Package ‘DTDA’

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DTDA-package

Description

Implementation of different algorithms for analyzing randomly truncated data, one-sided and two-sided (i.e. doubly) truncated data.

The package allows for the estimation of the distribution function of the doubly truncated (target) variable. The package provides estimators for the distribution of the truncation variables too. Pointwise confidence limits based on bootstrap methods are included. Automatic plots of cumulative distributions and survival functions are provided. The package also implements the kernel density estimator for doubly truncated data with different bandwidth selectors. The hazard rate function with least square cross-validation selector type is also included. Real datasets are provided within the package. Besides the right-truncated AIDS data, eight doubly truncated datasets are available: Childhood Cancer Data, AIDS Blood Transfusion Data, Equipment-S Rounded Failure Time Data, Quasars Data, Parkinson’s Disease Data (early and late onset groups), and Acute Coronary Syndrome Data (complete and reduced).

Details

Package: DTDA
Type: Package
Version: 3.0
Date: 2021-04-08
License: GPL-2
LazyLoad: yes

Missing data are allowed. Registries with missing data are simply removed. This package incorporates the functions efron.petrosian, lynden, shen to implement the iterative methods to compute the NPMLE for doubly truncated data; densityDT, hazardDT to calculate the kernel estimators of the density and hazard functions, respectively. The function rsim.DT, allows to simulate doubly truncated data in two different settings. For a complete list of functions, use library(help="DTDA").
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Author(s)

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ACS

Acute Coronary Syndrome data

Description

The data include information of 939 patients with confirmed diagnosis of type 1 (primary spontaneous) acute coronary syndrome (ACS). Patients were consecutively admitted to the Cardiology Department of two tertiary hospitals in Portugal between August 2013 and December 2014. The age at diagnosis is doubly truncated because of the interval sampling.

Usage

data(ACS)

Format

A data frame with 939 observations on the following 5 variables.

X  a numeric vector, age at diagnosis (in years).
U  a numeric vector, the elapsed time (in years) between birth and the beginning of the study (August 2013).
V  a numeric vector, the elapsed time (in years) between birth and end of the study (December 2014).
Sex  a numeric vector, sex of the participants (0 = female, 1 = male).
diagnosis  a numeric vector, type of diagnosis at discharge 1 - STEMI (ST elevation myocardial infarction) and 2 - NSTEMI (all others diagnoses).

Details

The age at diagnosis X is doubly truncated due to the interval sampling. The length of the sampling interval (V-U) is 1.42 years. The NPMLE of the cumulative distribution function of X does not exist or is not unique for this dataset. The necessary and sufficient graphical condition presented by Xiao and Hudgens (2020) to determine the existence and uniqueness of the NPMLE is not satisfied.
References


See Also

ACSred

Examples

```r
data(ACS)
str(ACS)
```

ACSred Acute Coronary Syndrome reduced data

Description

The data include information of 917 patients with confirmed diagnosis of type 1 (primary spontaneous) acute coronary syndrome (ACS). Patients were consecutively admitted to the Cardiology Department of two tertiary hospitals in Portugal between August 2013 and December 2014. The age at diagnosis is doubly truncated because of the interval sampling.

Usage

```r
data(ACSred)
```

Format

A data frame with 917 observations on the following 5 variables.

- **X**: a numeric vector, age at diagnosis (in years).
- **U**: a numeric vector, the elapsed time (in years) between birth and the beginning of the study (August 2013).
- **V**: a numeric vector, the elapsed time (in years) between birth and the end of the study (December 2014).
- **Sex**: a numeric vector, sex of the participants (0 = female, 1 = male).
- **diagnosis**: a numeric vector, type of diagnosis at discharge 1 - STEMI (ST elevation myocardial infarction) and 2 - NSTEMI (all others diagnoses).
Details

The age at diagnosis $X$ is doubly truncated due to the interval sampling. The length of the sampling interval ($V-U$) is 1.42 years. The NPMLE of the cumulative distribution function of $X$ for the complete data does not exist or is not unique for this dataset. This dataset is a reduced sample of the original ACS data, guaranteeing the existence and uniqueness of the NPMLE, according to Xiao and Hudgens (2020).

References


See Also

ACS

Examples

data(ACSred)
str(ACSred)

AIDS Blood Transfusion Data

Description

The data include information on the infection and induction times for 258 adults who were infected with HIV virus and developed AIDS by June 30, 1996. The data consist on the time in years, measured from April 1, 1978, when adults were infected by the virus from a contaminated blood transfusion, and the waiting time to development of AIDS, measured from the date of infection.

Usage

data(AIDS)

Format

A data frame with 258 observations on the following 3 variables.

INFTime a numeric vector, the infection time.
INDTime a numeric vector, the induction time.
V a numeric vector, the time from HIV infection to the end of the study.
Source
J.P. Klein and M.L.Moeschberger.

References

Examples
data(AIDS)
str(AIDS)

AIDS.DT  
* AIDSBlood Transfusion Data

Description
The data include information of transfusions cases of transfusion-related AIDS, corresponding to individuals diagnosed prior to July 1, 1986. Only 295 patients with consistent data, for which the infection could be attributed to a single transfusion or short series were included. Since HIV was unknown before 1982, this implies that cases developing AIDS prior to this date were not reported, leading to a doubly truncated data. The incubation time is doubly truncated due to the interval sampling.

Usage
data(AIDS.DT)

Format
A data frame with 295 observations on the following 4 variables.

- **X**: a numeric vector, the induction or incubation time: time elapsed from HIV infection to AIDS (in months).
- **U**: a numeric vector, time from 1982 to HIV infection (in months).
- **V**: a numeric vector, time from HIV infection to July 1, 1986 (in months).
- **AGE**: a numeric vector, age of the individual at diagnosis (in months).

Source
Kalbfleisch JD and Lawless JF

References
**ChildCancer**

**Examples**

```r
data(AIDS.DT)
str(AIDS.DT)
```

**Description**

This dataset corresponds to all children diagnosed from cancer between January 1, 1999 and December 31, 2003 in the region of North Portugal. The database includes information of 406 children with complete records on the age at diagnosis. Because of the interval sampling, the age at diagnosis is doubly truncated by the time from birth to the end of the study, and time from birth to the beginning of the study (time in days). The age at diagnosis is doubly truncated due to the interval sampling.

**Usage**

```r
data("ChildCancer")
```

**Format**

A data frame with 406 observations on the following 8 variables.

- **X** a numeric vector, age at diagnosis (time in days).
- **U** a numeric vector, time from birth to the beginning of the study (time in days).
- **V** a numeric vector, time from birth to the end of the study (time in days).
- **ICCGroup** a numeric vector, cancer types identified according to the International Classification of Childhood Cancer (ICCC). 1=Leukaemias, 2=Lymphoma, 3=Nervous System Tumour, 4=Neuroblastoma, 5=Retinoblastoma, 6=Renal, 7=Hepatic, 8=Bone, 9=Soft Tissues, 10=Germ Cell, 11=Melanoma and other epithelial tumours, 12=Other Tumours.
- **Status** a numeric vector, the status indicator at the end of the study: 0=alive, 1=dead.
- **SurvTime** a numeric vector, the survival time (time from birth to death or end of the study).
- **Residence** a numeric vector, districts of residence. 1=Braga, 2=Bragança, 3=Porto, 4=Viana do Castelo, 5=Vila Real.
- **Sex** a numeric vector, sex of the participants (1 = female, 2 = male).

**Source**

The childhood cancer data were gathered from the IPO (Registo Oncológico do Norte) service, kindly provided by Doctor Maria José Bento.

**References**

Examples

```r
data(ChildCancer)
str(ChildCancer)
```

---

**densityDT**  
*Estimation of a kernel density function under random double truncation*

---

**Description**

This function provides the nonparametric kernel density estimation of a doubly truncated random variable. A bandwidth value is required.

**Usage**

```r
densityDT(X, U, V, bw = "DPI2", from, to, n, wg = NA)
```

**Arguments**

- `X` numeric vector with the values of the target variable.
- `U` numeric vector with the values of the left truncation variable.
- `V` numeric vector with the values of the right truncation variable.
- `bw` The smoothing bandwidth to be used, but can also be a character string giving a rule to choose the bandwidth. This must be one of "NR", "DPI1", "DPI2", "LSCV" or "SBoot" with default "DPI2".
- `from` the left point of the grid at which the density is to be estimated.
- `to` the right point of the grid at which the density is to be estimated.
- `n` number of evaluation points on a equally spaced grid.
- `wg` Numeric vector of non-negative initial solution, with the same length as `X`. Default value is the solution obtained with Efron and Petrosian algorithm.

**Details**

The nonparametric kernel density estimation for a variable which is observed under random double truncation is computed as proposed in Moreira and de Uña-Álvarez (2012). As usual in kernel smoothing, the estimator is obtained as a convolution between a kernel function and an appropriate estimator of the cumulative df. Gaussian kernel is used. The automatic bandwidth selection procedures for the kernel density estimator are those proposed in Moreira and Van Keilegom (2013). The automatic bandwidth selection alternatives are appropriate modifications, i.e, taking into account the double truncation issue, of the normal reference rule, two types of plug-in procedures, the least squares cross-validation and a bootstrap based method proposed in Cao et al. (1994) and Sheater and Jones (1991) for the complete data.
Value

A list containing the following values:

- `x` the n coordinates of the points where the density is estimated.
- `y` the estimated density values.
- `bw` the bandwidth used.

Author(s)

Carla Moreira, de de Uña-Álvarez and Rosa Crujeiras

References


See Also

hazardDT

Examples

```r
n <- 50
X <- runif(n, 0, 1)
U <- runif(n, -1/3, 1)
V <- U + 1/3
for (i in 1:n){
  while (U[i] > X[i] | V[i] < X[i]){X[i] <- runif(1, 0, 1)
    U[i] <- runif(1, -1/3, 1)
    V[i] <- U[i] + 1/3
  }
}

vxDens1 <- densityDT(X, U, V, bw = "DPI1", 0, 1, 500)
vxDens2 <- densityDT(X, U, V, bw = "DPI2", 0, 1, 500)
vxDens3 <- densityDT(X, U, V, bw = 0.5, 0, 1, 500)
```
efron.petrosian

Doubly truncated data analysis with the first Efron-Petrosian algorithm

Description

This function computes the NPMLE of a lifetime distribution function observed under one-sided (right or left) and two-sided (double) truncation. It provides bootstrap pointwise confidence limits too.

Usage

efron.petrosian(X, U = NA, V = NA, wt = NA, error = NA, nmaxit = NA, boot = TRUE, B = NA, alpha = NA, display.F = FALSE, display.S = FALSE)

Arguments

X Numeric vector with the times of ultimate interest.
U Numeric vector with the left truncation times. If there are no truncation times from the left, put U=NA.
V Numeric vector with the right truncation times. If there are no truncation times from the left, put V=NA.
wt Numeric vector of non-negative initial solution, with the same length as X. Default value is set to 1/n, being n the length of X.
error Numeric value. Maximum pointwise error when estimating the density associated to X (f) in two consecutive steps. If this is missing, it is $1e-06$.
nmaxit Numeric value. Maximum number of iterations. If this is missing, it is set to nmaxit =100.
boot Logical. If TRUE (default), the simple bootstrap method is applied to lifetime distribution estimation. Pointwise confidence bands are provided.
B Numeric value. Number of bootstrap resamples . The default NA is equivalent to B =500.
alpha Numeric value. (1-alpha) is the nominal coverage for the pointwise confidence intervals.
display.F Logical. Default is FALSE. If TRUE, the estimated cumulative distribution function associated to X, (F) is plotted.
display.S Logical. Default is FALSE. If TRUE, the estimated survival function associated to X, (S) is plotted.
Details

The NPMLE of the lifetime is computed by the first algorithm proposed in Efron and Petrosian (1999). This is an alternative algorithm which converges to the NMPLE after a number of iterations. If the second (respectively third) argument is missing, computation of the Lynden-Bell estimator for right-truncated (respectively left-truncated) data is obtained. Note that individuals with NAs in the three first arguments will be automatically excluded.

Value

A list containing the following values:

- time: The timepoint on the curve.
- n.event: The number of events that occurred at time t.
- events: The total number of events.
- density: The estimated density values.
- cumulative.df: The estimated cumulative distribution values.
- truncation.probs: The probabilities of truncation values, in each region.
- S0: Error reached in the algorithm.
- Survival: The estimated survival values.
- n.iterations: The number of iterations used by this algorithm.
- B: Number of bootstrap resamples computed.
- alpha: The nominal level used to construct the confidence intervals.
- upper.df: The estimated upper limits of the confidence intervals for F.
- lower.df: The estimated lower limits of the confidence intervals for F.
- upper.Sob: The estimated upper limits of the confidence intervals for S.
- lower.Sob: The estimated lower limits of the confidence intervals for S.
- sd.boot: The bootstrap standard deviation.
- Boot.Repeat: The number of resamples done in each bootstrap call to ensure the existence and uniqueness of the bootstrap NPMLE.

Author(s)

Carla Moreira, Jacobo de Uña-Álvarez and Rosa Crujeiras

References

## Generating data which are doubly truncated

```r
n <- 25
X <- runif(n, 0, 1)
U <- runif(n, 0, 0.5)
V <- runif(n, 0.5, 1)
for (i in 1:n)
  while (X[i] < U[i] | X[i] > V[i]){
    U[i] <- runif(1, 0, 0.5)
    X[i] <- runif(1, 0, 1)
    V[i] <- runif(1, 0.5, 1)
  }
efron.petrosian(X = X, U = U, V = V, boot = FALSE, display.F = TRUE, display.S = TRUE)
```

### Description

Digitized data from Figure X in Ye and Tang 2016. The dataset contains (rounded) observations of 174 failure times of certain devices, observed under interval sampling. Right truncation is years between installation and 2011 and left truncation corresponds to right truncation time minus 15 years. The failure time is doubly truncated due to the interval sampling.

### Usage

```r
data("EqSRounded")
```

### Format

A data frame with 174 observations on the following 3 variables.

- **X**: a numeric vector, time to failure in years.
- **U**: a numeric vector, the number of years between installation and 2011 minus 15 years.
- **V**: a numeric vector, the number of years between installation and 2011.

### Source

Digitalization of the data plot in the original paper of Ye and Tang 2016.
References

Examples
data(EqSRounded)
str(EqSRounded)

```r
hazardDT
```

## Description
This function provides the nonparametric kernel hazard estimation for a variable which is observed under random double truncation, which is defined as a convolution of a kernel function with the NPMLE of the cumulative df. Least square cross validation bandwidth selection procedure is implemented too.

## Usage
```r
hazardDT(X, U, V, bw = "LSCV", from, to, n, wg = NA)
```

## Arguments
- **X**: numeric vector with the values of the target variable.
- **U**: numeric vector with the values of the left truncation variable.
- **V**: numeric vector with the values of the right truncation variable.
- **bw**: The smoothing bandwidth to be used, but can also be a character string giving a rule to choose the bandwidth. This must be "LSCV".
- **from**: the left point of the grid at which the density is to be estimated.
- **to**: the right point of the grid at which the density is to be estimated.
- **n**: number of evaluation points on a equally spaced grid.
- **wg**: numeric vector of non-negative initial solution, with the same length as X. Default value is the solution obtained with Efron and Petrosian algorithm.

## Details
The nonparametric kernel density estimation for a variable which is observed under random double truncation is computed as proposed in *Moreira et al. (2021)*. As usual in kernel smoothing, the estimator is obtained as a convolution between a kernel function and an appropriate estimator of the cumulative df. Gaussian kernel is used. The automatic bandwidth selection procedures for the kernel hazard estimator is the least square cross validation, presented in *Moreira et al. (2021)*.
Value

A list containing the following values:

- x: the n coordinates of the points where the density is estimated.
- y: the estimated density values.
- bw: the bandwidth used.

Author(s)

Carla Moreira, Jacobo de Uña-Álvarez and Rosa Crujeiras

References


See Also

densityDT

Examples

set.seed(4321)

n<-100
X <- runif(n, 0, 1)
U <- runif(n,-1/3, 1)
V <- U + 1/3
for (i in 1:n){
    while (U[i] > X[i] | V[i] < X[i]){
        X[i] <- runif(1, 0, 1)
        U[i] <- runif(1, -1/3, 1)
        V[i] <- U[i] + 1/3
    }
}

vxhazard1<-hazardDT(X,U,V,bw=0.3,0,1,500)
vxhazard2<-hazardDT(X,U,V,bw="LSCV",0,1,500)
Doubly truncated data analysis with the second Efron-Petrosian algorithm

Description
This function computes the NPMLE of a lifetime distribution function observed under one-sided (right or left) and two-sided (double) truncation. It provides bootstrap pointwise confidence limits too.

Usage
lynden(X, U = NA, V = NA, error = NA, nmaxit = NA, boot = TRUE, B = NA, alpha = NA, display.F = FALSE, display.S = FALSE)

Arguments
X
Numeric vector with the times of ultimate interest.
U
Numeric vector with the left truncation times. If there are no truncation times from the left, put U=NA.
V
Numeric vector with the right truncation times. If there are no truncation times from the left, put V=NA.
error
Numeric value. Maximum pointwise error when estimating the density associated to X (f) in two consecutive steps. If this is missing, it is $1e-06$.
nmaxit
Numeric value. Maximum number of iterations. If this is missing, it is set to nmaxit = 100.
boot
Logical. If TRUE (default), the simple bootstrap method is applied to lifetime distribution estimation. Pointwise confidence bands are provided.
B
Numeric value. Number of bootstrap resamples. The default NA is equivalent to B = 500.
alpha
Numeric value. (1-alpha) is the nominal coverage for the pointwise confidence intervals.
display.F
Logical. Default is FALSE. If TRUE, the estimated cumulative distribution function associated to X, (F) is plotted.
display.S
Logical. Default is FALSE. If TRUE, the estimated survival function associated to X, (S) is plotted.

Details
The NPMLE of the lifetime is computed by the second algorithm proposed in Efron and Petrosian (1999). This is an alternative algorithm which converges to the NMLE after a number of iterations. If the second (respectively third) argument is missing, computation of the Lynden-Bell estimator for right-truncated (respectively left-truncated) data is obtained. Note that individuals with NAs in the three first arguments will be automatically excluded.
Value

A list containing the following values:

time The timepoint on the curve.
n.event The number of events that occurred at time \( t \).
events The total number of events.
NJ The number of individuals in risk considering the left truncation times.
density The estimated density values.
cumulative.df The estimated cumulative distribution values.
truncation.probs The probabilities of truncation values, in each region.
hazard The estimated hazard values.
S0 error reached in the algorithm.
Survival The estimated survival values.
n.iterations The number of iterations used by this algorithm.
B Number of bootstrap resamples computed.
alpha The nominal level used to construct the confidence intervals.
upper.df The estimated upper limits of the confidence intervals for F.
lower.df The estimated lower limits of the confidence intervals for F.
upper.Sob The estimated upper limits of the confidence intervals for S.
lower.Sob The estimated lower limits of the confidence intervals for S.
sd.boot The bootstrap standard deviation.
Boot.Repeat The number of resamples done in each bootstrap call to ensure the existence and uniqueness of the bootstrap NPMLE.

Author(s)

Carla Moreira, Jacobo de Uña-Álvarez and Rosa Crujeiras

References


See Also

efron.petrosian
# Examples

```r
# Generating data which are doubly truncated
n<-25
X<-runif(n,0,1)
U<-runif(n,0,0.25)
V<-runif(n,0.75,1)
for (i in 1:n){
  while (X[i]<U[i]|X[i]>V[i]){ 
    U[i]<-runif(1,0,0.25)
    X[i]<-runif(1,0,1)
    V[i]<-runif(1,0.75,1)
  }
}
res<-lynden(X=X, U=U, V=V, boot=FALSE, display.F=TRUE, display.S=TRUE)

# Generating data which are right truncated
n<-25
X<-runif(n,0,1)
V<-runif(n,0.75,1)
for (i in 1:n){
  while (X[i]>V[i]){ 
    X[i]<-runif(1,0,1)
    V[i]<-runif(1,0.75,1)
  }
}
res<-lynden(X=X, U=NA, V=V, boot=FALSE)
```

---

**Description**

The sample consists of DNA from 99 Caucasian Parkinson’s Disease (PD) patients with earlier onset PD (age 35-55 years). To remove the selection bias related to survival, the study was limited to patients diagnosed from PD who had their DNA sample taken within eight years after onset. Consequently, the age of onset is doubly truncated by the age at blood sampling and this time minus 8 years. This is a situation of interval sampling, the sampling interval being subject-specific.

**Usage**

```r
data("PDearly")
```
Format

A data frame with 99 observations on the following 5 variables.

- \( X \) a numeric vector, age at onset of PD (in years).
- \( U \) a numeric vector, age at blood sampling minus 8 years.
- \( V \) a numeric vector, age at blood sampling.
- \( \text{SNP}\_A10398G \) a factor with alleles levels A and G.
- \( \text{SNP}\_\text{PGC}1a \) a factor with alleles levels A, AG and G.

Details

Clark et al., 2011 hypothesized that the rs8192678 PGC-1a single nucleotide polymorphism (SNP) and the A10398G mitochondrial SNP may influence risk or age of onset of PD. To test these hypotheses, genomic DNA samples from human blood samples were obtained from the National Institute of Neurological Disorders and Stroke (NINDS) Human Genetics DNA and Cell Line Repository at the Coriell Institute for Medical Research (Camden, New Jersey).

Source


References


See Also

PDlate

Examples

data(PDearly)
str(PDearly)

---

PDlate Parkinson’s Disease Data: late onset

Description

The sample consists of DNA from 100 Caucasian Parkinson’s Disease (PD) patients with late onset PD (age 63-87 years). To remove the selection bias related to survival, the study was limited to patients diagnosed from PD who had their DNA sample taken within eight years after onset. Consequently, the age of onset is doubly truncated by the age at blood sampling and this time minus 8 years. This is a situation of interval sampling, the sampling interval being subject-specific.
Usage

data("PDlate")

Format

A data frame with 99 observations on the following 5 variables.

X  a numeric vector, age at onset of PD (in years).
U  a numeric vector, age at blood sampling minus 8 years.
V  a numeric vector, age at blood sampling.
SNP_A10398G  a factor with alleles levels A and G.
SNP_PGC1a  a factor with alleles levels A, AG and G.

Details

Clark et al., 2011 hypothesized that the rs8192678 PGC-1a single nucleotide polymorphism (SNP) and the A10398G mitochondrial SNP may influence risk or age of onset of PD. To test these hypotheses, genomic DNA samples from human blood samples were obtained from the National Institute of Neurological Disorders and Stroke (NINDS) Human Genetics DNA and Cell Line Repository at the Coriell Institute for Medical Research (Camden, New Jersey).

Source


References


See Also

PDearly

Examples

data(PDlate)
str(PDlate)
Quasars Data

Description

The original dataset studied by Efron and Petrosian (1999) comprised independently collected quadruplets of the redshift and the apparent magnitude of a quasar object. Due to experimental constraints, the distribution of each luminosity in a log-scale is truncated to a known interval.

Usage

data(Quasars)

Format

A data frame with 210 observations on the following 3 variables.

- y (adj lum) a numeric vector, the log luminosity values.
- u (lower) a numeric vector, lower truncation limits.
- v (upper) a numeric vector, upper truncation limits.

Details

Quadruplets in the original data set studied by Efron and Petrosian (1999) are of the form \((z_i; m_i; a_i; b_i)\), \(i = 1, \ldots, n\), where \(z_i\) is the redshift of the \(i\)th quasar and \(m_i\) is the apparent magnitude. Due to experimental constraints, the distribution of each luminosity in the log-scale \((y_i = t(z_i, m_i))\) is truncated to a known interval \([a_i; b_i]\), where \(t\) represents a transformation which depends on the cosmological model assumed (see Efron and Petrosian (1999) for details). Quasars with apparent magnitude above \(b_i\) were too dim to yield dependent redshifts, and hence they were excluded from the study. The lower limit \(a_i\) was used to avoid confusion with non quasar stellar objects.

Source

Vahé Petrosian and Bradley Efron.

References


Examples

data(Quasars)
str(Quasars)
rsim.DT

Random generation functions of doubly truncated data

Description

Random generation functions of doubly truncated data with two different patterns of observational bias.

Usage

rsim.DT(n, tau, model=NULL)

Arguments

- \( n \)  number of observations to generate.
- \( \tau \)  length of the observational window.
- \( \text{model} \)  model to be simulated. Number 1 or 2 corresponding to different patterns of observational bias.

Details

If \( \text{model}=1 \), \( U \sim \text{Unif}(-\tau, 1) \) and \( V=U+\tau \). If \( \text{model}=2 \), \( U \sim \text{Unif}(0, 1)^2 \times (\tau+1) - \tau \) and \( V=U+\tau \). In \( \text{model}=1 \) there is no observational bias due double truncation while in \( \text{model}=2 \) double truncation induces observational bias.

Value

A matrix with \( n \) unit length rows representing the generated values from a doubly truncated data with triplets \((X, U, and V)\), in which \((U \leq X \leq V)\).

Author(s)

Carla Moreira, Jacobo de Uña-Álvarez and Rosa Crujeiras

Examples

rsim.DT(500, 1/2, model=2)
Doubly truncated data analysis with the Shen algorithm

Description

This function computes the NPMLE of a lifetime distribution function observed under one-sided (right or left) and two-sided (double) truncation. The NPMLE of the joint distribution of the truncation times along with its marginal distributions are also computed. It provides bootstrap pointwise confidence limits too.

Usage

```r
shen(X, U = NA, V = NA, wt = NA, error = NA,
      nmaxit = NA, boot = TRUE, boot.type = "simple",
      B = NA, alpha = NA, display.FS = FALSE,
      display.UV = FALSE, plot.joint = FALSE, plot.type = NULL)
```

Arguments

- `X` Numeric vector with the times of ultimate interest.
- `U` Numeric vector with the left truncation times. If there are no truncation times from the left, put U=NA.
- `V` Numeric vector with the right truncation times. If there are no truncation times from the left, put V=NA.
- `wt` Numeric vector of non-negative initial solution, with the same length as X. Default value is set to 1/n, being n the length of X.
- `error` Numeric value. Maximum pointwise error when estimating the density associated to X (f) in two consecutive steps. If this is missing, it is $1e-06$.
- `nmaxit` Numeric value. Maximum number of iterations. If this is missing, it is set to nmaxit =100.
- `boot` Logical. If TRUE (default), the simple bootstrap method is applied to lifetime and truncation times distributions estimation. Pointwise confidence bands are provided.
- `boot.type` A character string giving the bootstrap type to be used. This must be one of "simple" or "obvious", with default "simple".
- `B` Numeric value. Number of bootstrap resamples. The default NA is equivalent to B =500.
- `alpha` Numeric value. (1-alpha) is the nominal coverage for the pointwise confidence intervals.
- `display.FS` Logical. Default is FALSE. If TRUE, the estimated cumulative distribution function and the estimated survival function associated to X, (F) and (S) respectively, are plotted.
- `display.UV` Logical. Default is FALSE. If TRUE, the marginal distributions of U (fU) and V (fV), are plotted.
plot.joint Logical. Default is FALSE. If TRUE, the joint distribution of the truncation times is plotted.

plot.type A character string giving the plot type to be used to represent the joint distribution of the truncation times. This must be one of "image" or "persp", with default NULL.

Details

The NPMLE of the lifetime is computed by a single algorithm proposed in Shen (2008). This is an alternative algorithm which converges to the NMPLE after a number of iterations. Initial solutions are given by the ordinary empirical distribution functions. If the second (respectively third) argument is missing, computation of the Lynden-Bell estimator for right-truncated (respectively left-truncated) data is obtained. Note that individuals with NAs in the three first arguments will be automatically excluded.

Value

A list containing the following values:

time The timepoint on the curve.
n.event The number of events that occurred at time t.
events The total number of events.
density The estimated density values associated to X.
cumulative.df The estimated cumulative distribution values of X.
truncation.probs The probabilities of truncation values, in each region.
S0 error reached in the algorithm.
Survival The estimated survival values.
density.joint The estimated joint densities values associated to (U,V).
marginal.U The estimated cumulative univariate marginal values of the U.
marginal.V The estimated cumulative univariate marginal values of the V.
cumulative.joint The estimated joint cumulative distribution values.
n.iterations The number of iterations used by this algorithm.
biasf The estimated probabilities of observing the lifetimes.
Boot The type of bootstrap method applied.
B Number of bootstrap resamples computed.
alpha The nominal level used to construct the confidence intervals.
upper.df The estimated upper limits of the confidence intervals for F.
lower.df The estimated lower limits of the confidence intervals for F.
upper.Sob The estimated upper limits of the confidence intervals for S.
lower.Sob The estimated lower limits of the confidence intervals for S.
upper.fU The estimated upper limits of the confidence intervals for fU.
The estimated lower limits of the confidence intervals for \( f_U \).

The estimated upper limits of the confidence intervals for \( f_V \).

The estimated lower limits of the confidence intervals for \( f_V \).

The bootstrap standard deviation.

The number of resamples done in each bootstrap call to ensure the existence and uniqueness of the bootstrap NPMLE.

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**References**


**See Also**

*lynden*

**Examples**

```r
## Generating data which are doubly truncated

n<-25
X<-runif(n,0,1)
U<-runif(n,0,0.67)
V<-runif(n,0.33,1)
for (i in 1:n){
  while (X[i]<U[i]|X[i]>V[i]){  
    U[i]<-runif(1,0,0.67)
    X[i]<-runif(1,0,1)
    V[i]<-runif(1,0.33,1)
  }
}
res<-shen(X,U,V,boot=FALSE, plot.joint=TRUE, plot.type="persp")
```
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