# maxlogL: A general computational procedure for Maximum Likelihood estimation in R

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#### Abstract

Maximum likelihood (ML) method is preferred among others because it produces consistent and efficient estimators. However, likelihood optimization processes frequently involve unwieldy mathematical expressions and it is necessary in some cases to implement distributions and constantly build (log-)likelihood functions in computing languages in order to get numerical solutions. We present maxlogL, a function contained in the EstimationTools package for ML parameter estimation of any probability function loaded in R with no need of special structures, given a data set. We show via simulation that the routine has good performance in estimating the parameters of three distributions (normal, ZIP and user-defined). Finally, we present two application examples with real data.

Keywords: Maximum likelihood estimation, parameter estimation, R, EstimationTools.

#### 1. Introduction

Parameter estimation for probability density functions or probability mass functions is a central problem in statistical analysis and applied sciences because it allows to build predictive models and make inferences. Traditionally this problem has been tackled by means of likelihood maximization, as it was introduced by Fisher (1912). The method consists on performing an optimization through first derivative of the log-likelihood function and solve the outcoming system of equations. Despite its validity for any probability distribution, there exist a vast variety of them with cumbersome derivatives which produce non-linear systems of equations, therefore it is necessary to implement numerical methods and develop computational algorithms to find a solution.

R (R Core Team 2019) is a free language developed for statistical computing and equipped with unconstrained and box-constrained general-purpose optimization tools in **base** package: Fox, Hall, and Schryer (1978) developed the function nlminb for optimization using PORT (portable Fortran programs for numerical computation) routines; Nash (1979) implemented optim, a function that performs optimization based on three algorithms: (1) Nelder and Mead (1965), (2) quasi-Newton (BFGS) and (3) conjugate-gradient (CG). Either the former or the latter is implemented, users must take a distribution included in any package or create their own function otherwise and then write the likelihood.

On the other hand, R posses an extensive number of libraries (add-ons) in order to enhance its capabilities, e.g gamlss package (Stasinopoulos and Rigby 2007) for fitting generalized additive models for location, scale and shape. It is possible to carry out parameter estimation with

an empty regression model of any distribution implemented as a gamlss.family structure. Visit Stasinopoulos, Rigby, Heller, Voudouris, and De Bastiani (2017) for more details.

In this paper, we introduce the function maxlogL, which is capable of applying maximum likelihood estimation based on optimization through optim or nlminb only with the density/mass function defined as usual in R. The user could define its own distribution or use whichever existing in any package. The remainder of the article defines the maximization problem mathematically and computationally. Then, we present a simulation study to evaluate the performance of maxlogL with data generated from normal, ZIP and user-defined distributions. Finally, we give an application with a real data set and present some conclusions.

## 2. Maximum Likelihood estimation

Let be  $\mathbf{y}^{\top} = (y_1, y_2, ..., y_n)$  a random sample with n observations drawn from a population with distribution  $f(\cdot|\boldsymbol{\theta})$ , with  $\boldsymbol{\theta}$  a vector of parameters. The likelihood function of  $\boldsymbol{\theta}$  is

$$L(\boldsymbol{\theta}|\boldsymbol{y}) = \prod_{i=1}^{n} f(y_i|\boldsymbol{\theta}). \tag{1}$$

The method of ML finds the parameter values that makes data more probable. It is achieved by computing a vector  $\hat{\boldsymbol{\theta}}$  such that

$$\hat{\boldsymbol{\theta}} = \underset{\boldsymbol{\theta} \in \Theta}{\operatorname{arg\,max}} \ L(\boldsymbol{\theta}|\boldsymbol{y}). \tag{2}$$

It is usual to perform maximization of log-likelihood function, i.e.  $l(\theta|y) = \log L(\theta|y)$ . The variance-covariance matrix of ML estimators is given by

$$Var(\hat{\boldsymbol{\theta}}) = I^{-1}(\hat{\boldsymbol{\theta}}) = C(\hat{\boldsymbol{\theta}}), \tag{3}$$

where  $I(\hat{\theta})$  is the Fisher Information Matrix. The standard errors can be calculated as the square root of the diagonal elements of matrix C (Pawitan 2013)

$$S.E(\hat{\boldsymbol{\theta}}) = \sqrt{C_{jj}(\hat{\boldsymbol{\theta}})}.$$
 (4)

The R function presented here calculates  $l(\boldsymbol{\theta}|\boldsymbol{y})$  computationally, and computes standard errors from Hessian matrix.

# 3. Basic usage and features

Our maxlogL function is a S3 object of class maxlogL. It is included in **EstimationTools**, a package available from the Comprehensive R Archive Netwrok (CRAN) https://cran.r-project.org/package=EstimationTools. It can be downloaded and loaded in global environment typing the following instructions in the console:

R> install.packages("EstimationTools")
R> library(EstimationTools)

With maxlogL we provide a flexible implementation of ML estimation. It cab be executed stating its most important arguments

```
R> maxlogL(x, dist, optimizer, lower = NULL, upper = NULL)
```

where the argument x is a vector with data to be fitted, dist corresponds to the probability density/mass function of the working distribution, whereas upper and lower are limits used when user selects box-constrained algorithms. maxlogL is a wrapper function specifically developed for ML estimation, which allows to implement any of optim algorithms for optimization or nlminb routine for unconstrained or box-constrined optimization through the argument optimizer.

EstimationTools package provides a summary method for class maxlogL, which displays AIC (Akaike Information Criterion), BIC (Bayesian Information Criterion), ML estimates and its standard error. The method also reports the optimization routine selected by the user and the method used for computation of standard error. There are three methods available: hessian function from numDeriv package, calculation with optim (setting the argument hessian = TRUE and bootstrap calculation, with boot function of boot package (Davison and Hinkley 1997; Canty and Ripley 2017).

Hence, for non-censorship fitting the user must pass a vector with data and specify a probability distribution function available in R. For example, fitting a sample generated from a normal distribution,  $Z \sim NO(\mu = 10, \sigma^2 = 1)$ , could be done with the next command lines:

```
R> set.seed(1000)
R > z < -rnorm(n = 1000, mean = 10, sd = 1)
R > fit1 \leftarrow maxlogL(x = z, dist = 'dnorm', start=c(2, 3),
                 lower=c(-15, 0), upper=c(15, 10))
R> summary(fit1)
______
Optimization routine: nlminb
Standard Error calculation: Hessian from optim
      ATC
              BTC
 2804.033 2800.033
    Estimate Std. Error
      9.98752
                 0.0310
mean
      0.98126
                 0.0219
sd
```

The link argument is a list with entries fun and over, which specify the link functions applied and the name of linked parameters in probability function implemented in R respectively. The estimation performed above can be carried out applying logarithm link function to  $\sigma$ , avoiding problems of estimation in the boundary of parametric space. The usage is illustrated in the following code snippet:

```
R> fit2 <- maxlogL(x = z, dist = 'dnorm',</pre>
                link = list(over = "sd", fun = "log_link"))
R> summary(fit2)
Optimization routine: nlminb
Standard Error calculation: Hessian from optim
      AIC
             BIC
 2804.033 2800.033
_____
    Estimate Std. Error
mean 9.98752 0.0310
                0.0219
sd
      0.98126
The user can apply link functions to more than one parameter of the distribution:
R > fit3 < - maxlogL(x = z, dist = 'dnorm',
                link = list(over = c("mean", "sd"),
                           fun = c("log_link", "log_link")))
R> summary(fit3)
Optimization routine: nlminb
Standard Error calculation: Hessian from optim
      AIC
             BIC
 2804.033 2800.033
_____
    Estimate Std. Error
```

Other link functions available are logit and negative inverse, which must be specified as "logit\_link" and "NegInv\_link". On the other hand, maxlogL allows to define fixed known parameters, e.g., the sample size n in estimation of success proportion in a binomial distribution  $N \sim \mathrm{BI}(p=0.3,\,n=10)$ . This parameters can be specified with fixed argument, which is a list that stores the fixed parameters value specified by their names

mean 9.98752 0.0310

0.0219

0.98126

sd

```
Optimization routine: nlminb
Standard Error calculation: numDeriv::hessian

AIC BIC
334.9805 334.9805

Estimate Std. Error
prob 0.312 0.0147
```

As can be seen, the procedure applies the inverse of the link function to return the parameter to the original scale.

## 4. Simulation Study

In this section we simulated ML estimation with some distributions with maxlogL.

#### 4.1. Existing distributions

We simulated the following two types of random variables:  $X \sim \text{NO}(\mu = 5, \sigma^2 = 4)$  with sample sizes  $n \in \{3, ..., 400\}$  and  $Y \sim \text{ZIP}(\lambda = 6, \pi = 0.08)$  with  $n \in \{3, ..., 500\}$ . We performed 1000 replications of each sample size.

#### 4.2. User-defined distribution

Additionally, we simulated  $W \sim \text{EEB}(\mu = 0.5, \sigma = 1, \nu = 1.5, m = 10)$  with  $n \in \{3, ..., 1000\}$  (an exponentiated exponential binomial distribution, proposed by Bakouch, Ristić, Asgharzadeh, Esmaily, and Al-Zahrani (2012)). As the previous case, we performed 1000 replications of each sample size. Probability density function of EEB distribution is given by the following expression:

$$f(w|\mu, \sigma, \nu, m) = \frac{\sigma \nu \mu m}{1 - (1 - \mu)^m} \left( 1 - e^{-\sigma w} \right)^{\nu - 1} \left[ 1 - \mu \left( 1 - e^{-\sigma w} \right)^{\nu} \right]^{m - 1}, \tag{5}$$

with w > 0,  $\mu \in (0, 1)$ ,  $\sigma, \nu > 0$  and m an integer such that  $m \ge 1$ . We took this distribution as the user-defined by implementing it as an proglangR function, as is customary:

```
R> dEEB <- function(x, mu = 0.5, sigma = 1, nu = 1.5, size = 10, log = FALSE)\{
+ if (any(x<0))
+ stop(paste("x must greater than zero", "\n", ""))
+ if (any(mu < 0) \mid any(mu > 1))
+ stop(paste("mu must be between 0 and 1", "\n", ""))
+ if (any(sigma<=0))
+ stop(paste("sigma must be greater than zero", "\n", ""))
+ if (any(nu<=0))
```

```
# stop(paste("nu must be greater than zero", "\n", ""))
# 
# loglik <- log(sigma * nu * size * mu) - sigma * x +
# (nu - 1) * log(1 - exp(-sigma * x)) -
# log(1 - (1 - mu)^size) +
# (size - 1) * log(1 - mu * (1 - exp(-sigma * x))^nu)
# if (log == FALSE)
# density <- exp(loglik) else density <- loglik
# return(density)
# }</pre>
```

Note that argument size corresponds to parameter m in equation (5).

## 5. Results

The results of Monte Carlo study for the three distributions mentioned are presented in Figures 1, 2 and 3 below. All the plots show that the mean estimates tend to the true value of the parameter as the sample size increases, as is expected under regularity conditions.

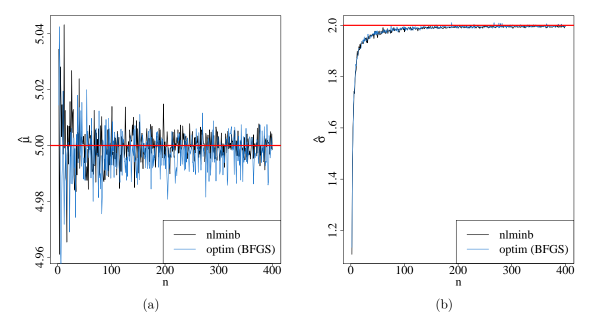


Figure 1: Mean value for (a) location parameter  $\hat{\mu}$  and (b) scale parameter  $\hat{\sigma}$  versus sample size n in normal distribution based on 1000 replications. Horizontal red lines are the true value of the parameters.

Variance of estimated parameters decreases as sample size increases, according to the efficiency property of ML estimators (Gurland 1954; Daniels 1961). This results are presented in figures 6, 7 and 8 of Appendix A.

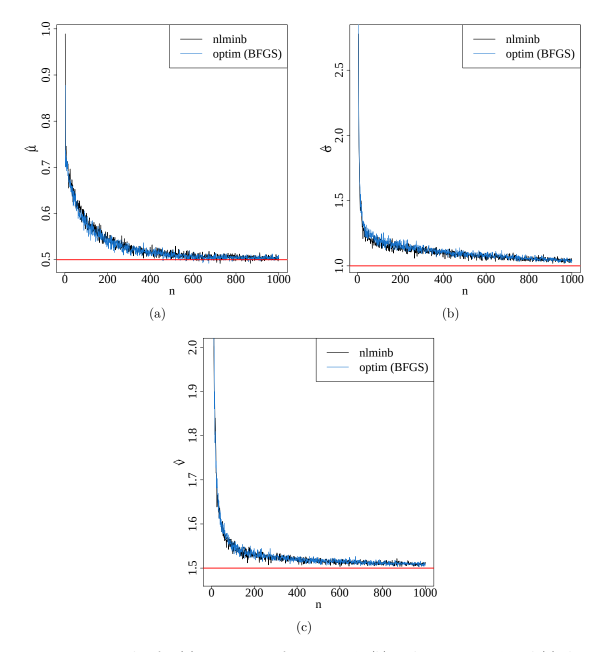


Figure 2: Mean value for (a) proportion of successes  $\hat{\mu}$ , (b) scale parameter  $\hat{\sigma}$  and (c) shape parameter  $\hat{\tau}$  versus sample size n in EEB distribution based on 1000 replications. Horizontal red lines are the true value of the parameters.

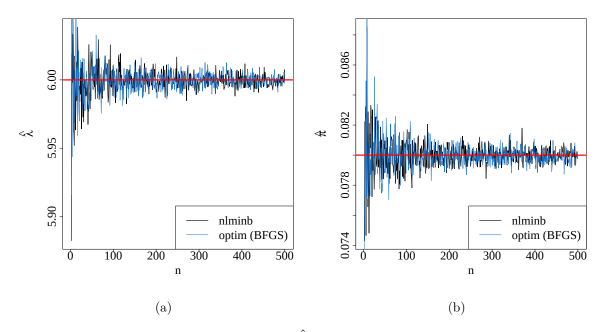


Figure 3: Mean value for (a) rate parameter  $\hat{\lambda}$  and (b) extra zeros proportion  $\hat{\pi}$  versus sample size n in ZIP distribution based on 1000 replications. Horizontal red lines are the true value of the parameters.

# 6. Illustrative examples

In the following examples we replicate maximum likelihood method with maxlogL in two applications: fitting of a power Lindley distribution to model tensile strength of carbon fibers and parameter estimation in two stage hierarchical model of retention proportions in memory tests.

## 6.1. Tensile strength data: power Lindley distribution

Data presented in Table 1 correspond to the tensile strength T (in GPa) of 69 specimens of carbon fiber tested under tension at gauge lengths of 20 mm.

1.312	1.314	1.479	1.552	1.700	1.803	1.861	1.865	1.944	1.958	1.966
1.997	2.006	2.027	2.055	2.063	2.098	2.14	2.179	2.224	2.240	2.253
2.270	2.272	2.274	2.301	2.301	2.359	2.382	2.382	2.426	2.434	2.435
2.478	2.490	2.511	2.514	2.535	2.554	2.566	2.57	2.586	2.629	2.633
2.642	2.648	2.684	2.697	2.726	2.770	2.773	2.800	2.809	2.818	2.821
2.848	2.88	2.954	3.012	3.067	3.084	3.090	3.096	3.128	3.233	3.433
3.585	3.585									

Table 1: Tensile strength of 69 fibers (Devendra and Rangaswamy 2013).

Ghitany, Al-Mutairi, Balakrishnan, and Al-Enezi (2013) fitted their power Lindley (PL) dis-

tribution

$$f(u|\mu,\sigma) = \frac{\mu\sigma^2}{\sigma+1}(1+u^{\mu})u^{\mu-1}e^{-\sigma u^{\mu}}, \quad u > 0, \ \mu,\sigma > 0.$$
 (6)

We implemented this density function in the R function dPL displayed below:

```
R> dPL <- function(x, mu, sigma, log=FALSE){</pre>
     if (any(x < 0))
       stop(paste("x must be positive", "\n", ""))
     if (any(mu \le 0))
       stop(paste("mu must be positive", "\n", ""))
     if (any(sigma <= 0))</pre>
       stop(paste("sigma must be positive", "\n", ""))
     loglik <- log(mu) + 2*log(sigma) - log(sigma+1) +</pre>
       log(1+(x^mu)) + (mu-1)*log(x) - sigma*(x^mu)
     if (log == FALSE)
       density <- exp(loglik)
     else density <- loglik
     return(density)
   }
```

Then, we estimate parameters with maxlogL taking the vector of strengths from data set fibers, as follows:

```
R> # Fitting of tensile strenght data
R> st <- Fibers$Strenght</pre>
R > theta <- maxlogL(x = st, dist = "dPL",
               link = list(over = c("mu", "sigma"),
                        fun = c("log_link", "log_link"))
R> summary(theta)
______
Optimization routine: nlminb
Standard Error calculation: Hessian from optim
_____
         BTC
    AIC
 102.119 98.119
    Estimate Std. Error
     3.867777
              0.3137
sigma 0.049671
               0.0160
```

Estimations are  $\hat{\mu} = 3.8678$  and  $\hat{\sigma} = 0.0497$ . Essentially, we get the same values computed by Ghitany et al. (2013). In Figure 4 we showed the performance of parameter estimation plotting corresponding density along with the histogram in the left panel and estimated survival function along with Kaplan-Meier estimator in the right panel.

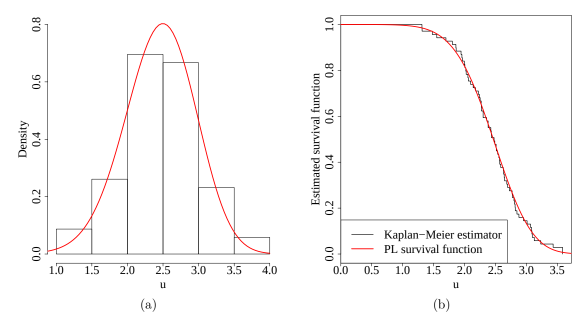


Figure 4: Fitting of tensile strength data: (a) estimated density and histogram; (b) estimated power Lindley survival function and Kaplan-Meier estimator.

#### 6.2. Forgetting data: hierarchical binomial distribution

maxlogL is capable of fit hierarchical models with the proper function definition of input variable. To illustrate the estimation, we replicated forgetting data example presented in Myung (2003), which used data from Murdock, Bennet B. (1961).

The retention function is a probability function that models the proportion of correct recall at time  $t_i$  in each trial in memory tests. Myung (2003) studied the following two models in their application example:

power model: 
$$p(a, b, t) = at^{-b}$$
,  
exponential model:  $p(a, b, t) = ae^{-bt}$ ,  $a, b > 0$ . (7)

m = 100 (number of trials)										
Retention Interval (sec.)	1	3	6	9	12	18				
Observed proportion	0.94	0.77	0.40	0.26	0.24	0.16				

Table 2: Observed proportion of recalls at each time.

Each observation in the data set corresponds to a proportion obtained as the quotient of correct responses  $(x_i)$  and the total number of independent trials (replications of each memory

test, represented by m). This kind of experiment can be modeled with a binomial distribution:

$$f(x_i|a,b,m) = \frac{m!}{(m-x_i)! x_i!} p(a,b,t_i) \left[1 - p(a,b,t_i)\right]^{m-x_i}$$
(8)

The usefulness of each retention equation is given by their goodness of fit.

#### Power model implementation

The hierarchical model with power retention function is implemented in an R function as follows:

Conditional density in equation (8) depends on  $x_i$ , but proportion of successes depends on t, as equation (7) shows. For this reason, the input argument  $\mathbf{x}$  must be a  $n \times 2$  matrix, where n is the sample size. In forgetting data, n=6. Then, the estimation is performed as usual with maxlogL. Note line six in the following chunk of code, which corresponds to the matrix definition of the input data aforementioned:

```
R> # Power model estimation
R> m <- 100 # Independent trials
R > t < c(1,3,6,9,12,18) # time intervals
R > p.ob < -c(0.94, 0.77, 0.40, 0.26, 0.24, 0.16) # Observed proportion
R> x <- p.ob*m # Correct responses
R > x <- as.integer(x)
R> Xi <- matrix(c(t,x), ncol=2, nrow=6)
R > retention.pwr < -maxlogL(x = Xi, dist = "power.logL", lower = c(0.01,0.01),
                          upper = c(1,1), start = c(0.1,0.1))
R> summary(retention.pwr)
______
Optimization routine: nlminb
Standard Error calculation: Hessian from optim
     AIC
             BTC
 57.4522 53.4522
 Estimate Std. Error
   0.95312
              0.0186
   0.49793
              0.0324
```

In this application, convergence was achieved by solving a box-constrained likelihood optimization, whose boundaries were specified in arguments lower and upper. Furthermore, tuning initial values was necessary with argument start. Computed values are  $\hat{\boldsymbol{\theta}} = (\hat{a}_{\text{pwr}}, \hat{b}_{\text{pwr}}) = (0.953, 0.498)$ , which are the same estimates of Myung (2003). Power fitting is illustrated in Figure 5.

### Exponential model implementation

Similarly as before, exponential retention function is implemented and input data is defined as a matrix, in this fashion:

```
R> # Exponential model implementation
R> exp.logL <- function(x, a, b, log = FALSE){</pre>
    p <- a * exp(-x[,1]*b)
    f \leftarrow dbinom(x = x[,2], size = m, prob = p)
    if (log == TRUE)
      density <- log(f) else density <- f
    return(density)
+ }
R> # Exponential model estimation
R> m <- 100 # Independent trials
R > t < c(1,3,6,9,12,18) # time intervals
R > p.ob < -c(0.94, 0.77, 0.40, 0.26, 0.24, 0.16) \# Observed proportion
R> x <- p.ob*m # Correct responses
R > x < -as.integer(x)
R> Xi <- matrix(c(t,x), ncol=2, nrow=6)
R> retention.exp <- maxlogL(x = Xi, dist = 'exp.logL', lower = c(0.01, 0.01),
                         upper = c(2,2), start = c(0.5,0.5))
R> summary(retention.exp)
______
Optimization routine: nlminb
Standard Error calculation: Hessian from optim
     AIC
 41.3329 37.3329
_____
 Estimate Std. Error
  1.07011
              0.0313
  0.13083
              0.0093
```

Again, we obtain the same estimates of Myung (2003):  $\hat{\theta} = (\hat{a}_{\text{exp}}, \hat{b}_{\text{exp}}) = (1.070, 0.131)$ . Exponential fitting is shown in Figure 5. According to the Akaike information criterion, the exponential model has better fitness ( $AIC_{\text{exp}} = 41.33$  against  $AIC_{\text{pwr}} = 57.45$ ).

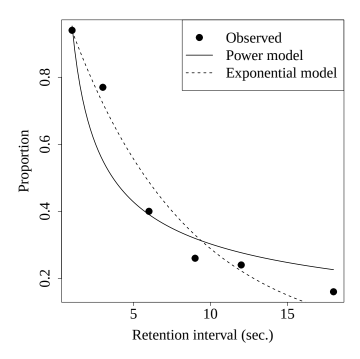


Figure 5: Observed proportion of recalls and models fitted.

#### 7. Conclusions

We have implemented classic estimation via maximization of log-likelihood function through basic optimization routines in R such as optim and nlminb. With maxlogL, we enable researchers, developers and users in general to compute ML estimators of any distribution in a fast and reliable way. With our summary method, it is possible to calculate standard error of estimates through Hessian matrix or bootstrap algorithm. In some cases, it is possible to implement estimation in hierarchical models, although Bayesian methods are preferred.

In future revisions, we could take our routine to develop further work on log-likelihood estimation, such as developing new parametric regression models.

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### A. Variance of estimates

In this section we report results of variance versus sample size of ML estimation with distributions fitted with maxlogL in section 5.

#### A.1. Normal distribution

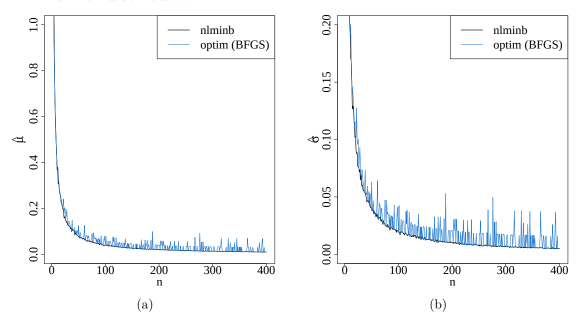


Figure 6: Variance for (a) location parameter  $\hat{\mu}$  and (b) scale parameter  $\hat{\sigma}$  versus sample size n in normal distribution based on 1000 replications..

## A.2. ZIP distribution

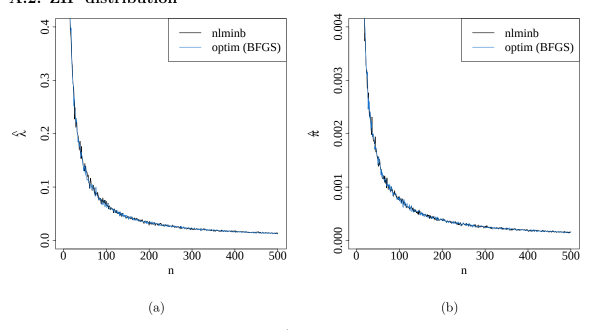


Figure 7: Variance for (a) rate parameter  $\hat{\lambda}$  and (b) extra zeros proportion  $\hat{\pi}$  versus sample size n in ZIP distribution based on 1000 replications.

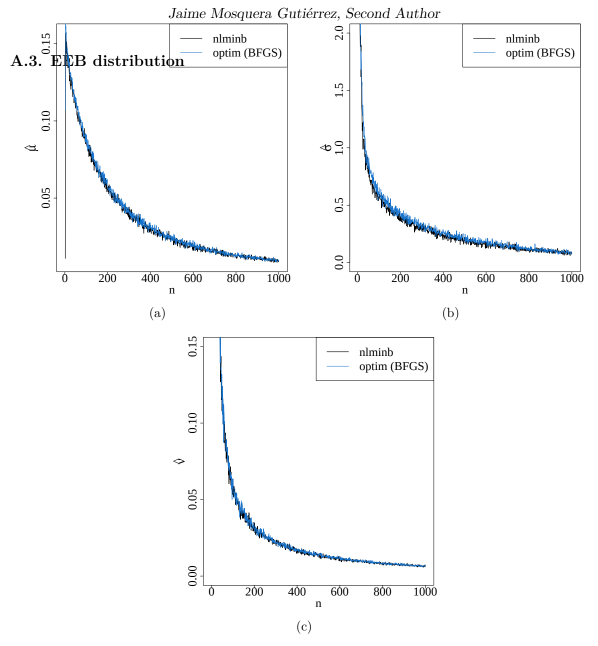


Figure 8: Variance for (a) proportion of successes  $\hat{\mu}$ , (b) scale parameter  $\hat{\sigma}$  and (c) shape parameter  $\hat{\tau}$  versus sample size n in EEB distribution based on 1000 replications.

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