Package 'gamlss.tr'

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Depends R ($>= 2.2.1$), gamlss.dist, gamlss ($>= 5.0-0$), methods
Description This is an add on package to GAMLSS. The purpose of this package is to allow users to defined truncated distributions in GAMLSS models. The main function gen.trun() generates truncated version of an existing GAMLSS family distribution.
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gamlss.tr-package fitTail gen.trun trun trun.d trun.p trun.q trun.r
Index 23

2 gamlss.tr-package

gamlss.tr-package Generating and Fitting Truncated 'gamlss.family' Distributions

Description

This is an add on package to GAMLSS. The purpose of this package is to allow users to defined truncated distributions in GAMLSS models. The main function gen.trun() generates truncated version of an existing GAMLSS family distribution.

Details

The DESCRIPTION file:

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Author: Mikis Stasinopoulos <d.stasinopoulos@gre.ac.uk>, Bob Rigby <r.rigby@gre.ac.uk>

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URL: https://www.gamlss.com/

Index of help topics:

fitTail For fitting truncated distribution to the tails

of data

gamlss.tr-package Generating and Fitting Truncated

'gamlss.family' Distributions

gen.trun Generates a truncated distribution from a

gamlss.family

trun Fits a Truncate Distribution from a

gamlss.family

trun.d Truncated Probability Density Function of a

gamlss.family Distribution

trun.p Truncated Cumulative Density Function of a

gamlss.family Distribution

trun.q Truncated Inverse Cumulative Density Function

of a gamlss.family Distribution

trun.r Generates Random Values from a Truncated

Density Function of a gamlss.family

Distribution

fitTail 3

Author(s)

Mikis Stasinopoulos <d.stasinopoulos@gre.ac.uk>, Bob Rigby <r.rigby@gre.ac.uk>

Maintainer: Mikis Stasinopoulos <d.stasinopoulos@gre.ac.uk>

References

Rigby, R. A. and Stasinopoulos D. M. (2005). Generalized additive models for location, scale and shape, (with discussion), *Appl. Statist.*, **54**, part 3, pp 507-554.

Rigby, R. A., Stasinopoulos, D. M., Heller, G. Z., and De Bastiani, F. (2019) *Distributions for modeling location, scale, and shape: Using GAMLSS in R*, Chapman and Hall/CRC. An older version can be found in https://www.gamlss.com/.

Stasinopoulos D. M. Rigby R.A. (2007) Generalized additive models for location scale and shape (GAMLSS) in R. *Journal of Statistical Software*, Vol. **23**, Issue 7, Dec 2007, https://www.jstatsoft.org/v23/i07/.

Stasinopoulos D. M., Rigby R.A., Heller G., Voudouris V., and De Bastiani F., (2017) *Flexible Regression and Smoothing: Using GAMLSS in R*, Chapman and Hall/CRC.

```
(see also https://www.gamlss.com/).
```

Examples

```
# generating a t-distribution from 0 to 100
gen.trun(par=c(0,100),family="TF", name="0to100", type="both")
op<-par(mfrow=c(2,2))
plot(function(x) dTF0to100(x, mu=80 ,sigma=20, nu=5), 0, 100, ylab="pdf")
plot(function(x) pTF0to100(x, mu=80 ,sigma=20, nu=5), 0, 100, ylab="cdf")
plot(function(x) qTF0to100(x, mu=80 ,sigma=20, nu=5), 0.01, .999, ylab="invcdf")
hist(s1<-rTF0to100(1000, mu=80 ,sigma=20, nu=5), ylab="hist", xlab="x", main="generated data")
par(op)</pre>
```

fitTail

For fitting truncated distribution to the tails of data

Description

There are two functions here. The function fitTail() which fits a truncated distribution to certain percentage of the tail of a response variable and the function fitTailAll() which does a sequence of truncated fits. Plotting the results from those fits is analogous to the Hill plot, Hill (1975).

Usage

4 fitTail

Arguments

The variable of interest family a gamlsss. family distribution what percentage of the tail need to be modelled, default is 10% percentage how many observations in the tail needed. This is an alternative to percentage. howmany If it specified it take over from the percentage argument otherwise percentage is used. which tall needs checking the right (default) of the left type plot whether to plot with default equal TRUE whether to print the coefficients with default equal TRUE print whether to save the fitted linear model with default equal FALSE save start where to start fitting from the tail of the data 0: no output 1: minimal 2: print estimates trace

Details

. . .

The idea here is to fit a truncated distribution to the tail of the data. Truncated log-normal and Weibull distributions could be appropriate distributions. More details can be found in Chapter 6 of "The Distribution Toolbox of GAMLSS" book which can be found in https://www.gamlss.com/).

for further argument to the fitting function

Value

A fitted gamlss model

Author(s)

Bob Rigby <r.rigby@gre.ac.uk>, Mikis Stasinopoulos <d.stasinopoulos@gre.ac.uk> and Vlassios Voudouris

References

Hill B. M. (1975) A Simple General Approach to Inference About the Tail of a Distribution *Ann. Statist.* Volume 3, Number 5, pp 1163-1174.

Rigby, R. A. and Stasinopoulos D. M. (2005). Generalized additive models for location, scale and shape, (with discussion), *Appl. Statist.*, **54**, part 3, pp 507-554.

Rigby, R. A., Stasinopoulos, D. M., Heller, G. Z., and De Bastiani, F. (2019) *Distributions for modeling location, scale, and shape: Using GAMLSS in R*, Chapman and Hall/CRC. An older version can be found in https://www.gamlss.com/.

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Stasinopoulos D. M., Rigby R.A., Heller G., Voudouris V., and De Bastiani F., (2017) *Flexible Regression and Smoothing: Using GAMLSS in R*, Chapman and Hall/CRC.

(see also https://www.gamlss.com/).

gen.trun 5

See Also

```
loglogSurv, logSurv
```

Examples

```
data(film90)
F90 <- exp(film90$lborev1)# original scale
# trucated plots
# 10%
w403<- fitTail(F90, family=WEI3)
qqnorm(resid(w403))
abline(0,1, col="red")
## Not run:
# hill -sequential plot 10
w1<-fitTailAll(F90)
# plot sigma
plot(w1[,2])
#LOGNO
1403<- fitTail(F90, family=LOGNO)
plot(1403)
qqnorm(resid(1403))
abline(0,1)
# hill -sequential plot 10
11<-fitTailAll(F90, family=LOGNO)</pre>
plot(11[,2])
## End(Not run)
```

gen.trun

Generates a truncated distribution from a gamlss.family

Description

The gen.trun() function allows the user to generate d, p, q, and r distribution functions plus an extra gamlss.family function for fitting a truncated distribution with gamlss.

For continuous distributions left truncation at 3 means that the random variable can take the value 3. For discrete distributions left truncation at 3 means that the random variable can take values from 4 onwards. This is the same for right truncation. Truncation at 15 for a discrete variable means that 15 and greater values are not allowed but for continuous variable it mean values greater that 15 are not allowed (so 15 is a possible value).

If the user want a different link (rather the default) for any of the parameters she/he has to declare at the generation of the functions, see example.

gen.trun

Usage

Arguments

par	a vector with one (for "left" or "right" truncation) or two elements for "both". When the argument varying = TRUE then par can be a vector or a matrix with two columns respectively.
family	a gamlss.family object, which is used to define the distribution and the link functions of the various parameters. The distribution families supported by gamlss() can be found in gamlss.family.
name	the extra characters to be added to the name of new truncated distribution, by default it adds tr
type	whether "left", "right" or in "both" sides truncation is required
varying	whether the truncation varies for different observations. This can be useful in regression analysis. If varying = TRUE then par should be an n-length vector for type equal "left" and "right" and an n by 2 matrix for type="both"
print	whether to print the names of the created distribution
	for extra arguments

Value

Returns the d, the p, the q, the r and the fitting functions of a truncated gamlss.family distribution.

Author(s)

Mikis Stasinopoulos <d. stasinopoulos@gre.ac.uk> and Bob Rigby

References

Rigby, R. A. and Stasinopoulos D. M. (2005). Generalized additive models for location, scale and shape, (with discussion), *Appl. Statist.*, **54**, part 3, pp 507-554.

Rigby, R. A., Stasinopoulos, D. M., Heller, G. Z., and De Bastiani, F. (2019) *Distributions for modeling location, scale, and shape: Using GAMLSS in R*, Chapman and Hall/CRC. An older version can be found in https://www.gamlss.com/.

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Stasinopoulos D. M., Rigby R.A., Heller G., Voudouris V., and De Bastiani F., (2017) *Flexible Regression and Smoothing: Using GAMLSS in R*, Chapman and Hall/CRC.

```
(see also https://www.gamlss.com/).
```

See Also

```
trun.d, trun.p, trun.q, trun.r
```

trun 7

Examples

```
# generating a t-distribution from 0 to 100
gen.trun(par=c(0,100),family="TF", name="0to100", type="both")
op<-par(mfrow=c(2,2))</pre>
plot(function(x) dTF0to100(x, mu=80 ,sigma=20, nu=5), 0, 100, ylab="pdf")
plot(function(x) pTF0to100(x, mu=80 ,sigma=20, nu=5), 0, 100, ylab="cdf")
plot(function(x) qTF0to100(x, mu=80 ,sigma=20, nu=5), 0.01, .999, ylab="invcdf")
hist(s1<-rTF0to100(1000, mu=80 ,sigma=20, nu=5), ylab="hist", xlab="x",
            main="generated data")
par(op)
m1<-histDist(s1, family=TF0to100, xlim=c(0,100))# fitting the data
# using the argumnt varying
# left part varies right part equal 100
leftPAR <- rPO(100)
gen.trun(par=cbind(leftPAR,rep(100, 100)),family="TF", name="0to100Varying",
            type="both", varying=TRUE)
YY<- rTF0to100Varying(100, mu=80, sigma=20, nu=5)
m1<-gamlss(YY~1, family=TF0to100Varying)
m1
```

trun

Fits a Truncate Distribution from a gamlss.family

Description

This function can be used to fit truncated distributions. It takes as an argument an existing GAMLSS family distribution and a parameter vector, of the type c(left.value, right.value), and generates a gamlss.family object which then can be used to fit a truncated distribution.

Usage

Arguments

par	a vector with one (for "left" or "right" truncation) or two elements for "both". When the argument varying = TRUE then par can be a vector or a matrix with two columns respectively.
family	an existing gamlss.family distribution
type	what type of truncation is required, left, right or both. If both the par should be a vector of length two. (the default is left truncation)
name	a character string to be added to name of the created object i.e. with family=TF and name=trZero the gamlss.family object will be called TFtrZero
local	if TRUE the function will try to find the environment of gamlss to generate the d and p functions required for the fitting, if FALSE the functions will be generated in the global environment

8 trun

delta the delta increment used in the numerical derivatives

varying whether the truncation varies for different observations. This can be usefull in

regression analysis. If varying = TRUE then par should be an n-length vector for type equal "left" and "right" and an n by 2 matrix for type="both"

... for extra arguments

Details

This function is created to help the user to fit a truncated form of existing gamlss distribution. It does this by taking an existing gamlss.family and changing some of the components of the distribution to help the fitting process. It particular it i) creates a pdf (d) and a cdf (p) function within gamlss, ii) changes the global deviance function G.dev.incr, the first derivative functions (see note below) and the quantile residual function.

Value

It returns a gamlss.family object which has all the components needed for fitting a distribution in gamlss.

Note

This function is experimental and could be changed. The function trun changes the first derivatives of the original gamlss family d function to numerical derivatives for the new truncated d function. The default increment delta, for this numerical derivatives function, is eps * pmax(abs(x), 1) where eps<-sqrt(.Machine\$double.eps). The default delta could be inappropriate for specific applications and can be overwritten by using the argument delta.

Author(s)

Mikis Stasinopoulos <d. stasinopoulos@gre.ac.uk> and Bob Rigby

References

Rigby, R. A. and Stasinopoulos D. M. (2005). Generalized additive models for location, scale and shape, (with discussion), *Appl. Statist.*, **54**, part 3, pp 507-554.

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```
(see also https://www.gamlss.com/).
```

See Also

```
trun.d, trun.p, trun.q, trun.r, gen.trun
```

Examples

```
# generate a left truncated zero t family
gen.trun(0,family="TF")
# take a random sample of 1000 observations
sam<-rTFtr(1000,mu=10,sigma=5, nu=5)
hist(sam)
# fit the distribution to the data
mod1<-gamlss(sam~1, family=trun(0,TF))
mod1
# now create a gamlss.family object before the fitting
Ttruc.Zero<- trun(par=0,family=TF, local=FALSE)
mod2<-gamlss(sam~1, family=Ttruc.Zero)
# now check the sensitivity of delta
Ttruc.Zero<- trun(par=0,family=TF, local=FALSE, delta=c(0.01,0.01, 0.01))
mod3<-gamlss(sam~1, family=Ttruc.Zero)</pre>
```

trun.d

Truncated Probability Density Function of a gamlss.family Distribution

Description

Creates a truncated probability density function version from a current GAMLSS family distribution For continuous distributions left truncation at 3 means that the random variable can take the value 3. For discrete distributions left truncation at 3 means that the random variable can take values from 4 onwards. This is the same for right truncation. Truncation at 15 for a discrete variable means that 15 and greater values are not allowed but for continuous variable it mean values greater that 15 are not allowed (so 15 is a possible value).

Usage

Arguments

par	a vector with one (for "left" or "right" truncation) or two elements for "both". When the argument varying = TRUE then par can be a vector or a matrix with two columns respectively.
family	a gamlss.family object, which is used to define the distribution and the link functions of the various parameters. The distribution families supported by gamlss() can be found in gamlss.family. Functions such as BI() (binomial) produce a family object.
type	whether left, right or in both sides truncation is required, (left is the default).
varying	whether the truncation varies for different observations. This can be usefull in regression analysis. If varying = TRUE then par should be an n-length vector for type equal "left" and "right" and an n by 2 matrix for type="both"
	for extra arguments

Value

Returns a d family function

Author(s)

Mikis Stasinopoulos <d. stasinopoulos@gre.ac.uk> and Bob Rigby

References

Rigby, R. A. and Stasinopoulos D. M. (2005). Generalized additive models for location, scale and shape, (with discussion), *Appl. Statist.*, **54**, part 3, pp 507-554.

Rigby, R. A., Stasinopoulos, D. M., Heller, G. Z., and De Bastiani, F. (2019) *Distributions for modeling location, scale, and shape: Using GAMLSS in R*, Chapman and Hall/CRC. An older version can be found in https://www.gamlss.com/.

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Stasinopoulos D. M., Rigby R.A., Heller G., Voudouris V., and De Bastiani F., (2017) *Flexible Regression and Smoothing: Using GAMLSS in R*, Chapman and Hall/CRC.

```
(see also https://www.gamlss.com/).
```

See Also

```
trun.p, trun.q, trun.r, gen.trun
```

Examples

```
#------
# continuous distribution
# left truncation
test1<-trun.d(par=c(0), family="TF", type="left")
test1(1)
dTF(1)/(1-pTF(0))
if(abs(test1(1)-(dTF(1)/pTF(0)))>0.00001) stop("error in left trucation")
test1(1, log=TRUE)
log(dTF(1)/(1-pTF(0)))
if(abs(test1(1, log=TRUE)-log(dTF(1)/pTF(0)))>0.00001)
                 stop("error in left trucation")
integrate(function(x) test1(x, mu=-2, sigma=1, nu=1),0,Inf)
# the pdf is defined even with negative mu
integrate(function(x) test1(x, mu=0, sigma=10, nu=1),0,Inf)
integrate(function(x) test1(x, mu=5, sigma=5, nu=10),0,Inf)
plot(function(x) test1(x, mu=-3, sigma=1, nu=1), 0, 10)
plot(function(x) test1(x, mu=3, sigma=5, nu=10),0,10)
# right truncation
test2<-trun.d(par=c(10), family="BCT", type="right")
test2(1)
dBCT(1)/(pBCT(10))
```

```
#if(abs(test2(1)-(dBCT(1)/pBCT(10)))>0.00001) stop("error in right trucation")
test2(1, log=TRUE)
log(dBCT(1)/(pBCT(10)))
if(abs(test2(1, log=TRUE)-log(dBCT(1)/(pBCT(10))))>0.00001)
                   stop("error in right trucation")
integrate(function(x) test2(x, mu=2, sigma=1, nu=1),0,10)
integrate(function(x) test2(x, mu=2, sigma=.1, nu=1),0,10)
integrate(function(x) test2(x, mu=2, sigma=.1, nu=10),0,10)
plot(function(x) test2(x, mu=2, sigma=.1, nu=1),0,10)
plot(function(x) test2(x, mu=2, sigma=1, nu=1),0,10)
# both left and right truncation
test3<-trun.d(par=c(-3,3), family="TF", type="both")</pre>
test3(0)
dTF(0)/(pTF(3)-pTF(-3))
if(abs(test3(0)-dTF(0)/(pTF(3)-pTF(-3)))>0.00001)
              stop("error in right trucation")
test3(0, log=TRUE)
log(dTF(0)/(pTF(3)-pTF(-3)))
if(abs(test3(0, log=TRUE)-log(dTF(0)/(pTF(3)-pTF(-3))))>0.00001)
            stop("error in both trucation")
plot(function(x) test3(x, mu=0, sigma=1, nu=1),-3,3)
integrate(function(x) test3(x, mu=2, sigma=1, nu=1),-3,3)
# discrete distribution
# left
# Poisson truncated at zero means zero is excluded
test4<-trun.d(par=c(0), family="PO", type="left")</pre>
test4(1)
dPO(1)/(1-pPO(0))
if(abs(test4(1)-dPO(1)/(1-pPO(0)))>0.00001) stop("error in left trucation")
test4(1, log=TRUE)
log(dPO(1)/(1-pPO(0)))
if(abs(test4(1, log=TRUE)-log(dPO(1)/(1-pPO(0))))>0.00001)
               stop("error in left trucation")
sum(test4(x=1:20, mu=2)) #
sum(test4(x=1:200, mu=80)) #
plot(function(x) test4(x, mu=20), from=1, to=51, n=50+1, type="h") # pdf
# right truncated at 10 means 10 is excluded
test5<-trun.d(par=c(10), family="NBI", type="right")</pre>
test5(2)
dNBI(2)/(pNBI(9))
if(abs(test5(1)-dNBI(1)/(pNBI(9)))>0.00001) stop("error in right trucation")
test5(1, log=TRUE)
log(dNBI(1)/(pNBI(9)))
if(abs(test5(1, log=TRUE)-log(dNBI(1)/(pNBI(9))))>0.00001) stop("error in right trucation")
sum(test5(x=0:9, mu=2, sigma=2)) #
sum(test5(x=0:9, mu=300, sigma=5)) # can have mu > parameter
plot(function(x) test5(x, mu=20, sigma=3), from=0, to=9, n=10, type="h") # pdf
plot(function(x) test5(x, mu=300, sigma=5), from=0, to=9, n=10, type="h") # pdf
# both
```

```
test6<-trun.d(par=c(0,10), family="NBI", type="both")
test6(2)
dNBI(2)/(pNBI(9)-pNBI(0))
if(abs(test6(2)-dNBI(2)/(pNBI(9)-pNBI(0)))>0.00001)
       stop("error in right trucation")
test6(1, log=TRUE)
log(dNBI(1)/(pNBI(9)-pNBI(0)))
if(abs(test6(1, log=TRUE)-log(dNBI(1)/(pNBI(9)-pNBI(0))))>0.00001)
 stop("error in right trucation")
sum(test6(x=1:9, mu=2, sigma=2)) #
sum(test6(x=1:9, mu=100, sigma=5)) # can have mu > parameter
plot(function(x) test6(x, mu=20, sigma=3), from=1, to=9, n=9, type="h") # pdf
plot(function(x) test6(x, mu=300, sigma=.4), from=1, to=9, n=9, type="h") # pdf
# now try when the trucated points varies for each observarion
# this will be appropriate for regression models only
# continuous
# left truncation
test7<-trun.d(par=c(0,1,2), family="TF", type="left", varying=TRUE)
test7(c(1,2,3))
dTF(c(1,2,3))/(1-pTF(c(0,1,2)))
test7(c(1,2,3), log=TRUE)
# right truncation
test8<-trun.d(par=c(10,11,12), family="BCT", type="right", varying=TRUE)
test8(c(1,2,3))
dBCT(c(1,2,3))/(pBCT(c(10,11,12)))
test8(c(1,2,3), log=TRUE)
# both left and right truncation
test9<-trun.d(par=cbind(c(0,1,2),c(10,11,12)), family="TF", type="both",
             varying=TRUE)
test9(c(1,2,3))
dTF(c(1,2,3))/(pTF(c(10,11,12))-pTF(c(0,1,2)))
test3(c(1,2,3), log=TRUE)
# discrete
# left
test10<-trun.d(par=c(0,1,2), family="PO", type="left", varying=TRUE)
test10(c(1,2,3))
dPO(c(1,2,3))/(1-pPO(c(0,1,2)))
# right
test11<-trun.d(par=c(10,11,12), family="NBI", type="right", varying=TRUE)
test11(c(1,2,3))
dNBI(c(1,2,3))/pNBI(c(9,10,11))
test12<-trun.d(par=rbind(c(0,10), c(1,11), c(2,12)), family="NBI", type="both", varying=TRUE)
test12(c(2,3,4))
dNBI(c(2,3,4))/(pNBI(c(9,10,11))-pNBI(c(0,1,2)))
```

trun.p

trun.p	Truncated Cumulative Density Function of a gamlss.family Distribution

Description

Creates a truncated cumulative density function version from a current GAMLSS family distribution.

For continuous distributions left truncation at 3 means that the random variable can take the value 3. For discrete distributions left truncation at 3 means that the random variable can take values from 4 onwards. This is the same for right truncation. Truncation at 15 for a discrete variable means that 15 and greater values are not allowed but for continuous variable it mean values greater that 15 are not allowed (so 15 is a possible value).

Usage

Arguments

par	a vector with one (for "left" or "right" truncation) or two elements for "both". When the argument varying = TRUE then par can be a vector or a matrix with two columns respectively.
family	a gamlss.family object, which is used to define the distribution and the link functions of the various parameters. The distribution families supported by gamlss() can be found in gamlss.family. Functions such as BI() (binomial) produce a family object.
type	whether left, right or in both sides truncation is required, (left is the default)
varying	whether the truncation varies for different observations. This can be usefull in regression analysis. If varying = TRUE then par should be an n-length vector for type equal "left" and "right" and an n by 2 matrix for type="both"
	for extra arguments

Value

Return a p family function

Author(s)

Mikis Stasinopoulos <d.stasinopoulos@gre.ac.uk> and Bob Rigby

14 trun.p

References

Rigby, R. A. and Stasinopoulos D. M. (2005). Generalized additive models for location, scale and shape, (with discussion), *Appl. Statist.*, **54**, part 3, pp 507-554.

Rigby, R. A., Stasinopoulos, D. M., Heller, G. Z., and De Bastiani, F. (2019) *Distributions for modeling location, scale, and shape: Using GAMLSS in R*, Chapman and Hall/CRC. An older version can be found in https://www.gamlss.com/.

Stasinopoulos D. M. Rigby R.A. (2007) Generalized additive models for location scale and shape (GAMLSS) in R. *Journal of Statistical Software*, Vol. **23**, Issue 7, Dec 2007, https://www.jstatsoft.org/v23/i07/.

Stasinopoulos D. M., Rigby R.A., Heller G., Voudouris V., and De Bastiani F., (2017) *Flexible Regression and Smoothing: Using GAMLSS in R*, Chapman and Hall/CRC.

```
(see also https://www.gamlss.com/).
```

See Also

```
trun.d, trun.q, trun.r, gen.trun
```

Examples

```
# trucated p continuous function
# continuous
# left
test1<-trun.p(par=c(0), family="TF", type="left")
test1(1)
(pTF(1)-pTF(0))/(1-pTF(0))
if(abs(test1(1)-(pTF(1)-pTF(0))/(1-pTF(0)))>0.00001)
                    stop("error in left trucation of p")
plot(function(x) test1(x, mu=2, sigma=1, nu=2),0,10)
#-----
# right
test2 <- trun.p(par=c(10), family="BCT", type="right")</pre>
test2(1)
pBCT(1)/pBCT(10)
if(abs(test2(1)-pBCT(1)/pBCT(10))>0.00001) stop("error in right trucation")
test2(1, lower.tail=FALSE)
1-pBCT(1)/pBCT(10)
if(abs(test2(1, lower.tail=FALSE)-(1-pBCT(1)/pBCT(10)))>0.00001)
               stop("error in right trucation")
test2(1, log.p=TRUE)
log(pBCT(1)/pBCT(10))
if(abs(test2(1, log.p=TRUE)-log(pBCT(1)/pBCT(10)))>0.00001)
              stop("error in right trucation")
plot(function(x) test2(x, mu=2, sigma=1, nu=2, tau=2), 0, 10)
plot(function(x) test2(x, mu=2, sigma=1, nu=2, tau=2,
                     lower.tail=FALSE),0,10)
# both
test3<-trun.p(par=c(-3,3), family="TF", type="both")
test3(1)
```

trun.p

```
(pTF(1)-pTF(-3))/(pTF(3)-pTF(-3))
if(abs(test3(1)-(pTF(1)-pTF(-3))/(pTF(3)-pTF(-3)))>0.00001)
             stop("error in right trucation")
test3(1, lower.tail=FALSE)
1-(pTF(1)-pTF(-3))/(pTF(3)-pTF(-3))
if(abs(test3(0,lower.tail=FALSE)-
            (1-(pTF(0)-pTF(-3))/(pTF(3)-pTF(-3)))>0.00001)
      stop("error in right trucation")
plot(function(x) test3(x, mu=2, sigma=1, nu=2, ),-3,3)
plot(function(x) test3(x, mu=2, sigma=1, nu=2, lower.tail=FALSE),-3,3)
# Discrete
# trucated p function
# left
test4<-trun.p(par=c(0), family="PO", type="left")</pre>
test4(1)
(pPO(1)-pPO(0))/(1-pPO(0))
if(abs(test4(1)-(pPO(1)-pPO(0))/(1-pPO(0)))>0.00001)
               stop("error in left trucation of p")
plot(function(x) test4(x, mu=2), from=1, to=10, n=10, type="h")
cdf <- stepfun(1:40, test4(1:41, mu=5), f = 0)
plot(cdf, main="cdf", ylab="cdf(x)", do.points=FALSE )
# right
test5<-trun.p(par=c(10), family="NBI", type="right")</pre>
test5(2)
pNBI(2)/(pNBI(9))
if(abs(test5(2)-(pNBI(2)/(pNBI(9))))>0.00001)
          stop("error in right trucation of p")
plot(function(x) test5(x, mu=2), from=0, to=9, n=10, type="h")
cdf \leftarrow stepfun(0:8, test5(0:9, mu=5), f = 0)
plot(cdf, main="cdf", ylab="cdf(x)", do.points=FALSE )
# both
test6<-trun.p(par=c(0,10), family="NBI", type="both")</pre>
(pNBI(2)-pNBI(0))/(pNBI(9)-pNBI(0))
if(abs(test6(2)-(pNBI(2)-pNBI(0))/(pNBI(9)-pNBI(0)))>0.00001)
               stop("error in the both trucation")
test6(1, log=TRUE)
log((pNBI(1)-pNBI(0))/(pNBI(9)-pNBI(0)))
if(abs(test6(1, log=TRUE)-log((pNBI(1)-pNBI(0))/(pNBI(9)-pNBI(0))))>0.00001)
             stop("error in both trucation")
plot(function(y) test6(y, mu=20, sigma=3), from=1, to=9, n=9, type="h")
plot(function(y) test6(y, mu=300, sigma=.4), from=1, to=9, n=9, type="h")
cdf < - stepfun(1:8, test6(1:9, mu=5), f = 0)
plot(cdf, main="cdf", ylab="cdf(x)", do.points=FALSE )
# varying truncation
# coninuous
# left
```

16 trun.q

```
test6<-trun.p(par=c(0,1,2), family="TF", type="left", varying=TRUE)
test6(c(2,3,4))
(pTF(c(2,3,4))-pTF(c(0,1,2)))/(1-pTF(c(0,1,2)))
test6(c(2,3,4), log.p=TRUE)
# right
test7 <- trun.p(par=c(10,11,12), family="BCT", type="right", varying=TRUE)
test7(c(1,2,3))
pBCT(c(1,2,3))/pBCT(c(10,11,12))
test7(c(1,2,3), lower.tail=FALSE)
1-pBCT(c(1,2,3))/pBCT(c(10,11,12))
test7(c(1,2,3), log.p=TRUE)
# both
test8<-trun.p(par=cbind(c(0,1,2), c(10,11,12)), family="TF",
           type="both", varying=TRUE)
test8(c(1,2,3))
(pTF(c(1,2,3))-pTF(c(0,1,2)))/(pTF(c(10,11,12))-pTF(c(0,1,2)))
test8(c(1,2,3), lower.tail=FALSE)
1-(pTF(c(1,2,3))-pTF(c(0,1,2)))/(pTF(c(10,11,12))-pTF(c(0,1,2)))
# discrete
# left
test9<-trun.p(par=c(0,1,2), family="PO", type="left", varying=TRUE)</pre>
test9(c(1,2,3))
(pPO(c(1,2,3))-pPO(c(0,1,2)))/(1-pPO(c(0,1,2)))
# right
test10<-trun.p(par=c(10,11,12), family="NBI", type="right", varying=TRUE)</pre>
test10(c(2,3,4))
pNBI(c(2,3,4))/(pNBI(c(9,10,11)))
test11<-trun.p(par=rbind(c(0,10), c(1,11), c(2, 12)), family="NBI",
              type="both", varying=TRUE)
test11(c(2,3,4))
(pNBI(c(2,3,4))-pNBI(c(0,1,2)))/(pNBI(c(9,10,11))-pNBI(c(0,1,2)))
```

trun.q

Truncated Inverse Cumulative Density Function of a gamlss.family Distribution

Description

Creates a function to produce the inverse of a truncated cumulative density function generated from a current GAMLSS family distribution.

For continuous distributions left truncation at 3 means that the random variable can take the value 3. For discrete distributions left truncation at 3 means that the random variable can take values from

trun.q 17

4 onwards. This is the same for right truncation. Truncation at 15 for a discrete variable means that 15 and greater values are not allowed but for continuous variable it mean values greater that 15 are not allowed (so 15 is a possible value).

Usage

Arguments

par	a vector with one (for "left" or "right" truncation) or two elements for "both". When the argument varying = TRUE then par can be a vector or a matrix with two columns respectively.
family	a gamlss.family object, which is used to define the distribution and the link functions of the various parameters. The distribution families supported by gamlss() can be found in gamlss.family. Functions such as BI() (binomial) produce a family object.
type	whether left, right or in both sides truncation is required, (left is the default)
varying	whether the truncation varies for different observations. This can be usefull in regression analysis. If varying = TRUE then par should be an n-length vector for type equal "left" and "right" and an n by 2 matrix for type="both"
	for extra arguments

Value

Returns a q family function

Author(s)

Mikis Stasinopoulos <d. stasinopoulos@gre.ac.uk> and Bob Rigby

References

Rigby, R. A. and Stasinopoulos D. M. (2005). Generalized additive models for location, scale and shape, (with discussion), *Appl. Statist.*, **54**, part 3, pp 507-554.

Rigby, R. A., Stasinopoulos, D. M., Heller, G. Z., and De Bastiani, F. (2019) *Distributions for modeling location, scale, and shape: Using GAMLSS in R*, Chapman and Hall/CRC. An older version can be found in https://www.gamlss.com/.

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Stasinopoulos D. M., Rigby R.A., Heller G., Voudouris V., and De Bastiani F., (2017) *Flexible Regression and Smoothing: Using GAMLSS in R*, Chapman and Hall/CRC.

```
(see also https://www.gamlss.com/)..
```

18 trun.q

See Also

```
trun.d, trun.q, trun.r, gen.trun
```

Examples

```
# trucated q continuous function
# continuous
# left
test1<-trun.q(par=c(0), family="TF", type="left")</pre>
test1(.6)
qTF(pTF(0)+0.6*(1-pTF(0)))
# right
test2 <- trun.q(par=c(10), family="BCT", type="right")</pre>
test2(.6)
qBCT(0.6*pBCT(10))
#-----
# both
test3<-trun.q(par=c(-3,3), family="TF", type="both")
test3(.6)
qTF(0.6*(pTF(3)-pTF(-3))+pTF(-3))
# varying par
# left
test7<-trun.q(par=c(0,1,2), family="TF", type="left", varying=TRUE)
test7(c(.5,.5,.6))
qTF(pTF(c(0,1,2))+c(.5,.5,.6)*(1-pTF(c(0,1,2))))
# right
test9 <- trun.q(par=c(10,11,12), family="BCT", type="right", varying=TRUE)</pre>
test9(c(.5,.5,.6))
qBCT(c(.5,.5,.6)*pBCT(c(10,11,12)))
# both
test10<-trun.q(par=cbind(c(0,1,2), c(10,11,12)), family="TF", type="both", varying=TRUE)
test10(c(.5, .5, .7))
qTF(c(.5, .5, .7)*(pTF(c(10,11,12))-pTF(c(0,1,2)))+pTF(c(0,1,2)))
#-----
# FOR DISCRETE DISTRIBUTIONS
# trucated q function
# left
test4<-trun.q(par=c(0), family="PO", type="left")</pre>
test4(.6)
qPO(pPO(0)+0.6*(1-pPO(0)))
# varying
test41<-trun.q(par=c(0,1,2), family="PO", type="left", varying=TRUE)
test41(c(.6,.4,.5))
qPO(pPO(c(0,1,2))+c(.6,.4,.5)*(1-pPO(c(0,1,2))))
# right
```

```
test5 <- trun.q(par=c(10), family="NBI", type="right")</pre>
test5(.6)
qNBI(0.6*pNBI(10))
test5(.6, mu=10, sigma=2)
qNBI(0.6*pNBI(10, mu=10, sigma=2), mu=10, sigma=2)
test51 <- trun.q(par=c(10, 11, 12), family="NBI", type="right", varying=TRUE)
test51(c(.6,.4,.5))
qNBI(c(.6,.4,.5)*pNBI(c(10, 11, 12)))
test51(c(.6,.4,.5), mu=10, sigma=2)
qNBI(c(.6,.4,.5)*pNBI(c(10, 11, 12), mu=10, sigma=2), mu=10, sigma=2)
# both
test6<-trun.q(par=c(0,10), family="NBI", type="both")</pre>
test6(.6)
qNBI(0.6*(pNBI(10)-pNBI(0))+pNBI(0))
# varying
test61 < -trun.q(par=cbind(c(\emptyset,1,2),\ c(10,11,12)),\ family="NBI",\ type="both",\ varying=TRUE)
test61(c(.6,.4,.5))
qNBI(c(.6,.4,.5)*(pNBI(c(10,11,12))-pNBI(c(0,1,2)))+pNBI(c(0,1,2)))
```

trun.r

Generates Random Values from a Truncated Density Function of a gamlss.family Distribution

Description

Creates a function to generate randon values from a truncated probability density function created from a current GAMLSS family distribution

For continuous distributions left truncation at 3 means that the random variable can take the value 3. For discrete distributions left truncation at 3 means that the random variable can take values from 4 onwards. This is the same for right truncation. Truncation at 15 for a discrete variable means that 15 and greater values are not allowed but for continuous variable it mean values greater that 15 are not allowed (so 15 is a possible value).

Usage

Arguments

par

a vector with one (for "left" or "right" truncation) or two elements for "both". When the argument varying = TRUE then par can be a vector or a matrix with two columns respectively.

family	a gamlss.family object, which is used to define the distribution and the link functions of the various parameters. The distribution families supported by gamlss() can be found in gamlss.family. Functions such as BI() (binomial) produce a family object.
type	whether left, right or in both sides truncation is required, (left is the default)
varying	whether the truncation varies for different observations. This can be usefull in regression analysis. If varying = TRUE then par should be an n-length vector for type equal "left" and "right" and an n by 2 matrix for type="both"
	for extra arguments

Value

Returns a r family function

Author(s)

Mikis Stasinopoulos <d. stasinopoulos@gre.ac.uk> and Bob Rigby

References

Rigby, R. A. and Stasinopoulos D. M. (2005). Generalized additive models for location, scale and shape, (with discussion), *Appl. Statist.*, **54**, part 3, pp 507-554.

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```
(see also https://www.gamlss.com/).
```

See Also

```
trun.p, trun.q, trun.d, gen.trun
```

Examples

```
rr<-test2(1000)
hist(rr)
# both
test3<-trun.r(par=c(-3,3), family="TF", type="both")</pre>
rr<-test3(1000)
hist(rr)
# discrete
# trucated r function
# left
test4<-trun.r(par=c(0), family="PO", type="left")</pre>
tN <- table(Ni <- test4(1000))
r <- barplot(tN, col='lightblue')</pre>
# right
test5 <- trun.r(par=c(10), family="NBI", type="right")</pre>
tN <- table(Ni <- test5(1000))
r <- barplot(tN, col='lightblue')</pre>
tN <- table(Ni <- test5(1000,mu=5))
r <- barplot(tN, col='lightblue')</pre>
tN <- table(Ni <- test5(1000,mu=10, sigma=.1))
r <- barplot(tN, col='lightblue')</pre>
# both
test6<-trun.r(par=c(0,10), family="NBI", type="both")</pre>
tN <- table(Ni <- test6(1000,mu=5))
r <- barplot(tN, col='lightblue')</pre>
# varying = TRUE
# continuous
test7<-trun.r(par=c(0,1,2), family="TF", type="left", varying=TRUE)</pre>
test7(3)
#-----
test8 <- trun.r(par=c(10,11,12), family="BCT", type="right", varying=TRUE)</pre>
test8(3)
# both
test9 < -trun.r(par=rbind(c(-3,3),\ c(-1,5),\ c(\emptyset,6)),\ ,\ family="TF",\ type="both",\ varying=TRUE)
#-----
# discrete
# trucated r function
test10<-trun.r(par=c(0,1,2), family="PO", type="left", varying=TRUE)</pre>
test10(3)
# right
```

Index

```
* distribution
    fitTail, 3
    gamlss.tr-package, 2
    gen.trun,5
    trun, 7
    trun.d,9
    trun.p, 13
    trun.q, 16
    trun.r, 19
* package
    gamlss.tr-package, 2
* regression
    fitTail, 3
    {\tt gamlss.tr-package}, {\tt 2}
    gen.trun,5
    trun, 7
    trun.d,9
    trun.p, 13
    trun.q, 16
    trun.r, 19
fitTail, 3
fitTailAll (fitTail), 3
gamlss.tr(gamlss.tr-package), 2
gamlss.tr-package, 2
gen.trun, 5, 8, 10, 14, 18, 20
loglogSurv, 5
logSurv, 5
trun, 7
trun.d, 6, 8, 9, 14, 18, 20
trun.p, 6, 8, 10, 13, 20
trun.q, 6, 8, 10, 14, 16, 18, 20
trun.r, 6, 8, 10, 14, 18, 19
```