Package ‘lmboot’

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Imports evd (>= 2.3.0), stats (>= 3.6.0)
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R topics documented:

lmboot-package .................................................. 2
ANOVA.boot .......................................................... 4
bayesian.boot .......................................................... 6
jackknife ................................................................. 7
paired.boot .............................................................. 8
residual.boot ........................................................... 10
wild.boot ................................................................. 11
Description


Details

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Title: Bootstrap in Linear Models
Version: 0.0.1
Date: 2019-05-13
Authors@R: person("Megan", "Heyman", email="heyman@rose-hulman.edu", role=c("aut","cre"))
Description: Various efficient and robust bootstrap methods are implemented for linear models with least squares estimation.
Depends: R (>= 3.5.0)
Imports: evd (>= 2.3.0), stats (>= 3.6.0)
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Author: Megan Heyman [aut, cre]
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Index of help topics:

- ANOVA.boot: Residual and wild bootstrap in 1-way and 2-way ANOVA
- bayesian.boot: Bayesian Bootstrap in Linear Models
- jackknife: Delete-1 Jackknife in Linear Models
- lmboot-package: Bootstrap in Linear Models
- paired.boot: Paired Bootstrap in Linear Models
- residual.boot: Residual bootstrap in linear models
- wild.boot: Wild Bootstrap in Linear Models

This package is useful to users who wish to perform bootstrap in linear models. The package contains functions to create the sampling distributions for linear model parameters using either efficient or robust bootstrap methods.
As classified by Liu and Singh (1992), efficient bootstrap types include the residual bootstrap \( \text{residual.boot}(\)\). These types of bootstrap are useful when it is not reasonable to assume that errors come from a normal distribution, but you may make other classical assumptions: errors are independent, have mean 0, and have constant variance.

Robust bootstrap types include the paired bootstrap \( \text{paired.boot}(\)\), wild bootstrap \( \text{wild.boot}(\)\), and the jackknife \( \text{jackknife}(\)\). These types of bootstrap are useful when it is not reasonable to assume that errors have constant variance, but you may make other classical assumptions: errors are independent and have mean 0.

The package also contains a function for Bayesian bootstrap \( \text{bayesian.boot}(\)\) and a function to perform bootstrap in the ANOVA hypothesis test \( \text{ANOVA.boot}(\)\). The ANOVA bootstrap function has options to use the wild or residual bootstrap techniques and has been tested to work in 2-way ANOVA. Its functionality allows K-way ANOVA, however those capabilities have not been fully tested.

Currently, the user must manipulate the output of the function to conduct hypothesis tests and create confidence intervals for the predictor coefficients. More convenient/streamlined output is expected in future package versions.

**Author(s)**

NA

Maintainer: NA

**References**


**Examples**

```r
Seed <- 14
set.seed(Seed)
y <- rnorm(20) # randomly generated response
x <- rnorm(20) # randomly generated predictor

ResidObj <- residual.boot(y~x, B=100, seed=Seed) # perform the residual bootstrap
WildObj <- wild.boot(y~x, B=100, seed=Seed) # perform the wild bootstrap

# residual bootstrap 95% CI for slope parameter (percentile method)
quantile(ResidObj$bootEstParam[,2], probs=c(.025, .975))

# bootstrap 95% CI for slope parameter (percentile method)
quantile(WildObj$bootEstParam[,2], probs=c(.025, .975))
```
ANOVA.boot
Residual and wild bootstrap in 1-way and 2-way ANOVA

Description
This function performs the residual bootstrap as described by Efron (1979) and wild bootstrap as described by Wu (1986) for ANOVA hypothesis testing. Linear models incorporating categorical and/or quantitative predictor variables with a quantitative response are allowed. The function output creates the bootstrap null distribution for each term to be tested. Estimation is performed via least squares and only Type I sum of squares are calculated.

Usage
ANOVA.boot(formula, B = 1000, type = "residual", wild.dist = "normal", seed = NULL, data = NULL, keep.boot.resp = FALSE)

Arguments
formula
input a linear model formula of the form response~predictors as you would in the lm() function. All variables must contain non-missing entries.

B
number of bootstrap samples. This should be a large, positive integer value.

type
type of bootstrap to perform. Select either "residual" for residual bootstrap or "wild" for wild bootstrap.

wild.dist
distribution used to create the wild bootstrap weights for the residuals. Allowed distributions include "normal", "uniform", "exponential", "laplace", "lognormal", "gumbel", "t5", "t8", and "t14". The numbers after the t-distributions indicate the degrees of freedom. Any selected distribution creates weights with mean 0 and variance 1 from the named distribution.

seed
optionally, set a value for the seed for the bootstrap sample generation. The default NULL will pick a random value for the seed.

data
optionally, input the name of the dataset where variables appearing in the model are stored.

keep.boot.resp
a boolean indicating whether the list of returns includes raw bootstrap responses. Setting this to TRUE may not be possible for larger datasets or too many bootstrap samples due to memory usage.

Details
Currently, the user must manipulate the output of the function manually to view the bootstrap ANOVA table components and visualize the null distribution. More convenient/streamlined output is expected in future package versions.

Thanks to Bochuan Lyu who helped to coding to this function.
Value

terms  names of the terms/rows of the ANOVA table. These correspond to each predictor variable input to the formula.

df  degrees of freedom associated with each term/row in the ANOVA table. These correspond to the number of categories in each predictor variable (or are 1 for quantitative predictors)

origFStats  original F-statistic value. Same value as obtained by aov() using type I sum of squares.

origSSE  original sum of squares, error. Same value as obtained by aov() using type I sum of squares.

origSSTr  original sum of squares, treatment. Vector containing the sum of squares for each term in the ANOVA model. These are the same values as obtained by aov() using type I sum of squares.

bootFStats  matrix containing the bootstrap F statistics. Each column corresponds to a term in the ANOVA table. There are B rows.

bootSSE  matrix containing the bootstrap sum of squares, error. Each column corresponds to a term in the ANOVA table. There are B rows. These are calculated using type I sum of squares.

bootSSTr  matrix containing the bootstrap sum of squares, treatment. Each column corresponds to a term in the ANOVA table. There are B rows. These are calculated using type I sum of squares.

‘p-values‘  vector containing the bootstrap p-values for each predictor term in the ANOVA model. These are calculated by counting the number of bootstrap test statistics which are greater than the original observed test statistic and dividing by B

Author(s)

Megan Heyman, heyman@rose-hulman.edu

References


See Also

wild.boot, residual.boot

Examples

data(mtcars)  #load an example dataset
myANOVA2 <- ANOVA.boot(mpg~as.factor(cyl)*as.factor(am), data=mtcars)
myANOVA2$p-values'  #bootstrap p-values for 2-way interactions model
myANOVA1 <- ANOVA.boot(mpg~as.factor(cyl), data=mtcars)
myANOVA1$`p-values` #bootstrap p-values for 1-way model

myANOVA2a <- ANOVA.boot(mpg~as.factor(cyl)+as.factor(am), data=mtcars)
myANOVA2a$`p-values` #bootstrap p-values for 1-way additive model

**bayesian.boot**  
*Bayesian Bootstrap in Linear Models*

**Description**

This function performs the bayesian bootstrap in linear models as described by Rubin (1981) <doi:10.1214/aos/1176345338>. Linear models incorporating categorical and/or quantitative predictor variables with a quantitative response are allowed. The function output creates the bootstrap sampling distribution for each coefficient. Estimation is performed via least squares.

**Usage**

```r
bayesian.boot(formula, B = 1000, seed = NULL, data = NULL)
```

**Arguments**

- `formula` input a linear model formula of the form `response~predictors` as you would in the `lm()` function. All variables must contain non-missing entries.
- `B` number of bootstrap samples. This should be a large, positive integer value.
- `seed` optionally, set a value for the seed for the bootstrap sample generation. The default NULL will pick a random value for the seed.
- `data` optionally, input the name of the dataset where variables appearing in the model are stored.

**Details**

Currently, the user must manipulate the output of the function to conduct hypothesis tests and create confidence intervals for the predictor coefficients. More convenient/streamlined output is expected in future package versions.

**Value**

- `bootEstParam` matrix containing the bootstrap parameter estimates. Each column corresponds to a coefficient. There are B rows, each corresponding to a bootstrap sample.
- `origEstParam` vector containing the least squares parameter estimates. These are the same as estimates obtained from `lm`.
- `seed` numerical value set for the seed. This is associated with the set of bootstrap parameter estimates and helps the process to be reproducible.
Author(s)
Megan Heyman, heyman@rose-hulman.edu

References

Examples

```r
Seed <- 14
set.seed(Seed)
y <- rnorm(20) # randomly generated response
x <- rnorm(20) # randomly generated predictor
BayesObj <- bayesian.boot(y~x, B=100, seed=Seed) # perform the Bayesian bootstrap

# plot the sampling distribution of the slope coefficient
hist(BayesObj$bootEstParam[,2], main="Bayesian Bootstrap Sampling Distn.", xlab="Slope Estimate")

# bootstrap 95% CI for slope parameter (percentile method)
quantile(BayesObj$bootEstParam[,2], probs=c(.025, .975))
```

---

**jackknife**  
*Delete-1 Jackknife in Linear Models*

**Description**

This function performs the delete-1 jackknife in linear models as described by Quenouille (1956)  
<doi:10.2307/2332914>. Linear models incorporating categorical and/or quantitative predictor  
variables with a quantitative response are allowed. The function output creates the jackknife sam-  
ppling distribution for each coefficient. Estimation is performed via least squares.

**Usage**

```r
jackknife(formula, data = NULL)
```

**Arguments**

- `formula` input a linear model formula of the form `response~predictors` as you would in the `lm()` function. All variables must contain non-missing entries.
- `data` optionally, input the name of the dataset where variables appearing in the model are stored.

**Details**

Currently, the user must manipulate the output of the function to conduct hypothesis tests and create confidence intervals for the predictor coefficients. More convenient/streamlined output is expected in future package versions.
Value

bootEstParam matrix containing the jackknife parameter estimates. Each column corresponds to a coefficient. There are n-1 rows, each corresponding to a jackknife sample.

origEstParam vector containing the least squares parameter estimates. These are the same as estimates obtained from lm.

Author(s)

Megan Heyman, heyman@rose-hulman.edu

References


Examples

Seed <- 14
set.seed(Seed)
y <- rnorm(20) # randomly generated response
x <- rnorm(20) # randomly generated predictor
JackObj <- jackknife(y~x) # perform the jackknife

# plot the sampling distribution of the slope coefficient
hist(JackObj$bootEstParam[,2], main="Jackknife Sampling Distn.",
     xlab="Slope Estimate")

# jackknife 95% CI for slope parameter (percentile method)
quantile(JackObj$bootEstParam[,2], probs=c(.025, .975))

Description

This function performs the paired bootstrap in linear models as described by Efron (1979, ISBN: 978-1-4612-4380-9). Linear models incorporating categorical and/or quantitative predictor variables with a quantitative response are allowed. The function output creates the bootstrap sampling distribution for each coefficient. Estimation is performed via least squares.

Usage

paired.boot(formula, B = 1000, seed = NULL, data = NULL)
Arguments

formula: input a linear model formula of the form response~predictors as you would in the lm() function. All variables must contain non-missing entries.

B: number of bootstrap samples. This should be a large, positive integer value.

seed: optionally, set a value for the seed for the bootstrap sample generation. The default NULL will pick a random value for the seed.

data: optionally, input the name of the dataset where variables appearing in the model are stored.

Details

Currently, the user must manipulate the output of the function to conduct hypothesis tests and create confidence intervals for the predictor coefficients. More convenient/streamlined output is expected in future package versions.

Value

bootEstParam: matrix containing the bootstrap parameter estimates. Each column corresponds to a coefficient. There are B rows, each corresponding to a bootstrap sample.

origEstParam: vector containing the least squares parameter estimates. These are the same as estimates obtained from lm.

seed: numerical value set for the seed. This is associated with the set of bootstrap parameter estimates and helps the process to be reproducible.

Author(s)

Megan Heyman, heyman@rose-hulman.edu

References


Examples

Seed <- 14
set.seed(Seed)
y <- rnorm(20) # randomly generated response
x <- rnorm(20) # randomly generated predictor
PairObj <- paired.boot(y=x, B=100, seed=Seed) # perform the paired bootstrap

# plot the sampling distribution of the slope coefficient
hist(PairObj$bootEstParam[,2], main="Paired Bootstrap Sampling Distr.",
     xlab="Slope Estimate")

# bootstrap 95% CI for slope parameter (percentile method)
quantile(PairObj$bootEstParam[,2], probs=c(.025, .975))
residual.boot  Residual bootstrap in linear models

Description
This function performs the residual bootstrap in linear models as described by Efron (1979, ISBN:978-1-4612-4380-9). Linear models incorporating categorical and/or quantitative predictor variables with a quantitative response are allowed. The function output creates the bootstrap sampling distribution for each coefficient. Estimation is performed via least squares.

Usage
residual.boot(formula, B = 1000, data = NULL, seed = NULL)

Arguments
- formula: input a linear model formula of the form response~predictors as you would in the `lm()` function. All variables must contain non-missing entries.
- B: number of bootstrap samples. This should be a large, positive integer value.
- data: optionally, input the name of the dataset where variables appearing in the model are stored.
- seed: optionally, set a value for the seed for the bootstrap sample generation. The default NULL will pick a random value for the seed.

Details
Currently, the user must manipulate the output of the function to conduct hypothesis tests and create confidence intervals for the predictor coefficients. More convenient/streamlined output is expected in future package versions.

Value
- bootEstParam: matrix containing the bootstrap parameter estimates. Each column corresponds to a coefficient. There are B rows, each corresponding to a bootstrap sample.
- origEstParam: vector containing the least squares parameter estimates. These are the same as estimates obtained from `lm`.
- seed: numerical value set for the seed. This is associated with the set of bootstrap parameter estimates and helps the process to be reproducible.

Author(s)
Megan Heyman, heyman@rose-hulman.edu

References
Examples
Seed <- 14
set.seed(Seed)
y <- rnorm(20) # randomly generated response
x <- rnorm(20) # randomly generated predictor
ResidObj <- residual.boot(y-x, B=100, seed=Seed) # perform the residual bootstrap

# plot the sampling distribution of the slope coefficient
hist(ResidObj$bootEstParam[,2], main="Residual Bootstrap Sampling Distn.", xlab="Slope Estimate")

# bootstrap 95% CI for slope parameter (percentile method)
quantile(ResidObj$bootEstParam[,2], probs=c(.025, .975))

Description
This function performs the wild/external bootstrap in linear models as described by Wu (1986) <doi:10.1214/aos/1176350142>. Linear models incorporating categorical and/or quantitative predictor variables with a quantitative response are allowed. The function output creates the bootstrap sampling distribution for each coefficient. Estimation is performed via least squares.

Usage
wild.boot(formula, B = 1000, data = NULL, seed = NULL, bootDistn = "normal")

Arguments
formula input a linear model formula of the form response~predictors as you would in the `lm()` function. All variables must contain non-missing entries.
B number of bootstrap samples. This should be a large, positive integer value.
data optionally, input the name of the dataset where variables appearing in the model are stored.
seed optionally, set a value for the seed for the bootstrap sample generation. The default NULL will pick a random value for the seed.
bootDistn distribution used to create the wild bootstrap weights for the residuals. Allowed distributions include "normal", "uniform", "exponential", "laplace", "lognormal", "gumbel", "t5", "t8", and "t14". The numbers after the t-distributions indicate the degrees of freedom. Any selected distribution creates weights with mean 0 and variance 1 from the named distribution.

Details
Currently, the user must manipulate the output of the function to conduct hypothesis tests and create confidence intervals for the predictor coefficients. More convenient/streamlined output is expected in future package versions.
Value

- `bootEstParam` - matrix containing the bootstrap parameter estimates. Each column corresponds to a coefficient. There are `B` rows, each corresponding to a bootstrap sample.
- `origEstParam` - vector containing the least squares parameter estimates. These are the same as estimates obtained from `lm`.
- `seed` - numerical value set for the seed. This is associated with the set of bootstrap parameter estimates and helps the process to be reproducible.
- `bootDistn` - type of distribution used to generate the wild bootstrap weights for the residuals

Author(s)

Megan Heyman, heyman@rose-hulman.edu

References


Examples

```r
Seed <- 14
set.seed(Seed)
y <- rnorm(20) # randomly generated response
x <- rnorm(20) # randomly generated predictor
WildObj <- wild.boot(y~x, B=100, seed=Seed) # perform the wild bootstrap

# plot the sampling distribution of the slope coefficient
hist(WildObj$bootEstParam[,2], main="Wild Bootstrap Sampling Distn.", xlab="Slope Estimate")

# bootstrap 95% CI for slope parameter (percentile method)
quantile(WildObj$bootEstParam[,2], probs=c(.025, .975))
```
Index

*Topic distribution
  bayesian.boot, 6
  wild.boot, 11
*Topic models regression
  ANOVA.boot, 4
  bayesian.boot, 6
  jackknife, 7
  paired.boot, 8
  residual.boot, 10
  wild.boot, 11
*Topic nonparametric
  ANOVA.boot, 4
  bayesian.boot, 6
  jackknife, 7
  paired.boot, 8
  residual.boot, 10
  wild.boot, 11
*Topic package
  lmboot-package, 2

ANOVA.boot, 4
bayesian.boot, 6
jackknife, 7
lmboot (lmboot-package), 2
lmboot-package, 2
paired.boot, 8
residual.boot, 5, 10
wild.boot, 5, 11