# Use of the library fitdistrplus to specify a distribution from data

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Here you will find some easy examples of use of the functions of the library fitdistrplus. The aim is to show you by examples how to use these functions to help you to specify a parametric distribution from data corresponding to a random sample drawn from a theoretical distribution that you want to describe. For details, see the documentations of each function, using the R help command (ex.: ?fitdist). Do not forget to load the library using the function library before testing following examples.

#### > library(fitdistrplus)

### Contents

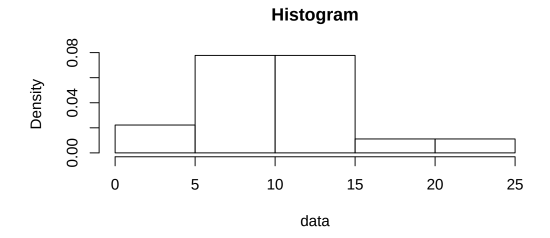
1	Specification of a distribution from non-censored continuous data	2
	1.1 Graphical display of the observed distribution	2
	1.2 Characterization of the observed distribution	
	1.3 Fitting of a distribution	4
	1.4 Simulation of the uncertainty by boostrap	
2	Specification of a distribution from non-censored discrete data	13
3	Specification of a distribution from censored data	17
	3.1 Graphical display of the observed distribution	17
	3.2 Fitting of a distribution	10

## 1 Specification of a distribution from non-censored continuous data

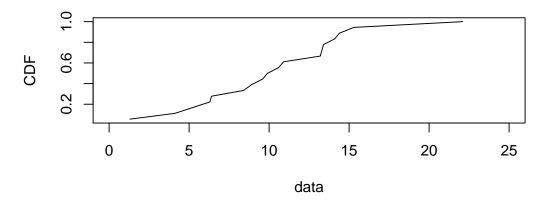
### 1.1 Graphical display of the observed distribution

First of all, the observed distribution may be plotted using the function plotdist.

```
> x1 <- c(6.4, 13.3, 4.1, 1.3, 14.1, 10.6, 9.9, 9.6, 15.3, 22.1,
+ 13.4, 13.2, 8.4, 6.3, 8.9, 5.2, 10.9, 14.4)
> plotdist(x1)
```



## **Cumulative distribution plot**



### 1.2 Characterization of the observed distribution

Descriptive parameters of the empirical distribution may be computed using the function descdist. This function will also provide by default a skewness-kurtosis plot which may help you to select which distribution(s) to fit among the potential candidates.

#### > descdist(x1)

summary statistics

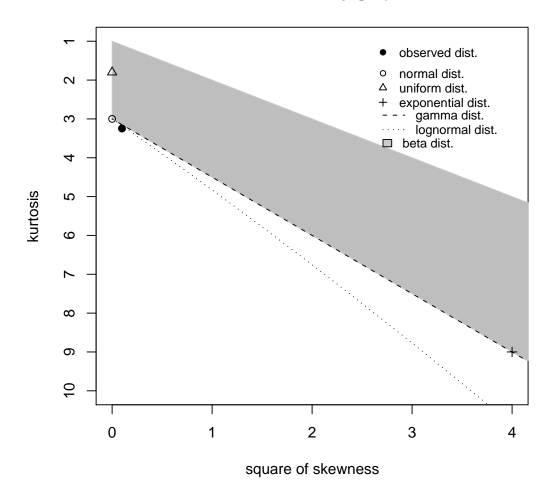
\_\_\_\_

min: 1.3 max: 22.1

median: 10.2 mean: 10.4 sample sd: 4.75

sample skewness: 0.314 sample kurtosis: 3.25

## **Cullen and Frey graph**



In order to take into account the uncertainty of the estimated values of kurtosis and skewness, the data set may be boostrapped by fixing the argument boot to an integer above 10 in descdist. boot values of skewness and kurtosis corresponding to the boot nonparametric bootstrap samples are then computed and reported in blue color on the skewness-kurtosis plot.

#### > descdist(x1, boot = 1000)

#### summary statistics

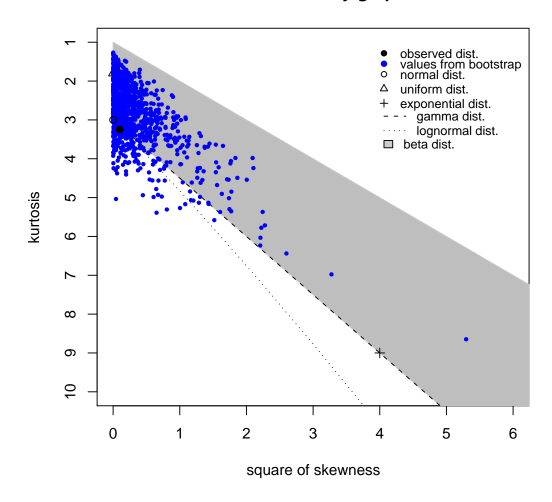
-----

min: 1.3 max: 22.1

median: 10.2 mean: 10.4 sample sd: 4.75

sample skewness: 0.314
sample kurtosis: 3.25

## **Cullen and Frey graph**



#### 1.3 Fitting of a distribution

One or more parametric distributions may then be fitted to the data set, one at a time, using the fonction fitdist. This function uses the maximum likelihood method if the argument method="mle" (or if it is omitted) or the matching moments method if the argument method="mom". When fitting continuous distributions, Kolmogorov-Smirnov and Anderson-Darling statistics are computed and corresponding tests are performed when possible. Even if less appropriate for continuous distributions, the Chi-squared statistic is also computed when possible. For this calculation, cells are defined by the argument chisqbreaks or automatically defined from the data set and from the argument meancount (the approximate mean count per cell) which is fixed to  $(4n)^{2/5}$  if omitted (with n the length of the data set). For more details, see the help of the function fitdist. Four goodness of fit plots are also provided.

Below is the result of a fit of a gamma distribution by maximum likelihood.

```
> f1g <- fitdist(x1, "gamma")</pre>
> plot(f1g)
> summary(f1g)
FITTING OF THE DISTRIBUTION ' gamma ' BY MAXIMUM LIKELIHOOD
PARAMETERS
      estimate Std. Error
         3.575
shape
                    1.140
         0.343
                    0.118
rate
Loglikelihood:
                -54.4
GOODNESS-OF-FIT STATISTICS
    _____ Chi-squared__
Chi-squared statistic: 7.93
Degree of freedom of the Chi-squared distribution: 3
Chi-squared p-value: 0.0475
```

```
!!! the p-value may be wrong with some theoretical counts < 5 !!!
```

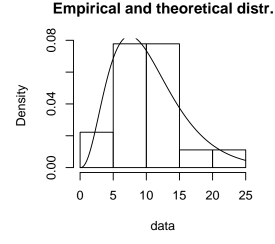
!!! For continuous distributions, Kolmogorov-Smirnov and Anderson-Darling statistics should be prefered !!!

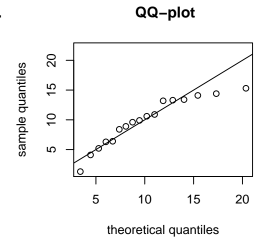
\_\_\_\_\_ Kolmogorov-Smirnov\_\_\_\_\_

Kolmogorov-Smirnov statistic: 0.138 Kolmogorov-Smirnov test: not calculated

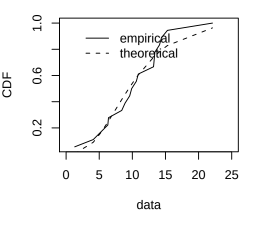
\_\_\_\_\_\_ Anderson-Darling\_\_\_\_\_

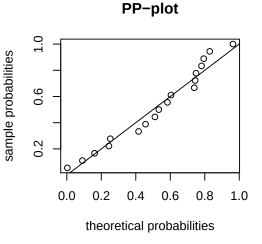
Anderson-Darling statistic: 0.457 Anderson-Darling test: not rejected





## **Empirical and theoretical CDFs**





Below is the result of another fit of the same distribution by matching moments.

- > f1gbis <- fitdist(x1, "gamma", method = "mom")</pre>
- > summary(f1gbis)

FITTING OF THE DISTRIBUTION ', gamma ', BY MATCHING MOMENTS PARAMETERS

estimate

shape 4.810 rate 0.462

----

GOODNESS-OF-FIT STATISTICS

\_\_\_\_\_\_ Chi-squared\_\_\_\_\_

Chi-squared statistic: 7.27

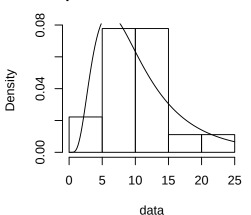
Degree of freedom of the Chi-squared distribution: 3

Chi-squared p-value: 0.0637

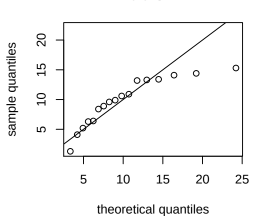
```
!!! the p-value may be wrong with some theoretical counts < 5 !!!
!!! For continuous distributions, Kolmogorov-Smirnov and
      Anderson-Darling statistics should be prefered !!!
_____Kolmogorov-Smirnov_____
Kolmogorov-Smirnov statistic: 0.144
Kolmogorov-Smirnov test: not calculated
_____ Anderson-Darling_____
Anderson-Darling statistic: 0.471
Anderson-Darling test: not rejected
  As can be seen in this returned summary, the automatic definition of the cells required to calculate the Chi-squared
statistic does not give theoretical counts large enough to validate the use of the test in this example. It is often the
case for small data sets. The observed and theoretical counts may be printed as below:
> f1g$chisqtable
       obscounts theocounts
<= 5.2 3.0000000 2.8950753
<= 8.4 3.0000000 4.5964712
<= 9.9 3.0000000 2.1080076
<= 13.2 3.0000000 3.7064776
<= 14.1 3.0000000 0.7577383
> 14.1 3.0000000 3.9362300
  Below is the fit of a lognormal distribution.
> f11 <- fitdist(x1, "lnorm")
> plot(f11)
> summary(f11)
FITTING OF THE DISTRIBUTION ' lnorm ' BY MAXIMUM LIKELIHOOD
PARAMETERS
      estimate Std. Error
meanlog 2.197
                   0.147
         0.622
sdlog
                     0.104
Loglikelihood: -56.5
GOODNESS-OF-FIT STATISTICS
______ Chi-squared_____
Chi-squared statistic: 11.1
Degree of freedom of the Chi-squared distribution: 3
Chi-squared p-value: 0.0110
!!! the p-value may be wrong with some theoretical counts < 5 !!!
\verb|!!!| For continuous distributions, Kolmogorov-Smirnov and
      Anderson-Darling statistics should be prefered !!!
 _____Kolmogorov-Smirnov_____
Kolmogorov-Smirnov statistic: 0.178
Kolmogorov-Smirnov test: not calculated
_____ Anderson-Darling_____
```

Anderson-Darling statistic: 0.793 Anderson-Darling test: rejected

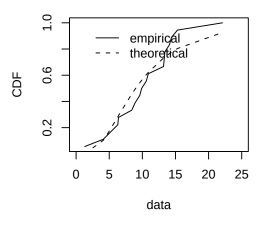




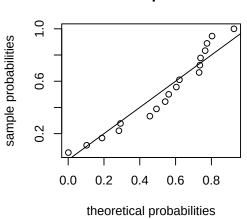
## QQ-plot



### **Empirical and theoretical CDFs**



### PP-plot



Below is the fit of a normal distribution.

- > f1n <- fitdist(x1, "norm")
- > plot(f1n)
- > summary(f1n)

FITTING OF THE DISTRIBUTION ' norm ' BY MAXIMUM LIKELIHOOD PARAMETERS

estimate Std. Error
mean 10.41 1.119
sd 4.75 0.791
Loglikelihood: -53.6

-----

GOODNESS-OF-FIT STATISTICS

\_\_\_\_\_\_ Chi-squared\_\_\_\_\_

Chi-squared statistic: 4.83

Degree of freedom of the Chi-squared distribution: 3

Chi-squared p-value: 0.185

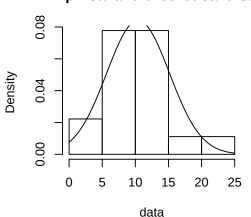
!!! the p-value may be wrong with some theoretical counts < 5 !!!

!!! For continuous distributions, Kolmogorov-Smirnov and Anderson-Darling statistics should be prefered !!!

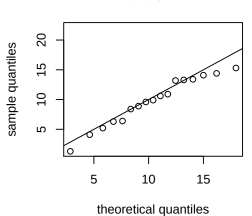
\_\_\_\_\_ Kolmogorov-Smirnov \_\_\_\_\_ Kolmogorov-Smirnov statistic: 0.110 Kolmogorov-Smirnov test: not calculated

Anderson-Darling

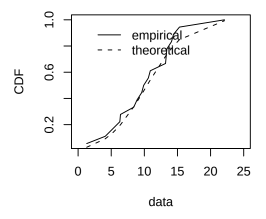
Anderson-Darling statistic: 0.226 Anderson-Darling test: not rejected



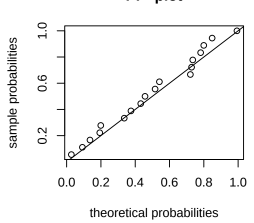
## QQ-plot



#### **Empirical and theoretical CDFs**



### PP-plot



Below is the fit of a Weibull distribution.

- > f1w <- fitdist(x1, "weibull")</pre>
- > plot(f1w)
- > summary(f1w)

FITTING OF THE DISTRIBUTION 'weibull 'BY MAXIMUM LIKELIHOOD PARAMETERS

 $\begin{array}{cccc} & \text{estimate Std. Error} \\ \text{shape} & 2.29 & 0.426 \\ \text{scale} & 11.70 & 1.264 \\ \text{Loglikelihood:} & -53.5 \end{array}$ 

-----

GOODNESS-OF-FIT STATISTICS

\_\_\_\_\_\_ Chi-squared\_\_\_\_\_

Chi-squared statistic: 5.87

Degree of freedom of the Chi-squared distribution: 3

Chi-squared p-value: 0.118

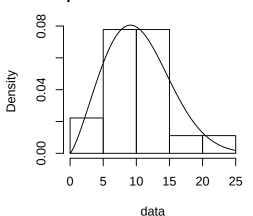
!!! the p-value may be wrong with some theoretical counts < 5 !!!

!!! For continuous distributions, Kolmogorov-Smirnov and Anderson-Darling statistics should be prefered !!!

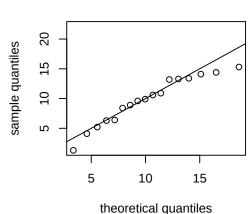
\_\_\_\_\_ Kolmogorov-Smirnov \_\_\_\_\_ Kolmogorov-Smirnov statistic: 0.121 Kolmogorov-Smirnov test: not calculated

Anderson-Darling

Anderson-Darling statistic: 0.282 Anderson-Darling test: not rejected

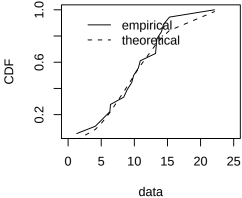


### QQ-plot

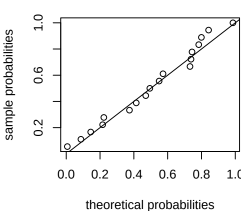


## **Empirical and theoretical CDFs**









The values of the Anderson-Darling statistic (or another result of the fit: see the help of fitdist for details) for the different fittings may be extracted and compared to help the selection of a distribution:

```
> anderson <- list(lnorm = f11$ad, gamma = f1g$ad, norm = f1n$ad,
```

+ weibull = f1w\$ad)

> anderson

#### \$lnorm

[1] 0.7925666

#### \$gamma

[1] 0.4567361

#### \$norm

[1] 0.225598

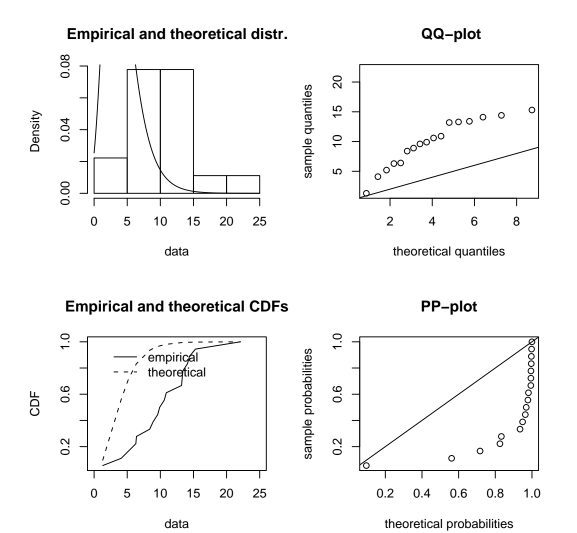
#### \$weibull

[1] 0.2821827

For some distributions (see the help of fitdist for details), it is necessary to specify initial values for the distribution parameters in the argument start when using the maximum likelihood method. start must be a named list of parameters initial values. The names of the parameters in start must correspond exactly to their definition in R or to their definition in a previous R code. The function plotdist may help to find correct initial values for the distribution parameters in non trivial cases, by an manual iterative use if necessary.

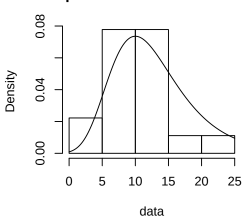
For example, below is the definition of the Gumbel distribution (also named extreme value distribution) and a first plot of the data set with the Gumbel distribution with arbitrary values for parameters.

```
> dgumbel \leftarrow function(x, a, b) 1/b * exp((a - x)/b) * exp(-exp((a - x)/b))
+ x)/b)
> pgumbel \leftarrow function(q, a, b) exp(-exp((a - q)/b))
> pgumbel \leftarrow function(p, a, b) a - b * log(-log(p))
> plotdist(x1, "gumbel", para = list(a = 3, b = 2))
```

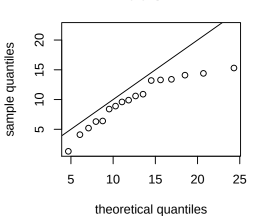


The same data set may be plotted with a Gumbel distribution with modified values for parameters.

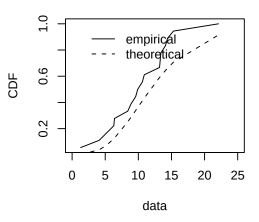
> plotdist(x1, "gumbel", para = list(a = 10, b = 5))



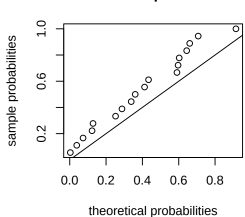
### QQ-plot



#### **Empirical and theoretical CDFs**



#### PP-plot



And a Gumbel distribution may be fitted to data with these values for initial parameter values.

- $> fgu \leftarrow fitdist(x1, "gumbel", start = list(a = 10, b = 5))$
- > plot(fgu)
- > summary(fgu)

FITTING OF THE DISTRIBUTION ' gumbel ' BY MAXIMUM LIKELIHOOD PARAMETERS

estimate Std. Error

a 8.09 1.092 b 4.38 0.766

Loglikelihood: -54.1

-----

GOODNESS-OF-FIT STATISTICS

\_\_\_\_\_ Chi-squared\_\_\_\_\_

Chi-squared statistic: 7.56

Degree of freedom of the Chi-squared distribution: 3

Chi-squared p-value: 0.056

!!! the p-value may be wrong with some theoretical counts < 5 !!!

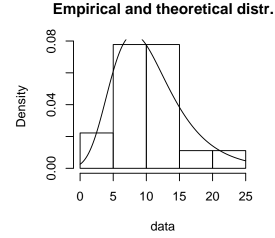
!!! For continuous distributions, Kolmogorov-Smirnov and Anderson-Darling statistics should be prefered !!!

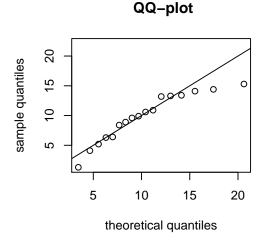
\_\_\_\_\_ Kolmogorov-Smirnov Statistic: 0 121

Kolmogorov-Smirnov statistic: 0.121 Kolmogorov-Smirnov test: not calculated

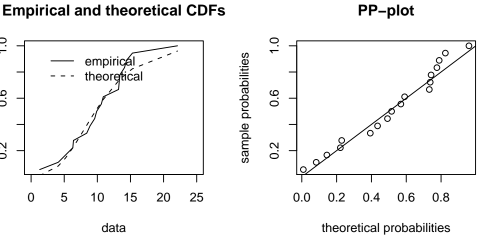
\_\_\_\_\_ Anderson-Darling\_\_\_\_\_

Anderson-Darling statistic: 0.34 Anderson-Darling test: not calculated





## empirical theoretical 9.0 CDF 0 5 10 15 20 25 data



#### Simulation of the uncertainty by boostrap

The uncertainty in the parameters of the fitted distribution may be simulated by parametric or nonparametric boostrap using the function boodist. This function returns the boostrapped values of parameters which may be plotted to visualize the bootstrap region. It also calculates the 95 percent confidence intervals for each parameter from the 2.5 and 97.5 percentiles of the boostrap values of each parameter (see the help of the function bootdist for details).

Below is an example of the use of this function with the previous fit of the gamma distribution.

- > b1g <- bootdist(f1g)
- > plot(b1g)
- > summary(b1g)

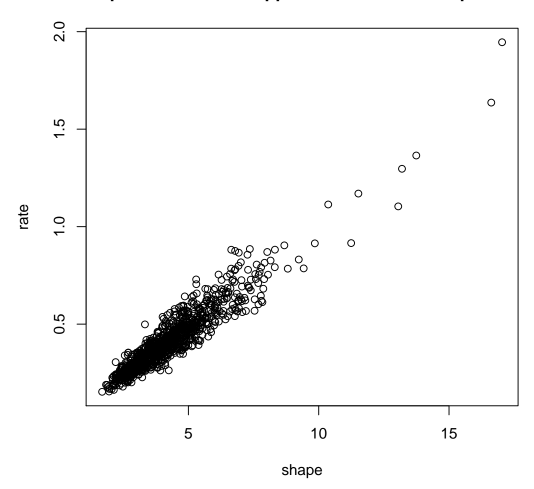
Parametric bootstrap medians and 95% CI  $\,$ 

Median 2.5% 97.5% 3.90 2.173 8.410 shape

0.38 0.204 0.842 rate

Maximum likelihood method converged for 999 among

## Scatterplot of the boostrapped values of the two parameters

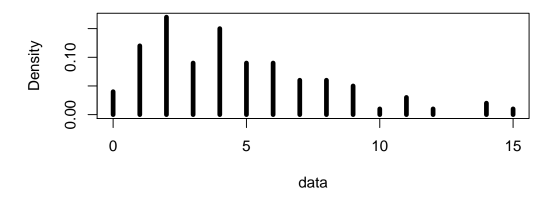


# 2 Specification of a distribution from non-censored discrete data

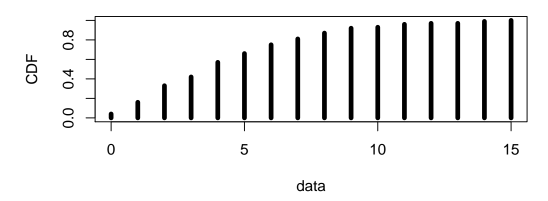
A discrete data set may be considered as a continuous one for example for a large data set from a binomial distribution converging to a normal one. A discrete plot of the distribution may also be provided, fixing the argument discrete of the function plotdist to TRUE.

```
> x2 <- rnbinom(n = 100, size = 2, prob = 0.3)
> plotdist(x2, discrete = TRUE)
```

## **Empirical distribution**



## **Empirical CDFs**



As for continuous distributions, descriptive parameters of the empirical distribution may be computed using the function descdist which also provides a skewness-kurtosis plot which may help you to choose which distribution(s) to fit.

#### > descdist(x2, discrete = T)

#### summary statistics

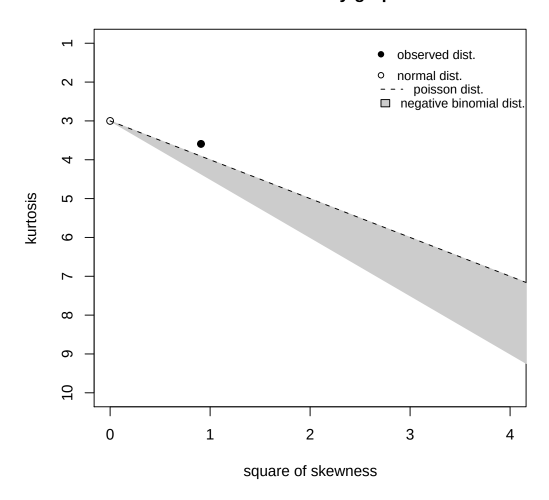
-----

min: 0 max: 15

median: 4
mean: 4.65
sample sd: 3.31

sample skewness: 0.953
sample kurtosis: 3.59

## **Cullen and Frey graph**



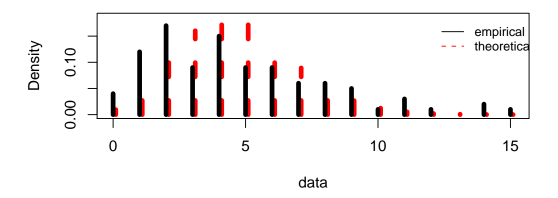
As for continuous distributions, one or more parametric distributions may then be fitted to the data set by maximum likelihood or matching moments.

Below is the result of the fit of a Poisson distribution with the bootstrap simulations.

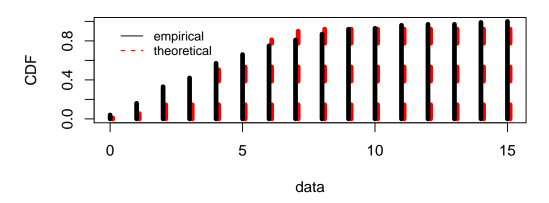
```
> f2p <- fitdist(x2, "pois")</pre>
> plot(f2p)
> summary(f2p)
FITTING OF THE DISTRIBUTION ' pois ' BY MAXIMUM LIKELIHOOD
PARAMETERS
       estimate Std. Error
lambda
           4.65
                     0.216
Loglikelihood: -273
GOODNESS-OF-FIT STATISTICS
_____ Chi-squared_
Chi-squared statistic: 48.9
Degree of freedom of the Chi-squared distribution: 6
Chi-squared p-value: 7.84e-09
!!! the p-value may be wrong with some theoretical counts < 5 !!!
> b2p <- bootdist(f2p)</pre>
> summary(b2p)
Parametric bootstrap medians and 95\% CI
     Median 2.5% 97.5%
2.5%
      4.64 4.23 5.08
```

Maximum likelihood method converged for 999 among 999

iterations



## **Empirical and theoretical CDFs**



Below is the result of the fit of a negative binomial distribution with the boostrap simulations.

```
> f2n <- fitdist(x2, "nbinom")</pre>
```

> summary(f2n)

FITTING OF THE DISTRIBUTION ' nbinom ' BY MAXIMUM LIKELIHOOD

PARAMETERS

estimate Std. Error size 3.38 0.856 mu 4.65 0.332

Loglikelihood: -249

----

GOODNESS-OF-FIT STATISTICS

\_\_\_\_\_\_ Chi-squared\_\_\_\_\_\_

Chi-squared statistic: 3.46

Degree of freedom of the Chi-squared distribution: 5

Chi-squared p-value: 0.63

> b2n <- bootdist(f2n)

> summary(b2n)

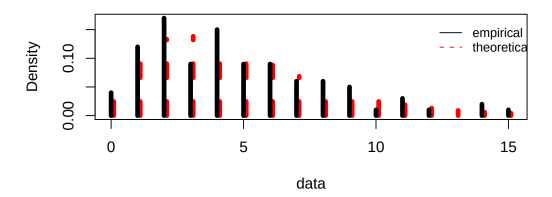
Parametric bootstrap medians and 95% CI

Median 2.5% 97.5% size 3.48 2.23 6.31

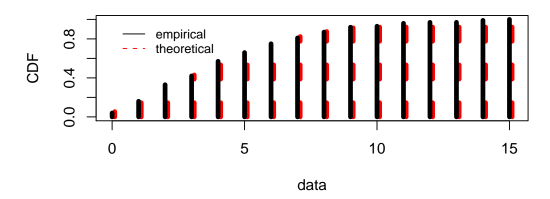
mu 4.63 4.01 5.29

Maximum likelihood method converged for 999 among 999 iterations

<sup>&</sup>gt; plot(f2n)



## **Empirical and theoretical CDFs**

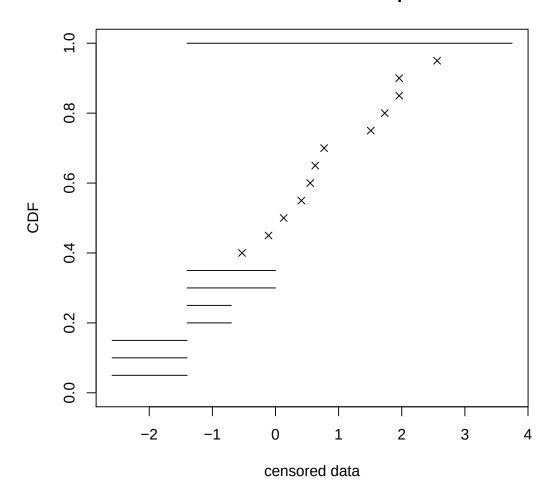


## 3 Specification of a distribution from censored data

Censored data may contain left censored, right censored and interval censored values, with several lower and upper bounds. Data must be coded into a dataframe with two columns, respectively named left and right, describing each observed value as an interval. The left column contains either NA for left censored observations, the left bound of the interval for interval censored observations, or the observed value for non-censored observations. The right column contains either NA for right censored observations, the right bound of the interval for interval censored observations, or the observed value for non-censored observations.

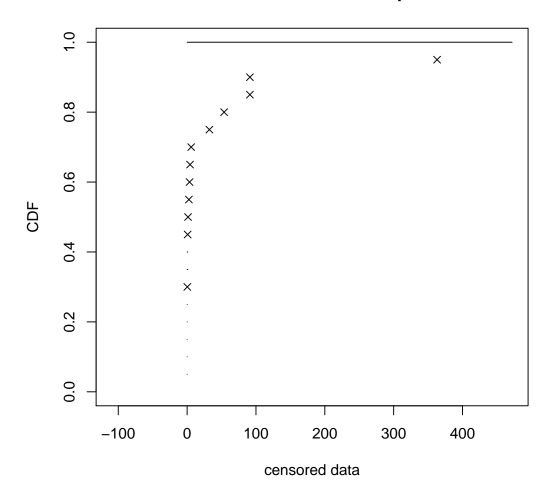
#### 3.1 Graphical display of the observed distribution

First of all, the observed distribution may be plotted using the function plotdistcens. Data are reported directly as segments for interval, left and right censored data, and as points for non-censored data. For more details, see the help of the function plotdistcens.



When left or right NA-values correspond to finite value (for example 0 for left NA-values of positive data), the arguments leftNA (or rightNA) must be affected to this finite value to ensure a correct plot of left (or right) censored observations, as in the example below.

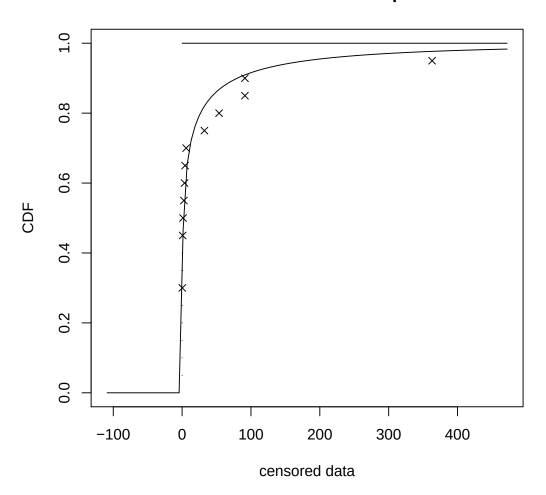
```
> d2 <- data.frame(left = 10^(d1\$left), right = 10^(d1\$right)) > plotdistcens(d2, leftNA = 0)
```



#### 3.2 Fitting of a distribution

One or more parametric distributions may then be fitted to the censored data set, one at a time, using the fonction fitdistcens. This function always uses the maximum likelihood method. For more details, see the help of the function fitdistcens. Only one goodness of fit plot is provided for censored data, in cumulative frequencies. The uncertainty in the parameters of the fitted distribution may be simulated by nonparametric boostrap only, using the function boodistcens

Below is the result of a fit of a Weibull distribution by maximum likelihood and the results of the corresponding boostrap simulations.



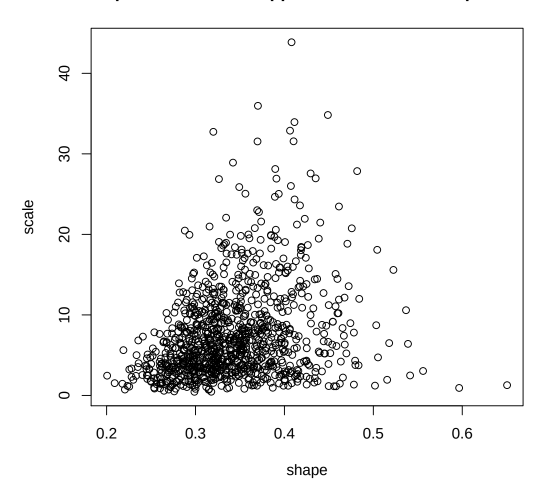
- > b2w <- bootdistcens(f2w)
- > summary(b2w)

> plot(b2w)

Nonparametric bootstrap medians and 95% CI Median 2.5% 97.5% shape 0.336 0.244 0.477 scale 6.004 1.255 21.896

Maximum likelihood method converged for 999 among 999 iterations

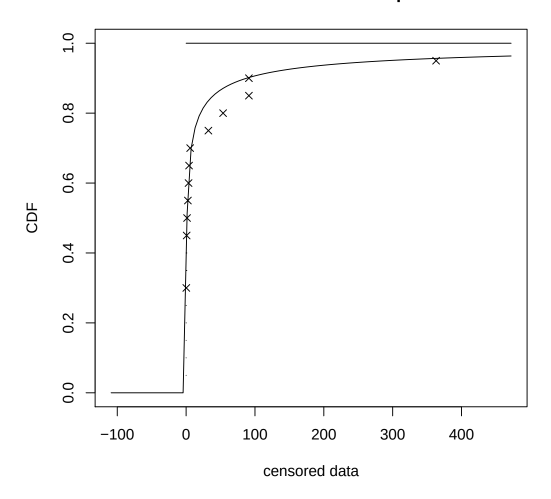
## Scatterplot of the boostrapped values of the two parameters



Goodness of fit statistics are not computed for fit on censored data, so the quality of fit may only be estimated from the loglikelihood and the goodness of fit plot.

Below is the fit of a lognormal distribution to the same censored data set.

> plot(f21, leftNA = 0)



Below is the fit of an exponential distribution.

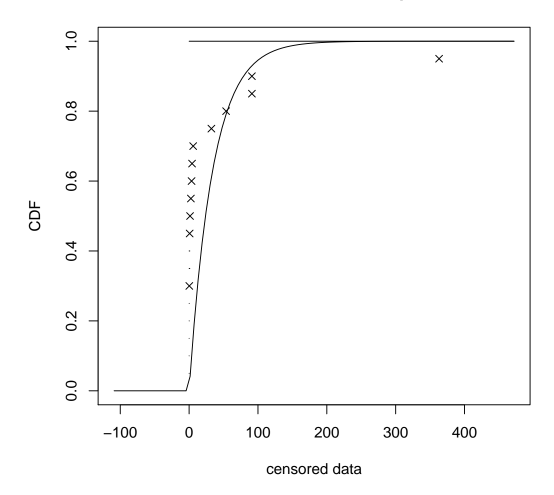
```
> f2e <- fitdistcens(d2, "exp")</pre>
```

> summary(f2e)

FITTING OF THE DISTRIBUTION '  $\exp$  ' BY MAXIMUM LIKELIHOOD ON CENSORED DATA PARAMETERS

estimate Std. Error rate 0.0292 0.00668 Loglikelihood: -99.6

> plot(f2e, leftNA = 0)



As with fitdist, for some distributions (see the help of fitdistcens for details), it is necessary to specify initial values for the distribution parameters in the argument start. start must be a named list of parameters initial values. The names of the parameters in start must correspond exactly to their definition in R or to their definition in a previous R code. The function plotdistcens may help to find correct initial values for the distribution parameters in non trivial cases, by an manual iterative use if necessary, as explained previously for non-censored continuous data.