Introduction to the garma Package

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Introduction

This is an introduction to the garma package. The package is designed to simulate, estimate and generate Moving Block Bootstrap resamples and diagnosis for the GARMA process of Benjamin et al. (2003). It has four core classes and respective methods, all related to the evaluation of simulated GARMA process. The S4 classes are: GarmaSpec, GarmaSim, GarmaFit and GarmaSimBoot. The syntax is straightforward, in the firt class 'Spec' stands for Specification of a GARMA process, while 'Sim' for Simulation of the specified model in the second, 'Fit' for estimating in the third, and 'SimBoot' for a Moving Block Bootstrap evaluation of the simulated process. All classes have the built-in methods print, summary and plot, and their behaviour depends on some factors as the number of Monte Carlo simulations being specified. ##How to install the package: The garma package depends on the dboot package. So the first step in the installation process is to certify that you have the package devtools installed:

```
if (!require("devtools")) install.packages("devtools")
```

Now the **dboot** and **garma** pkgs:

```
if (!require("dboot")) install_github("matheusbarroso/dboot")
if (!require("garma"))
install_github("matheusbarroso/garma")
```

The next section gives a formal defintion of the forementioned process followed by a section for the classes and method explaining their usage and providing examples.

The GARMA model

Let Y be a stochastic process, i.e. a collection of random variables $Y = Y_t(\omega), t \in \mathcal{T}, \omega \in \Omega$, defined in the filtered probability space $(\Omega, \mathcal{F}, \mathcal{F}_{t \geq 0}, \mathcal{P})$, where Ω is the set of all possible states, \mathcal{F} is the σ – field of all subsets of Ω , $\mathcal{F}_{t \geq 0}$ is a filtration, \mathcal{P} is a probability measure unter \mathcal{F} and \mathcal{T} an arbitrary set. In the garma model we have that each realization of $Y_t, t = 1, ...n$, has conditional distribution belonging to the exponential family of distributions, where the conditioning is w.r.t. to \mathcal{F}_{t-1} . $\mathcal{F}_{t-1} = \{\mathbf{x_1}, ..., \mathbf{x_{t-1}}; y_1, ..., y_{t-1}; \mu_1, ..., \mu_{t-1}\}$ such that the conditional density $f_{Y_t|\mathcal{F}_{t-1}}$ is of the form:

$$f_{Y_t|\mathcal{F}_{t-1}} = exp\left(\frac{y_t v_t - a(v_t)}{\varphi} + b(y_t, \varphi)\right)$$
(1)

where $a(\cdot)$ and $b(\cdot)$ are specific functions definining the particular member of the exponential familiy, with v_t as the canonical and φ as the scale parameter of the member, \mathbf{x} is a r dimensional vector of explanatory variables and μ is the mean vector. Additionally, the predictor η is such that $\eta = g(\mu_t)$ where g is the link function.

$$\eta_t = \mathbf{x}_t' \boldsymbol{\beta} + \sum_{j=1}^p \phi_j \{ g(y_{t-j}) - \mathbf{x}_{t-j}' \boldsymbol{\beta} \} + \sum_{j=1}^q \theta_j g(y_{t-j}) - \eta_{t-j}$$
(2)

where $\beta' = (\beta_1, \beta_2, \dots, \beta_r)$ is the vector of parameters of the linear predictor, $\phi' = (\phi_1, \phi_2, \dots, \phi_p)$ the vector of autoregressive parameters and $\theta' = (\theta_1, \theta_2, \dots, \theta_p)$ the vector of moving average parameters. Eq.1 and Eq.2 define the GARMA model.

In this package only the conditional *Gamma* and *Poisson* were implemented, though the application to all the exponential family is already in the pipeline. Thus, the next two subsctions deal with the specifics of the aforementioned models.

Poisson-GARMA model

If $Y_t|F_{t-1}$ follows a Poisson distribution with mean parameter μ_t then its p.m.f. is:

$$f_{Y_t|\mathcal{F}_{t-1}} = \frac{e^{-\mu_t} \mu_t^{y_t}}{y_t!} = \exp\left(y_t \log(\mu_t) - \mu_t - \log(y_t!)\right)$$
(3)

Consequently, we have that $y_t|\mathcal{F}_{t-1}$ belongs to the exponential distribution, $v_t = log(\mu_t)$, $a(v_t) = e^{-\mu_t}$, $b(y_t, \varphi) = -log(y_t!)$ and $\varphi = 1$. The canonical link function is $log \equiv ln$. Hence η_t is such that:

$$\eta_{t} = log(\mu_{t}) = \mathbf{x}'_{t}\boldsymbol{\beta} + \sum_{j=1}^{p} \phi_{j} \{ log(y_{t-j}^{*}) - \mathbf{x}'_{t-j}\boldsymbol{\beta} \} + \sum_{j=1}^{q} \theta_{j}g(y_{t-j}) - \eta_{t-j}$$
(4)

where $y_{t-i}^* = max(y_{t-i}, \alpha), 0 < \alpha < 1$. α is the offset parameter and its default value in the package is 0.1

Gamma-GARMA model

If $Y_t|F_{t-1}$ follows a Gamma distribution with shape parameter δ and scale parameter γ (so a $\Gamma(\delta, \gamma)$ distribution) its p.d.f is given by:

$$f_{Y_t|\mathcal{F}_{t-1}} = \frac{y_t^{\delta-1} e^{-y_t/\delta}}{\Gamma(\delta)\gamma^{\delta}} \tag{5}$$

with $E_{Y_t|\mathcal{F}_{t-1}} = \delta \gamma$ and $Var_{Y_t|\mathcal{F}_{t-1}} = \delta \gamma^2$. However, here we adopt a more useful parametrization of the gamma density given by this table transformation: $delta = 1/\sigma^2$ and $\gamma = \sigma^2 \mu_t$. With these new parameters at hand we have that $E_{Y_t|\mathcal{F}_{t-1}} = \mu_t$ and $Var_{Y_t|\mathcal{F}_{t-1}} = \sigma^2 \mu_t^2$. In the Gamma case the canonical link function is the reciprocal function, though, we make usage of the logarithmic function. A new version of the package with all reasonable link functions will be developed but keep in mind that the log is currently being used in the Gamma case so that the equation for η_t is the same for the Poisson case, that is, Eq. 4.

The garma package

After a formal definition of the *GARMA* model now we are able to understand better each class and its respective methods in the **garma** package. This section contains four subsections, one for each main class in the package.

GarmaSpec

The GarmaSpec class is the more general class for specificating the *GARMA* model and contain the slots that are common to all members of the exponential family. For those unfamiliar with S4 classes, slots might be seen as the parameters that specify the class, and are defined as follows: * family: A character vector specifying the family of the specification object. Accepted values are: "Po" for poisson and "GA" for gamma families.

- beta.x:A numeric vector with length of the desired specification, in the current version the intercept term must be includer here. The default behaviour is beta.x = 1L.
- phi: A numeric vector with length of the desired autoregressive term order specification. The default behaviour is phi = OL.
- theta: A numeric vector with length of the desired moving-average term order specification. The default behaviour is theta = 0L.
- X: A $n \times m$ matrix, where n = nsteps + burnin + max.order and m = length(beta.x). Where nsteps is the number of simulations desired, burnin the number of burnin observations and max.order is equal to the highest order of the GARMA model (i.e. max(length(phi),length(theta).)

This class has two child classes: PoissonSpec and GammaSpec. Each one specifies the additional slots required for a correc specification of a *Poisson* or *Gamma GARMA* models.

PoissonSpec

Inherits from GarmaSpec and has the slots:

- family: A character with the tag "PO".
- alpha: Numeric, specifying the offset (α) parameter. The default value is $\alpha = 0.01$
- mu0: A numeric vector with length equal to the max.order of the Poisson-GARMA model. The default value is $\mu_0 = 10$.

GammaSpec

Inherits from GarmaSpec and has the slots:

- family: A character with the tag "PO".
- sigma2: Numeric, specifying the σ^2 parameter.
- mu0: A numeric vector with length equal to the max.order of the Poisson-GARMA model. The default value is $\mu_0 = 10$.

Methods

To avoid problems in the model specification the easiest way to create an object of the GarmaSpec(Gamma/Poisson) is through the construtor function (method) GarmaSpec. At least the family argument must be provided in order to create an object of this class. If no further arguments are passed the default behaviour is used:

```
library(garma)
GarmaSpec("PO") # This results in no ARMA terms.
#> An object of class "PoissonSpec"
#> Slot "alpha":
#> [1] 0.1
#>
#> Slot "mu0":
#> [1] 10
#> Slot "family":
#> [1] "PO"
#>
#> Slot "beta.x":
#> [1] 1
#>
#> Slot "phi":
#> [1] 0
#> Slot "theta":
#> [1] 0
#>
#> Slot "X":
#> <0 x 0 matrix>
GarmaSpec("GA") # This results in no ARMA terms.
#> An object of class "GammaSpec"
#> Slot "sigma2":
#> [1] 1
#>
#> Slot "mu0":
#> [1] 10
#>
#> Slot "family":
#> [1] "GA"
#> Slot "beta.x":
#> [1] 1
#>
#> Slot "phi":
#> [1] 0
#>
#> Slot "theta":
#> [1] 0
#>
#> Slot "X":
#> <0 x 0 matrix>
```

If we wish to specify a second order poisson model we would need at least:

```
GarmaSpec("PO",phi=c(0.3,0.2),mu0=c(1,1))
#> An object of class "PoissonSpec"
#> Slot "alpha":
#> [1] 0.1
#>
#> Slot "mu0":
#> [1] 1 1
#> Slot "family":
#> [1] "PO"
#>
#> Slot "beta.x":
#> [1] 1
#>
#> Slot "phi":
#> [1] 0.3 0.2
#> Slot "theta":
#> [1] 0
#>
#> Slot "X":
#> <0 x 0 matrix>
```

Now we are ready to move to some more elaborate examples:

1. A GA-GARMA(1,0) with an intercept term, note that the length of x1 is 101, as we will simulate 100 steps, and the other 1 is for the initial value of Phi. The initial value for μ_0 is the default.

```
spec1 <- GarmaSpec("GA",
phi = 0.5,
beta.x = 1,
sigma2 =2,
X = as.matrix(
   data.frame(
    x1 = rep(10,101))))</pre>
```

2. A PO-GARMA(1,3) with an intercept term, here the default behaviour of μ_0 will not suffice, so we need three initial values. The intercept will be generated automatically

```
spec2 <- GarmaSpec("PO",
phi = 0.2,
theta = c(0.1, 0.3, 0.5),
mu0 = 1:3)</pre>
```

3. A PO-GARMA(2,0) with an intercept term and one covariate. A series of length 100 will be generated, though, 1000 burnin observations will be deleted and the covariates must take into consideration these obs and the order of the process, so that the nrow of X should be 100 (desired length)+1000 (burnin)+2 (order of the process)

```
spec3 <- GarmaSpec("PO",
phi = c(0.5,0.15),
beta.x = c(1,1),
mu0 = c(2,2),
X = as.matrix(
data.frame(
intercept = rep(1,1102),</pre>
```

```
x1 = c(rep(7,100), rep(2,1002))))
```

4. A PO-GARMA(0,0), with an intercept and covariate:

```
spec4 <- GarmaSpec("PO",
beta.x = c(0.1,1),
X = as.matrix(
  data.frame(
  intercept = rep(1,1100),
     x1 = c(rep(7,100),
         rep(2,1000)))))</pre>
```

GarmaSim

The GarmaSim class is used to simulate an object of the class GarmaSpec. The methods GarmaSim, print, plot and summary are implemented. The GarmaSim parameters are:

- spec: An object of the GarmaSpec class, as provided by the method GarmaSpec
- nmonte: A positive integer, specifying the number of Monte Carlo simulations to perform, the default value is 1000.
- nsteps: A numeric vector with the number of steps in the Garma model simulation, that is, the length of the time series to simulate. The default value is 100.
- burnin: A numeric vector indicating the number of burn in observations. If you want to generate only nsteps this argument should be set equal to zero. Otherwise, provide a positive integer. The default value is 1000.
- allow.parallel: Logical TRUE/FALSE indicating whether parallel computation via the *foreach* package should be used. The default value is TRUE. OBS:parallel backend must be registered prior to calling GarmaSim.
- seed: Numeric, the seed to set.seed() for replicable examples. Default value is 123. Before moving to the examples it is useful to register the parallel back end:

```
library(doParallel)
no_cores <- detectCores() - 1
registerDoParallel(no_cores)</pre>
```

Now we can move to some examples: 1. Using the specifications build in the session Garma Spec:

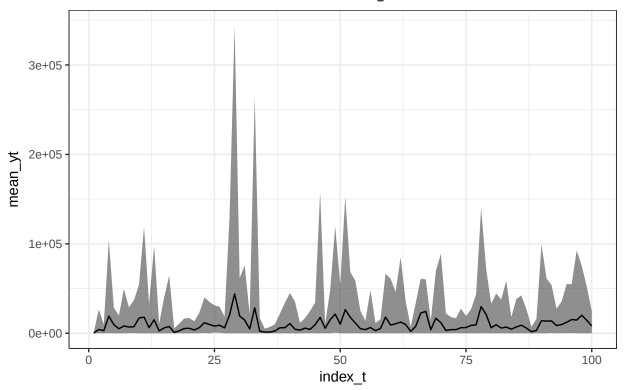
The print method give us a tidy output in the monte carlo setting, with a brief descritpion of the model and each generated series in the columns.

```
print(sim1)
```

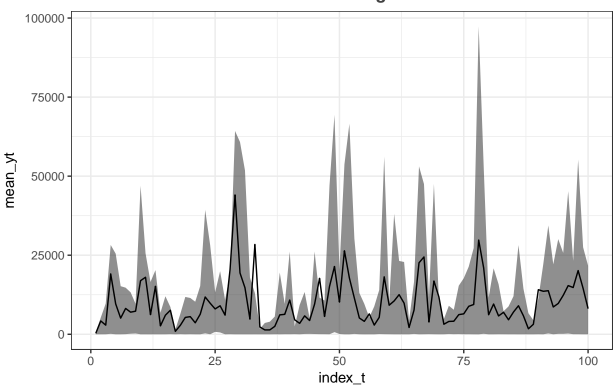
When nmonte > 1, the plot comes with th 95% empirical confidence interval for the mean of each observation. The confidence level can be changed through the confint parameter:

```
plot(sim1)
```

Simulated GA-Garma(1,0): Mean values and 95% confidence interval through time



Simulated GA-Garma(1,0): Mean values and 80% confidence interval through time



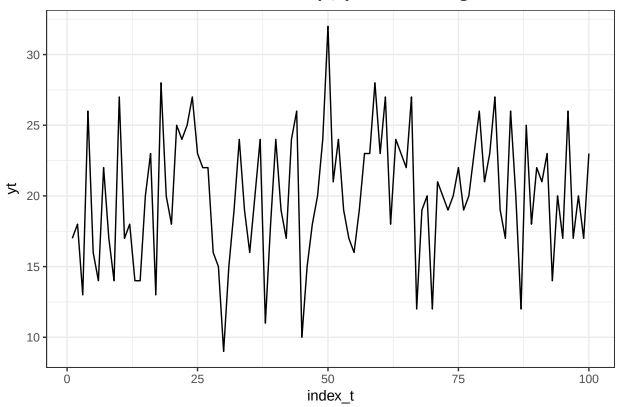
The summary(GarmaSim) method has a different behaviour for nmonte > 1 and nmonte = 1

```
summary(sim1)
#> -----
#> A GA-Garma(1,0) simulation object:
#> Number of Monte Carlo Simulations ('nmonte') = 10
#> Time Series Length ('nsteps') = 100
\#> Burn-in ('burnin') = 0
#> parallel = TRUE
#> Mean Monte Carlo mean values = 9932.798 with distribution:
      2.5%
               50%
                       97.5%
#> 2 5638.614 8282.88 18799.36
#>
#> Mean Monte Carlo min values = 0.4406 with distribution:
#> 2.5% 50% 97.5%
#> 1 0 0.027 1.512
#> Mean Monte Carlo max values = 134052.1 with distribution:
#> 2.5% 50% 97.5%
#> 3 53313.97 92538.95 344191.5
#>
                      2.5%
                                 50%
                                          97.5%
             mean
          0.4406
                      0.000
                               0.027
                                          1.512
#> min
#> mean 9932.7980 5638.614 8282.880 18799.355
#> max 134052.0926 53313.974 92538.953 344191.509
```

2. An Poisson-GARMA(0,0), with only one Monte Carlo simulation:

```
sim2 <-
 GarmaSim(
   GarmaSpec("PO",
             beta.x = c(0.1,1),
             X = as.matrix(
             data.frame(
              intercept = rep(10,1100),
              x1 = c(rep(7,100), rep(2,1000)))),
 nmonte = 1,
 allow.parallel = TRUE)
print(sim2)
#> ---
#> A PO-Garma(0,0) simulation object:
#>
#> beta.intercept = 0.1
\#> beta.x1 = 1
#> mu0[1] = 10
#> Number of Monte Carlo Simulations ('nmonte') = 1
#> Time Series Length ('nsteps') = 100
#> Burn-in ('burnin') = 1000
#> parallel = TRUE
#> -----
                       _____
#>
   [1] 17 18 13 26 16 14 22 17 14 27 17 18 14 14 20 23 13 28 20 18 25 24 25
#>
#> [24] 27 23 22 22 16 15 9 15 19 24 19 16 20 24 11 18 24 19 17 24 26 10 15
#> [47] 18 20 24 32 21 24 19 17 16 19 23 23 28 23 27 18 24 23 22 27 12 19 20
#> [70] 12 21 20 19 20 22 19 20 23 26 21 23 27 19 17 26 20 12 25 18 22 21 23
#> [93] 14 20 17 26 17 20 17 23
plot(sim2)
```

Simulated PO-Garma(0,0) values through time



```
summary(sim2)
#>
    A PO-Garma(0,0) simulation object:
#>
#>
   Number of Monte Carlo Simulations ('nmonte') = 1
#>
   Time Series Length ('nsteps') = 100
#>
   Burn-in ('burnin') = 1000
#>
#>
    parallel = TRUE
#>
      Min. 1st Qu.
                   Median
                              Mean 3rd Qu.
                                              Max.
      9.00 17.00
                   20.00
                            20.07 23.00
                                             32.00
```

Further specification examples can be found at the GarmaSim-class help menu.

GarmaFit

Given a GarmaSim object we can simultaneously estimate all the nmonte series (if parallel = TRUE and the parallel back-end registered). The methods GarmaFit, print, plot and summary are implemented. The GarmaSim method has the following parameters:

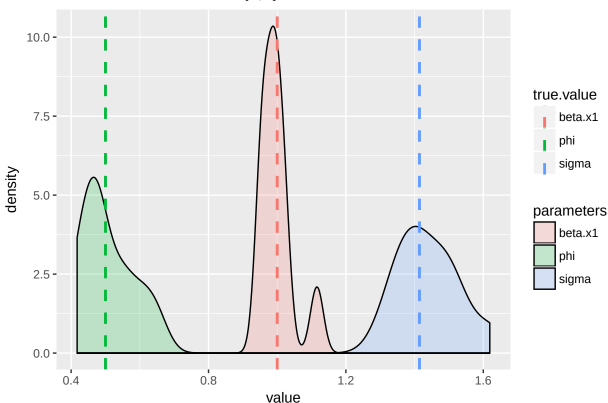
- garma: An object of the GarmaSim class, as provided by GarmaSim.
- allow.parallel: Logical TRUE/FALSE indicating whether parallel computation via the foreach package should be used. The default value is TRUE. OBS:parallel backend must be registered prior to calling GarmaSim.
- seed: Numeric, the seed to set.seed() for replicable examples. Default value is 123.

- errorhandling: Character, either 'try' or 'pass'
- n.try: Positive integer. If errorhandling = 'try', this specifies the number of attempts in the algorithm.
- control: List. This is passed to the garmaFit2 function. The options are given by garmaFit function in the gamlss.util package.

The examples: 1.

```
fit1 <- GarmaFit(sim1)</pre>
print(fit1)
#>
#> A GA-Garma(1,0) simulation fitted object:
#>
#> phi 0.5
\#> beta.x1 = 1
\#> muO[1] = 10
\#> siqma2 = 2
#> Number of Monte Carlo Simulations ('nmonte') = 10
#> Time Series Length ('nsteps') = 100
#> Burn-in ('burnin') = 0
#> parallel = TRUE
#> errorhandling = try
\#> n.try = 5
#>
#>
#> Estimated parameters:
              [,1]
                       [,2]
                                [,3]
                                         [,4]
                                                  [,5]
#>
                                                           [,6]
#> beta.x1 0.9512578 1.1156606 0.9952164 0.9638448 1.0227713 0.9951840
#> phi 0.4214639 0.6464954 0.4174278 0.4640413 0.6021564 0.4757756
[,9]
              [,7]
                                        [,10]
                       [,8]
#> beta.x1 0.9919034 0.9755214 1.0169597 0.9530157
         0.5313133 0.5566795 0.4767271 0.4704674
#> phi
#> siqma
         1.6189999 1.3584369 1.5088017 1.4912128
plot(fit1)
```

Simulated GA-Garma(1,0): Coefficients Distribution

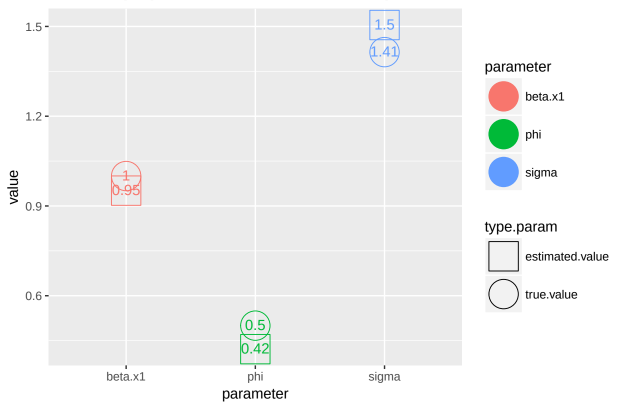


```
summary(fit1)
#> A GA-Garma(1,0) simulation object:
#>
#> Number of Monte Carlo Simulations ('nmonte') = 10
#> Time Series Length ('nsteps') = 100
\#> Burn-in ('burnin') = 0
#> parallel = TRUE
#> parameters
                  Min. 1st Qu. Median.
                                             Mean. 3rd Qu.
#> 1 beta.x1 0.9512578 0.9638448 0.9919034 0.9981335 1.0169597 1.1156606
        phi 0.4174278 0.4640413 0.4757756 0.5062548 0.5566795 0.6464954
       sigma 1.3231409 1.3715135 1.4208644 1.4424751 1.5047123 1.6189999
#> true.value
#> 1 1.000000
#> 2 0.500000
#> 3 1.414214
```

2. An example with nmonte = 1:

```
x1 = rep(10,101))),
 nmonte = 1,
 burnin = 0)
fit2 <- GarmaFit(sim2)</pre>
print(fit2)
#> -----
#> A GA-Garma(1,0) simulation fitted object:
#>
#> phi 0.5
\#> beta.x1 = 1
#> mu0[1] = 10
#> sigma2 = 2
#> Number of Monte Carlo Simulations ('nmonte') = 1
#> Time Series Length ('nsteps') = 100
#> Burn-in ('burnin') = 0
#> parallel = TRUE
#> errorhandling = try
\#> n.try = 5
#> -----
#>
#> Estimated parameters:
#> [,1]
#> beta.x1 0.9512578
plot(fit2)
```





```
summary(fit2)
#>
   A GA-Garma(1,0) simulation object:
#>
  Number of Monte Carlo Simulations ('nmonte') = 1
#>
#> Time Series Length ('nsteps') = 100
  Burn-in ('burnin') = 0
#>
#>
   parallel = TRUE
#>
    parameters true.value estimate
#> 1
      beta.x1 1.000000 0.9512578
#> 2
        phi 0.500000 0.4214639
         sigma 1.414214 1.5047123
#> 3
```

GarmaSimBoot

Given a GarmaSim object we can simultaneously perform the Moving Block Bootstrap for all the nmonte series (if parallel = TRUE and the parallel back-end registered). The methods GarmaSimBoot, print, plot and summary are implemented. The GarmaSimBoot method has the following parameters:

- sim: An object of the GarmaSim class, as provided by GarmaSim.
- 1: 1 is the fixed block length used in generating the replicate time series.
- R: A positive integer giving the number of bootstrap replicates required.
- allow.parallel: Logical TRUE/FALSE indicating whether parallel computation via the foreach package

should be used. The default value is TRUE. OBS:paralllel backend must be registered prior to calling GarmaSim.

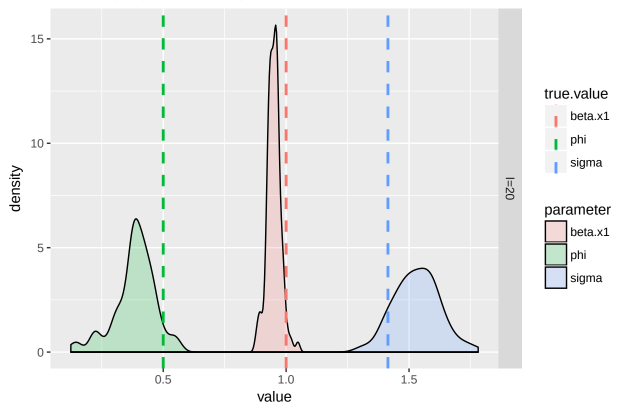
- seed: Numeric, the seed to set.seed() for replicable examples. Default value is 123.
- errorhandling: Character, either 'try' or 'pass'
- n.try: Positive integer. If errorhandling = 'try', this specifies the number of attempts in the algorithm.
- boot.function: A function to summarise the bootstrap replicates. The default function returns 0. Be aware that this is not a problem, as by default the mean values is already being returned. This is useful if the user wants to specify a quantity not being reported, as an example consider the 0.2 quantile. Warning: this feature is only enabled if nmonte > 1.
- control: List. This is passed to the garmaFit2 function. The options are given by garmaFit function in the gamlss.util package.

The examples:

1.nmonte = 1 and only one block length for the MBB:

```
# specification and simulation:
Sim1 <- GarmaSim(</pre>
  GarmaSpec("GA",
            phi = 0.5,
            beta.x = 1,
            sigma2 = 2,
            X = as.matrix(
              data.frame(
                 x1 = rep(10, 101))),
  nmonte = 1.
  burnin = 0)
#MBB
mbb1 <- GarmaSimBoot(Sim1,1 = 20)</pre>
#> deviance of linear model= 1853.087
#> deviance of garma model= 1805.009
#print(mbb1)
plot(mbb1)
```

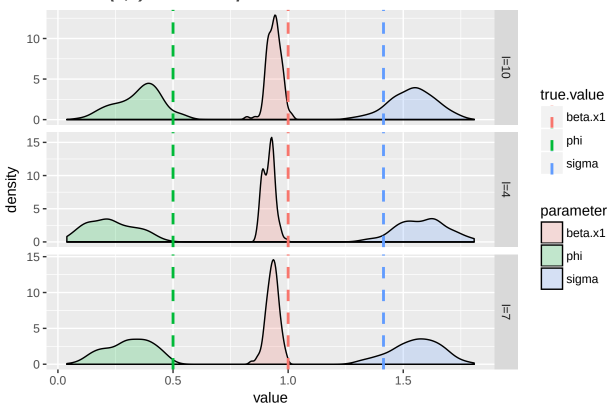
ed GA-Garma(1,0): Bootstrap Distribution from 1 Monte Carlo Simulation



```
summary(mbb1)
#> A GA-Garma(1,0) bootstrap object:
#>
#> Number of Monte Carlo Simulations ('nmonte') = 1
#> Time Series Length ('nsteps') = 100
\#> Burn-in ('burnin') = 0
#> parallel = TRUE
#> errorhandling = try
\#> n.try = 5
#> block length = 20
\#>R=100
#>
#>
#> Bootstrap Statistic Summary:
#> $`True values`
#> parameter true.value
#> 1 beta.x1 1.000000
       phi 0.500000
#> 2
#> 3
       sigma 1.414214
#>
#> $Estimates
#> length parameter variable Mean Est.
#> 1  l=20  beta.x1 original 0.9502413
#> 2  l=20  phi original 0.3853386
#> 3  l=20  sigma original 1.5316826
```

2. nmonte = 10, multiple block lengths

ed GA-Garma(1,0): Bootstrap Distribution from 1 Monte Carlo Simulation

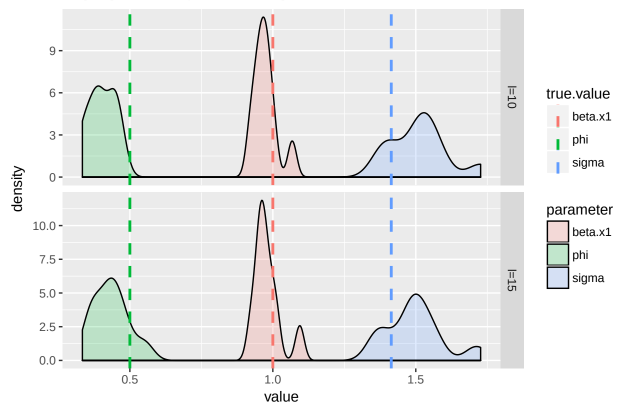


```
#> Bootstrap Statistic Summary:
#> $`True values`
#> parameter true.value
#> 1 beta.x1 1.000000
#> 2
       phi
              0.500000
#> 3
      sigma 1.414214
#>
#> $Estimates
#> length parameter variable Mean Est.
#> 8 l=7
             phi original 0.3079951
#> 9
    l=7 sigma original 1.5559196
#> 4
     l=4 beta.x1 original 0.9161941
#> 5
     l=4
              phi original 0.2294262
#> 6 l=4
            sigma original 1.5817752
#> 1  l=10  beta.x1 original 0.9389050
#> 2
     l=10
             phi original 0.3516731
#> 3 l=10
            sigma original 1.5401262
```

3. nmonte = 10, multiple block lengths + a user defined function to apply in the bootstrap resamples (the 0.1 and 0.9 quantiles)

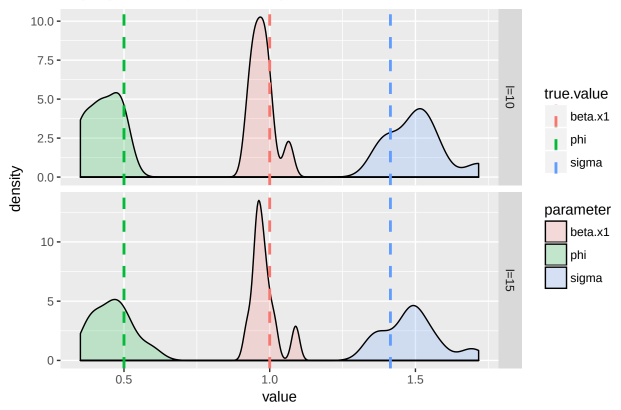
```
#print(mbb3)
plot(mbb3)
```

A-Garma(1,0): Average Bootstrap Estimates from 10 Monte Carlo Simulati



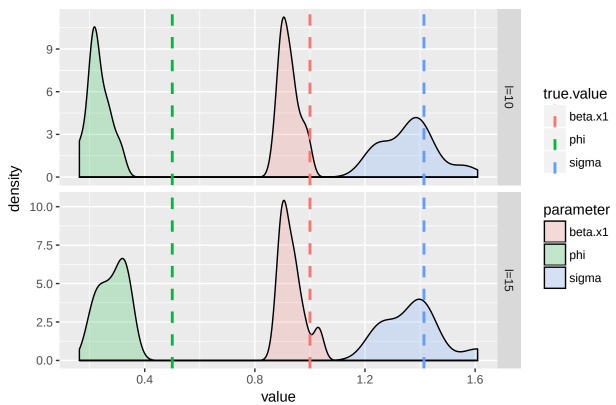
plot(mbb3, variable = "Median.")

A-Garma(1,0): Average Bootstrap Estimates from 10 Monte Carlo Simulati



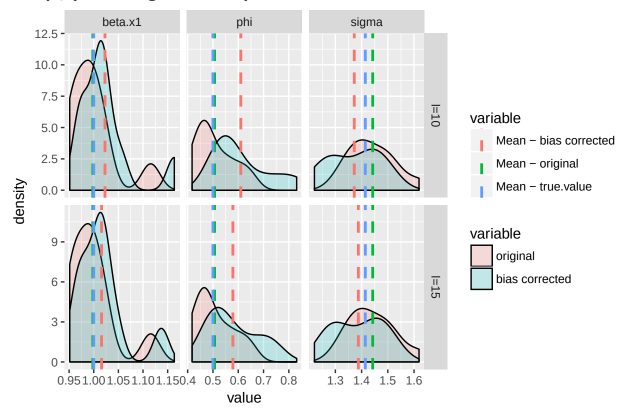
plot(mbb3, variable = "boot.func.10%")

A-Garma(1,0): Average Bootstrap Estimates from 10 Monte Carlo Simulati



#plot(mbb3, variable = "test") # if you type a variable that #does not exists you can see all available
plot(mbb3, type = "original-bias", scales = 'free')

arma(1,0): Average Bootstrap Estimates from 10 Monte Carlo Simulations



```
summary(mbb3)
   A GA-Garma(1,0) bootstrap object:
#>
  Number of Monte Carlo Simulations ('nmonte') = 10
#>
#> Time Series Length ('nsteps') = 100
\#> Burn-in ('burnin') = 0
  parallel = TRUE
#>
  errorhandling = try
  n.try = 5
#>
#>
   block length = 10 15
#>
   R = 100
#>
#>
#> Bootstrap Statistic Summary:
#> $`True values`
    parameter true.value
              1.000000
     beta.x1
          phi
#> 2
              0.500000
#> 3
        sigma
              1.414214
#>
#> $Estimates
#> $Estimates$original
     length parameter variable summarise
       l=10 beta.x1 original 0.9981340
#> 1
#> 13  l=10  phi original 0.5062552
```

```
#> 25
     l=10 sigma original 1.4424760
#> 37 l=15
            beta.x1 original 0.9981340
            phi original 0.5062552
#> 49
      l=15
#> 61
       l=15
             sigma original 1.4424760
#> $Estimates$bias
#> length parameter variable summarise
     l=10 beta.x1 bias -0.02468396
#> 2
#> 14 l=10
            phi
                      bias -0.10392664
     l=10
                     bias 0.07047019
#> 26
             siqma
#> 38
     l=15 beta.x1
                      bias -0.01776717
#> 50 l=15
             phi
                    bias -0.07256006
#> 62 l=15
                    bias 0.05485127
            sigma
#>
#> $Estimates$`bias corrected`
#> length parameter variable summarise
#> 3
      l=10 beta.x1 bias corrected 1.0228179
#> 15
      l=10
            phi bias corrected 0.6101819
#> 27
     l=10
            sigma bias corrected 1.3720058
#> 39
     l=15 beta.x1 bias corrected 1.0159012
             phi bias corrected 0.5788153
#> 51 l=15
#> 63
     l=15
             sigma bias corrected 1.3876247
#>
#> $Estimates$` std. error`
#> length parameter variable summarise
      l=10 beta.x1 std. error 0.03779253
#> 4
#> 16
     l=10
            phi std. error 0.11520966
#> 28
     l=10
            sigma std. error 0.11798771
            beta.x1 std. error 0.03809448
#> 40 l=15
#> 52
     l=15
            phi std. error 0.10869360
#> 64
     l=15
             sigma std. error 0.10148352
#>
#> $Estimates$Min.
#> length parameter variable summarise
#> 5
      l=10 beta.x1 Min. 0.87451636
                      Min. 0.08138220
#> 17
     l=10
              phi
#> 29
      l=10
                      Min. 1.26085910
             sigma
#> 41  l=15  beta.x1
                    Min. 0.87138490
#> 53 l=15
            phi Min. 0.07255566
#> 65
     l=15
                    Min. 1.26234587
            sigma
#> $Estimates$`1st Qu.`
#> length parameter variable summarise
#> 6
      l=10 beta.x1 1st Qu. 0.9484606
#> 18
      l=10
               phi 1st Qu. 0.3232478
#> 30
     l=10
              sigma 1st Qu. 1.4278401
     l=15
            beta.x1 1st Qu. 0.9574524
#> 42
            phi 1st Qu. 0.3751582
#> 54
       l=15
#> 66
       l=15
            sigma 1st Qu. 1.4280658
#> $Estimates$Median.
#> length parameter variable summarise
#> 7  l=10 beta.x1 Median. 0.9745827
```

```
#> 19  l=10  phi  Median. 0.4298073
#> 31 l=10
              sigma Median. 1.5046131
            beta.x1 Median. 0.9808227
#> 43
       l=15
#> 55
            phi Median. 0.4603785
       l=15
             sigma Median. 1.4889247
#> 67
       l=15
#>
#> $Estimates$Mean.
#> length parameter variable summarise
#> 8
       l=10 beta.x1 Mean. 0.9734500
              phi Mean. 0.4023286
#> 20
      l=10
             sigma Mean. 1.5129462
#> 32
      l=10
#> 44
       l=15
            beta.x1 Mean. 0.9803668
#> 56
              phi Mean. 0.4336952
       l=15
                     Mean. 1.4973273
#> 68
       l=15
              siqma
#>
#> $Estimates$`3rd Qu.`
#> length parameter variable summarise
      l=10 beta.x1 3rd Qu. 0.9986573
#> 21
       l=10
               phi 3rd Qu. 0.4844474
            sigma 3rd Qu. 1.5866617
#> 33
      l=10
            beta.x1 3rd Qu. 1.0071224
#> 45
      l=15
#> 57
       l=15
             phi 3rd Qu. 0.5076496
#> 69
       l=15
             sigma 3rd Qu. 1.5592318
#> $Estimates$Max.
#> length parameter variable summarise
#> 10  l=10  beta.x1  Max. 1.0622121
#> 22
      l=10
              phi
                      Max. 0.5977572
                      Max. 1.8272891
#> 34
      l=10
              siqma
#> 46
                      Max. 1.0629719
      l=15
            beta.x1
#> 58
      l=15
            phi
                     Max. 0.5909486
#> 70
              sigma Max. 1.7863893
      l=15
#> $Estimates$`boot.func.10%`
#> length parameter variable summarise
      l=10 beta.x1 boot.func.10% 0.9238160
#> 11
              phi boot.func.10% 0.2322866
#> 23
       l=10
#> 35
       l=10
             sigma boot.func.10% 1.3632199
       l=15 beta.x1 boot.func.10% 0.9325777
#> 47
       l=15
            phi boot.func.10% 0.2830445
#> 59
#> 71
       l=15
             sigma boot.func.10% 1.3765794
#>
#> $Estimates$`boot.func.90%`
#> length parameter variable summarise
#> 12
      l=10 beta.x1 boot.func.90% 1.0208273
#> 24
       l=10
              phi boot.func.90% 0.5225375
#> 36
      l=10
              sigma boot.func.90% 1.6683108
#> 48
       l=15
            beta.x1 boot.func.90% 1.0261614
#> 60
       l=15
              phi boot.func.90% 0.5408858
#> 72
       l=15
            sigma boot.func.90% 1.6252759
```

ConfidenceInterval

Provided a MBB object, another useful thing to do is to examine the behaviour of the simulated confidence intervals. This task is easy, with built-in functions handling it and also the task of computing coverage rates and default plots.

Given a GarmaSimBoot object we can simultaneously perform confidence inteval computations for all the nmonte series (if parallel = TRUE and the parallel back-end registered). The methods ConfidenceInterval, print, plot, summary and coverage are implemented. The ConfidenceInterval method has the following parameters:

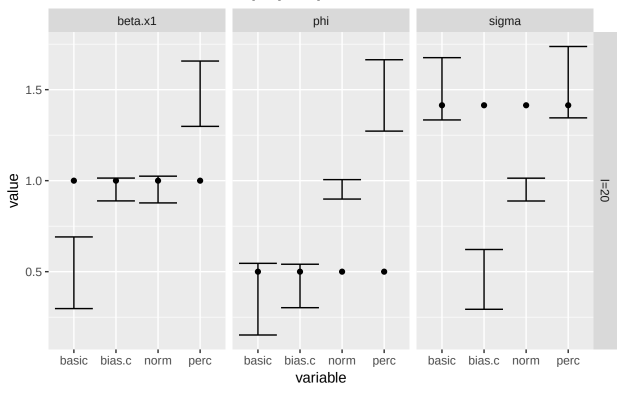
- garma.boot: An object of the GarmaSimBoot class, as provided by GarmaSimBoot.
- allow.parallel: Logical TRUE/FALSE indicating whether parallel computation via the foreach package should be used. The default value is TRUE. OBS:parallel backend must be registered prior to calling GarmaSim.
- conf: A scalar containing the confidence level of the required interval(s). The default value is 0.95.

The examples:

1. nmonte = 1:

```
coverage (ci)
#>
   A GA-Garma(1,0) MBB ci's coverage object:
#>
#>
#> Number of Monte Carlo Simulations ('nmonte') = 1
#> Time Series Length ('nsteps') = 100
#>
   parallel = TRUE
  block\ length = 20
#>
#>
  R = 100
#>
#>
#>
   Bootstrap Confidence Interval Coverage Rates:
    length parameter norm bias.c basic perc
#>
#> 1
     l=20 beta.x1
                     1
                            1 1
#> 2 l=20
                             1
                phi
                       1
                                   1
                                        1
#> 3 l=20
                                        1
               siqma 1
                             1
                                   1
plot(ci)
```

CI's Lower Bounds (LB) and Upper Bounds (LB) for parameters true values



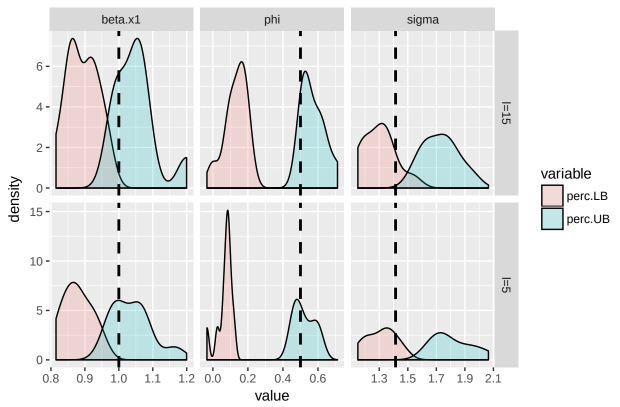
Note that we have information regarding the Normal, Basic, Bias Corrected and Percentile confidence intervals. In the case of nmonte > 1 we just have do provide the respective tag for each ci (actually type.ci argument to the plot method): 'norm', 'basic', 'bias.c' and 'perc'.

2. nmonte > 1:

```
spec2 <- GarmaSim(</pre>
GarmaSpec("GA",
         phi = c(0.5),
         beta.x = 1,
         sigma2 = 2,
         X = as.matrix(
           data.frame(
             x1 = rep(10,101))),
nmonte = 10, burnin = 0)
boot2 <- GarmaSimBoot(spec2,1 = c(5,15))
ci <- ConfidenceInterval(boot2)</pre>
print(ci)
summary(ci)
coverage (ci)
#>
    A GA-Garma(1,0)
                       MBB ci's coverage object:
#>
   Number of Monte Carlo Simulations ('nmonte') = 10
```

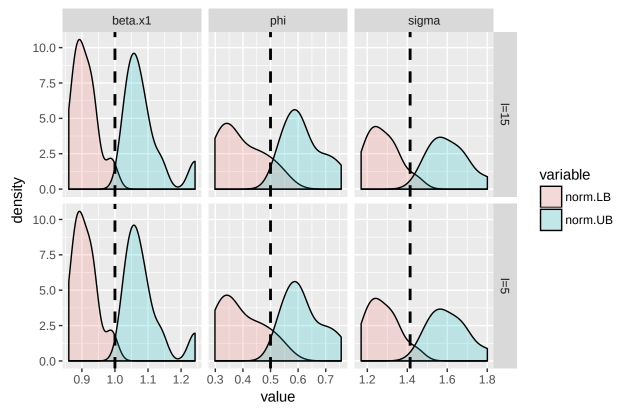
```
#> Time Series Length ('nsteps') = 100
#> parallel = TRUE
#> block length = 5 15
\#>R=100
#>
#>
#> Bootstrap Confidence Interval Coverage Rates:
  length parameter norm bias.c basic perc
#> 1  l=5  beta.x1 1.0
                          0.9
                               0.9 0.7
     l=5
#> 2
              phi 0.9
                          0.7
                              0.7 0.5
#> 3 l=5 sigma 0.9
                          1.0
                               1.0 0.8
#> 4 l=15 beta.x1 1.0
                          0.9
                               0.9 0.7
#> 5 l=15
                               0.8 1.0
               phi 0.9
                          1.0
#> 6 l=15
              sigma 0.9
                          1.0 1.0 0.9
plot(ci,type.ci='perc')
```

erc CI's Lower Bounds (LB) and Upper Bounds (LB) for 'nsims' = 10



plot(ci) #norm ci





Estimation of a general Garma model

This section deals with the estimation of any garma model, making usage of the garmaFit2 function from the dboot package (an adapted version of the garmaFit function from the gamlss.util pkg). This is a general purpose function, handling families other than the ones in the exponential family, for details check the gamlss.util package. The GarmaFit method is only a constructor from a GarmaSim object to the garmaFit function. The minimal structure for estimating a GARMA process is (as defined in the garmaFit function):

- formula: A formula for linear terms i.e. like in lm()
- order: the relevant data.frame.
- data: Numeric, the seed to set.seed() for replicable examples. Default value is 123.

So we can construc an example with the GarmaSim function or use the analogy into a real data example:

```
sim <- GarmaSim(spec,</pre>
              nmonte = 1,
              nsteps = 100)
#creating the required data.frame
db <- data.frame(cbind(yt = print(sim), intercept = rep(1,100)))</pre>
#> -----
#> A PO-Garma(2,0) simulation object:
#>
\#> phi1 = 0.5
\#> phi2 = 0.15
#> beta.intercept = 1
\#> beta.x1 = 1
\#> mu0[1] = 2
\#> mu0[2] = 2
#> Number of Monte Carlo Simulations ('nmonte') = 1
#> Time Series Length ('nsteps') = 100
#> Burn-in ('burnin') = 1000
#> parallel = TRUE
#> -----
#>
set.seed(123)
fit <- garmaFit2(yt~.-1 ,data= db,order = sim@order )</pre>
#> deviance of linear model= 637.7676
#> deviance of garma model= 571.322
library(gamlss.util)
#> Loading required package: zoo
#>
#> Attaching package: 'zoo'
#> The following objects are masked from 'package:base':
#>
      as.Date, as.Date.numeric
summary(fit)
#>
#> Family: c("NO", "Normal")
#> Fitting method: "nlminb"
#>
#> Call: qarmaFit2(formula = yt ~ . - 1, order = sim@order, data = db)
#>
#> Coefficient(s):
                 Estimate Std. Error t value Pr(>|t|)
#> beta.intercept 19.0463703     1.5215935 12.51738 < 2.22e-16 ***</pre>
          #> phi1
#> phi2
               4.4636610 0.3188329 14.00000 < 2.22e-16 ***
#> sigma
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#>
#> Degrees of Freedom for the fit: 4 Residual Deg. of Freedom 96
#> Global Deviance: 571.322
#>
             AIC:
                   579.322
#>
             SBC:
                    589.743
```

```
#the same task using the built-in intercept:
db <- data.frame(yt = print(sim))</pre>
#> -----
#> A PO-Garma(2,0) simulation object:
#>
\#> phi1 = 0.5
\#> phi2 = 0.15
#> beta.intercept = 1
\#> beta.x1 = 1
\#> mu0[1] = 2
\#> mu0[2] = 2
#> Number of Monte Carlo Simulations ('nmonte') = 1
#> Time Series Length ('nsteps') = 100
#> Burn-in ('burnin') = 1000
#> parallel = TRUE
#> -----
#>
set.seed(123)
fit <- garmaFit2(yt~. ,data= db,order = sim@order )
#> deviance of linear model= 637.7676
#> deviance of garma model= 571.322
summary(fit)
#>
#> Family: c("NO", "Normal")
#> Fitting method: "nlminb"
#>
#> Call: qarmaFit2(formula = yt ~ ., order = sim@order, data = db)
#>
#>
#> Coefficient(s):
#>
                   Estimate Std. Error t value Pr(>|t|)
#> beta.(Intercept) 19.0463703    1.5215935 12.51738 < 2.22e-16 ***
#> phi1
                 #> phi2
                  0.1976032 0.0973950 2.02889
                                                 0.04247 *
#> sigma
                  4.4636610 0.3188329 14.00000 < 2.22e-16 ***
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#> Degrees of Freedom for the fit: 4 Residual Deg. of Freedom
#> Global Deviance:
                     571.322
#>
             AIC:
                     579.322
#>
             SBC:
                     589.743
```

MBB of a general Garma model

In this section we deal with the MBB estimation of a *GARMA* process using the tsboot2 function from the **dboot** package. This is only a modified version of the tsboot function from the **boot** package, with similar arguments.

```
mu0 = c(2,2),
                  X = as.matrix(
                    data.frame(
                      intercept = rep(1,1102),
                      x1 = c(rep(7,100),
                             rep(2,1002)))))
sim <- GarmaSim(spec,</pre>
                nmonte = 1,
                nsteps = 100)
\#db \leftarrow data.frame(yt = print(sim))
db <- data.frame(cbind(yt = print(sim), intercept = rep(1,100)))</pre>
#> -----
#> A PO-Garma(2,0) simulation object:
#>
\#> phi1 = 0.5
\#> phi2 = 0.15
#> beta.intercept = 1
#> beta.x1 = 1
\#> mu0[1] = 2
\#> mu0[2] = 2
#> Number of Monte Carlo Simulations ('nmonte') = 1
#> Time Series Length ('nsteps') = 100
#> Burn-in ('burnin') = 1000
#> parallel = TRUE
#>
set.seed(123)
boot.function <- function(data, order, family) {</pre>
fit <- garmaFit2(yt~.-1,data = data ,order = order,
                  family = family)
return(fit$coef)
}
ord <- sim@order ; fam <- sim@spec@family
MBB <- tsboot2(db,
               statistic = boot.function,
               R = 10,
               1 = 5,
               order = ord,
               family = fam,
               export = "garmaFit2",
              package = "gamlss")
#> deviance of linear model= 655.8949
#> deviance of garma model= 568.2738
\#> Warning: <anonymous>: ... may be used in an incorrect context: 'statistic(ran.gen(ts.orig[inds,],n))
\#The\ original\ statistic\ (applied\ to\ the\ original\ series\ ):
MBB$t0
#> beta.intercept
                          phi1
                                           phi2
```

```
2.9929513
                       0.4841739
                                      0.2068798
#The resamples:
MBB$t
#>
             [,1]
                       [,2]
                                   [,3]
#>
    [1,] 3.037345 0.4416777 0.14509357
    [2,] 2.935912 0.3022353
                             0.12467772
   [3,] 3.114831 0.5144340 0.14757825
#> [4,] 2.981993 0.5308902 -0.03848191
#> [5,] 3.003604 0.5799550 -0.07059052
#> [6,] 2.982355 0.3891946 -0.04175624
#> [7,] 2.945518 0.3152008 0.07419220
#> [8,] 2.883092 0.4431917 0.12134359
#> [9,] 2.966544 0.4938093
                            0.06522629
#> [10,] 2.976808 0.2649635 0.28420021
#Quick statistic for the resamples:
apply(MBB$t,2, mean)
#> [1] 2.98280007 0.42755522 0.08114832
```

Here we only dealt with the *Poisson-GARMA* model, for all available families:

```
library(gamlss.dist)
?gamlss.family
```

References

Benjamin, Michael A., Rigby, Robert A. and Stasinopoulos, D. Mikis. 2003. Generalized Autoregressive Moving Average Models. Journal of the American Statistical Association. Mar, 2003, Vol. 98, 461, pp. 214-223.