method is very similar to the predict.lm method. We find the same arguments: object, interval, se.fit, newdata and level. The difference is that it returns the predicted outcome for the treated and control groups separately and the argument vcov., a function like vcovHC or vcovCL, can be used to compute robust standard errors. The function return a list of two elements, treated and control. Each element contains the prediction fit and the standard errors se.fit when se.fit is set to TRUE. When interval is set to "confidence", fit is a matrix containing the prediction, and the lower and upper bound of the confidence interval. Here is an example with the previous simplified model:

```
## $treated
## fit lower upper
## 1 6975.337 4960.082 8990.592
## 2 7064.591 5119.244 9009.937
##
## $control
## fit lower upper
## 3 5054.036 3455.978 6652.093
## 4 5081.834 3423.558 6740.110
```

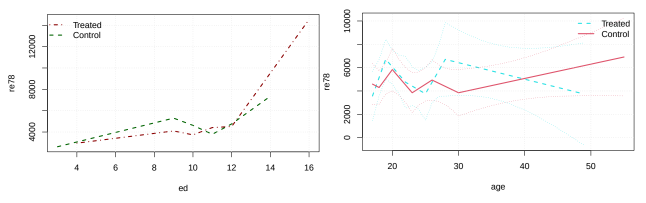
The predict method is called by the plot method to compare the predicted outcome for the treated and control group with respect to a given covariate. By default, all other covariates are fixed to their sample means. Consider the following model:

```
model1 <- setModel(re78~treat | ~age+re75+ed+married, data=nsw)
fit1 <- estModel(model1)</pre>
```

Suppose we want to compare the predicted income with respect to age or education, holding the other covariates fixed to their means. The following show some possible options.

Outcome versus ed using piecewise polynomials

Outcome versus age using piecewise polynomials

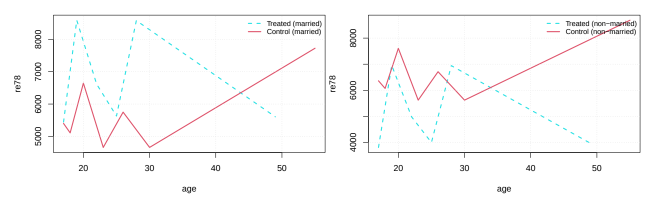


It is also possible to set some of the other covariates to a specific value by changing the argument newdata. This argument must be a named vector with the names corresponding to the variables you want to fix. You can also add a description to the legend with the argument addToLegend.

```
plot(fit1, "age", newdata=c(married=1, re75=10000), addToLegend="married", cex=0.8)
plot(fit1, "age", newdata=c(married=0, re75=10000), addToLegend="non-married", cex=0.8)
```

Outcome versus age using piecewise polynomials

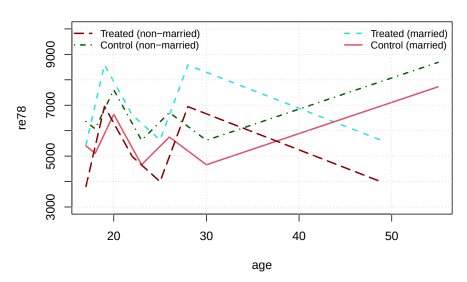
Outcome versus age using piecewise polynomials



To be better compare the two, it is also possible to have them plotted on the same graph by setting the argument add. to TRUE. We just to be careful and adjust the arguments correctly to avoid confusion.

```
plot(fit1, "age", newdata=c(married=1, re75=10000), addToLegend="married", cex=0.8,
        ylim.=c(3000,10000))
plot(fit1, "age", newdata=c(married=0, re75=10000), addToLegend="non-married", cex=0.8,
        legendPos='topleft', col0="darkgreen", col1="darkred", lty0=4, lty1=5,
        add.=TRUE,)
```

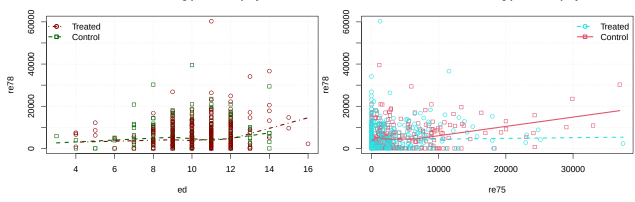
Outcome versus age using piecewise polynomials



Finally, it is also possible to add the observed points to the graph.

Outcome versus ed using piecewise polynomials

Outcome versus re75 using piecewise polynomials



The causal function

Once we have a model with knots, we can estimate the different causal effects. This is done by the causal function. The function assumes we are satisfied with the knots and estimate the causal effects and their standard errors. To define the different causal effect measures, let's redefine $U_0(X)$ and $U_1(X)$ as the spline bases using the knots of the control and treated group respectively, but with all data points. This differs from how it is defined in the introduction, because this $U_0(X_i)$ is not equal to 0 when $Z_i = 1$ and $U_1(X_i)$ is not equal to 0 when $Z_i = 0$. The regression estimated by estModel, or the one defined in the introduction, can be written as

$$Y = \beta_0(1-Z) + \beta_1 Z + \psi_0'[U_0(X)(1-Z)] + \psi_1'[U_1(X)Z] + u.$$

Let $\hat{\beta}_0$, $\hat{\beta}_1$, $\hat{\psi}_0$ and $\hat{\psi}_1$ be the least squares estimates. Then, the estimated causal effects are defined as:

$$\begin{split} & \text{ACE} &= \hat{\beta}_1 - \hat{\beta}_0 + \hat{\phi}_1' \overline{U_1(X)} - \hat{\phi}_0' \overline{U_0(X)} \\ & \text{ACT} &= \hat{\beta}_1 - \hat{\beta}_0 + \hat{\phi}_1' \overline{U_1(X)Z} - \hat{\phi}_0' \overline{U_0(X)Z} \\ & \text{ACN} &= \hat{\beta}_1 - \hat{\beta}_0 + \hat{\phi}_1' \overline{U_1(X)(1-Z)} - \hat{\phi}_0' \overline{U_0(X)(1-Z)} \,, \end{split}$$

where

$$\overline{U_j(X)} = \frac{1}{n} \sum_{i=1}^{n} U_j(X_i), \text{ for } j=0,1$$

$$\overline{U_j(X)Z} = \frac{1}{n_1} \sum_{i=1}^{n} U_j(X_i) Z_i, \text{ for } j=0,1$$

$$\overline{U_j(X)(1-Z)} = \frac{1}{n_0} \sum_{i=1}^{n} U_j(X_i) (1-Z_i), \text{ for } j=0,1$$

and n_0 and n_1 are the number of individuals in the control and treated groups. The function causal is a method registered for tlseFit and tlseModel objects. In other words, we can compute the causal effects directly from the model:

causal(model1)

Causal Effect using Thresholding Least Squares

We see that the selection method (for the knots) and the criteria used to select the knots are set to unknown. This is because it is not specified in the model object how the knots were selected. We will clarify this below. The method return an object of class causaltlse. We see above what its print method returns and the following show its summary method:

```
ce <- causal(model1)
summary(ce)
## Causal Effect using Thresholding Least Squares
## ************
## Selection Method: SLSE
##
      Estimate Std. Error t value Pr(>|t|)
         814.3
                   482.1
## ACE
                          1.689
                                  0.0912 .
## ACT
         831.9
                   499.5
                          1.665
                                  0.0958 .
## ACN
         802.0
                   498.9
                          1.608
                                  0.1079
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The standard errors are computed using an analytical expression derived in the paper (need to add a citation to our paper), which takes into account the variance of the sample means of the covariates. Asymptotically, these variances converge to 0, so it only makes a difference in small samples. Alternatively, we can set the argument seType to "lm" and use the least squares standard errors based on the asymptotic properties. By default, vcov.lm is used, but it is possible to modify it by changing the argument vcov.. In the following, we estimate the standard errors using the HC3 type of heteroskedasticity robust standard errors.

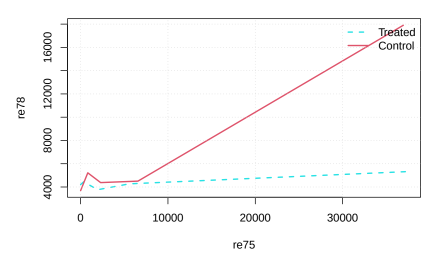
```
ce2 <- causal(model1, seType="lm", vcov.=vcovHC, type="HC3")
summary(ce2)</pre>
```

```
## Causal Effect using Thresholding Least Squares
## *************
## Selection Method: SLSE
##
      Estimate Std. Error t value Pr(>|t|)
## ACE
         814.3
                   506.1
                          1.609
                                  0.108
## ACT
         831.9
                   527.4
                                  0.115
                          1.577
## ACN
         802.0
                   514.2
                          1.560
                                  0.119
```

The object causaltlse inherits from the class tlseFit, so we can apply the plot (or the predict) method directly on this object.

```
plot(ce2, "re75")
```

Outcome versus re75 using piecewise polynomials



Optimal selection of the knots

We propose two methods for selecting the knots: a backward (BTLSE) and a forward (FTLSE) methods. For each method, we propose three criteria: the asymptotic (ASY), the Akaike Information (AIC) and the Bayesian Information (BIC). The two selection methods can be summarized as follows:

BTLSE:

- 1. We estimate the model with all knots included in the model.
- 2. For each knot, we test if the slope of the piecewise linear polynomial is the same before and after, and return the p-value.
- 3. The knots are selected using one of the following criteria
 - ASY: We remove all knots with a p-value greater than a specified threshold.
 - AIC or BIC: We order the p-values in descending order. Then, going from the largest to the smallest, we remove the knot associated with the p-value one by one, estimate the model and return the information criterion. We keep the model with the smallest information citerion.

FTLSE:

- 1. We estimate the model by including a subset of the knots one variable at the time. When we test a knot for one variable, the number of knots is set to 0 for all the others.
- 2. For each knot, we test if the slope of the piecewise linear polynomial is the same before and after, and return the p-value. The set of knots used for each test depends on the following:
 - Variables with 1 knot: we return the p-value of the test of equality before and after the
 - Variables with 2 knots: we include the two knots and return the p-values of the test of equality before and after for each knot.
 - Variables with p knots (p > 2): We test the equality before and after the knot i, for i = 1, ..., p, using the sets of knots $\{1, 2\}, \{1, 2, 3\}, \{2, 3, 4\}, ..., \{p 2, p 1, p\}$ and $\{p 1, p\}$ respectively.
- 3. The knots are selected using one of the following criteria
 - ASY: We remove all knots with a p-value greater than a specified threshold.

• **AIC** or **BIC**: We order the p-values in ascending order. Then, starting with a model with no knots and going from the smallest to the highest highest p-value, we add the knot associated with the p-value one by one, estimate the model and return the information criterion. We keep the model with the smallest information citerion.

The selection is done using the function selTLSE. The arguments are:

- model: An object of class tlseModel.
- method: This is the selection method. We have the choice between "FTLSE" (the default) and "BTLSE".
- **crit**: This is the criterion used by the selection method. We have the choice between "ASY" (the default), "AIC" or "BIC".
- minPV: This is a function that returns the p-value threshold. It is a function of one argument, the average number of knots per covariate. The default is function(p) 1/log(p). It is also possible to set it to a fix threshold. For example, function(p) 0.20 set the threshold to 0.2. This argument affects the result only when method is set to "ASY".
- vcov.: By default, the p-values are computed with the lm covariance matrix method vcov. Alternatively, we can use sandwich estimators like vcovHC.
- ...: This is used to pass arguments to the vcov. function.

The function returns a model of class tlseModel with the optimal selection of knots. For example, we can compare the starting knots of model1, with the model selected by the default arguments.

```
print(model1, knots=TRUE)
```

```
## Lists of knots for the treated group
## ************
## age:
## 20% 40% 60% 80%
## 19
        22
            25
                 28
## re75:
        40%
                            80%
                  60%
##
  357.9499
            1961.8640
## ed:
## 20%
       40%
            60%
                80%
##
    9
        10
            11
                 12
## married:
## None
##
## Lists of knots for the Control group
## ***********
## age:
## 16.66667% 33.33333%
                                 66.66667% 83.33333%
                             23
##
                   20
                                        26
                                                  30
         18
## re75:
        50% 66.66667% 83.33333%
##
## 823.2544 2292.1710
                      6567.3290
## ed:
## 16.66667%
            33.33333%
                       66.66667%
                                 83.33333%
##
          9
                   10
                             11
## married:
## None
model2 <- selTLSE(model1)</pre>
print(model2, knots=TRUE)
## Lists of knots for the treated group
## *******************
## age:
## 20% 40% 60% 80%
## 19
       22
            25
                 28
## re75:
```

```
##
       60%
## 1961.864
## ed:
## 20% 40% 60% 80%
  9 10
##
            11
                12
## married:
## None
##
## Lists of knots for the Control group
## ********************
## age:
## 33.3333%
                  50% 66.66667% 83.33333%
                   23
                            26
## re75:
       50% 66.66667% 83.33333%
##
## 823.2544 2292.1710 6567.3290
## ed:
## 16.66667% 66.66667%
##
         9
                   11
## married:
## None
```

For example, the method has removed all knots from re75 for the treated group and kept 2 knots for the control group. We can then compute the causality measures for the new model. Notice that the selection method and criterion reflects what was used to update the model. In this case, we see FTLSE as selection method and ASY as criterion.

```
causal(model2)
```

```
## Causal Effect using Thresholding Least Squares
## *******************************
## Selection Method: FTLSE
## Criterion: ASY
##
## ACE = 820.3279
## ACT = 837.1689
## ACN = 808.559
We can compare with other methods:
model3 <- selTLSE(model1, method="BTLSE", crit="BIC")</pre>
causal(model3)
## Causal Effect using Thresholding Least Squares
## ************
## Selection Method: BTLSE
## Criterion: BIC
##
## ACE = 818.8162
## ACT = 889.3806
## ACN = 769.5041
```

The extract method

The package comes with an extract method for objects of class causaltlse, which is a required method for creating Latex tables using the texreg package. For example, we can compare different methods in a single table.

```
library(texreg)
c1 <- causal(model1)
c2 <- causal(selTLSE(model1, method="BTLSE"))</pre>
```

```
c3 <- causal(selTLSE(model1, method="FTLSE"))
texreg(list(SLSE=c1, BTLSE=c2, FTLSE=c3), table=FALSE, digits=4)</pre>
```

	SLSE	BTLSE	FTLSE
ACE	814.3083	816.1558	820.3279
	(482.1393)	(483.7087)	(483.7472)
ACT	831.8856	827.8434	837.1689
	(499.4948)	(501.7417)	(500.7082)
ACN	802.0249	807.9883	808.5590
	(498.8671)	(495.9641)	(495.3648)
Num. knots (Control)	12	10	9
Num. knots (Treated)	11	8	9
Num. covariates	4	4	4
Num. obs.	722	722	722
\mathbb{R}^2	0.0869	0.0866	0.0863
R_{adj}^2	0.0445	0.0511	0.0507

***p < 0.001; **p < 0.01; *p < 0.05

The causalTLSE function

We just saw how to estimate the causal effects step by step. The function causalTLSE estimate them in one step, once the model has been created. It returns an object of class causaltlse like the causal method does, so we can apply the same print, summary and predict and plot method to it. The last two can be applied to the object, because it inherits from the tlseFit class. The arguments are almost like the ones from the selTLSE and causal functions.

- model: An object of class tlseModel.
- selType: This is the selection method. We have the choice between "SLSE" (the default), "FTLSE" and "BTLSE". The SLSE method implies no selection, so all knots from the model are kept. It is therefore identical to estimating the model using the causal method.
- selCrit: This is the criterion used by the selection method. We have the choice between "ASY" (the default), "AIC" or "BIC".
- causal: What causality measure should the function compute? We have the choice between "All" (the default), "ACT", "ACE" or "ACT".
- **seType**: The method to compute the standard error of the causality measures. We have the choice between "analytical" (the default) or "lm". We have explained the difference when we presented the causal method.
- minPV: This is a function that returns the p-value threshold. We explained this argument when we presented the sellTLSE function.
- vcov.: An alternative was to compute the covariance matrix of the least squares estimates.
- ...: This is used to pass arguments to the vcov. function.

For example, we can generate the previous table as follows:

```
c1 <- causalTLSE(model1, selType="SLSE")
c2 <- causalTLSE(model1, selType="BTLSE")
c3 <- causalTLSE(model1, selType="FTLSE")
texreg(list(SLSE=c1, BTLSE=c2, FTLSE=c3), table=FALSE, digits=4)</pre>
```

	SLSE	BTLSE	FTLSE
ACE	814.3083	816.1558	820.3279
	(482.1393)	(483.7087)	(483.7472)
ACT	831.8856	827.8434	837.1689
	(499.4948)	(501.7417)	(500.7082)
ACN	802.0249	807.9883	808.5590
	(498.8671)	(495.9641)	(495.3648)
Num. knots (Control)	12	10	9
Num. knots (Treated)	11	8	9
Num. covariates	4	4	4
Num. obs.	722	722	722
\mathbb{R}^2	0.0869	0.0866	0.0863
R _{adj}	0.0445	0.0511	0.0507

***p < 0.001; **p < 0.01; *p < 0.05

An example with simulated data

In the package, the data set datSim1 was generated using the following data generating process.

$$Y(0) = 1 + X + X^{2} + e$$

 $Y(1) = 1 - 2X + u$
 $Z = Ber[\Lambda(1 + X)]$
 $Y = Y(1)Z + Y(0)(1 - Z)$

where X, e and u are standard normal, $\Lambda(x)$ is the CDF of the standard logistic distribution and Ber(p) is the Bernoulli distribution. The true causal effects ACE, ACT and ACN are approximately equal to -1, -1.6903 and 0.5867. We can start by building starting model:

```
data(simDat1)
mod <- setModel(Y~Z | ~X, data=simDat1)</pre>
```

Then we can compare three different methods:

```
c1 <- causalTLSE(mod, selType="SLSE")
c2 <- causalTLSE(mod, selType="BTLSE", selCrit="BIC")
c3 <- causalTLSE(mod, selType="FTLSE", selCrit="BIC")
texreg(list(SLSE=c1, BTLSE=c2, FTLSE=c3), table=FALSE, digits=4)</pre>
```

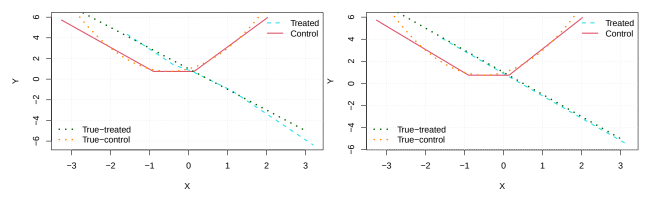
	SLSE	BTLSE	FTLSE	
ACE	-1.4396***	-1.4530***	-1.4530***	
	(0.2614)	(0.2605)	(0.2605)	
ACT	-1.9316***	-1.9316***	-1.9316***	
	(0.3030)	(0.3024)	(0.3024)	
ACN	-0.0865	-0.1369	-0.1369	
	(0.3263)	(0.3224)	(0.3224)	
Num. knots (Control)	2	2	2	
Num. knots (Treated)	4	0	0	
Num. covariates	1	1	1	
Num. obs.	300	300	300	
\mathbb{R}^2	0.7434	0.7386	0.7386	
R_{adj}^2	0.7354	0.7342	0.7342	
*** $p < 0.001$; ** $p < 0.01$; * $p < 0.05$				

We see that both selection methods choose to assign 0 knots for the treated group, which is not surprising since the true $f_1(x)$ is linear. We can compare the different fits (we ignore the FTLSE because the selected knots are the same):

```
plot(c1, "X")
curve(1-2*x, -3,3, col="darkgreen", lty=3, lwd=3, add=TRUE)
curve(1+x+x^2, -3,3, col="darkorange", lty=3, lwd=3, add=TRUE)
```

Outcome versus X using piecewise polynomials

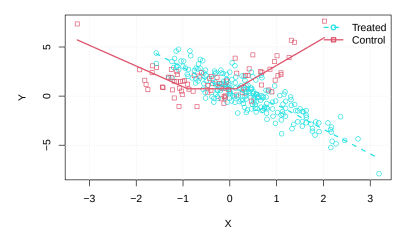
Outcome versus X using piecewise polynomials



We see that the piecewise polynomials are very close to the true $f_1(x)$ and $f_2(x)$. We can see from the following graph how the lines are fit through the observations by group.

plot(c1, "X", addPoints=TRUE)

Outcome versus X using piecewise polynomials



An example with another simulated data

The dataset datSim2 was generated using the following data generating process.

$$Y(0) = (1+X)I(X \le -1) + (-1-X)I(X > -1) + e$$

$$Y(1) = (1-2X)I(X \le 0) + (1+2X)I(X > 0) + e$$

$$Z = Ber[\Lambda(1+X)]$$

$$Y = Y(1)Z + Y(0)(1-Z)$$

where I(A) is the indicator function equal to 1 if A is true, X, e and u are standard normal, $\Lambda(x)$ is the CDF

of the standard logistic distribution and Ber(p) is the Bernoulli distribution. The true causal effects ACE, ACT and ACN are approximately equal to 3.763, 3.858 and 3.545. We can compare the SLSE, BTLSE with AIC and BTLSE with BIC.

```
data(simDat2)
mod <- setModel(Y~Z | ~X, data=simDat2)

c1 <- causalTLSE(mod, selType="SLSE")
c2 <- causalTLSE(mod, selType="BTLSE", selCrit="BIC")
c3 <- causalTLSE(mod, selType="BTLSE", selCrit="AIC")
texreg(list(SLSE=c1, BTLSE.BIC=c2, BTLSE.AIC=c3), table=FALSE, digits=4)</pre>
```

	SLSE	BTLSE.BIC	BTLSE.AIC	
ACE	3.9290***	3.9201***	3.9201***	
	(0.1703)	(0.1717)	(0.1717)	
ACT	3.9552***	3.9404***	3.9404***	
	(0.1891)	(0.1904)	(0.1904)	
ACN	3.8670***	3.8721***	3.8721***	
	(0.2371)	(0.2362)	(0.2362)	
Num. knots (Control)	2	1	1	
Num. knots (Treated)	3	2	2	
Num. covariates	1	1	1	
Num. obs.	300	300	300	
\mathbb{R}^2	0.7833	0.7829	0.7829	
R_{adj}^2	0.7774	0.7784	0.7784	
*** $p < 0.001$; ** $p < 0.01$; * $p < 0.05$				

The following illustrate the fit of BTLSE-AIC with the true $f_1(x)$ and $f_0(x)$, and the observations.

Outcome versus X using piecewise polynomials

Outcome versus X using piecewise polynomials

