

M3 Individual Claim Size Modelling

General Insurance Modelling : Actuarial Modelling III ¹

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17 May 2023

- 1 Introduction
- 2 Data analysis and descriptive statistics (MW 3.1)
- 3 Selected parametric claims size distributions (MW 3.2)
- 4 Fitting of distributions (MW3.2)
- 5 Model selection (MW 3.3)
- 6 ✕ Other advanced topics
- 7 Calculating within layers for claim sizes (MW 3.4)

- 1 Introduction
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Introduction

- How to fit loss models to insurance data?
- Peculiar characteristics of insurance data:
 - complete vs incomplete set of observations
 - left-truncated observations
 - right-censored observations
 - heavy tailed risks
- Parametric distribution models
 - model parameter estimation
 - judging quality of fit
 - model selection criteria
(graphical, score-based approaches, information criteria)
- Concepts and R functions are demonstrated with the help of some data sets

1 Introduction

- Steps in fitting loss models to data
- Insurance data
- Data set used for illustrations

Steps in fitting loss models to data

- ① explore and summarise data
 - graphical explorations
 - empirical moments and quantiles
- ② select a set of candidate distributions
 - Pareto, log-normal, inverse Gaussian, gamma, etc.
- ③ estimate the model parameters
 - method of moments
 - maximum likelihood (MLE)
 - maximum goodness (MGE)
- ④ evaluate the quality of a given model
 - graphical procedures (qq, pp plots, empirical cdf's)
 - score-based approaches (Kolmogorov-Smirnoff tests, AD tests, chi-square goodness-of-fit tests)
 - likelihood based information criteria (AIC, BIC)
- ⑤ determine which model to choose based on needs

1 Introduction

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Complete vs incomplete data

- complete, individual data
 - you observe the exact value of the loss
- incomplete data
 - exact data may not be available
 - in loss/claims data, these arise in the following situations:
 - ① observations may be grouped - observe only the range of values in which the data belongs
 - ② presence of censoring and/or truncation
 - ③ due to typical insurance and reinsurance arrangements such as deductibles and limits

Left-truncation and right-censoring

- left-truncated observation (e.g. excess / deductible)
 - observation is left-truncated at c if it is NOT recorded when it is below c and when it is above c , it is recorded at its exact value.
- right-censored observation (e.g. policy limit)
 - observation is right-censored at d if when it is above d it is recorded as being equal to d but when it is below d it is recorded at its observed value.
- similarly, we can define right-truncated, left-censored, ...
- of course, observations can be both left-truncated and right-censored; this is often the case in actuarial data

Zero claims

- Significant proportions of zero claims are frequent, for a number of reasons:
 - Data is policy-based, not claims-based;
 - Claim not exceeding deductible;
 - Mandatory reporting of accidents;
 - etc...
- This complicates the fitting (parametric distributions often don't have a flexible mass at 0, if at all)
- Several possible solutions
 - Adjust Y by mixing 0 with a parametric distribution
 - Adjust the frequency of claims accordingly (hence ignoring 0 claims), but this
 - may underestimate the volatility of claims (the proportion of 0's may also be random)
 - should be avoided if 0's are claims of no cost (rather than absence of claim)

Heavy tailed risks

- Essentially, these are risks that can be very large.
- This is translated by thick tails, that is, a density that goes to zero slowly in the tails.
- Typically, this means that the expectation of the excess over a threshold increases with that threshold.
- We will encounter such losses here, but a full treatment is deferred to Module 6 (Extreme Value Theory).

1 Introduction

- Steps in fitting loss models to data
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Data set used for illustrations

Bivariate data set of Swiss workers compensation medical and daily allowance costs:

- Example of real actuarial data; see Avanzi, Cassar, and Wong (2011)
- Data were provided by the SUVA (Swiss workers compensation insurer)
- Random sample of 5% of accident claims in construction sector with accident year 1999 (developped as of 2003)
- Claims are (joint) medical costs and daily allowance costs

```
SUVA <- read_excel("SUVA.xls")  
as_tibble(SUVA)
```

	medcosts	dailyallow
	407	0
	12591	13742
	269	0
	142	0
	175	0
	298	839
	47	0
	59	0
	191	7446
	159	0
	332	0
	226	21894
	453	1144
	182	410
	398	2989
	876	132
	462	4208
	3359	4687
	218	0

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Data analysis and descriptive statistics (MW 3.1)

It is essential, before any modelling is done, to make sure that one gets a good sense of what the data look like.

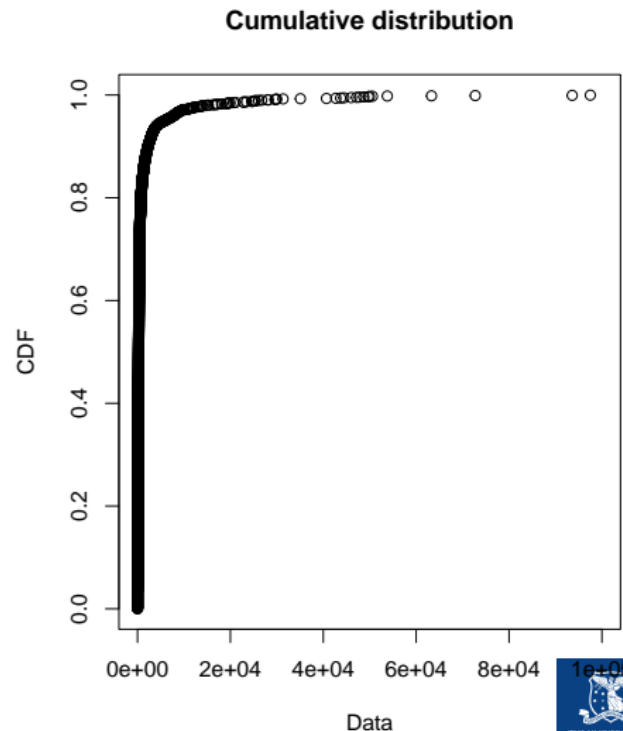
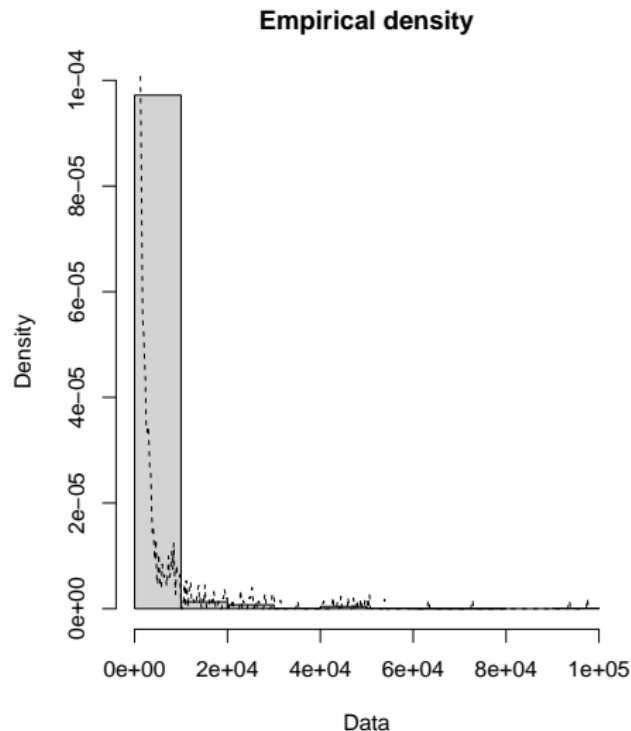
- For any type of data analysis, first thing to do is to summarise the data.
 - summary statistics: mean, median, standard deviation, coefficient of variation, skewness, quantiles, min, max, etc ...
 - gives a preliminary understanding of the data
- Visualising the data is often even more informative:
 - histogram, with associated kernel density
 - empirical cdf, which can be compared to that of a normal cdf via Q-Q or P-P plot
- When data are heavy tailed it often helps to perform the above on the log of the data (we can then compare the data to a lognormal)
- Data collection procedures and standards should be understood
- Any unusual feature (outliers, breaks, ...) should be investigated. If possible, ask the claims adjusters or data owners about them

2 Data analysis and descriptive statistics (MW 3.1)

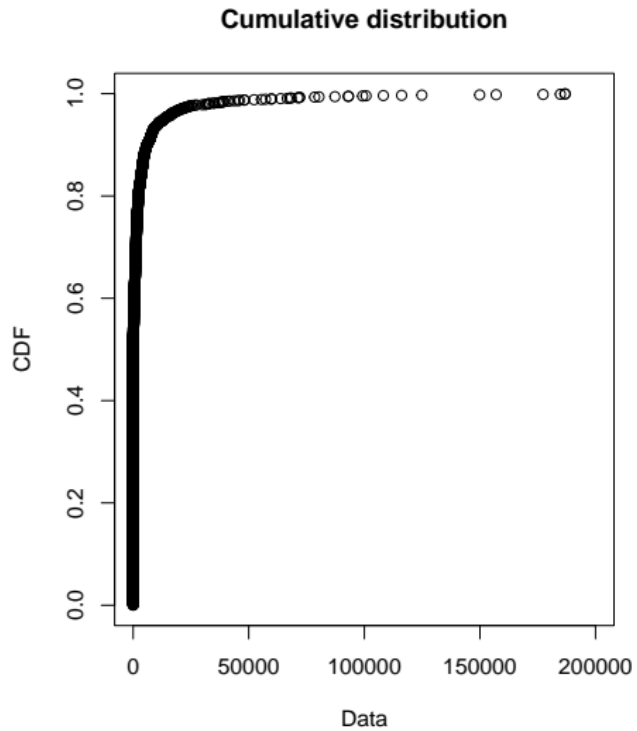
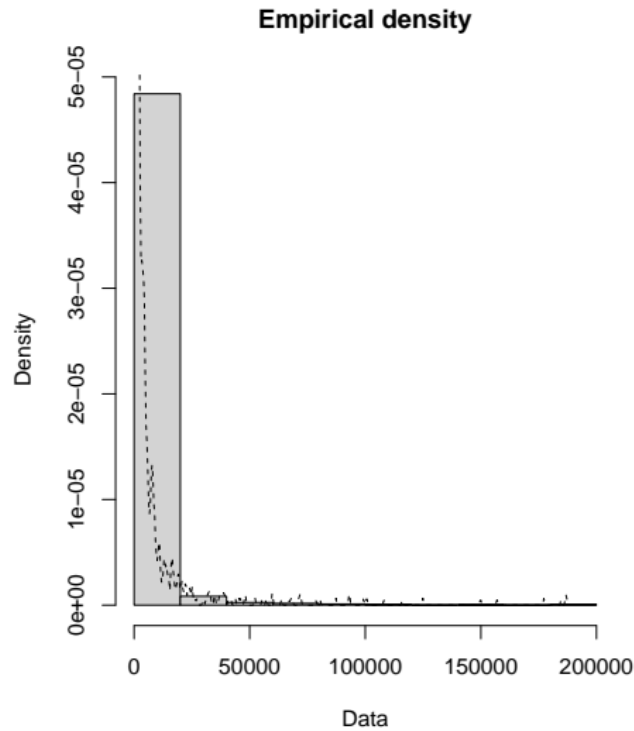
- Raw data
- Moments
- Quantiles
- Boxplots
- Log-log plots

Visualise the SUVA raw data

```
fitdistrplus::plotdist(SUVA$medcosts, histo = TRUE, demp = TRUE)
```

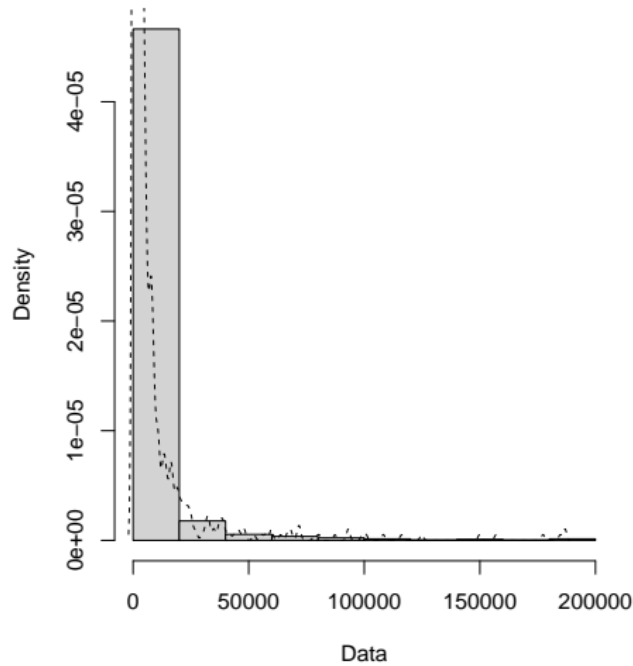


```
plotdist(SUVA$dailyyellow, histo = TRUE, demp = TRUE)
```

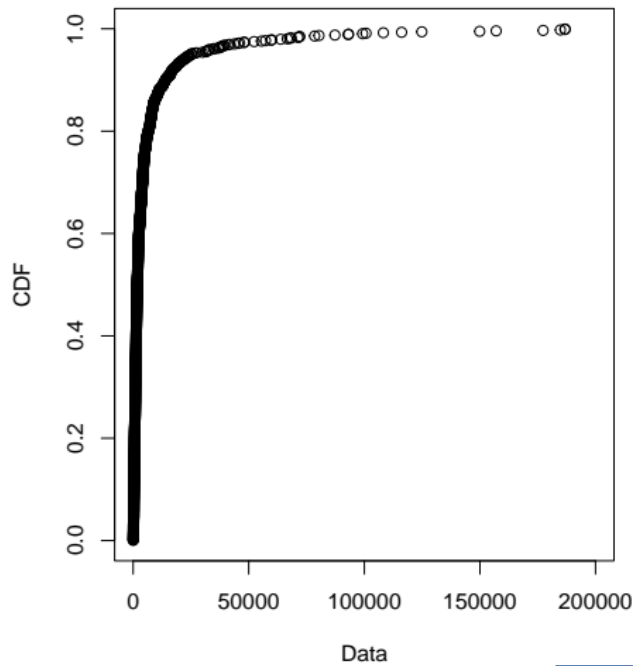


```
plotdist(SUVA$dailyallow[SUVA$dailyallow > 0], histo = TRUE,  
demp = TRUE)
```

Empirical density



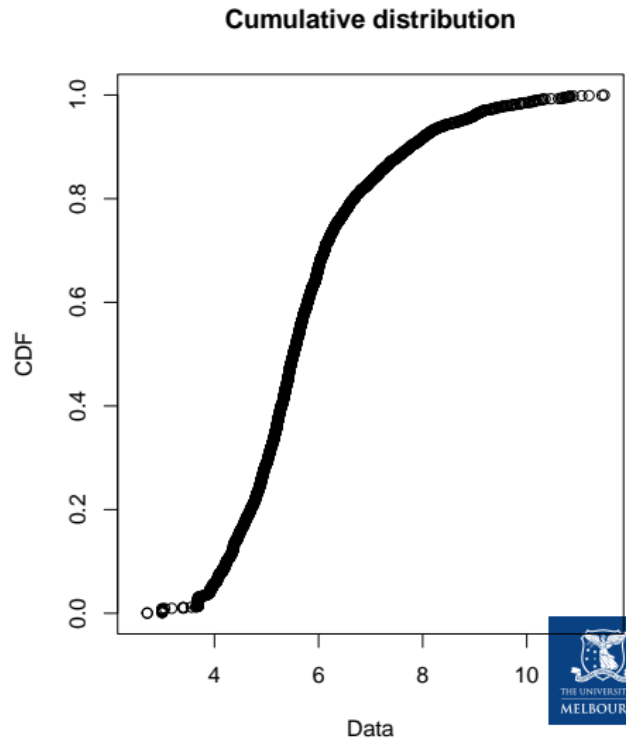
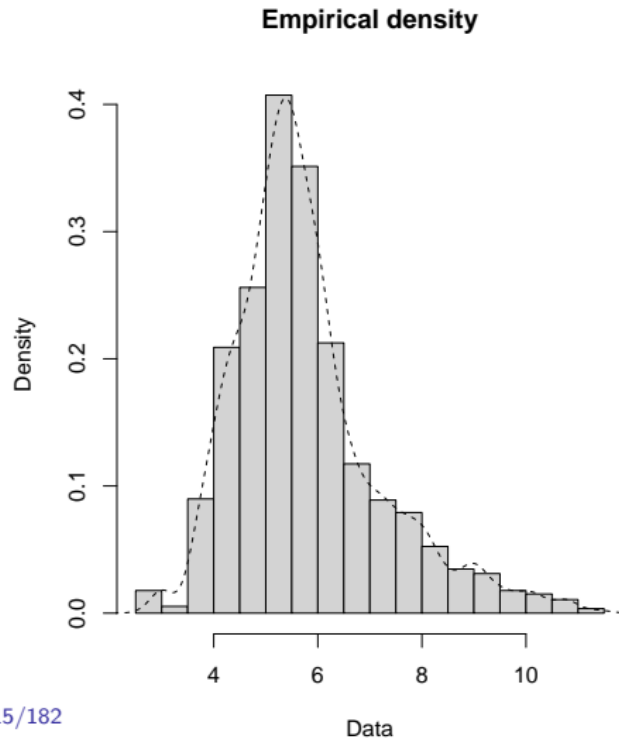
Cumulative distribution



A log transformation may help us see better what is happening.

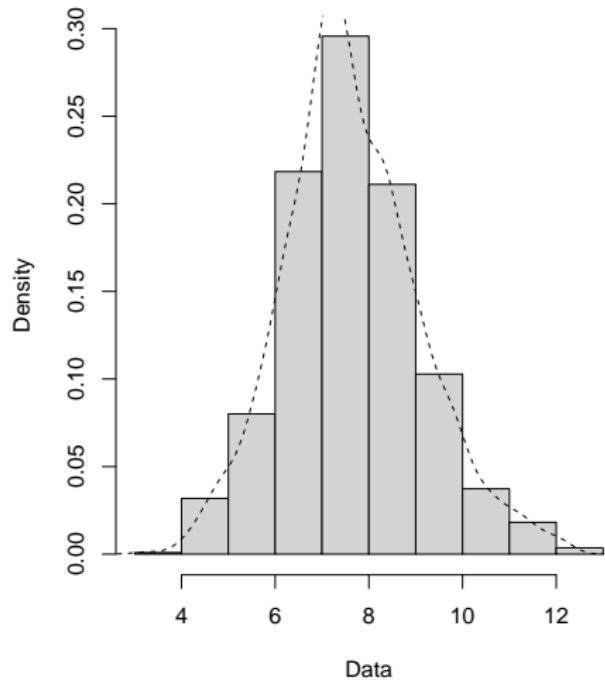
log of raw SUVA data

```
plotdist(log(SUVA$medcosts[SUVA$medcosts > 0]), histo = TRUE,  
demp = TRUE)
```

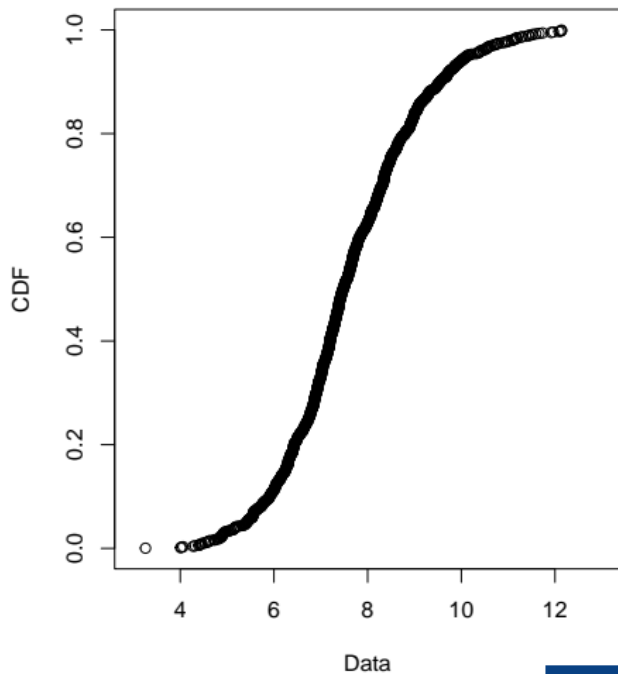


```
plotdist(log(SUVA$dailyyellow[SUVA$dailyyellow > 0]), histo = TRUE,  
demp = TRUE)
```

Empirical density



Cumulative distribution



- Medical costs are still skewed even after a log transformation, which suggests that a very heavy tailed distribution might be necessary.
- Daily allowance costs look symmetrical after the log transformation, which suggests a lognormal (or similar) distribution might be appropriate.
- Removing 0's is especially important for the daily allowance claims (and is necessary for taking the log anyway), as more than half of the claims are 0.

2 Data analysis and descriptive statistics (MW 3.1)

- Raw data
- Moments
- Quantiles
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Moments

Moments of a distribution provide information:

- The mean provides its location
- The second moment leads to the variance, and the coefficient of variation, which give an idea of the dispersion around the mean
- Skewness is self-explanatory
- Kurtosis provides an idea of how fat the tails are
 - “Excess Kurtosis” is with respect to a normal/lognormal distribution for raw/log claims, respectively

The following are also helpful:

- Loss size index function

$$\mathcal{I}(G(y)) = \frac{\int_0^y zdG(z)}{\int_0^\infty zdG(z)} \quad \text{and} \quad \widehat{\mathcal{I}}_n(\alpha) = \frac{\sum_{i=1}^{\lfloor n\alpha \rfloor} Y_{(i)}}{\sum_{i=1}^n Y_i}$$

for $\alpha \in [0, 1]$. Corresponds of the contribution of $[0, y]$ to the overall mean. (see also [Pareto principle](#), whereby 80% of overall cost would be due to 20% of (the most costly) claims)

- Mean excess function

$$e(u) = E[Y_i - u | Y_i > u] \quad \text{and} \quad \widehat{e}_n(u) = \frac{\sum_{i=1}^n (Y_i - u) \mathbf{1}_{\{Y_i > u\}}}{\sum_{i=1}^n \mathbf{1}_{\{Y_i > u\}}}$$

This is useful for the analysis of large claims, and for the analysis of reinsurance. Increasing values of the mean excess function indicate a heavy tailed distribution (see also Module 6) [Note $u = 0$ leads to the mean only when all claims are strictly positive.]

In R:

- To get numbers:
 - the function `actuar::emm` provides empirical moments up to any order
 - `mean`, `stats::var` and `stats::sd` provide mean, variance, and standard deviation (unbiased versions)
 - codes for the Loss size index function are provided in the illustration
 - codes for the Mean excess function are also provided, but the graph is most easily plotted with `extRemes::mrlplot` as will be demonstrated
- The function `fistdistrplus::descdist` provides a graph that shows where the couple "skewness/kurtosis" lies, in comparison with its theoretically possible locations for a certain number of distributions.
 - the parameter `boot` allows for nonparametric bootstrapping of that coordinate, which helps with the assessment of its potential variability (it is sensitive to outliers, that is, not "robust")
 - `method` can be "unbiased" or "sample" for the unbiased or sample versions of the moments

SUVA moments

Medical costs:

```
format(actuar::emm(SUVA$medcosts, order = 1:3), scientific = FALSE)

## [1] "1443.349" "34268506.007" "1791560934502.502"

sd(SUVA$medcosts)/mean(SUVA$medcosts)

## [1] 3.93143
```

Daily allowance:

```
format(actuar::emm(SUVA$dailyallow, order = 1:3), scientific = FALSE)

## [1] "3194.15" "172677852.63" "20364647975482.07"

sd(SUVA$dailyallow)/mean(SUVA$dailyallow)

## [1] 3.991459
```

Medical costs:

```
format(actuar::emm(SUVA$medcosts[SUVA$medcosts > 0], order = 1:3),  
       scientific = FALSE)  
  
## [1] "1492.765" "35441771.887" "1852899392464.571"  
  
sd(SUVA$medcosts[SUVA$medcosts > 0])/mean(SUVA$medcosts[SUVA$medcosts >  
0])  
  
## [1] 3.861552
```

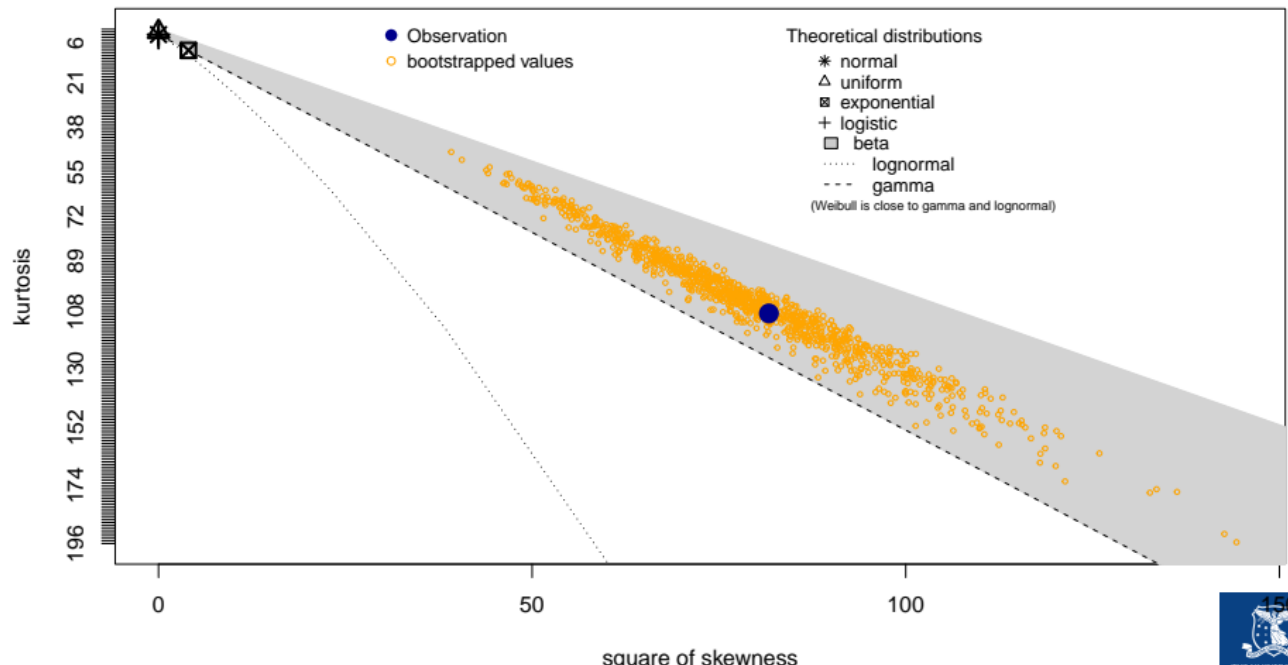
Daily allowance:

```
format(actuar::emm(SUVA$dailyallow[SUVA$dailyallow > 0], order = 1:3),  
       scientific = FALSE)  
  
## [1] "6760.322" "365467411.472" "43101156679682.695"  
  
sd(SUVA$dailyallow[SUVA$dailyallow > 0])/mean(SUVA$dailyallow[SUVA$dailyallow  
0])  
  
## [1] 2.646343
```

SUVA Medical Costs skewness and kurtosis

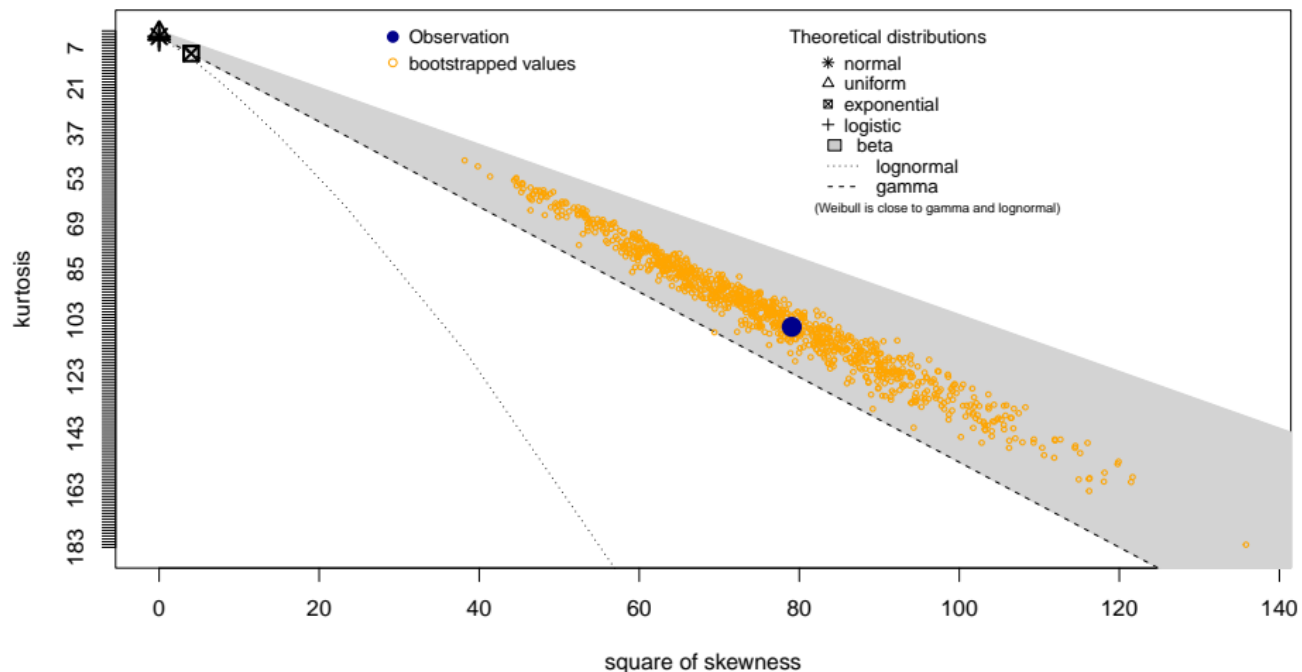
```
fitdistrplus::descdist(SUVA$medcosts, boot = 1000)
```

Cullen and Frey graph



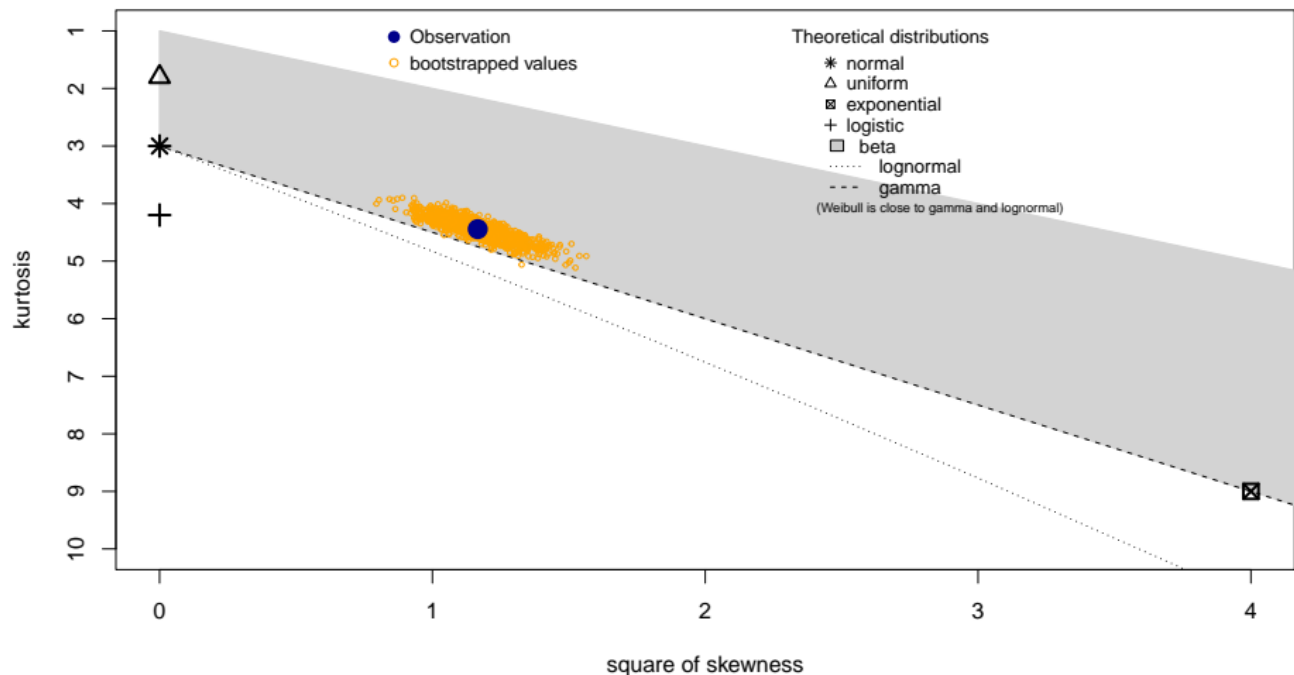
```
descdist(SUVA$medcosts [SUVA$medcosts > 0] , boot = 1000)
```

Cullen and Frey graph



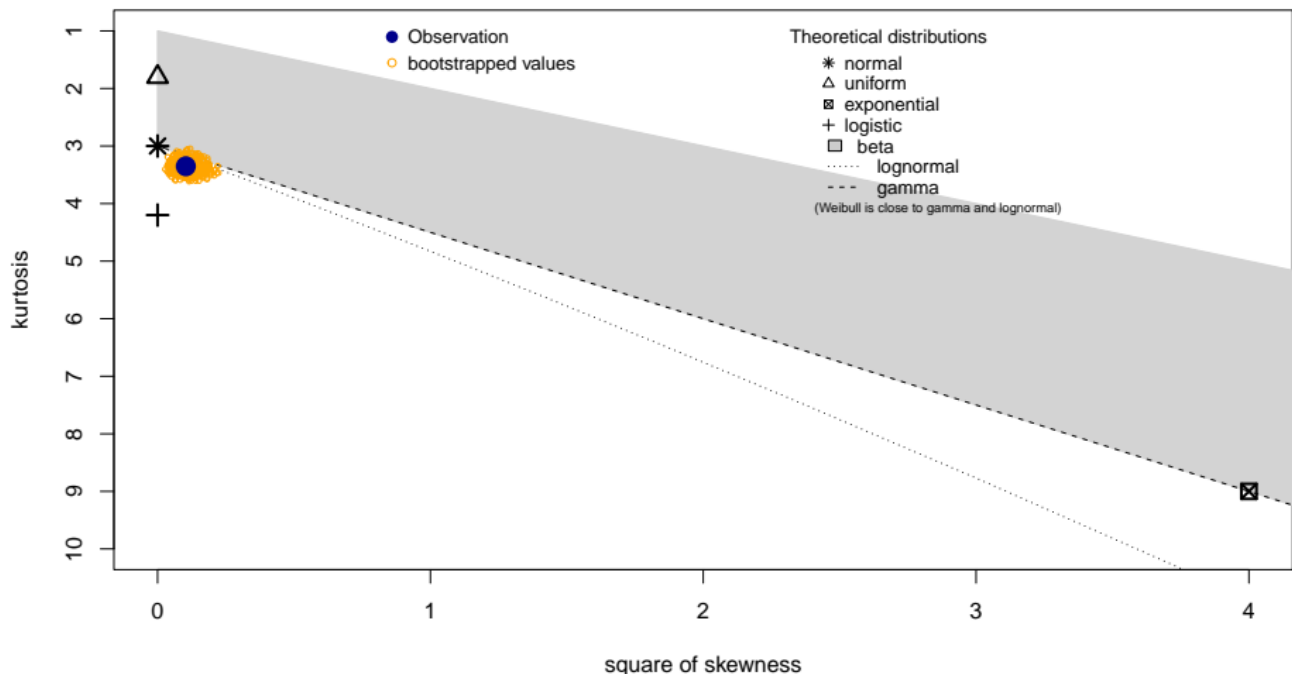
```
descdist(log(SUVA$medcosts[SUVA$medcosts > 0]), boot = 1000)
```

Cullen and Frey graph



```
descdist(log(log(SUVA$medcosts [SUVA$medcosts > 0])), boot = 1000)
```

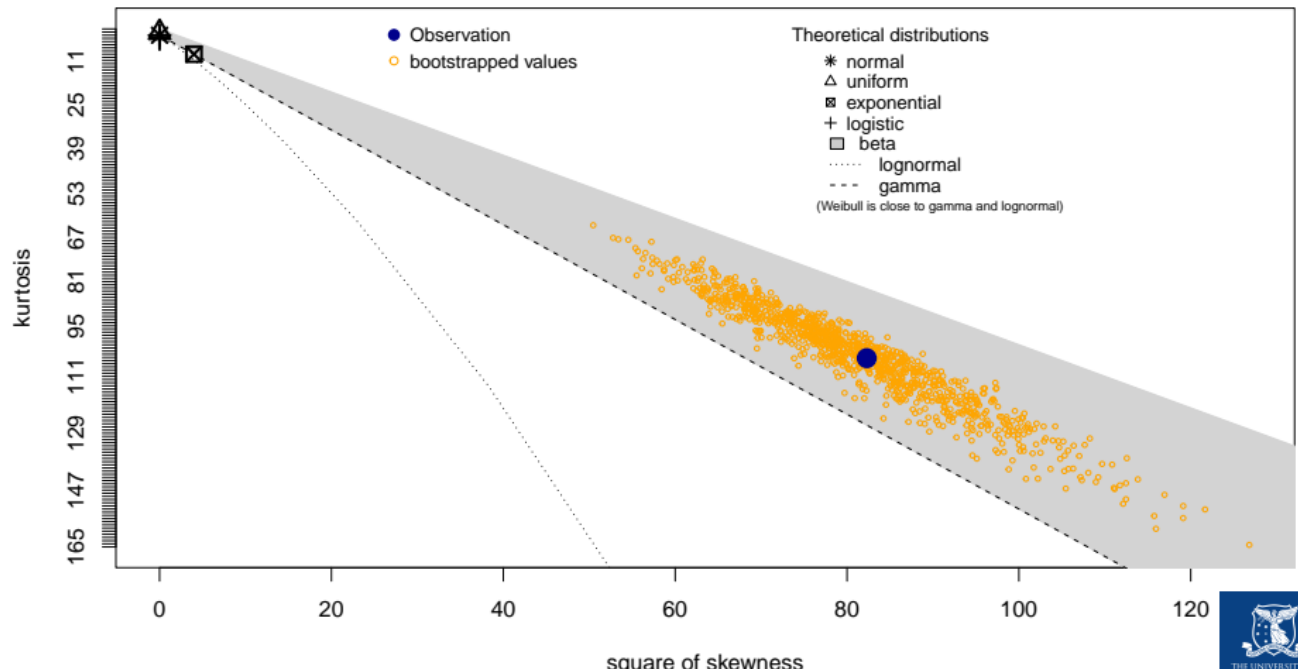
Cullen and Frey graph



SUVA Daily Allowance skewness and kurtosis

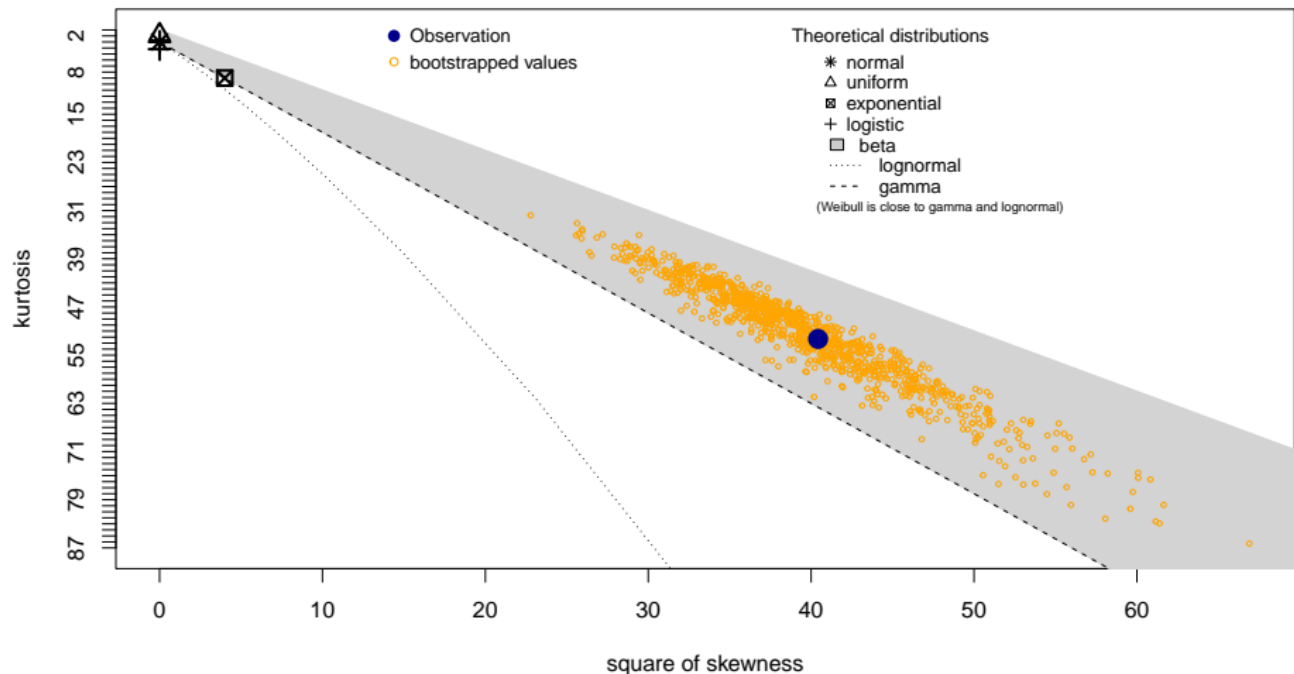
```
descdist(SUVA$dailyallow, boot = 1000)
```

Cullen and Frey graph



```
descdist(SUVA$dailyyellow[SUVA$dailyyellow > 0], boot = 1000)
```

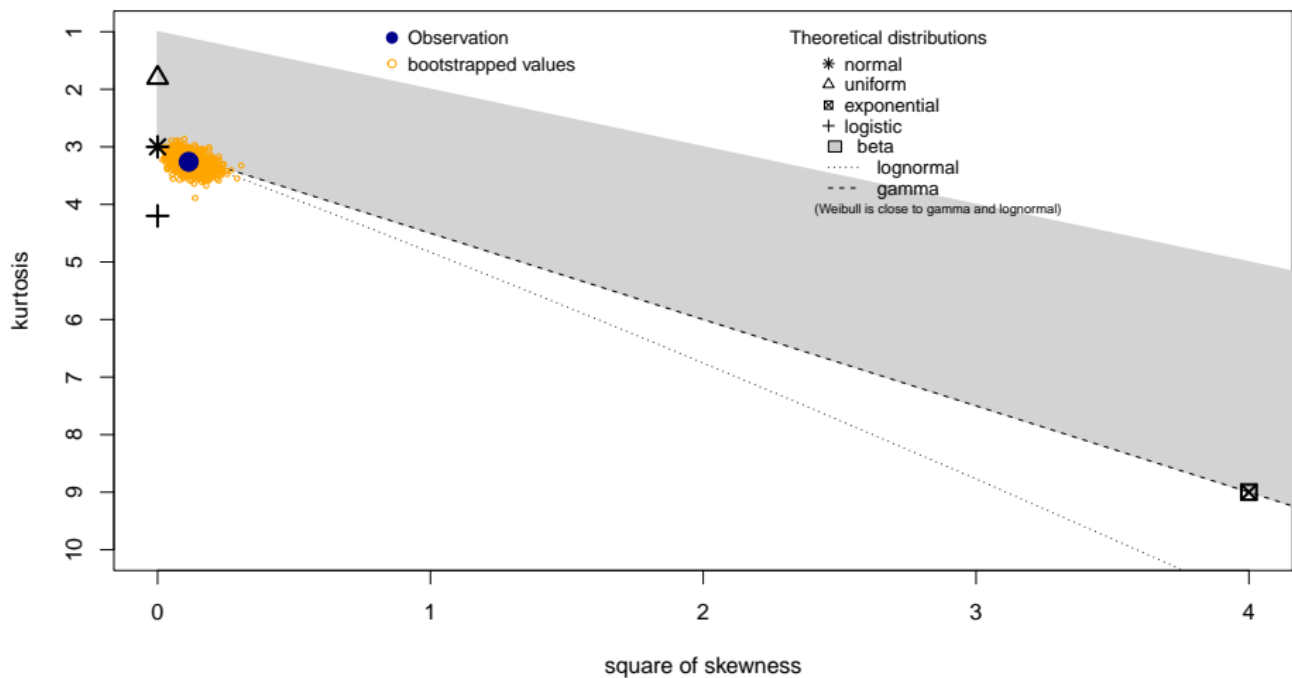
Cullen and Frey graph



```
descdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), boot = 1000)

## summary statistics
## -----
## min: 3.258097 max: 12.13806
## median: 7.474772
## mean: 7.63051
## estimated sd: 1.441014
## estimated skewness: 0.3378648
## estimated kurtosis: 3.259708
```

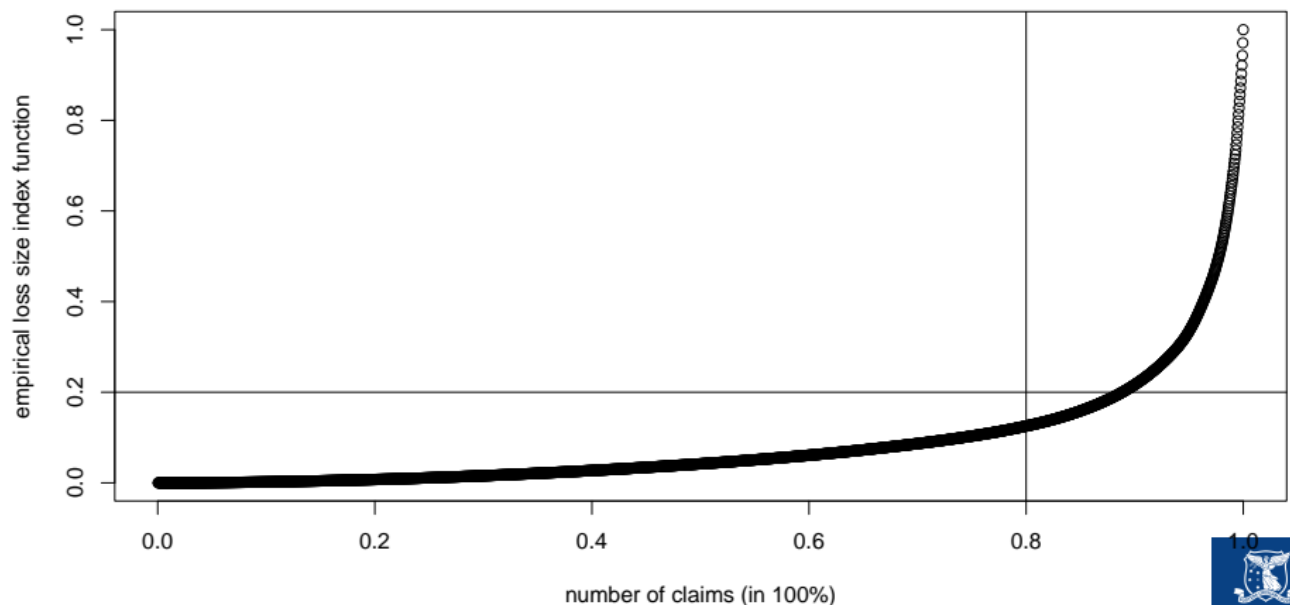
Note `descdist` also gives stats as above! (not shown on the previous slides)

Cullen and Frey graph

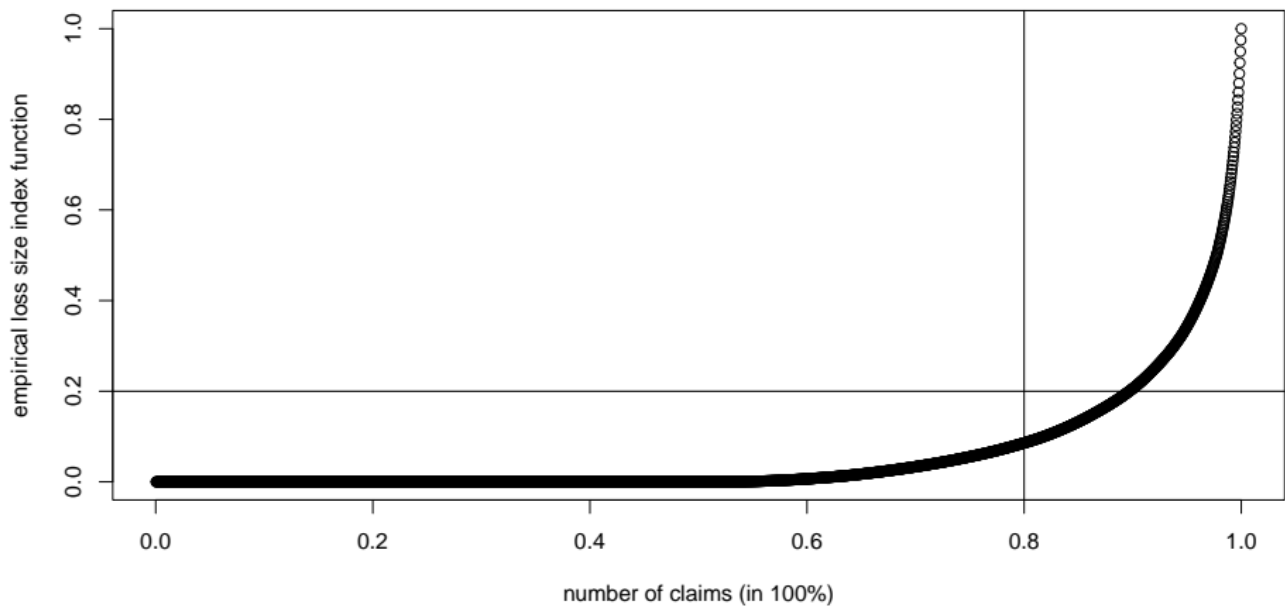
Good candidates seem to be lognormal, gamma, and potentially Weibull.

Loss index function

```
SUVA.MC.lif <- cumsum(sort(SUVA$medcosts))/sum(SUVA$medcosts)
plot(1:length(SUVA$medcosts)/length(SUVA$medcosts), SUVA.MC.lif,
     xlab = "number of claims (in 100%)", ylab = "empirical loss size index function",
     abline(h = 0.2, v = 0.8)
```



```
SUVA.DA.lif <- cumsum(sort(SUVA$dailyallow))/sum(SUVA$dailyallow)
plot(1:length(SUVA$dailyallow)/length(SUVA$dailyallow), SUVA.DA.lif,
     xlab = "number of claims (in 100%)", ylab = "empirical loss size index function",
     abline(h = 0.2, v = 0.8)
```



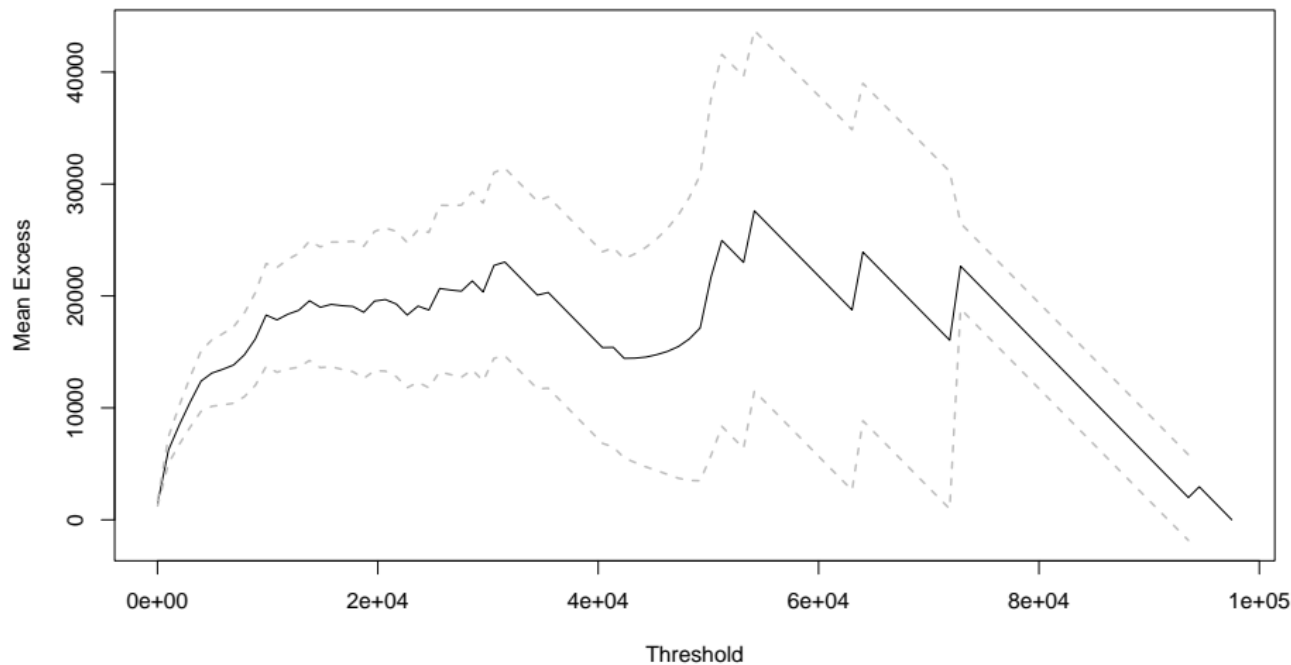
Mean excess function

This function will return the mean excess function for an arbitrary vector of thresholds u (for instance, 0, 100, 1000 and 10000 here)

```
mef <- function(x, u) {  
  mefvector <- c()  
  for (i in u) {  
    mefvector <- c(mefvector, sum(pmax(sort(x) - i, 0))/length(x[x >  
      i]))  
  }  
  return(mefvector)  
}  
mef(SUVA$medcosts, c(0, 100, 1000, 10000))  
## [1] 1492.765 1709.148 6233.296 18156.631  
mean(SUVA$medcosts)  
## [1] 1443.349  
mean(SUVA$medcosts[SUVA$medcosts > 0])  
## [1] 1492.765
```

The graph is best done with `extRemes::mrlplot`

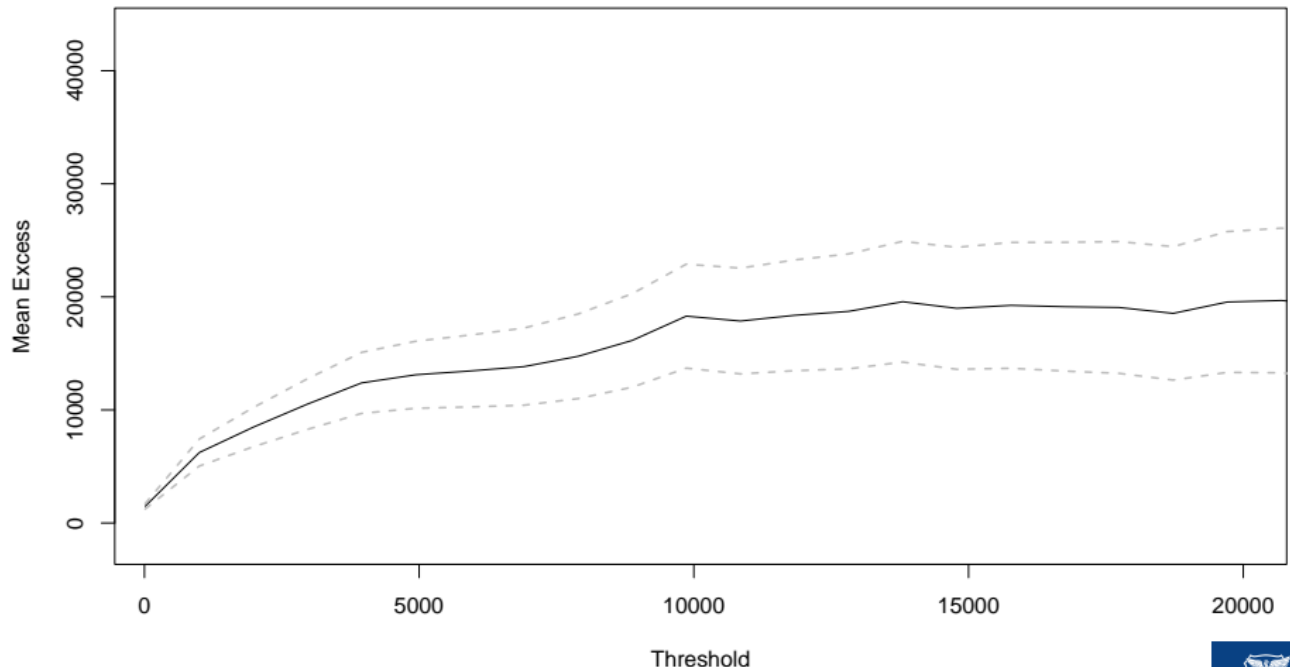
```
mrlplot(SUVA$medcosts [SUVA$medcosts > 0])
```



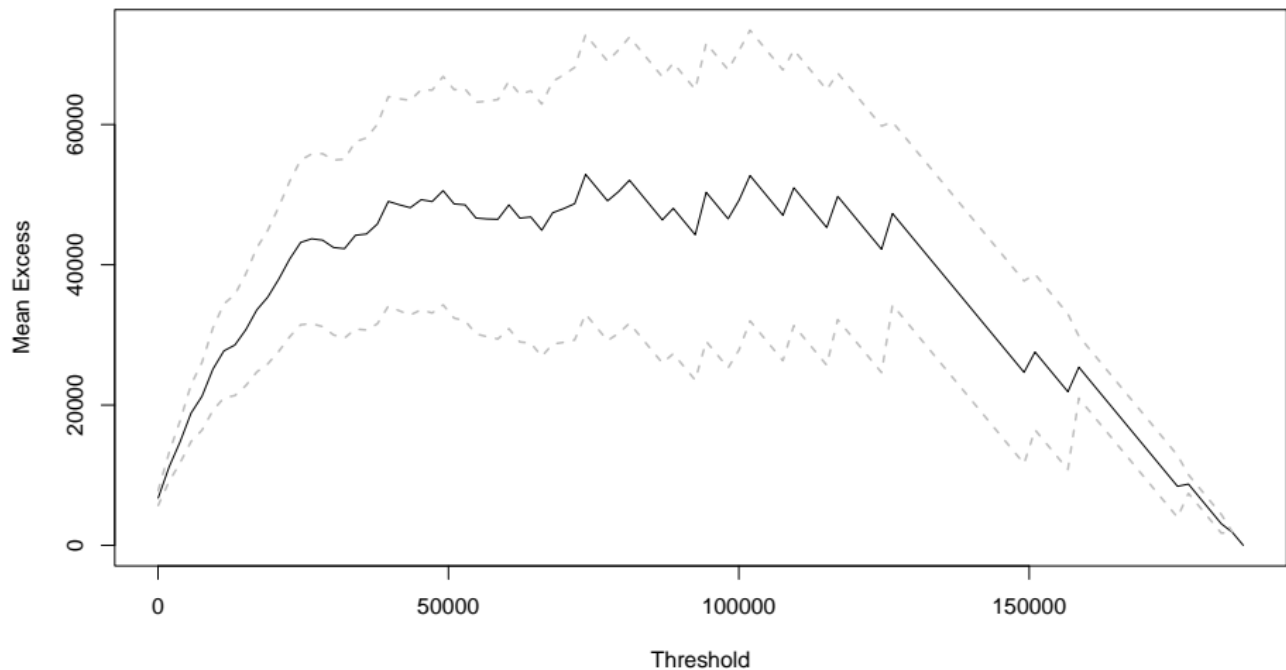
Linear increases point towards a heavy tailed distribution. Here the graph is biased because dominated by a few very large claims.

If we restrict the graph to range up to 20000 (which is roughly 99% of the data) we get:

```
mrlplot(SUVA$medcosts [SUVA$medcosts > 0], xlim = c(250, 20000))
```



```
mrlplot(SUVA$dailyyellow[SUVA$dailyyellow > 0])
```



This is not as heavy tailed.

2 Data analysis and descriptive statistics (MW 3.1)

- Raw data
- Moments
- Quantiles
- Boxplots
- Log-log plots

Quantiles

```
quantile(SUVA$dailyallow[SUVA$dailyallow > 0])
```

```
##      0%      25%      50%      75%     100%
## 26.0   842.0  1763.0  4841.5 186850.0
```

One can also focus on particular quantiles:

```
quantile(SUVA$dailyallow[SUVA$dailyallow > 0], probs = c(0.75,
  0.95, 0.99))
```

```
##      75%      95%      99%
## 4841.5 25140.5 93285.0
```

Note this “corresponds” to (crude) empirical versions of Values at Risk (“VaR”s).

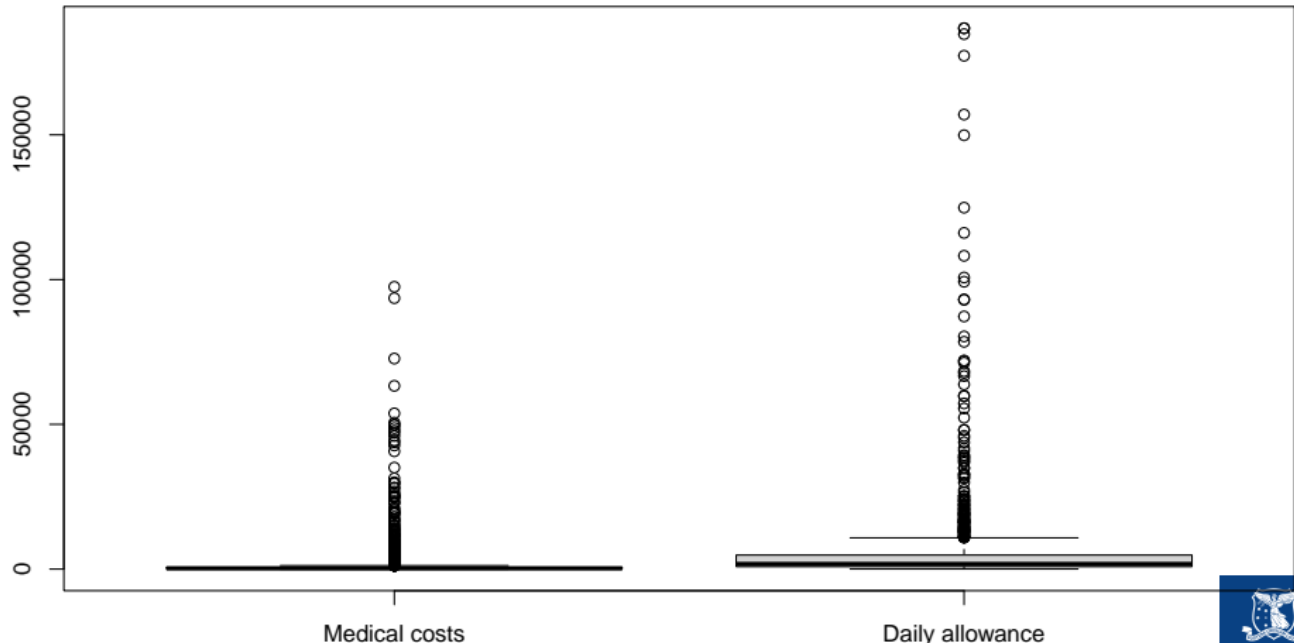
(There are better ways of estimating VaRs though.)

2 Data analysis and descriptive statistics (MW 3.1)

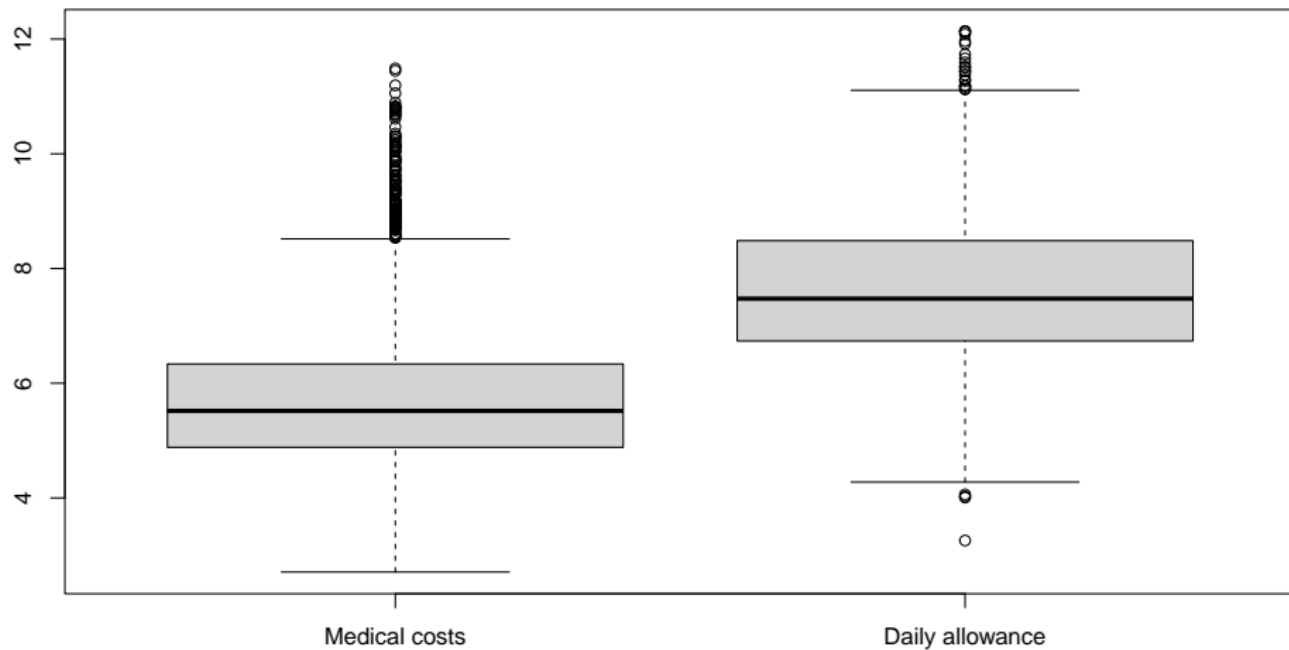
- Raw data
- Moments
- Quantiles
- Boxplots
- Log-log plots

Boxplots

```
boxplot(list(`Medical costs` = SUVA$medcosts[SUVA$medcosts > 0], `Daily allowance` = SUVA$dailyallow[SUVA$dailyallow > 0]))
```



```
boxplot(list(`Medical costs` = log(SUVA$medcosts[SUVA$medcosts > 0]), `Daily allowance` = log(SUVA$dailyallow[SUVA$dailyallow > 0])))
```



2 Data analysis and descriptive statistics (MW 3.1)

- Raw data
- Moments
- Quantiles
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- Log-log plots

Log-log plots

The log-log plot is defined as

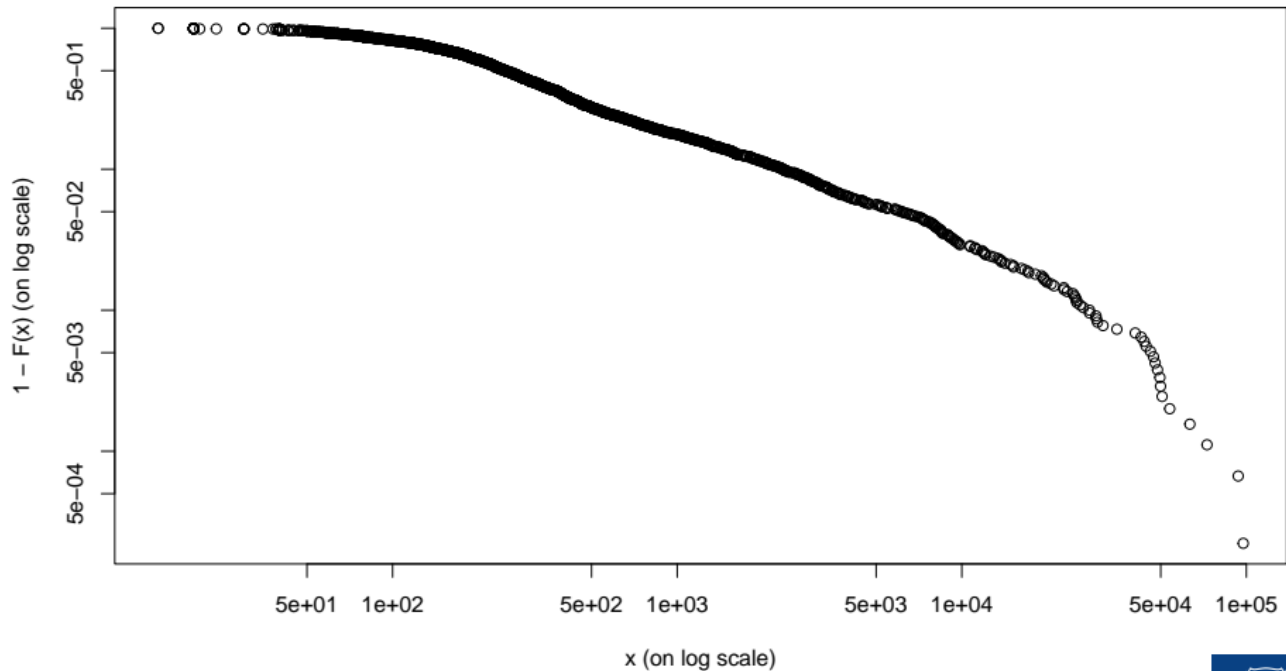
$$y \rightarrow [\log y, \log(1 - G(y))] ,$$

where G is simply replaced by \hat{G} for the empirical version.

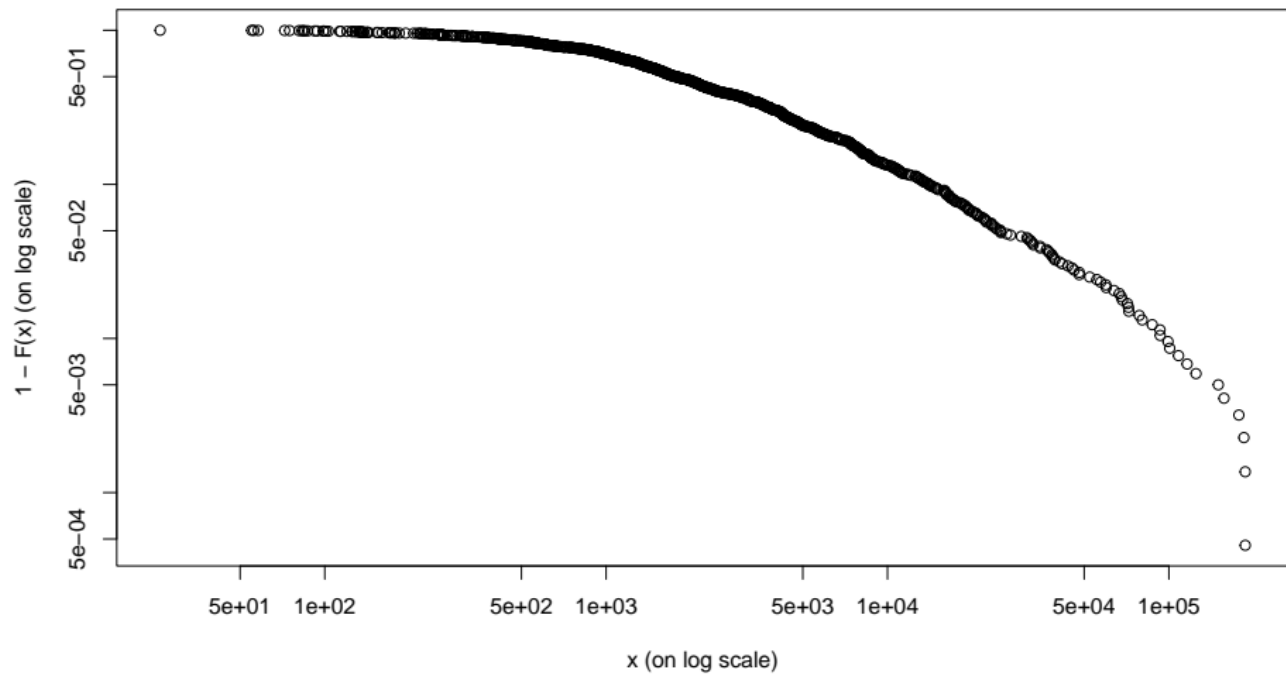
Just as for the (empirical) mean-excess plots, a linear behaviour (now decreasing) in the (empirical) log-log plot suggests a heavy-tailed distribution. Typical log-log plots are as in Figure 3.19 of Wuthrich (2022).

SUVA log-log plots

```
emplot(SUVA$medcosts[SUVA$medcosts > 0], alog = "xy", labels = TRUE)
```



```
emplot(SUVA$dailyallow[SUVA$dailyallow > 0], alog = "xy", labels = TRUE)
```



Again, medical costs are a good candidate for heavy tailed distributions (graph is more linear, except at the very end), whereas daily allowance is more reasonably behaved (graph is more concave).

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③ Selected parametric claims size distributions (MW 3.2)

- Parametric models for severity Y
- Extreme Value Theory

Gamma

- We shall write $Y \sim \Gamma(\alpha, \beta)$ if density has the form

$$g(y) = \frac{\beta^\alpha}{\Gamma(\alpha)} y^{\alpha-1} e^{-\beta y}, \text{ for } y > 0; \alpha, \beta > 0.$$

- Mean: $E(Y) = \alpha/\beta$
- Variance: $Var(Y) = \alpha/\beta^2$
- Skewness: $\varsigma_Y = 2/\sqrt{\alpha}$ [positively skewed distribution]
- Mgf: $M_Y(t) = \left(\frac{\beta}{\beta - t}\right)^\alpha$, provided $t < \beta$.
- Higher moments: $E(Y^k) = \frac{\Gamma(\alpha + k)}{\Gamma(\alpha) \beta^k}$
- Special case: When $\alpha = 1$, we have $Y \sim Exp(\beta)$

Inverse Gaussian

- We shall write $Y \sim IG(\alpha, \beta)$ if density has the form

$$g(y) = \frac{\alpha y^{-3/2}}{\sqrt{2\pi\beta}} \exp\left[-\frac{(\alpha - \beta y)^2}{2\beta y}\right], \text{ for } y > 0; \alpha, \beta > 0.$$

- Mean: $E(Y) = \alpha/\beta$
- Variance: $\text{Var}(Y) = \alpha/\beta^2$
- Skewness: $\varsigma_Y = 3/\sqrt{\alpha}$ [positively skewed distribution]
- Mgf: $M_Y(t) = e^{\alpha(1-\sqrt{1-2t/\beta})}$, provided $t < \beta/2$.
- The term “Inverse Gaussian” comes from the fact that there is an inverse relationship between its cgf and that of the Gaussian distribution, but NOT from the fact that the inverse is Gaussian!

Weibull

- We shall write $Y \sim \text{Weibull}(\tau, c)$ if density has the form

$$g(y) = (c\tau)(cy)^{\tau-1} \exp\{-(cy)^\tau\}, \text{ for } y \geq 0; \alpha, \beta > 0.$$

Note $G(u) = 1 - \exp\{-(cu)^\tau\}$.

- Mean: $E(Y) = \frac{\gamma(1+1/\tau)}{c}$
- Variance: $\text{Var}(Y) = \frac{\gamma(1+2/\tau)}{c^2} - \mu_Y^2$
- Skewness: $\varsigma_Y = \left[\frac{\gamma(1+2/\tau)}{c^2} - 3\mu_Y\sigma_Y^2 - \mu_Y^3 \right] / \sigma_Y^3$
- Mgf: does not exist for $\tau < 1$ and $t > 0$.
- Higher moments: $E(Y^k) = \frac{\gamma(1+k/\tau)}{c^k}$
- Note: if $Z \sim \text{expo}(1)$ then $Z^{1/\tau}/c \sim \text{Weibull}(\tau, c)$.

Lognormal

- We shall write $Y \sim \text{LN}(\mu, \sigma^2)$ and we have

$$\log Y \sim \mathcal{N}(\mu, \sigma^2).$$

- Mean: $E(Y) = \exp\{\mu + \sigma^2/2\}$
- Variance: $\text{Var}(Y) = \exp\{2\mu + \sigma^2\} (\exp\{\sigma^2\} - 1)$
- Skewness: $\varsigma_Y = (\exp\{\sigma^2\} + 2) (\exp\{\sigma^2\} - 1)^{1/2}$
- Mgf: does not exist for $t > 0$.

Log-gamma

- We shall write $\log Y \sim \Gamma(\gamma, c)$ and we have

$$g(y) = \frac{c^\gamma}{\Gamma(\gamma)} (\log y)^{\gamma-1} y^{-(c+1)}, \text{ for } y > 0; \alpha, \beta > 0.$$

- Mean: $E(Y) = \left(\frac{c}{c-1}\right)^\gamma$ for $c > 1$
- Variance: $Var(Y) = \left(\frac{c}{c-2}\right)^\gamma - \mu_Y^2$ for $c > 2$
- Skewness: $\varsigma_Y = \left[\left(\frac{c}{c-3}\right)^\gamma - 3\mu_Y\sigma_Y^2 - \mu_Y^3\right] / \sigma_Y^3$
- Mgf: does not exist for $t > 0$.
- Higher moments: $E(Y^k) = \left(\frac{c}{c-k}\right)^\gamma$ for $c > k$
- Special case: When $\alpha = 1$, we have $Y \sim Exp(\beta)$

Pareto Distribution

- We shall write $Y \sim \text{Pareto}(\theta, \alpha)$ if density has the form

$$g(y) = \frac{\alpha}{\theta} \left(\frac{y}{\theta} \right)^{-(\alpha+1)} \quad \text{for } y \geq \theta$$

- Mean: $E(Y) = \theta \frac{\alpha}{\alpha-1}$, $\alpha > 1$
- Variance: $Var(Y) = \theta^2 \frac{\alpha}{(\alpha-1)^2(\alpha-2)}$, $\alpha > 2$
- Skewness: $\varsigma_Y = \frac{2(1+\alpha)}{\alpha-3} \left(\frac{\alpha-2}{\alpha} \right)^{1/2}$, $\alpha > 3$
- Mgf: does not exist for $t > 0$
- Translated Pareto: distribution of $Y = Y - \beta$

③ Selected parametric claims size distributions (MW 3.2)

- Parametric models for severity Y
- Extreme Value Theory

Extreme Value Theory

Note that the following topics (which appear briefly in MW 3) will be covered in Module 6 on Extreme Value Theory:

- regular variation at infinity and tail index (page 58-59+)
- Hill plot (page 75+)
- Generalised Pareto (“GP”) and Generalised Extreme Value (“GEV”) distributions

- 1 Introduction
- 2 Data analysis and descriptive statistics (MW 3.1)
- 3 Selected parametric claims size distributions (MW 3.2)
- 4 Fitting of distributions (MW3.2)**
- 5 Model selection (MW 3.3)
- 6 ✕ Other advanced topics
- 7 Calculating within layers for claim sizes (MW 3.4)

4 Fitting of distributions (MW3.2)

- Introduction

- Moment matching estimation (MME)
- Maximum likelihood estimation (MLE)
- Parameter uncertainty
- ✕ Bootstrap
- Dealing with left-truncation and right-censoring

Introduction

- By now, the modeller should have identified some candidate parametric distributions for their data set.
 - Example: for SUVA\$dailyallow, based on data numerical and graphical explorations, we decided to try gamma, lognormal, and Weibull.
- In order to be able to compare them, that is, assess their goodness of fit (which will be discussed in the next section), one must find numerical values for the parameters of the candidates.
- The parameter values we will choose will depend on some criterion, which the modeller can choose. In some cases, they may even want to follow different approaches and choose which is the one they think works best for their purposes.

- Fitting criteria include:
 - moment matching: choose the m parameters such that the first m moments match
 - maximum likelihood: choose the parameters such that the overall likelihood that the model generated the data is maximal
 - ~~✗~~ maximum goodness: choose the parameters such that some goodness of fit criterion is maximised
- We will discuss the first two here, and the third after the goodness of fit section (for obvious reasons).

- There are other criteria which we will not discuss in details, such as:
 - quantile matching: choose parameters such that empirical quantiles match theoretical quantiles
 - least squares: choose parameters such that the “sum of squares” is minimal (this corresponds to maximum likelihood for Gaussian random variables)
 - so-called “minimum distance estimation”, which minimises distances between certain theoretical and empirical functions; see `actuar::mde`. For instance the `actuar` Cramér-von Mises method minimises the (weighted) distance between empirical and theoretical cdf's
 - score functions such as considered in probabilistic forecasting; see, e.g., package `scoringRules` in Jordan, Krüger, and Lerch (2019)

R and technical notes

- The function `MASS::fitdist` is standard, and uses by default MLE via the `optim` function.
- The package `fitdistrplus` allows other fitting criteria (such as method of moments and maximum goodness), and also allows for the user to supply their own optimisation function.
- Distributions are coded in the following way. For distribution `foo`:
 - `dfoo` is the density (pdf) of `foo`
 - `pfoo` is the distribution function (cdf) of `foo`
 - `qfoo` is the quantile function of `foo`
 - `rfoo` is the random number generator of `foo`
- This link provides a very comprehensive list of available distributions in R. As a user you can define any distribution yourself, and this syntax will be recognised by some functions.
- The package `actuar` provides additional distributions. It also automates the transformation of such distribution functions in presence of left-truncation and right-censoring (more later).

4 Fitting of distributions (MW3.2)

- Introduction
- Moment matching estimation (MME)
- Maximum likelihood estimation (MLE)
- Parameter uncertainty
- ✕ Bootstrap
- Dealing with left-truncation and right-censoring

Moment matching estimation (MME)

- This is quite straightforward; for a distribution of m parameters:
 - choose m moments you care about (usually the first m moments around the origin or the mean);
 - build a system of m equations (for the m parameters) matching empirical moments of your choice;
 - solve the system (might require a computer or some numerical techniques if the equations are not linear).
- In R, set the argument `method` to `mme` in the call to `fitdist`
 - For distributions of 1 and 2 parameters, mean and variances are matched
 - For a certain number of distributions a closed form formula is used. For the others, equations are solved numerically using `optim` and by minimising the sum of squared differences between theoretical and observed moments.

MME for SUVA

```
fit.lnorm.mme <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "lnorm", method = "mme", order = 1:2)
```

```
fit.lnorm.mme$estimate
```

```
## meanlog      sdlog
```

```
## 2.0146490 0.1871132
```

```
fit.lnorm.mme$loglik
```

```
## [1] -1959.06
```

```
fit.lnorm.mme2 <- fitdistrplus::mmedist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "lnorm", order = 1:2)
```

```
fit.lnorm.mme2$estimate
```

```
## meanlog      sdlog
```

```
## 2.0146490 0.1871132
```

```
fit.lnorm.mme2$loglik
```

```
## [1] -1959.06
```

```
fit.gamma.mme <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "gamma", method = "mme", order = 1:2)
fit.gamma.mme$estimate

##      shape      rate
## 28.065081  3.678009

fit.gamma.mme$loglik

## [1] -1951.838

# function to calculate sample raw moment
memp <- function(x, order) emm(x, order)
fit.weibull.mme <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "weibull", method = "mme", memp = memp, order = c(1, 2))
fit.weibull.mme$estimate

##      shape      scale
## 6.172910 8.212158

fit.weibull.mme$loglik

## [1] -2020.119
```

4 Fitting of distributions (MW3.2)

- Introduction
- Moment matching estimation (MME)
- Maximum likelihood estimation (MLE)
- Parameter uncertainty
- ✕ Bootstrap
- Dealing with left-truncation and right-censoring

Maximum likelihood estimation (MLE)

- The likelihood for a statistical model gives an indication of how likely it is that this data set was generated, should the model be correct.
- If there is no censoring or truncation, we have

$$L(\theta; \mathbf{x}) = \prod_{i=1}^n f(x_i; \theta).$$

Obviously, this is a function of the parameter (vector) $\theta = (\theta_1, \dots, \theta_m)$, for a given set of observations, denoted $\mathbf{x} = (x_1, \dots, x_n)$.

- The value $\hat{\theta}$ that maximises the likelihood is called the maximum likelihood estimator (MLE).
- Often, it is more convenient to maximise the log-likelihood function given by

$$\ell(\theta; \mathbf{x}) = \log L(\theta; \mathbf{x}).$$

This avoids issues of underflow for L when n is large.

Note:

- The formulation of a likelihood in presence of left-truncation and right-censoring will be covered in a later section.
- Sometimes, MLEs match MMEs (check book).
- MLEs have nice properties (such as asymptotic normality, which is one way to estimate standard errors; see later).

MLE for SUVA

```
fit.lnorm.mle <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "lnorm", method = "mle")
fit.lnorm.mle$estimate

##      meanlog      sdlog
## 2.0140792 0.1918442

fit.lnorm.mle$loglik

## [1] -1958.359

fit.gamma.mle <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "gamma")
fit.gamma.mle$estimate

##      shape      rate
## 27.825788  3.646718

fit.gamma.mle$loglik

## [1] -1951.818
```

```
fit.weibull.mle <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "weibull", method = "mle")
fit.weibull.mle$estimate
```

```
##      shape      scale
## 5.527347 8.231313
```

```
fit.weibull.mle$loglik
```

```
## [1] -2003.26
```

```
summary(fit.weibull.mle)
```

```
## Fitting of the distribution ' weibull ' by maximum likelihood
## Parameters :
## Loglikelihood: -2003.26    AIC: 4010.519    BIC: 4020.524
## Correlation matrix:
##           shape      scale
## shape 1.0000000 0.3301744
## scale 0.3301744 1.0000000
```

4 Fitting of distributions (MW3.2)

- Introduction
- Moment matching estimation (MME)
- Maximum likelihood estimation (MLE)
- Parameter uncertainty
- \times Bootstrap
- Dealing with left-truncation and right-censoring

Parameter uncertainty

- The estimation of parameters is not perfect.
- The mere fact that different methods lead to different estimates makes that point obvious.
- This will always be true even if we are fitting the right distribution, just because we have only a finite sample of data
- How far can we be from the “truth” ?
- There are different ways of answering that question, two of which we discuss here:
 - the Wald approximation: standard errors via the Hessian for MLEs
 - bootstrap

Hessian matrix

The score (or gradient) vector consists of first derivatives

$$S(\theta; \mathbf{x}) = \left(\frac{\partial \ell(\theta; \mathbf{x})}{\partial \theta_1}, \dots, \frac{\partial \ell(\theta; \mathbf{x})}{\partial \theta_m} \right)',$$

so that the MLE satisfies F.O.C. $S(\hat{\theta}; \mathbf{x}) = \mathbf{0} = (0, \dots, 0)'$. - The $m \times m$ Hessian matrix for $\ell(\theta; \mathbf{x})$ is defined by

$$H(\theta; \mathbf{x}) = \frac{\partial^2 \ell(\theta; \mathbf{x})}{\partial \theta \partial \theta'} = \begin{bmatrix} \frac{\partial^2 \ell(\theta; \mathbf{x})}{\partial \theta_1^2} & \dots & \frac{\partial^2 \ell(\theta; \mathbf{x})}{\partial \theta_1 \partial \theta_m} \\ \vdots & \ddots & \vdots \\ \frac{\partial^2 \ell(\theta; \mathbf{x})}{\partial \theta_m \partial \theta_1} & \dots & \frac{\partial^2 \ell(\theta; \mathbf{x})}{\partial \theta_m^2} \end{bmatrix}.$$

- This Hessian is used to estimate $\text{Var}(\hat{\theta})$.
- Minus the expected value of this is called the (expected) Fisher information.

The Wald approximation

It is well-known that a consistent estimator for the covariance matrix $\text{Var}(\hat{\theta})$ is given by the inverse of the negative of this Hessian matrix:

$$\text{Var}(\hat{\theta}) \geq \widehat{\text{Var}}(\hat{\theta}) = \left[-\mathbf{E}[H(\hat{\theta}; \mathbf{x})] \right]^{-1}.$$

The square root of the diagonal elements of this covariance estimate give the standard errors of the MLE estimates. Note:

- “Consistent” means it converges in probability to the value being estimated.
- When $n \rightarrow \infty$, the distribution of $\hat{\theta}$ is asymptotically normal.
- Note these are *asymptotic* results. Furthermore, their quality also strongly depends on the data, the distribution, and the parametrisation of the distribution.

Standard errors for SUVA data

Note these are obviously unavailable for MMEs. For MLEs:

```
fit.lnorm.mle$sd
```

```
##      meanlog      sdlog
## 0.005786951 0.004091492
summary(fit.lnorm.mle)
```

```
## Fitting of the distribution ' lnorm ' by maximum likelihood
## Parameters :
## Loglikelihood: -1958.359    AIC: 3920.717    BIC: 3930.722
## Correlation matrix:
##      meanlog  sdlog
## meanlog     1     0
## sdlog       0     1
```

```
fit.gamma.mle$sd

##      shape      rate
## 1.1799866 0.1560432
summary(fit.gamma.mle)

## Fitting of the distribution ' gamma ' by maximum likelihood
## Parameters :
## Loglikelihood: -1951.818    AIC: 3907.636    BIC: 3917.641
## Correlation matrix:
##      shape      rate
## shape 1.00000 0.99103
## rate  0.99103 1.00000
```

```
fit.weibull.mle$sd
```

```
##      shape      scale  
## 0.12127037 0.04760155
```

```
summary(fit.weibull.mle)
```

```
## Fitting of the distribution ' weibull ' by maximum likelihood
```

```
## Parameters :
```

```
## Loglikelihood: -2003.26    AIC: 4010.519    BIC: 4020.524
```

```
## Correlation matrix:
```

```
##      shape      scale
```

```
## shape 1.0000000 0.3301744
```

```
## scale 0.3301744 1.0000000
```

4 Fitting of distributions (MW3.2)

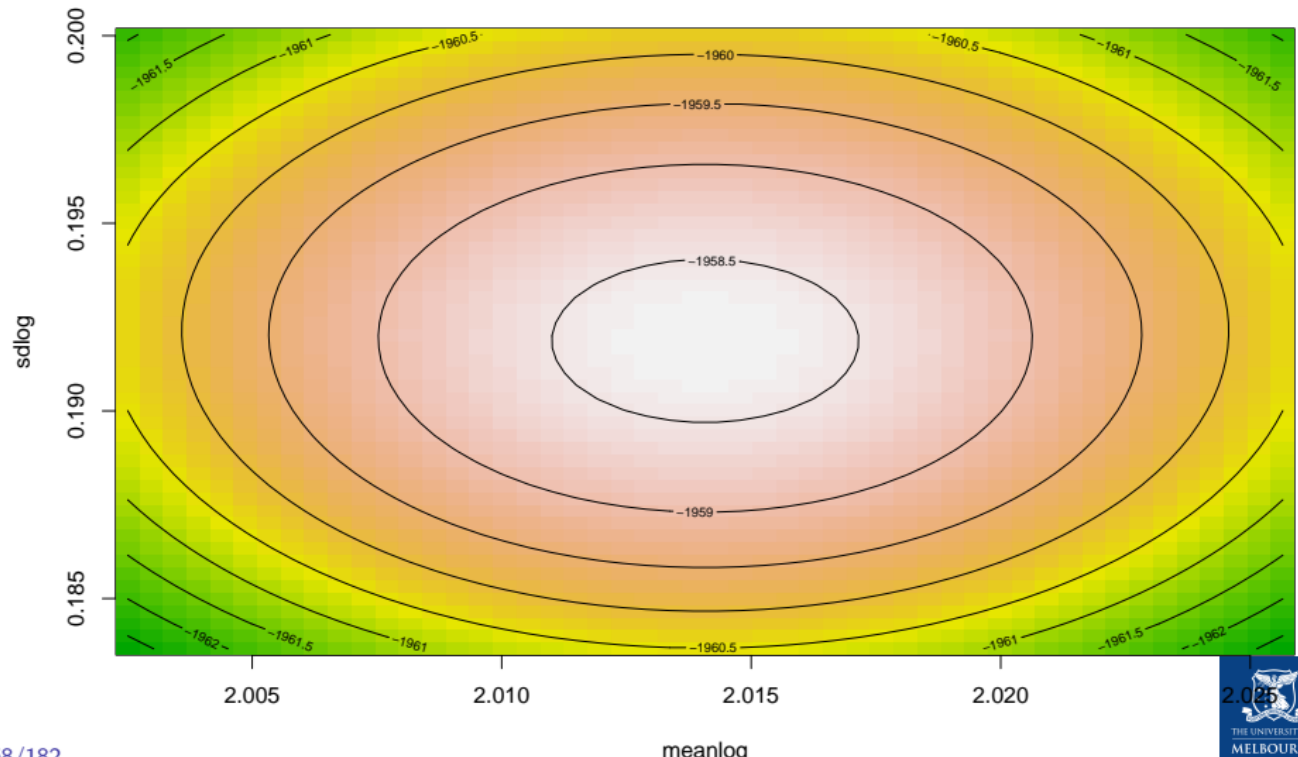
- Introduction
- Moment matching estimation (MME)
- Maximum likelihood estimation (MLE)
- Parameter uncertainty
- ✖ Bootstrap
- Dealing with left-truncation and right-censoring

Bootstrap

- It is advisable to compare the Wald approximation to the ones obtained using bootstrap procedures.
- Also, the Wald approximation assumes elliptical loglikelihood contours (related to the Gaussian asymptotic result), and hence having a look at the loglikelihood contours is also informative.
- More generally, one might want to simply use bootstrap to compute confidence intervals on parameters (or any function of those parameters).
- In R:
 - `llplot` will provide the loglikelihood contours; and
 - `bootdist` will provide bootstrap results on a fitted object.

✚ SUVA contours and bootstrap results

```
llplot(fit.lnorm.mle)
```



```
fit.lnorm.mle.boot <- bootdist(fit.lnorm.mle, niter = 1001)
fit.lnorm.mle.boot$fitpart # the Wald approximation

## Fitting of the distribution ' lnorm ' by maximum likelihood
## Parameters:
summary(fit.lnorm.mle.boot)

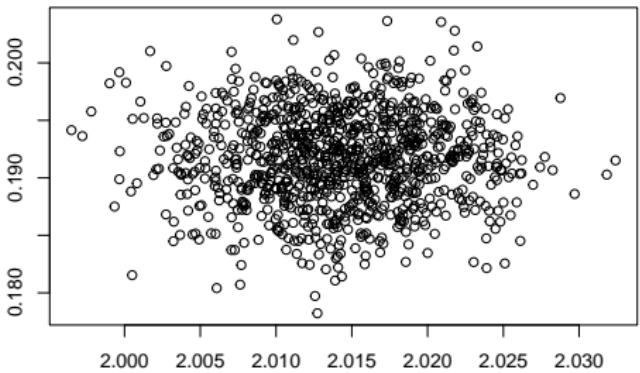
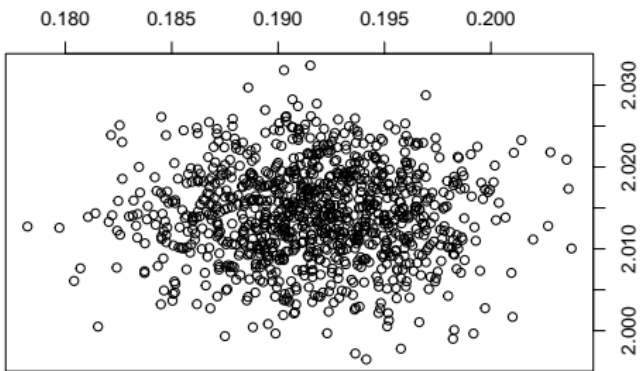
## Parametric bootstrap medians and 95% percentile CI
## Median      2.5%      97.5%
## meanlog 2.0140785 2.0027568 2.0248904
## sdlog   0.1917506 0.1837203 0.1991819

# CI to be compared with
fit.lnorm.mle$estimate + cbind(estimate = 0, `2.5%` = -1.96 *
  fit.lnorm.mle$sd, `97.5%` = 1.96 * fit.lnorm.mle$sd)

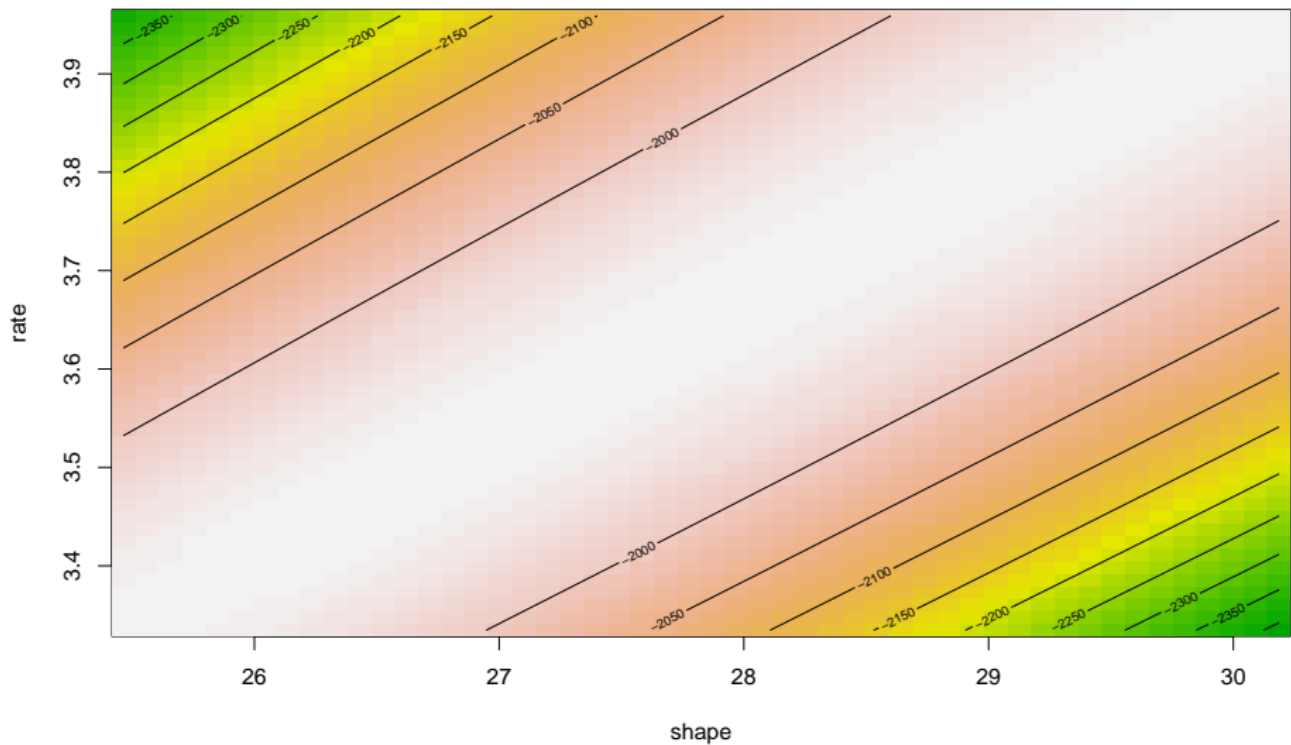
## estimate      2.5%      97.5%
## meanlog 2.0140792 2.0027368 2.0254216
## sdlog   0.1918442 0.1838249 0.1998635
```

```
plot(fit.lnorm.mle.boot)
```

Bootstrapped values of parameters



```
llplot(fit.gamma.mle)
```



```
fit.gamma.mle.boot <- bootdist(fit.gamma.mle, niter = 1001)
fit.gamma.mle.boot$fitpart # the Wald approximation

## Fitting of the distribution ' gamma ' by maximum likelihood
## Parameters:
summary(fit.gamma.mle.boot)

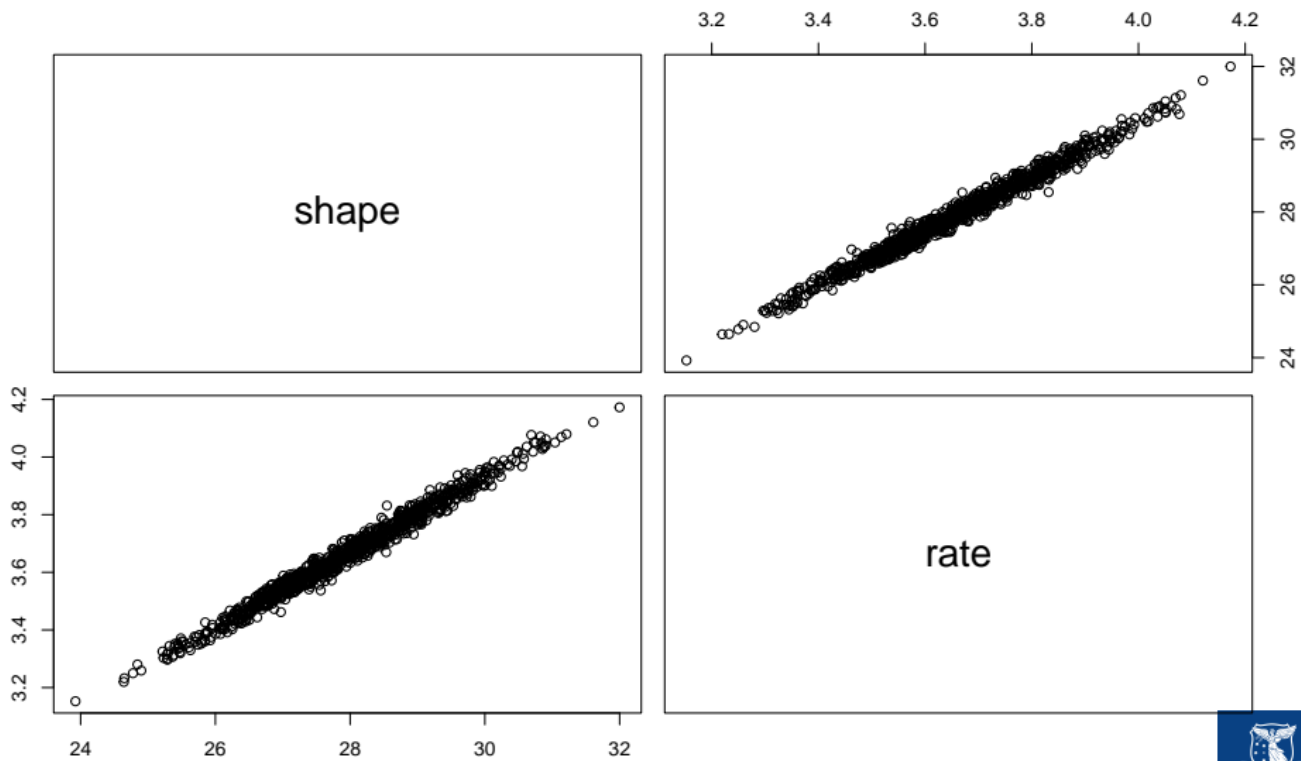
## Parametric bootstrap medians and 95% percentile CI
##           Median      2.5%     97.5%
## shape 27.844365 25.526058 30.341980
## rate   3.651136  3.352219  3.974696

# CI to be compared with
fit.gamma.mle$estimate + cbind(estimate = 0, `2.5%` = -1.96 *
  fit.gamma.mle$sd, `97.5%` = 1.96 * fit.gamma.mle$sd)

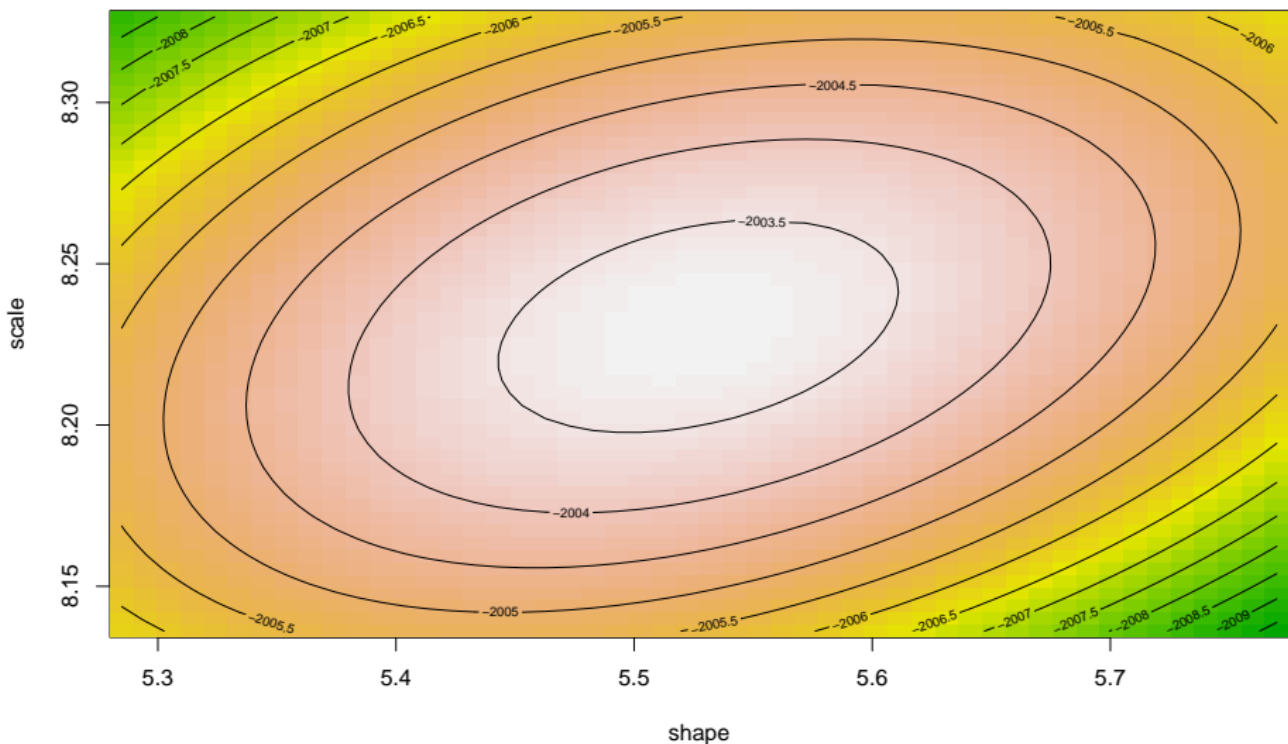
##           estimate      2.5%     97.5%
## shape 27.825788 25.513015 30.138562
## rate   3.646718  3.340873  3.952562
```

```
plot(fit.gamma.mle.boot)
```

Bootstrapped values of parameters



```
llplot(fit.weibull.mle)
```



```
fit.weibull.mle.boot <- bootdist(fit.weibull.mle, niter = 1001)
fit.weibull.mle.boot$fitpart # the Wald approximation

## Fitting of the distribution ' weibull ' by maximum likelihood
## Parameters:
summary(fit.weibull.mle.boot)

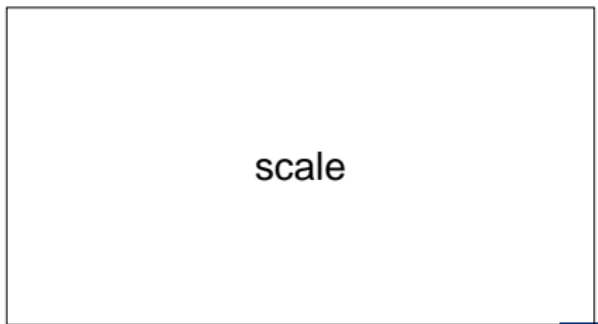
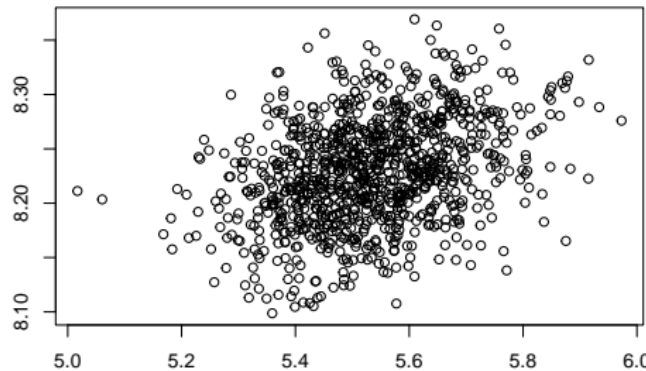
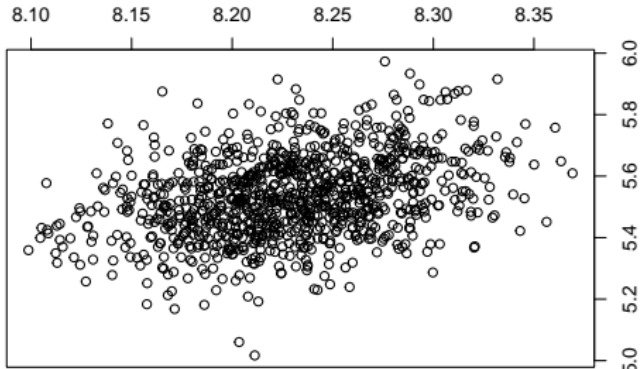
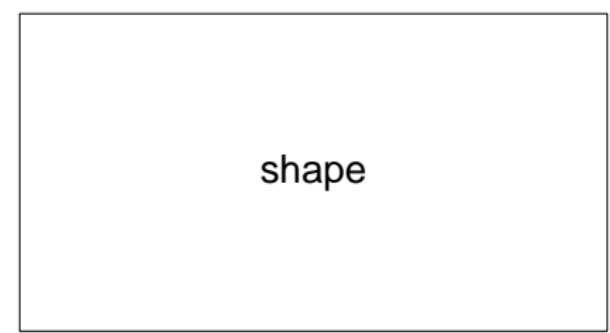
## Parametric bootstrap medians and 95% percentile CI
## Median      2.5%      97.5%
## shape 5.525935 5.283250 5.807544
## scale 8.229161 8.132619 8.322033

# CI to be compared with
fit.weibull.mle$estimate + cbind(estimate = 0, `2.5%` = -1.96 *
  fit.weibull.mle$sd, `97.5%` = 1.96 * fit.weibull.mle$sd)

## estimate      2.5%      97.5%
## shape 5.527347 5.289657 5.765037
## scale 8.231313 8.138014 8.324612
```

```
plot(fit.weibull.mle.boot)
```

Bootstrapped values of parameters



✖ Confidence intervals on other quantities

- One can readily obtain confidence intervals on quantiles with the `quantile` function and a `bootdist` object
- More generally, this can be done of any function; see Delignette-Muller and Dutang (2015).

```
quantile(fit.gamma.mle.boot)
```

```
## (original) estimated quantiles for each specified probability (non-censored)
## Median of bootstrap estimates
##          p=0.1    p=0.2    p=0.3    p=0.4    p=0.5    p=0.6
## estimate 5.845722 6.394506 6.811862 7.181263 7.538064 7.90777
##          p=0.7    p=0.8    p=0.9
## estimate 8.316399 8.811106 9.530596
##
## two-sided 95 % CI of each quantile
```

4 Fitting of distributions (MW3.2)

- Introduction
- Moment matching estimation (MME)
- Maximum likelihood estimation (MLE)
- Parameter uncertainty
- ✕ Bootstrap
- Dealing with left-truncation and right-censoring

Coding of censored data in R

- We need two columns named `left` and `right`.
- The `left` column contains:
 - NA for left censored observations
 - the left bound of the interval for interval censored observations
 - the observed value for non-censored observations
- The `right` column contains:
 - NA for right censored observations
 - the right bound of the interval for interval censored observations
 - the observed value for non-censored observations
- This corresponds to the coding names `interval2` in function `Surv` of the package `survival`

In other words, (left,right) corresponds to

- (a,a) for an exact observation at a ($= a$)
- (a,NA) for a right censored observation at a ($> a$)
- (NA,b) for a left censored observation at b ($\leq b$)
- (a,b) for an interval censored observation over (a, b) , where $b > a$
 $(> a \text{ and } \leq b)$

For example, consider twenty values from the canlifins of the CASdatasets package (see Delignette-Muller and Dutang (n.d.))

```
exitage <- c(81.1, 78.9, 72.6, 67.9, 60.1, 78.3, 83.4, 66.9,  
74.8, 80.5, 75.6, 67.1, 75.3, 82.8, 70.1, 85.4, 74, 70, 71.6,  
76.5)  
death <- c(0, 0, 1, 0, 0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 1, 1, 0,  
0, 0, 0)
```

- the first value is someone who exited the study at 81.1, but not via death, so it is a right-censored observation
- this needs to be coded:
 - left: 81.1
 - right: NA

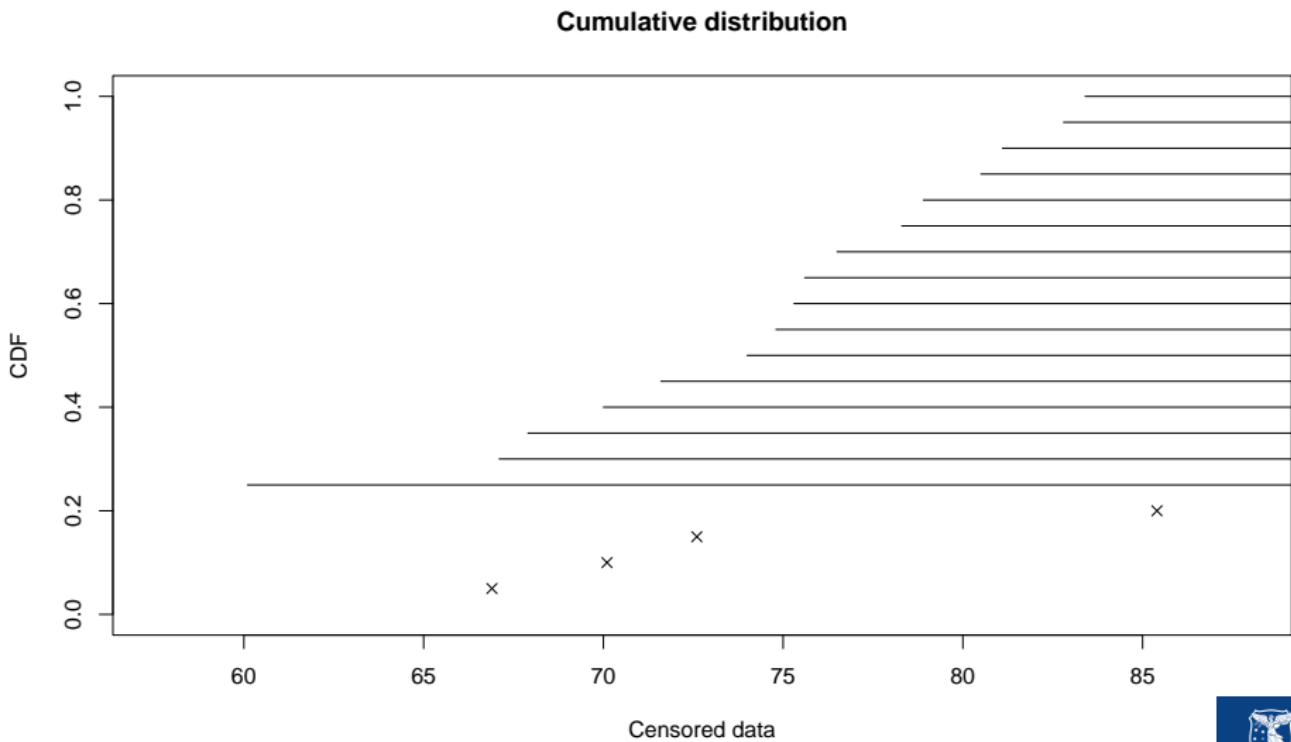
Overall, this becomes:

```
casdata <- cbind.data.frame(left = exitage, right = exitage)
casdata$right[death == 0] <- NA # the censored values
print(casdata)
```

(Note: How to use the function Surv() is explained in Delignette-Muller and Dutang, n.d.)

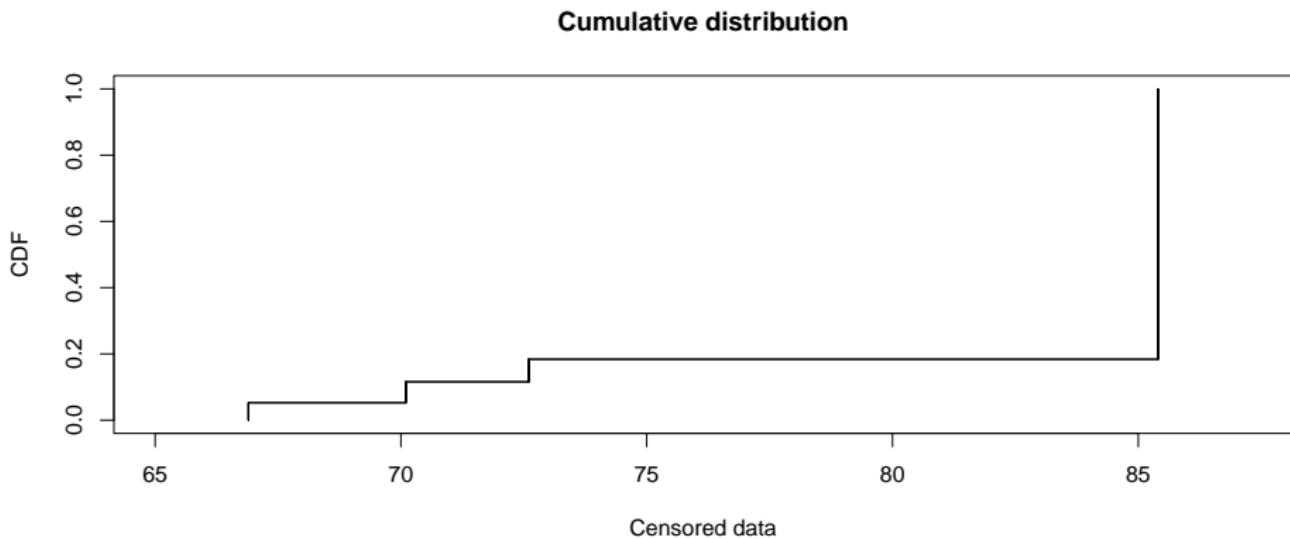
Censored data can be plotted raw...

```
plotdistcens(casdata, NPMLE = FALSE)
```



... or as an empirical distribution

```
plotdistcens(casdata)
```



Note that there are some technical subtleties with creating empirical distributions for censored data. This is out of scope, but details can be found in Delignette-Muller and Dutang (n.d.), Delignette-Muller and Dutang (2015), and references therein.

Fitting censored data

- Again, this is easily done with associated `foocens` functions

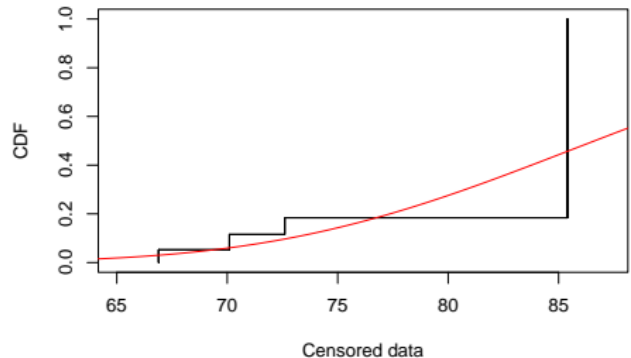
```
cas.gamma.fit <- fitdistcens(casdata, "gamma")
summary(cas.gamma.fit)
```

```
## Fitting of the distribution ' gamma ' By maximum likelihood on censored
## Parameters
## Loglikelihood: -20.0179    AIC: 44.0358    BIC: 46.02727
## Correlation matrix:
##           shape      rate
## shape 1.0000000 0.9983071
## rate   0.9983071 1.0000000
```

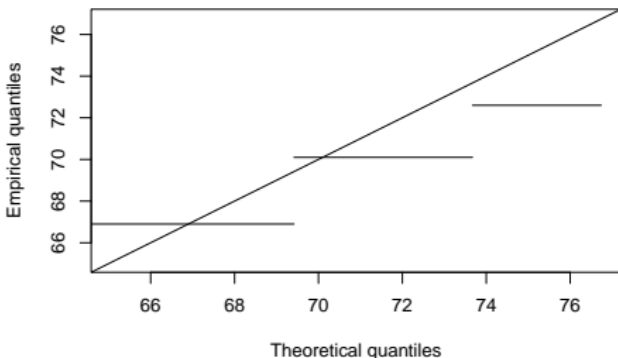
- Commands like `cdfcomp`, `cens`, `bootdist`, `cens` also exist.
- However, `gofstat` is not available as there do not exist results general enough to be coded in the package. Specific results for right-censored variables do exist, though.

```
plot(cas.gamma.fit)
```

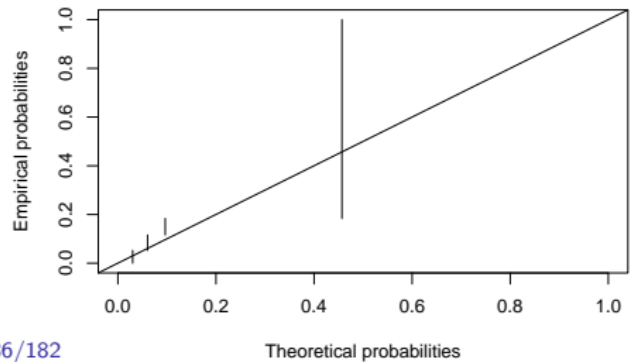
Empirical and theoretical CDFs



Q-Q plot



P-P plot



Left-truncation and R

- Unfortunately there is no pre-coded function for left-truncation.
- It can be done manually, with care.
- With left-truncation, the key (from definition) is that an observation will exist, if, *and only if*, it was beyond the truncation point. This means that the probability/lielihood associated to each observation is *conditional* on it being more than the truncation point.
- What follows is generalised to left- and right- truncation, and is taken from Delignette-Muller and Dutang (n.d.).
- For X before truncation, the l -left-truncated, u -right-truncated variable Y has density

$$g_Y(y) = \begin{cases} \frac{g_X(x)}{G_X(u)-G_X(l)} & \text{if } l < x < u \\ 0 & \text{otherwise} \end{cases}$$

As an example in R, the d and p functions of a truncated exponential can be coded as:

```
dtexp <- function(x, rate, low, upp) {  
  PU <- pexp(upp, rate = rate)  
  PL <- pexp(low, rate = rate)  
  dexp(x, rate)/(PU - PL) * (x >= low) * (x <= upp)  
}  
  
ptexp <- function(q, rate, low, upp) {  
  PU <- pexp(upp, rate = rate)  
  PL <- pexp(low, rate = rate)  
  (pexp(q, rate) - PL)/(PU - PL) * (q >= low) * (q <= upp) +  
  1 * (q > upp)  
}
```

If we generate 200 such truncated variables:

```
set.seed(22042021)
n <- 200 # number of observations
x <- rexp(n) # simulating sample of x's
y <- x[x > 0.5 & x < 3] # truncating to get sample of y's
```

and then fit them with either left- and right- truncation estimated from the data:

```
fit.texp.emp <- fitdist(y, "texp", method = "mle", start = list(rate = 3),
fix.arg = list(low = min(y), upp = max(y)))
```

... or in an informative way (i.e. we know what the bounds are):

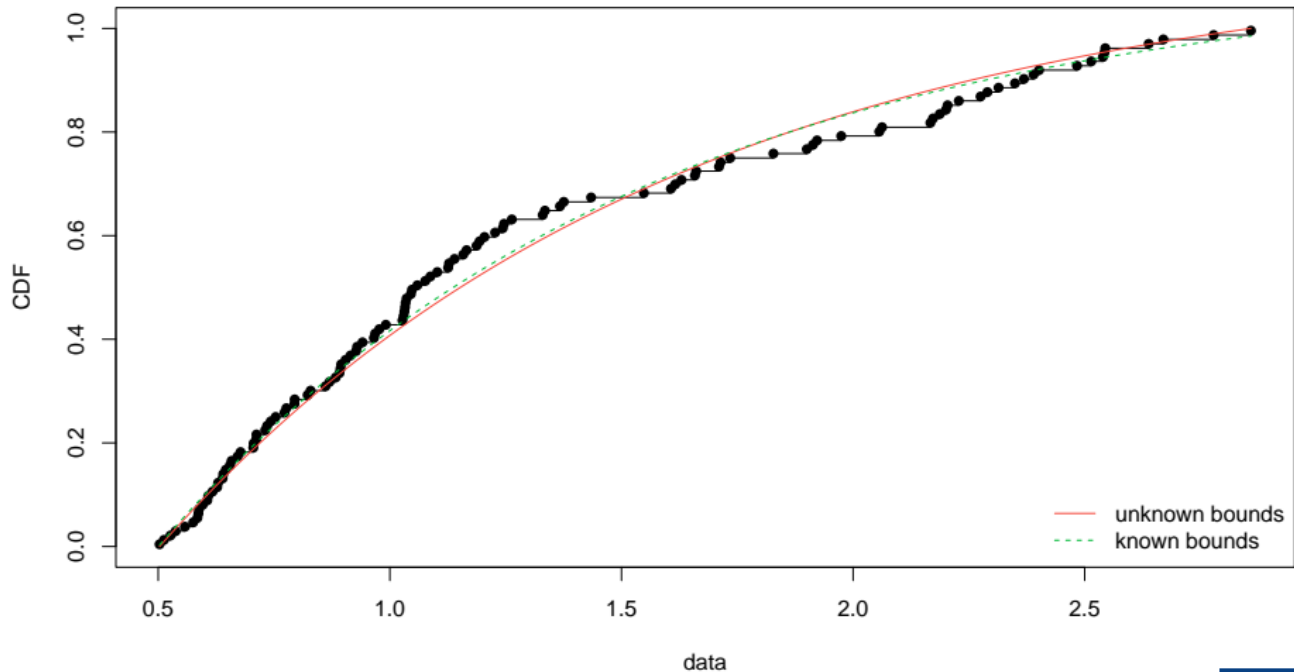
```
fit.texp.inf <- fitdist(y, "texp", method = "mle", start = list(rate = 3),
fix.arg = list(low = 0.5, upp = 3))
```

```
gofstat(list(fit.texp.emp, fit.texp.inf), fitnames = c("unknown bounds",
  "known bounds"))

## Goodness-of-fit statistics
##                                     unknown bounds known bounds
## Kolmogorov-Smirnov statistic      0.07546318  0.06747339
## Cramer-von Mises statistic       0.13211330  0.09999122
## Anderson-Darling statistic        Inf       0.59863722
##
## Goodness-of-fit criteria
##                                     unknown bounds known bounds
## Akaike's Information Criterion    165.3424    169.7301
## Bayesian Information Criterion   168.1131    172.5007
```

```
cdfcomp(list(fit.texp.emp, fit.texp.inf), legendtext = c("unknown bounds",  
"known bounds"))
```

Empirical and theoretical CDFs



Likelihood formulation with both left-truncation and right-censoring

- For our purposes, we shall represent our set of observations as

$$(t_j, x_j, \delta_j)$$

where

- t_j is the left truncation point;
 - x_j is the claim value that produced the data point; and
 - δ_j is indicator whether limit has been reached.
- For example:
 - (50, 250, 0)
 - (100, 1100, 1)

Densities will be as follows:

$$L(\theta; \mathbf{x}) = \prod_j \left[\frac{f(x_j; \theta)}{1 - F(t_j; \theta)} \right]^{1-\delta_j} \cdot \prod_j \left[\frac{1 - F(x_j; \theta)}{1 - F(t_j; \theta)} \right]^{\delta_j}.$$

The contribution to the likelihood function for a data point where the limit has not been reached is given by

$$\left[\frac{f(x_j)}{1 - F(t_j)} \right]^{1-\delta_j}.$$

The contribution to the likelihood function for a data point where the limit has been reached is given by

$$\left[\frac{1 - F(x_j)}{1 - F(t_j)} \right]^{\delta_j}.$$

Note here that the policy limit if reached would be equal to $x_j - t_j$.

When truncation and censoring levels are the same everywhere

- In R, the approach will be to code a left-truncated function, and then use the `foocens` functions.
- Let us do this on a gamma distribution:

```
dtgamma <- function(x, rate, shape, low) {  
  PL <- pgamma(low, rate = rate, shape = shape)  
  dgamma(x, rate = rate, shape = shape)/(1 - PL) * (x >= low)  
}  
  
ptgamma <- function(q, rate, shape, low) {  
  PL <- pgamma(low, rate = rate, shape = shape)  
  (pgamma(q, rate = rate, shape = shape) - PL)/(1 - PL) * (q >=  
  low)  
}
```

We initially assume that the truncation and censoring levels are the same everywhere.

For instance,

- truncated at 2
- censored at 20

for all data points.

Simulating one such dataset:

```
set.seed(22042021)
n <- 2000 # number of observations
x <- rgamma(n, shape = 2, rate = 0.2) # simulating sample of x's
x <- x[x > 2] # left-truncation at 2
n - length(x) # number of observations that were truncated

## [1] 123

censoring <- x > 20 # we will censor at 20
x[x > 20] <- 20 # censoring at 20
```

Transforming this into the right format:

```
censoring[censoring == FALSE] <- x[censoring == FALSE]
censoring[censoring == TRUE] <- NA
xcens <- cbind.data.frame(left = x, right = censoring)
```

And fitting:

```
# Not allowing for truncation:
fit.gamma.xcens <- fitdistcens(xcens, "gamma", start = list(shape = mean(xcens$left),
  rate = mean(xcens$left)/var(xcens$left)))
summary(fit.gamma.xcens)

...
## Fitting of the distribution ' gamma ' By maximum likelihood on censored
## Parameters
## Loglikelihood: -5412.521    AIC: 10829.04    BIC: 10840.12
## Correlation matrix:
##           shape      rate
## shape 1.0000000 0.9208904
##          ...
```



Allowing for truncation

```
fit.tgamma.xcens <- fitdistcens(xcens, "tgamma", start = list(shape = mean(xcens$left), rate = mean(xcens$left)/var(xcens$left)), fix.arg = list(low = min(xcens$left), high = max(xcens$right))), method = "Nelder-Mead")  
summary(fit.tgamma.xcens)
```

...

Fitting of the distribution ' tgamma ' By maximum likelihood on censored

Parameters

Fixed parameters:

Loglikelihood: -5340.151 AIC: 10684.3 BIC: 10695.38

Correlation matrix:

shape rate

shape 1.0000000 0.9401461

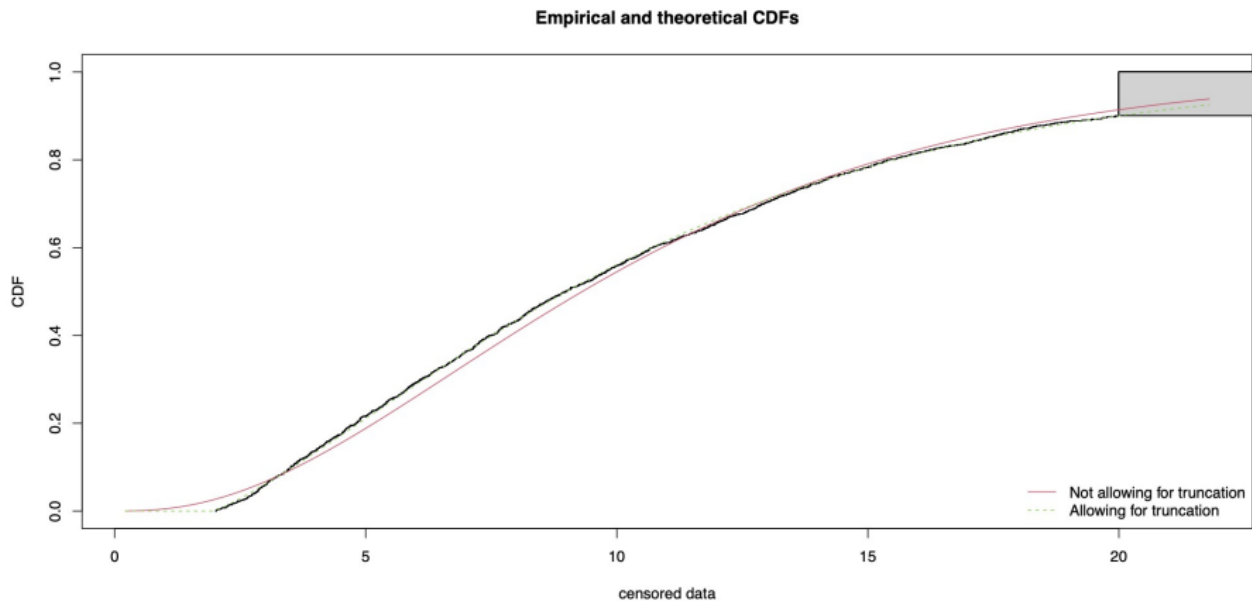
rate 0.9401461 1.0000000

NA

...



```
cdfcompcens(list(fit.gamma.xcens, fit.tgamma.xcens), legendtext = c("Not allowing for truncation"))
```



When truncation and censoring levels vary

In real life, an insurance product would have more than one level of deductibles and limits to suit different policyholders. Simulating another dataset:

```
set.seed(2022)
n <- 3006 # number of observations 9 x 334
orig_x <- rgamma(n, shape = 2, rate = 0.2) # simulating sample of x's
deductibles <- rep(c(rep(1, 3), rep(3, 3), rep(5, 3)), 334)
limits <- rep(c(15, 20, 30), 3 * 334) + deductibles
# limit is on payment, not raw loss
```

Manually applying the deductibles and limits:

```
x <- orig_x
censored <- x>limits # we will censor at the limits
x[censored] <- limits[censored] # censoring at the limits
# the above takes only elements of x which have TRUE
# in the vector censored
#
deducted <- x > deductibles
x <- x[deducted] # left-truncation at all points
# here the truncated observations disappear!
n-length(x) # observations that were truncated

## [1] 431
# that many were removed
#
claims <- data.frame(x = x, #
                      deduct = deductibles[deducted], #
                      limitI = censored[deducted])
```

Preliminary analysis:

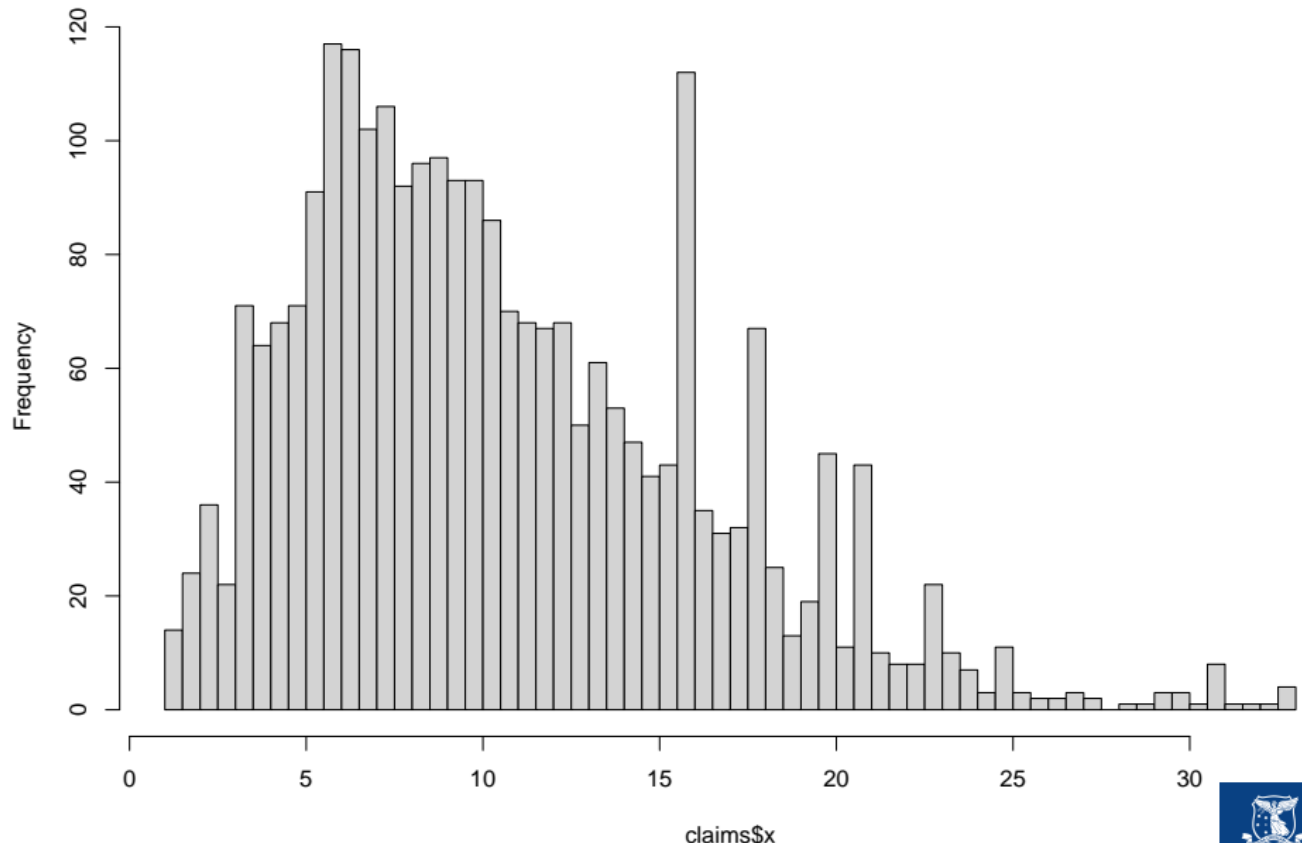
(note that there are no claims less than 1)

```
claims <- claims[sample(1:nrow(claims), nrow(claims)), ] # randomising data  
# we pretend we do not know how the data was generated  
head(claims)
```

	x	deduct	limitl
444	17.196879	1	FALSE
1733	13.964488	3	FALSE
1414	3.521077	1	FALSE
1407	15.488936	1	FALSE
1138	11.509672	3	FALSE
1977	4.875571	3	FALSE

```
hist(claims$x, breaks = 100)
```

Note this includes right-censored observations But not
the truncated values

Histogram of claims\$x

Preparing our joint log-likelihood function:

Here, we are minimising a negative log-likelihood instead of maximising a log-likelihood.

```
negloglik <- function(pdf, cdf, param, x, deduct, limitI) {  
  # Function returns the negative log likelihood of the  
  # censored and truncated dataset. Each data point's  
  # contribution to the log likelihood depends on the  
  # theoretical distribution pdf and cdf and also the  
  # deductible and limit values to adjust for truncation  
  # and censoring  
  PL <- do.call(cdf, c(list(q = deduct), param))  
  PX <- do.call(cdf, c(list(q = x), param))  
  fX <- do.call(pdf, c(list(x = x), param))  
  lik.contr <- ifelse(limitI, log(1 - PX), log(fX)) - log(1 -  
    PL)  
  return(-sum(lik.contr))  
}
```

Fitting the distribution

Let's try gamma. Note that our objective function needs starting values for the optimisation. What other starting values could we use?

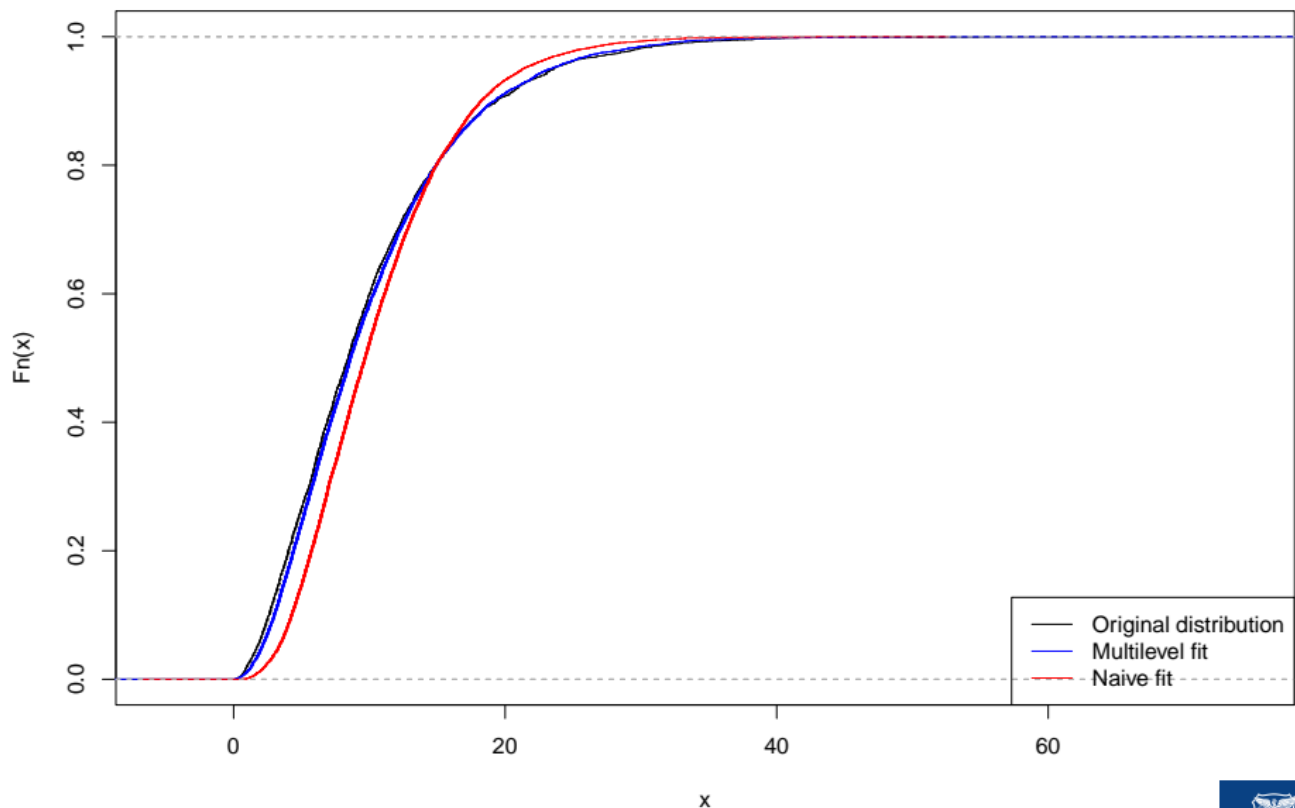
```
pdf <- dgamma
cdf <- pgamma
x <- claims$x
deduct <- claims$deduct
limitI <- claims$limitI
# MME for starting values
start <- list(shape = mean(x)^2/var(x), rate = mean(x)/var(x))
obj.fun <- function(shape, rate) {
  param <- list(shape = shape, rate = rate)
  return(negloglik(pdf, cdf, param, x, deduct, limitI))
} # we now have a function to minimise wrt shape and rate
gamma.ll.fit <- stats4::mle(obj.fun, start = start, lower = c(0,
  0))
summary(gamma.ll.fit)
param.g.ll <- stats4::coef(gamma.ll.fit)
param.g.ll
```

```
## Maximum likelihood estimation
##
## Call:
## stats4::mle(minuslogl = obj.fun, start = start, lower = c(0,
##           0))
##
## Coefficients:
##             Estimate Std. Error
## shape 2.1297568 0.089216920
## rate  0.2111871 0.008274053
##
## -2 log L: 14707.6
##
##      shape      rate
## 2.1297568 0.2111871
```

How did we go?

```
fit.tcens.param <- param.g.ll # from the proper fit
fit.param <- coef(fitdistrplus::fitdist(claims$x, "gamma", method = "mle"))
# this is a naive fit
sim.tcens.gamma <- rgamma(10000, shape = fit.tcens.param[1],
  rate = fit.tcens.param[2]) # sample from proper fit
sim.gamma <- rgamma(10000, shape = fit.param[1], rate = fit.param[2])
# sample from naive fit
# Comparing the proper fit (that accounts for l-trunc and
# r-cens) with a 'naive' fit (that does not account for
# those)
plot(ecdf(orig_x), main = "Empirical CDF plots", col = "black")
lines(ecdf(sim.tcens.gamma), col = "blue", type = "s")
lines(ecdf(sim.gamma), col = "red", type = "s")
legend("bottomright", legend = c("Original distribution", "Multilevel fit",
  "Naive fit"), lty = 1, col = c("black", "blue", "red"))
```

Empirical CDF plots



- 1 Introduction
- 2 Data analysis and descriptive statistics (MW 3.1)
- 3 Selected parametric claims size distributions (MW 3.2)
- 4 Fitting of distributions (MW3.2)
- 5 Model selection (MW 3.3)
- 6 ✕ Other advanced topics
- 7 Calculating within layers for claim sizes (MW 3.4)

5 Model selection (MW 3.3)

- Graphical approaches
- Hypothesis tests
- Information criteria
- SUVA GOF stats

Graphical approaches

For judging quality of model, do some graphical comparisons:

- histogram vs. fitted parametric density function;
- empirical CDF vs fitted parametric CDF;
- probability-probability (P-P) plot - theoretical vs empirical cumulative probabilities;
- quantile-quantile (Q-Q) plot - theoretical vs sample quantiles.

Let the (theoretical) fitted parametric distribution be denoted by $G(x; \hat{\theta})$.

P-P plot

To construct the P-P plot:

- order the observed data from smallest to largest: $x_{(1)}, x_{(2)}, \dots, x_{(n)}$.
- calculate the theoretical CDF at each of the observed data points:
 $G(x_{(i)}; \hat{\theta})$.
- for $i = 1, 2, \dots, n$, plot the points $\frac{i - 0.5}{n}$ against $G(x_{(i)}; \hat{\theta})$.

Note that using $\frac{i - 0.5}{n}$ is Hazen's rule, as recommended by Blom (1959).

see also the following video on YouTube:

- About the choice of $(i - 0.5)/n$ in Q-Q and P-P plots

Q-Q plot

To construct the Q-Q plot:

- order the observed data from smallest to largest: $x_{(1)}, x_{(2)}, \dots, x_{(n)}$.
- for $i = 1, 2, \dots, n$, calculate the theoretical quantiles: $G^{-1}\left(\frac{i - 0.5}{n}; \hat{\theta}\right)$.
- for $i = 1, 2, \dots, n$, plot the points $x_{(i)}$ against $G^{-1}\left(\frac{i - 0.5}{n}; \hat{\theta}\right)$.

These constructions hold only for the case where you have no censoring/truncation.

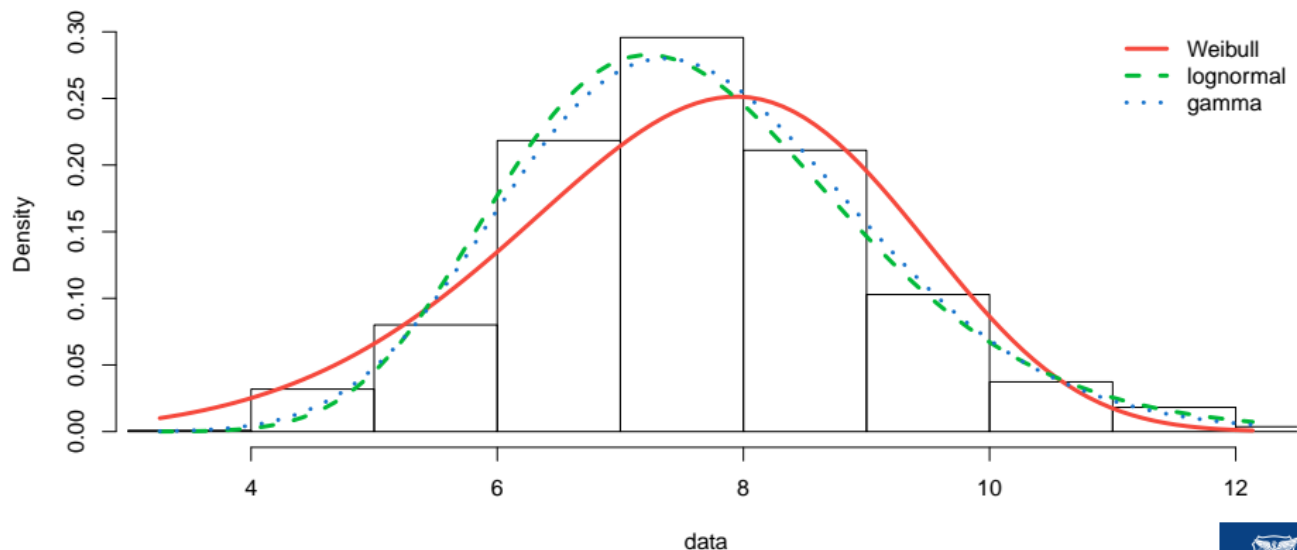
see also the following video on YouTube:

- How to build Q-Q plots and P-P plots

SUVA GOF graphs

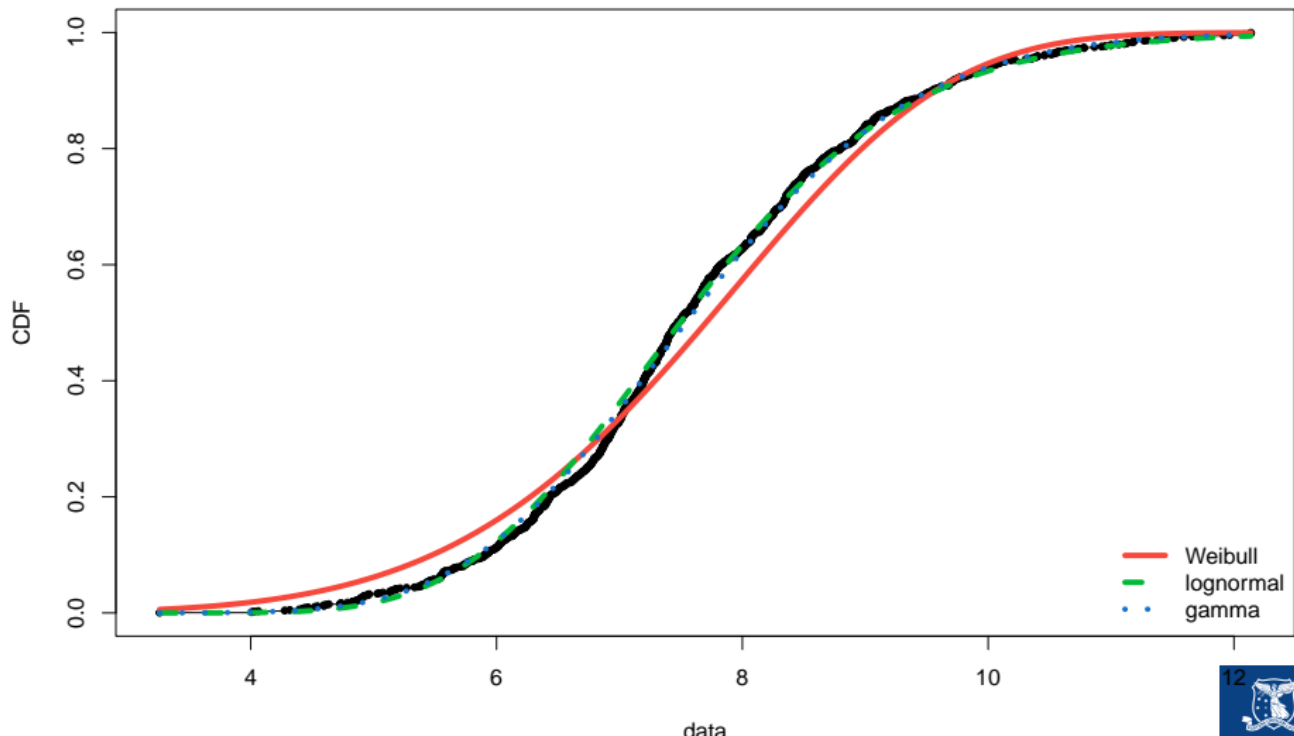
```
plot.legend <- c("Weibull", "lognormal", "gamma")
fitdistrplus::denscomp(list(fit.weibull.mle, fit.lnorm.mle, fit.gamma.mle),
  legendtext = plot.legend, fitlwd = 3)
```

Histogram and theoretical densities



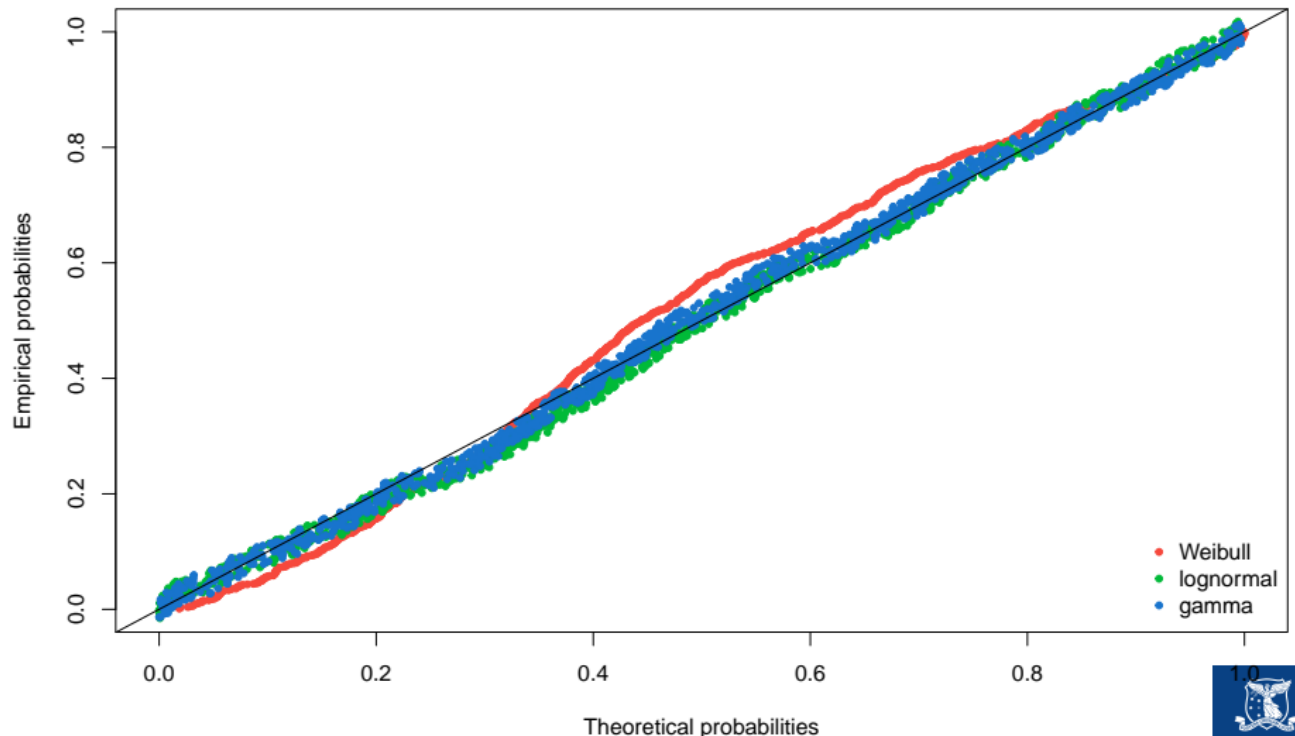
```
fitdistrplus::cdfcomp(list(fit.weibull.mle, fit.lnorm.mle, fit.gamma.mle),  
  legendtext = plot.legend, fitlwd = 4, datapch = 20)
```

Empirical and theoretical CDFs



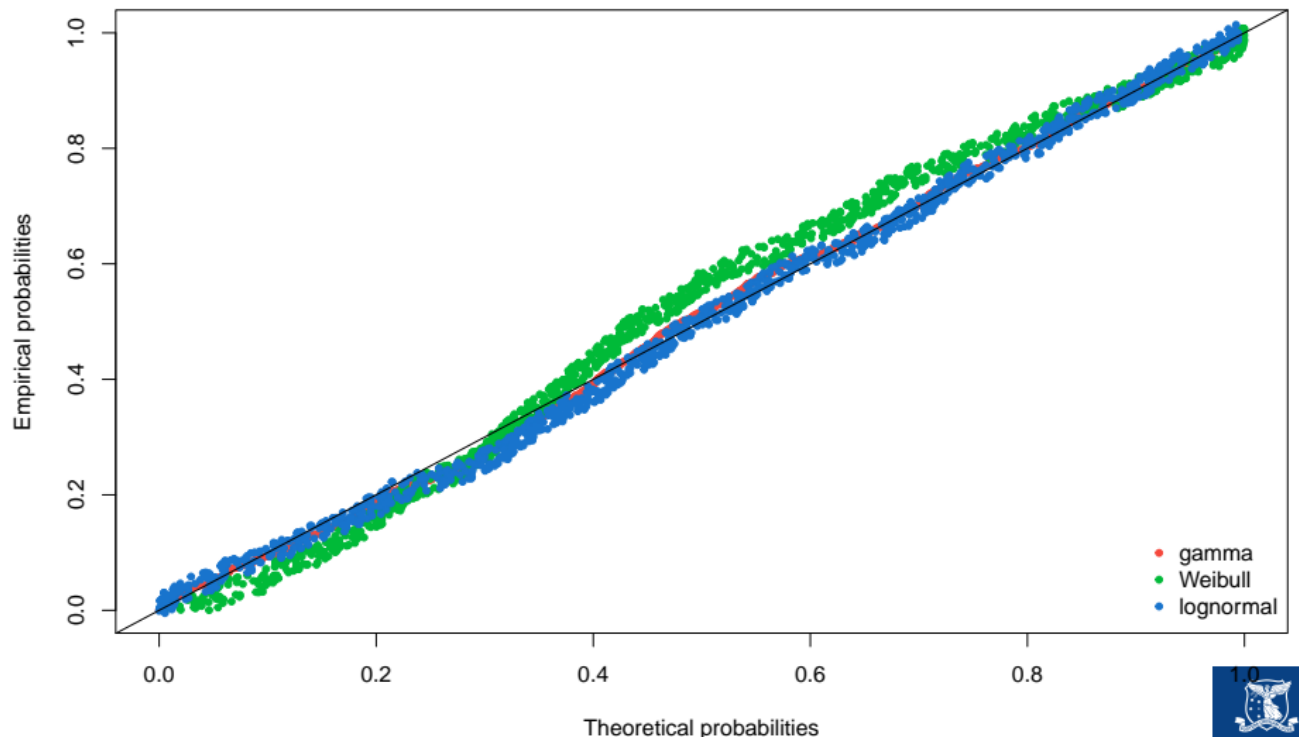
```
fitdistrplus::ppcomp(list(fit.weibull.mle, fit.lnorm.mle, fit.gamma.mle),  
  legendtext = plot.legend, fitpch = 20)
```

P-P plot

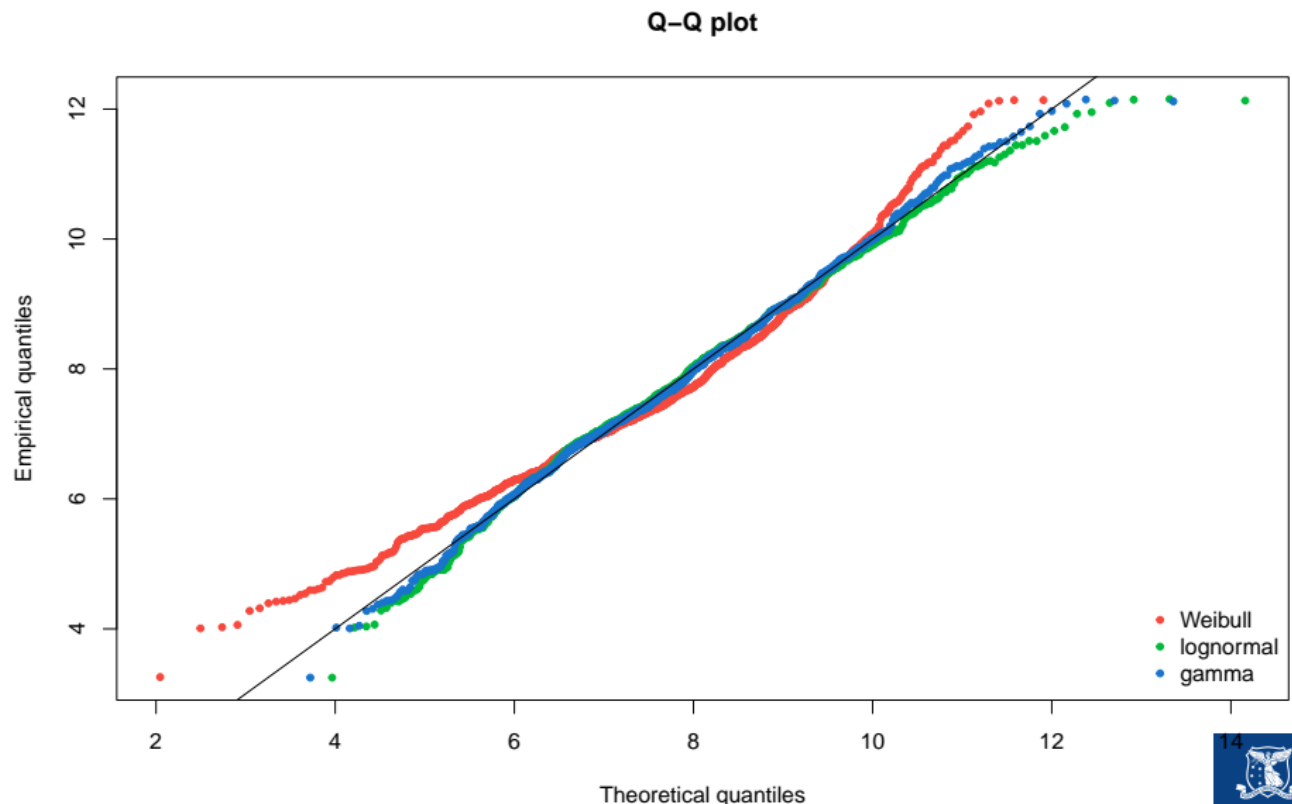


```
fitdistrplus::ppcomp(list(fit.gamma.mle, fit.weibull.mle, fit.lnorm.mle),  
  legendtext = c("gamma", "Weibull", "lognormal"), fitpch = 20) #the order
```

P-P plot



```
fitdistrplus::qqcomp(list(fit.weibull.mle, fit.lnorm.mle, fit.gamma.mle),  
  legendtext = plot.legend, fitpch = 20)
```



5 Model selection (MW 3.3)

- Graphical approaches
- Hypothesis tests
- Information criteria
- SUVA GOF stats

Hypothesis tests

We will test the null

- H_0 : data came from population with the specified model, against
- H_a : the data did not come from such a population.

Some commonly used tests, and their test statistics:

- Kolmogorov-Smirnoff: $K.S. = \sup_y |\hat{G}(y) - G(y; \hat{\theta})|$
- Anderson-Darling: $A.D. = n \int \frac{[\hat{G}(y) - G(y; \hat{\theta})]^2}{G(y; \hat{\theta}) [1 - G(y; \hat{\theta})]} dG(y)$
- χ^2 goodness-of-fit: $\chi^2 = \sum_j \frac{(observed - expected)^2}{expected}$

Computational formulas for those tests are available in Delignette-Muller and Dutang (2015).

Kolmogorov-Smirnoff test

$$K.S. = \sup_y |\hat{G}(y) - G(y; \hat{\theta})|,$$

where

- $\hat{G}(y)$ is the empirical distribution
- $G(y; \theta)$ is the assumed theoretical distribution in the null hypothesis
- $G(y; \theta)$ is assumed to be (must be) continuous
- $\hat{\theta}$ is the maximum likelihood estimate for θ under the null hypothesis

Note:

- There are tables for the critical values. There are several variations of these tables in the literature that use somewhat different scalings for the K-S test statistic and critical regions.
- The test (obviously) does not work for grouped data.

Anderson-Darling test

$$A.D. = n \int \frac{[\hat{G}(y) - G(y; \hat{\theta})]^2}{G(y; \hat{\theta}) [1 - G(y; \hat{\theta})]} dG(y),$$

where n is the sample size.

Note:

- The critical values for the Anderson-Darling test are dependent on the specific distribution that is being tested. There are tabulated values and formulas for a few specific distributions.
- The theoretical distribution is assumed to be (must be) continuous.
- The test does not work for grouped data.

χ^2 goodness-of-fit test

Procedure:

- Break down the whole range into k subintervals:
 $c_0 < c_1 < \cdots < c_k = \infty$
- χ^2 goodness-of-fit test: $\chi^2 = \sum_j^k \frac{(E_j - O_j)^2}{E_j}$
- Let $\hat{p}_j = G(c_j; \hat{\theta}) - G(c_{j-1}; \hat{\theta})$. Then, the number of expected observations in the interval $(c_{j-1}, c_j]$ assuming that the hypothesized model is true:

$$E_j = n\hat{p}_j \quad (\text{Here, } n \text{ is the sample size})$$

- Let $p_j = \hat{G}(c_j) - \hat{G}(c_{j-1})$. Then, the number of observations in the interval $(c_{j-1}, c_j]$:

$$O_j = np_j$$

The statistic has chi-square distribution with the degree of freedom equal to: $k - 1 - \text{number of parameters estimated}$

How these tests are used

- Besides testing whether data came from specified model or not, generally we would prefer models with:
 - lowest K-S test statistic
 - lowest A-D test statistic
 - lowest χ^2 goodness-of-fit test statistic (or equivalently highest p -value)
 - highest value of the likelihood function at the maximum
- Perform formal statistical test, or use as 'horse race'

Comparison

- K-S and A-D tests are quite similar - both look at the difference between the empirical and model distribution functions.
 - K-S in absolute value, A-D in squared difference.
 - But A-D is weighted average, with more emphasis on good fit in the tails than in the middle; K-S puts no such emphasis.
- For K-S and A-D tests, no adjustments are made to account for increase in the number of parameters, nor sample size. Result: more complex models often will fare better on these tests.
- The χ^2 test adjusts the degrees of freedom for increases in the number of parameters, but is easily manipulated as the choice of brackets is arbitrary.

5 Model selection (MW 3.3)

- Graphical approaches
- Hypothesis tests
- Information criteria
- SUVA GOF stats

Information criteria

- Information criteria penalise the log likelihood with a function that depends on the number of parameters.
- Akaike Information Criterion (AIC):

$$\text{AIC}^{(i)} = -2\ell_{\mathbf{Y}}^{(i)} + 2d^{(i)},$$

where $d^{(i)}$ denotes the number of estimated parameters in the density g_i that is considered, and where $\ell_{\mathbf{Y}}^{(i)}$ is the maximum log likelihood that can be achieved with that density.

- Bayesian Information Criterion (BIC)

$$\text{BIC}^{(i)} = -2\ell_{\mathbf{Y}}^{(i)} + \log(n)d^{(i)}.$$

This is -2SBC , where SBC is Schwarz Bayesian Criterion.

- Note that these can sometimes be presented in inverse sign. Here, because the penalty is added, it is obvious that smaller values of the IC are preferred.

5 Model selection (MW 3.3)

- Graphical approaches
- Hypothesis tests
- Information criteria
- SUVA GOF stats

GOF hypothesis test statistics

For MLE:

```
gofstat(list(fit.weibull.mle, fit.lnorm.mle, fit.gamma.mle),  
       fitnames = plot.legend)
```

```
## Goodness-of-fit statistics  
##                                     Weibull    lognormal      gamma  
## Kolmogorov-Smirnov statistic  0.07105097 0.04276791 0.03376236  
## Cramer-von Mises statistic   1.74049707 0.26568341 0.19438834  
## Anderson-Darling statistic   10.69572021 1.70209314 1.10385675  
##  
## Goodness-of-fit criteria  
##                                     Weibull    lognormal      gamma  
## Akaike's Information Criterion 4010.519   3920.717   3907.636  
## Bayesian Information Criterion 4020.524   3930.722   3917.641
```

For MME:

```
gofstat(list(fit.weibull.mme, fit.lnorm.mme, fit.gamma.mme),  
       fitnames = plot.legend)  
  
## Goodness-of-fit statistics  
##                                     Weibull    lognormal     gamma  
## Kolmogorov-Smirnov statistic  0.08023228 0.0374087 0.0327241  
## Cramer-von Mises statistic   1.66608962 0.1869752 0.1823197  
## Anderson-Darling statistic  10.53097690 1.4369047 1.0539353  
##  
## Goodness-of-fit criteria  
##                                     Weibull    lognormal     gamma  
## Akaike's Information Criterion 4044.239  3922.121 3907.677  
## Bayesian Information Criterion 4054.243  3932.125 3917.681
```

Note that when fitting discrete distributions, the chi-squared statistic is computed by the `gofstat` function (see Delignette-Muller and Dutang (2015) for details).

GOF hypothesis test results

R can also provide the results from the GOF hypothesis tests. For instance:

```
gammagof <- gofstat(list(fit.gamma.mle, fit.lnorm.mle), fitnames = c("gamma", "lognormal MLE"), chisqbreaks = c(10:20/2))
gammagof$chisqvalue

##      gamma MLE lognormal MLE
## 1.374256e-03 1.690633e-05

gammagof$adtest

##      gamma MLE lognormal MLE
## "rejected" "not computed"

gammagof$kstest

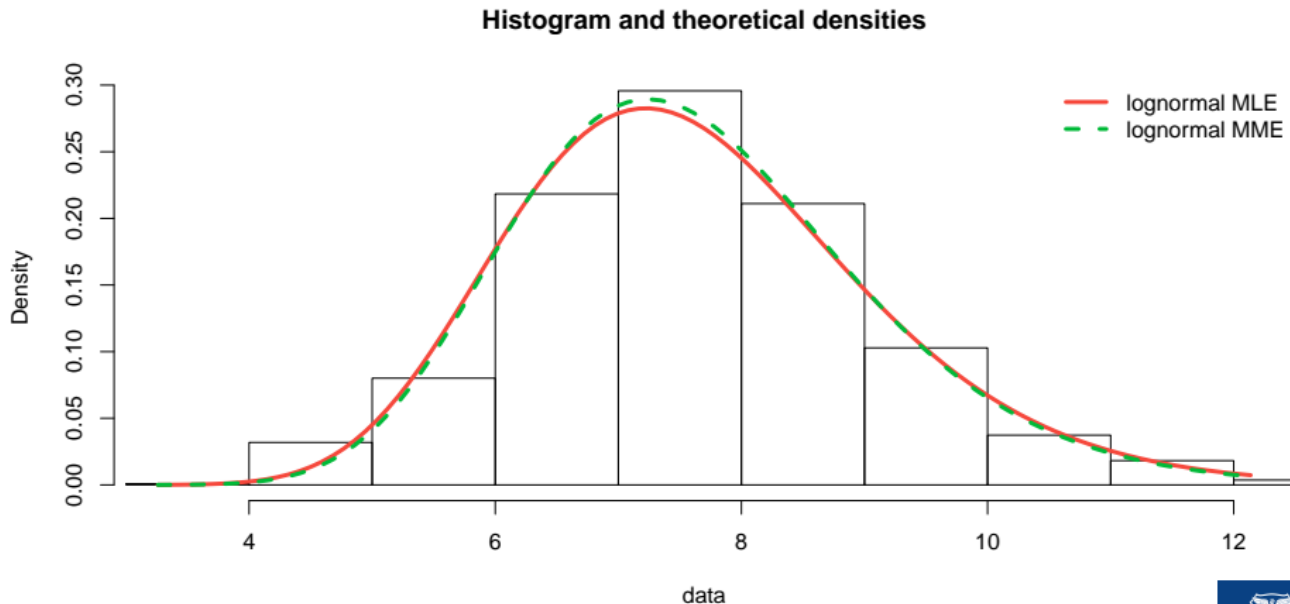
##      gamma MLE lognormal MLE
## "not rejected" "rejected"
```

gammagof\$chisqtable

	obscounts	theo gamma MLE	theo lognormal MLE
## <= 5	36	23.96251	19.19169
## <= 5.5	28	39.64865	39.53512
## <= 6	60	72.77895	76.73318
## <= 6.5	110	109.47716	116.38286
## <= 7	130	139.03193	145.08202
## <= 7.5	191	152.62949	154.45787
## <= 8	134	147.62688	144.65495
## <= 8.5	141	127.77796	121.96771
## <= 9	91	100.25469	94.30131
## <= 9.5	62	72.07629	67.84850
## <= 10	51	47.91524	45.97086
## > 10	65	65.82026	72.87394

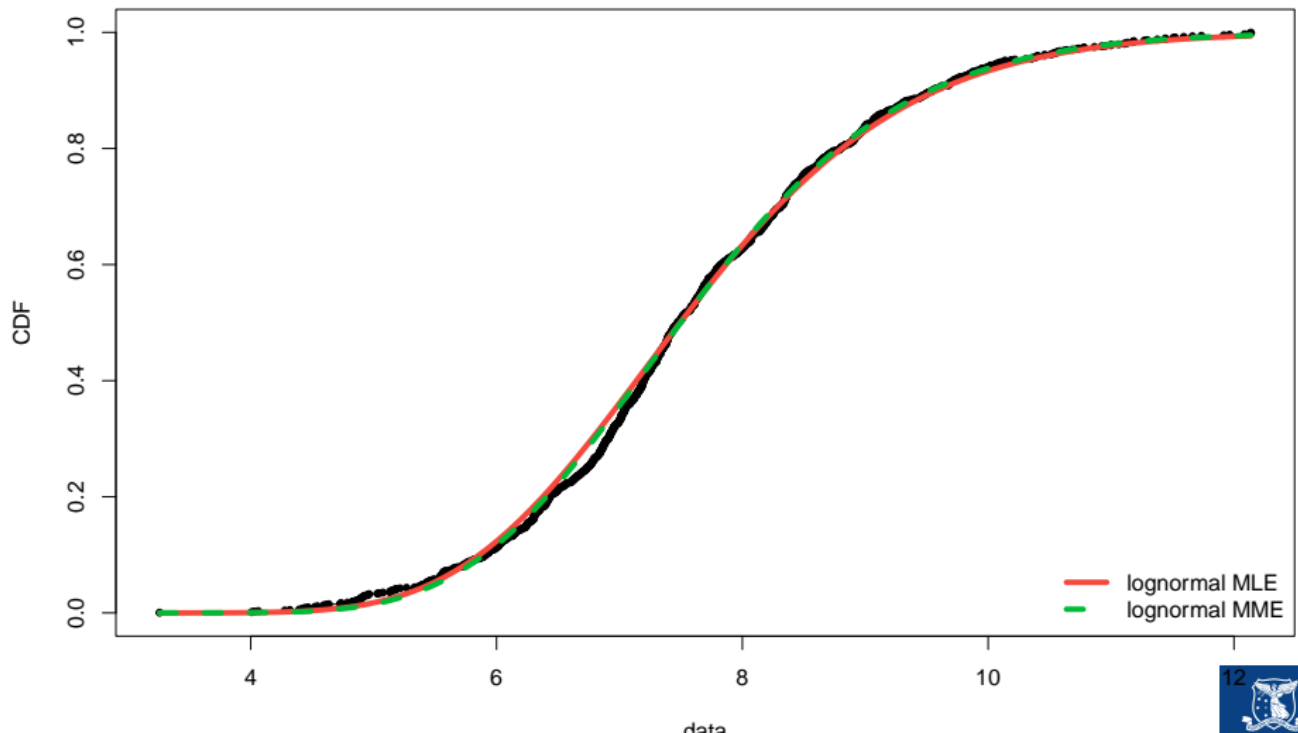
GOF graphical comparisons

```
plot.legend <- c("lognormal MLE", "lognormal MME")
fitdistrplus::denscomp(list(fit.lnorm.mle, fit.lnorm.mme), legendtext = plot.legend,
fitlwd = 3)
```

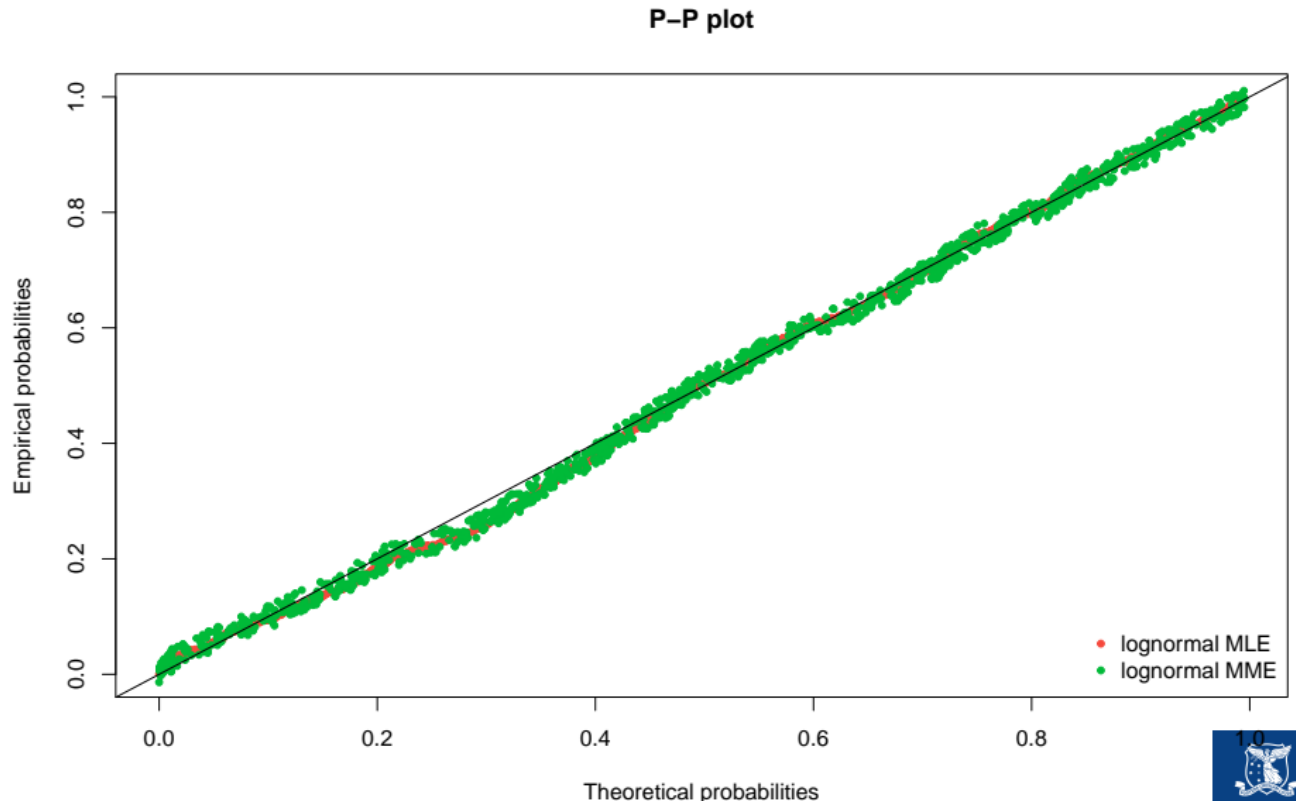


```
fitdistrplus::cdfcomp(list(fit.lnorm.mle, fit.lnorm.mme), legendtext = plot  
fitlwd = 4, datapch = 20)
```

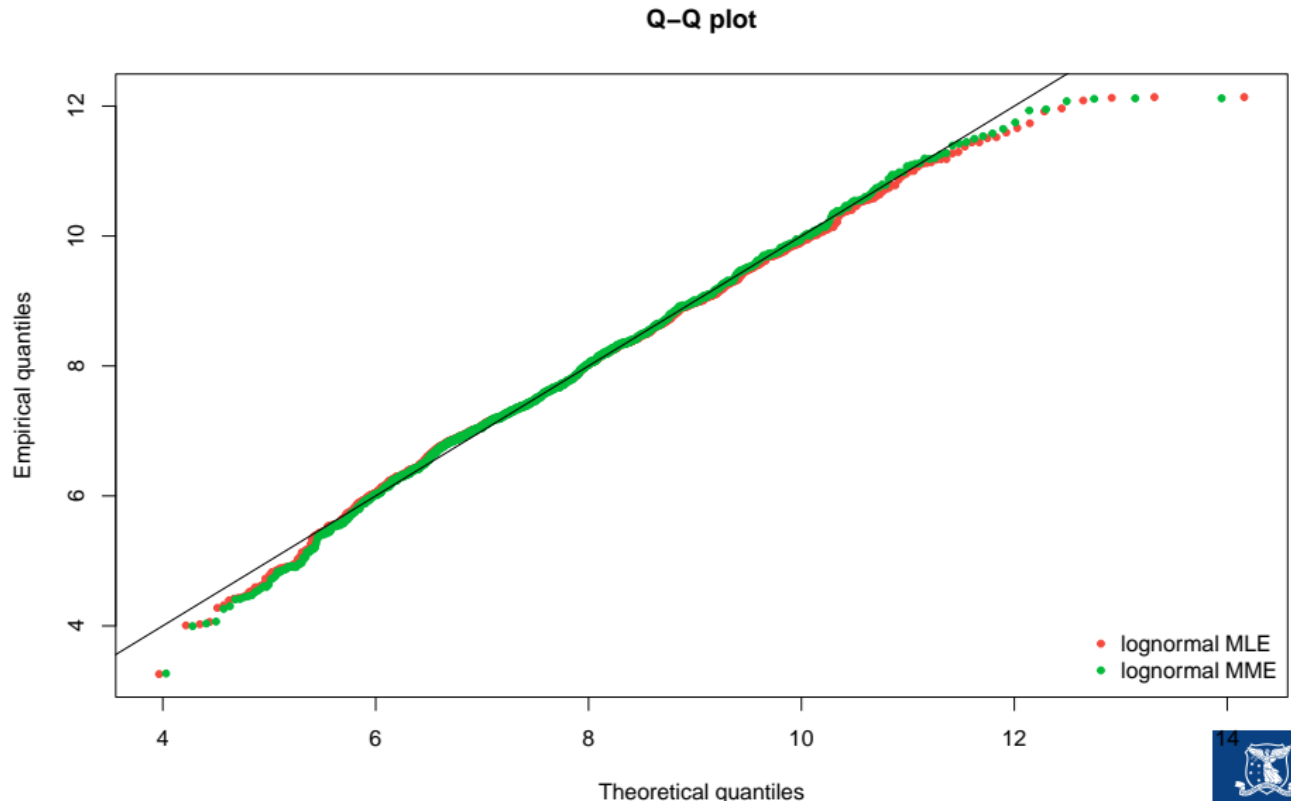
Empirical and theoretical CDFs



```
fitdistrplus::ppcomp(list(fit.lnorm.mle, fit.lnorm.mme), legendtext = plot.  
fitpch = 20)
```

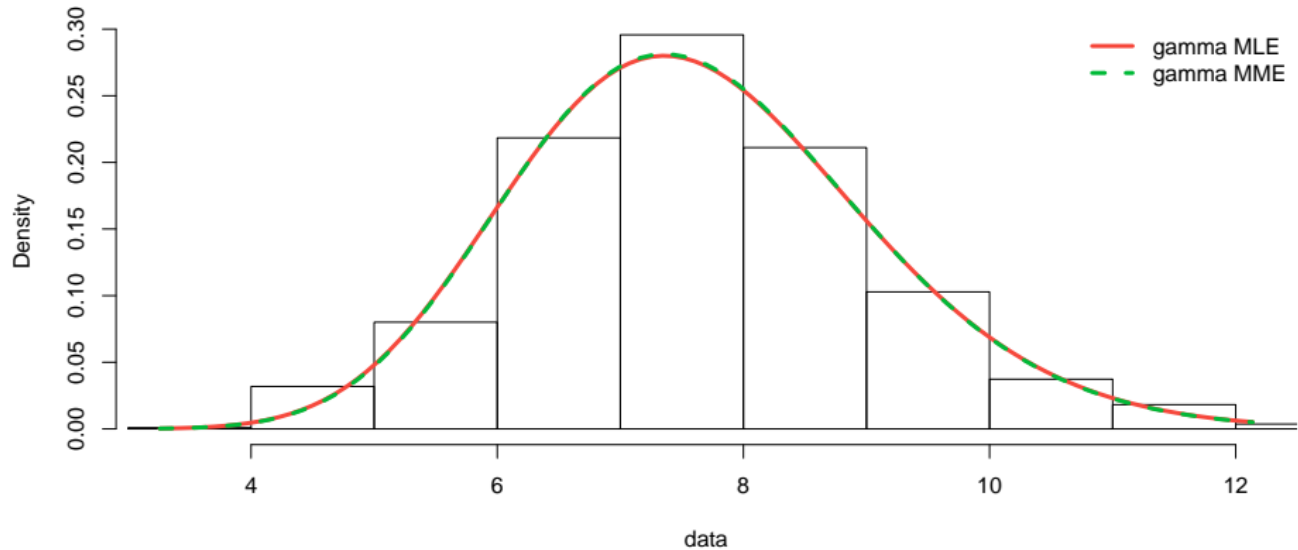


```
fitdistrplus::qqcomp(list(fit.lnorm.mle, fit.lnorm.mme), legendtext = plot.  
fitpch = 20)
```



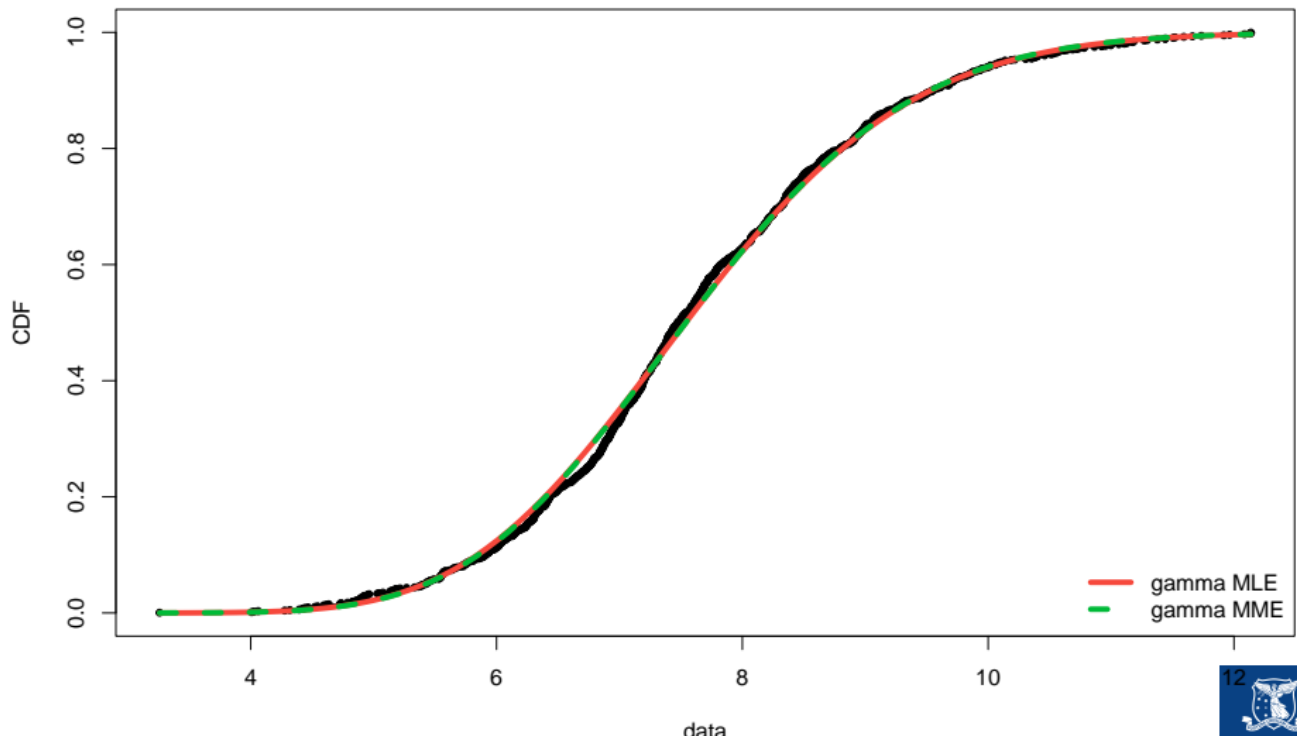
```
plot.legend <- c("gamma MLE", "gamma MME")
fitdistrplus::denscomp(list(fit.gamma.mle, fit.gamma.mme), legendtext = plot.legend,
fitlwd = 3)
```

Histogram and theoretical densities



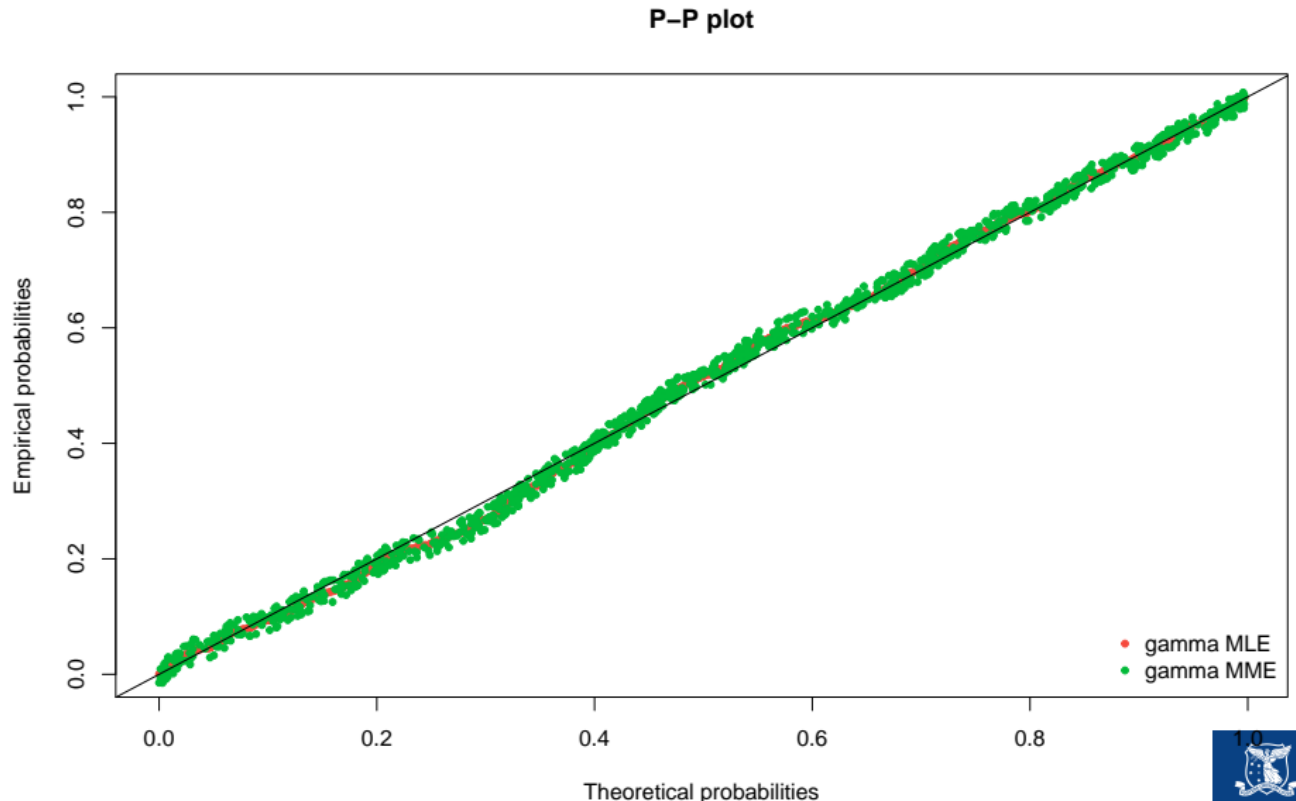
```
fitdistrplus::cdfcomp(list(fit.gamma.mle, fit.gamma.mme), legendtext = plot  
fitlwd = 4, datapch = 20)
```

Empirical and theoretical CDFs



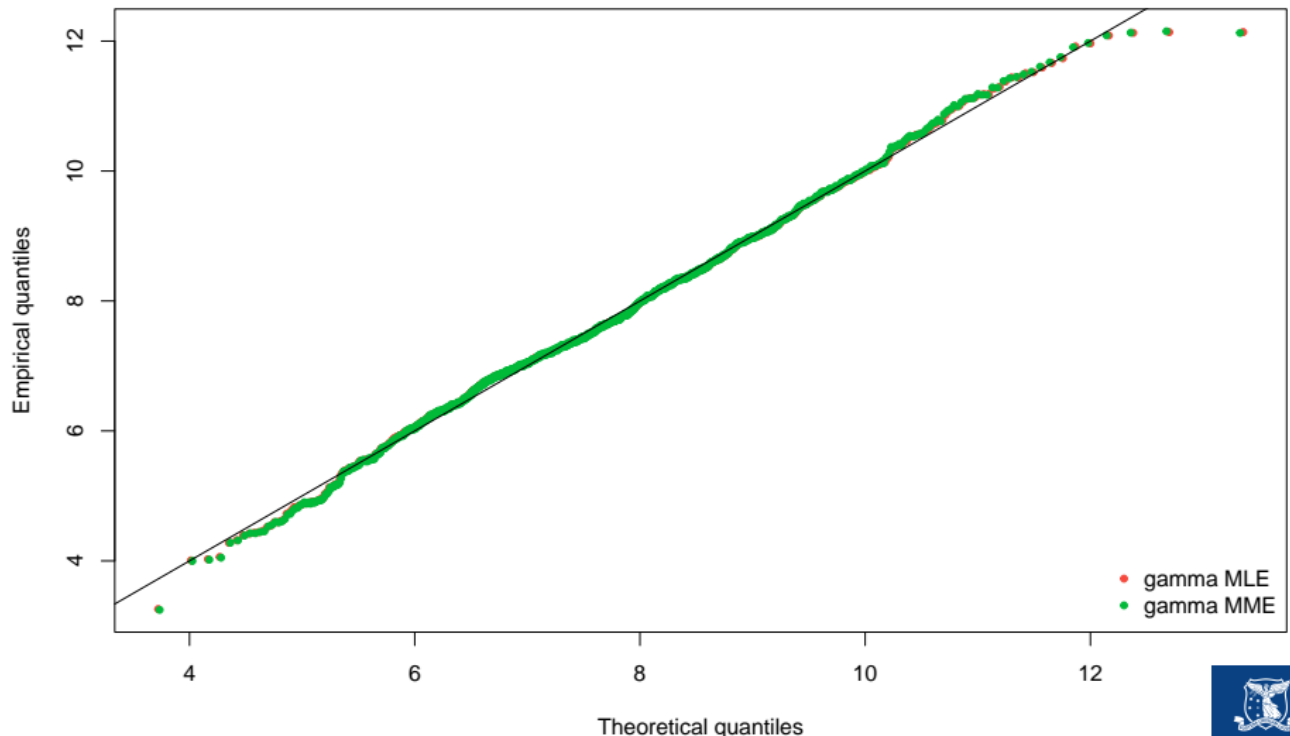
gamma MLE
gamma MME

```
fitdistrplus::ppcomp(list(fit.gamma.mle, fit.gamma.mme), legendtext = plot.  
fitpch = 20)
```



```
fitdistrplus::qqcomp(list(fit.gamma.mle, fit.gamma.mme), legendtext = plot.  
fitpch = 20)
```

Q-Q plot



- 1 Introduction
- 2 Data analysis and descriptive statistics (MW 3.1)
- 3 Selected parametric claims size distributions (MW 3.2)
- 4 Fitting of distributions (MW3.2)
- 5 Model selection (MW 3.3)
- 6 ✖ Other advanced topics
- 7 Calculating within layers for claim sizes (MW 3.4)

6 ✖ Other advanced topics

- ✖ Alternative methods for estimation
- ✖ Zero-inflated severity model $X = IB$

✖ Maximum Goodness Estimation (MGE)

- This is one form of “minimum distance estimation”, whereby parameters are chosen such that a distance (between empirical and theoretical) is minimised
- Here we focus on the GOF tests AD, CvM, and KS
- This can be readily chosen in R using `fitdist`, with `method="mge"` and where `gof=` one of AD, CvM, or KS.

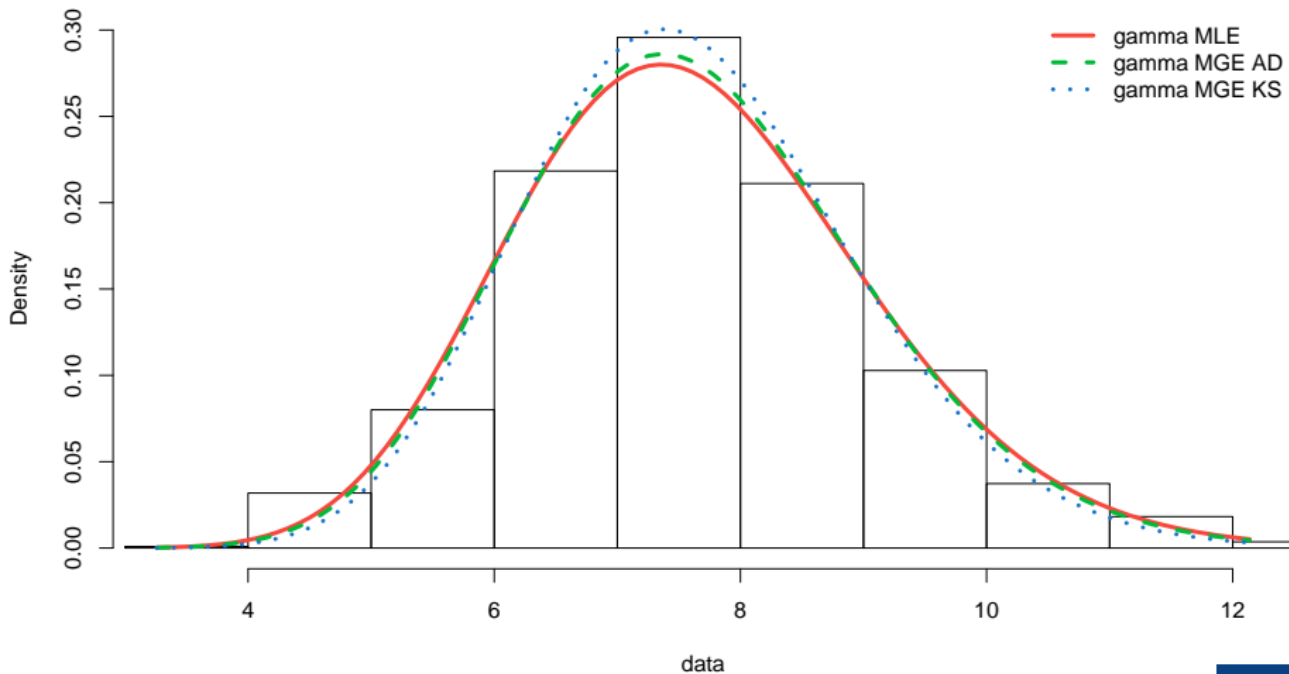
With the SUVA data:

```
fit.gamma.mge.ad <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "gamma", method = "mge", gof = "AD")
fit.gamma.mge.ks <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "gamma", method = "mge", gof = "KS")
gof.mge.legend <- c("gamma MLE", "gamma MGE AD", "gamma MGE KS")
gofstat(list(fit.gamma.mle, fit.gamma.mge.ad, fit.gamma.mge.ks),
       fitnames = gof.mge.legend)
```

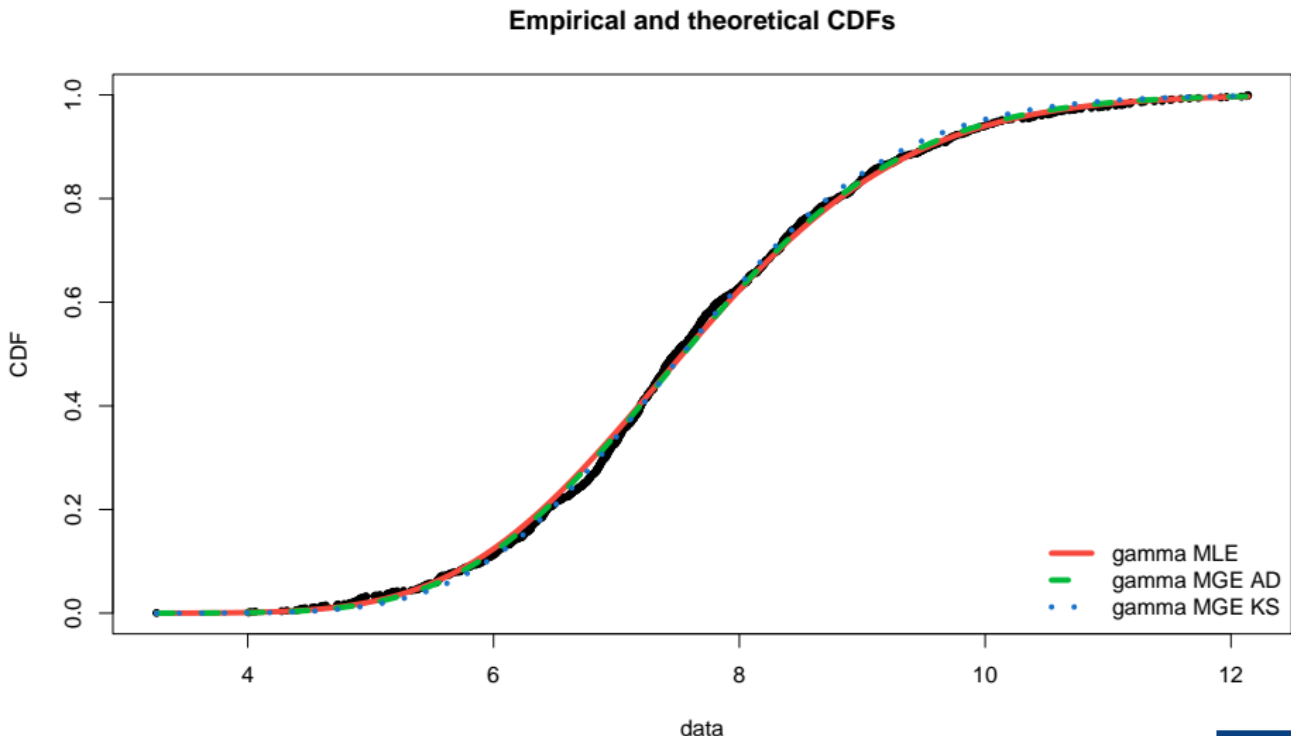
```
## Goodness-of-fit statistics
##                                     gamma MLE gamma MGE AD gamma MGE KS
## Kolmogorov-Smirnov statistic 0.03376236   0.02841676   0.0208791
## Cramer-von Mises statistic  0.19438834   0.14533386   0.1444858
## Anderson-Darling statistic  1.10385675   0.96063188   1.6991376
##
## Goodness-of-fit criteria
##                                     gamma MLE gamma MGE AD gamma MGE KS
## Akaike's Information Criterion 3907.636      3908.736     3919.475
## Bayesian Information Criterion 3917.641      3918.740     3929.480
```

```
denscomp(list(fit.gamma.mle, fit.gamma.mge.ad, fit.gamma.mge.ks),  
        legendtext = gof.mge.legend, fitlwd = 3)
```

Histogram and theoretical densities

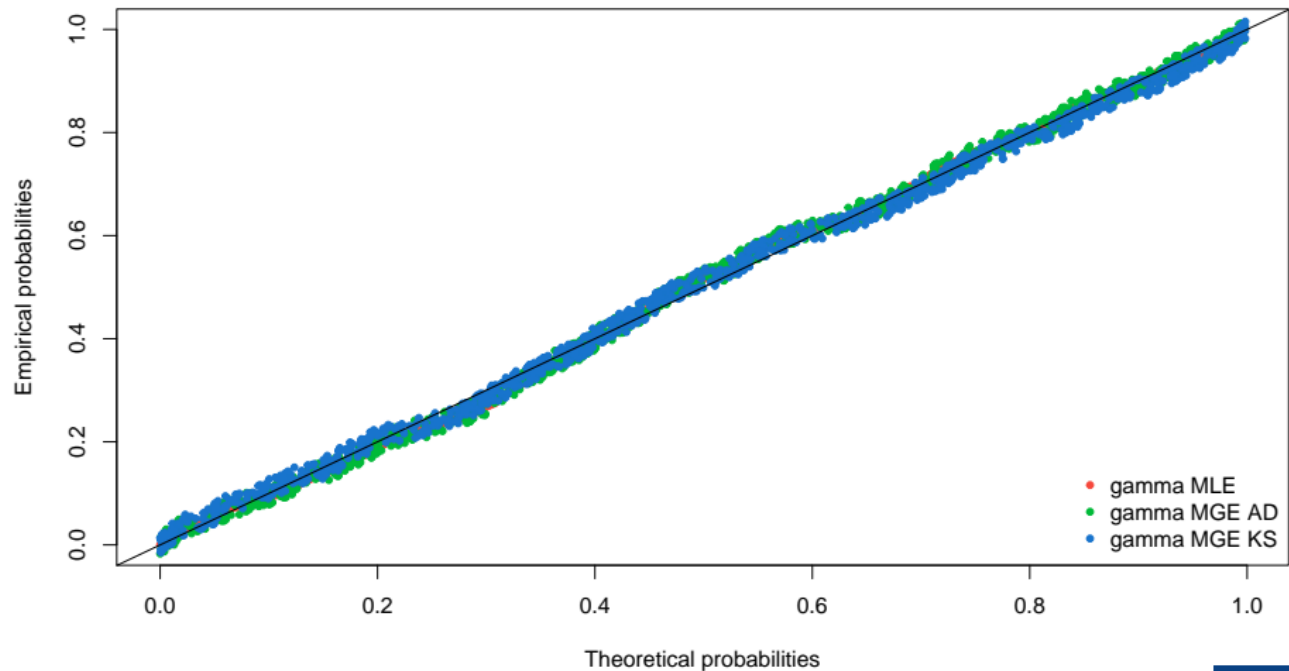


```
cdfcomp(list(fit.gamma.mle, fit.gamma.mge.ad, fit.gamma.mge.ks),  
        legendtext = gof.mge.legend, fitlwd = 4, datapch = 20)
```

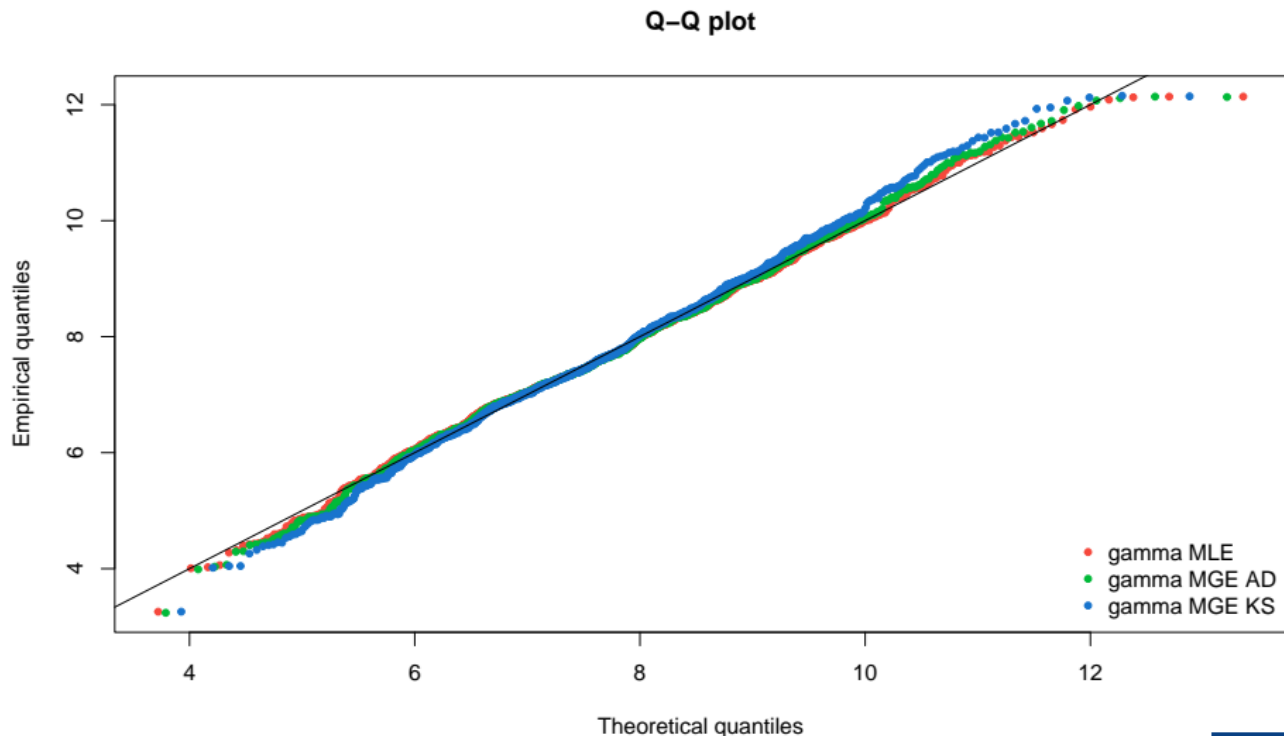


```
ppcomp(list(fit.gamma.mle, fit.gamma.mge.ad, fit.gamma.mge.ks),  
       legendtext = gof.mge.legend, fitpch = 20)
```

P-P plot



```
qqcomp(list(fit.gamma.mle, fit.gamma.mge.ad, fit.gamma.mge.ks),  
       legendtext = gof.mge.legend, fitpch = 20)
```



✖ AD based parameter estimation

- Recall the Anderson-Darling test statistic, which focused on the tails.
- What if we wanted to generalise the idea of AD to left tail or right tail only, or put even more weight on either of those tails?
- The AD test considers a weighted “sum” (integral) of the squared difference between empirical and theoretical cdf’s
- In its original formulation, the weight is of the form

$$w_i = \frac{1}{G(y)[1 - G(y)]},$$

which goes to infinity when $y \rightarrow 0$ or $y \rightarrow \infty$.

- There are 5 alternatives in `fitdistrplus`.

- ADR: Right-tail AD, where

$$w_i = \frac{1}{1 - G(y)};$$

- ADL: Left-tail AD, where

$$w_i = \frac{1}{G(y)};$$

- ADR2: Right-tail AD, 2nd order, where

$$w_i = \frac{1}{[1 - G(y)]^2};$$

- ADL2: Left-tail AD, 2nd order, where

$$w_i = \frac{1}{[G(y)]^2};$$

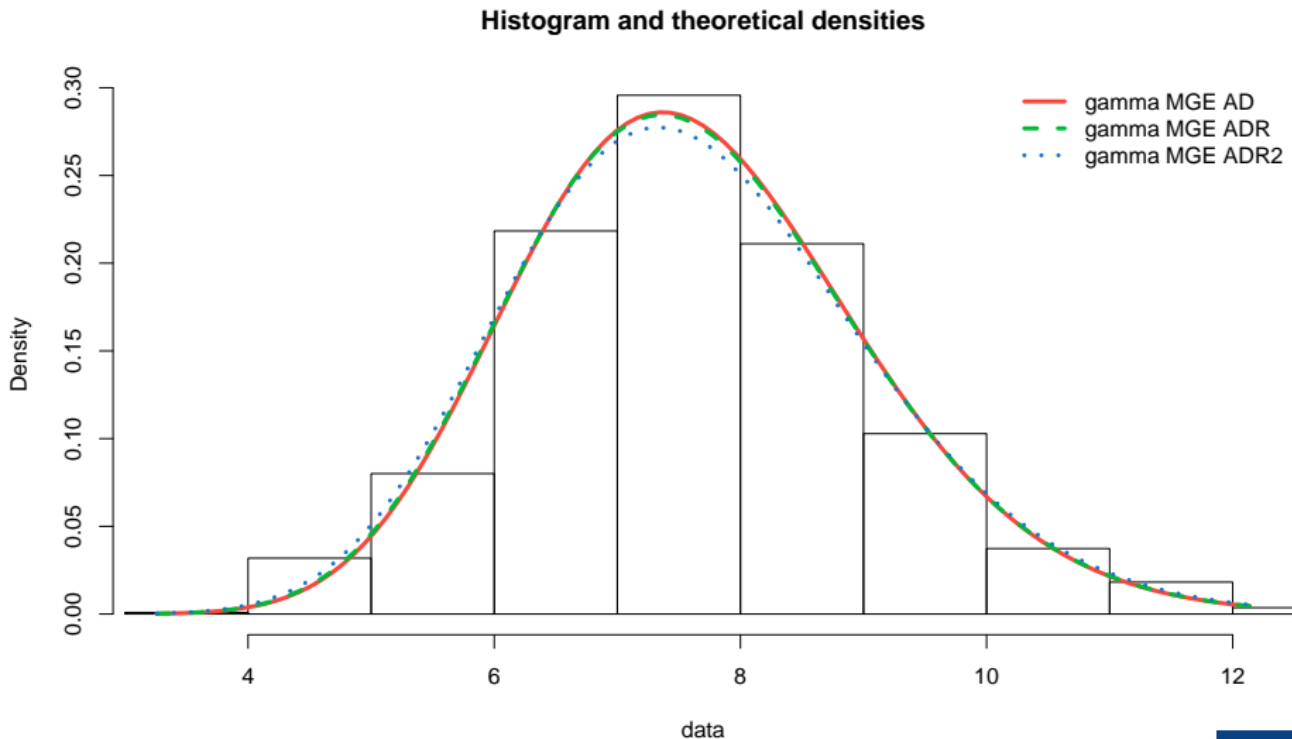
- AD2: AD, 2nd order. Here, R will minimise ADR2 + ADL2.

With the SUVA data:

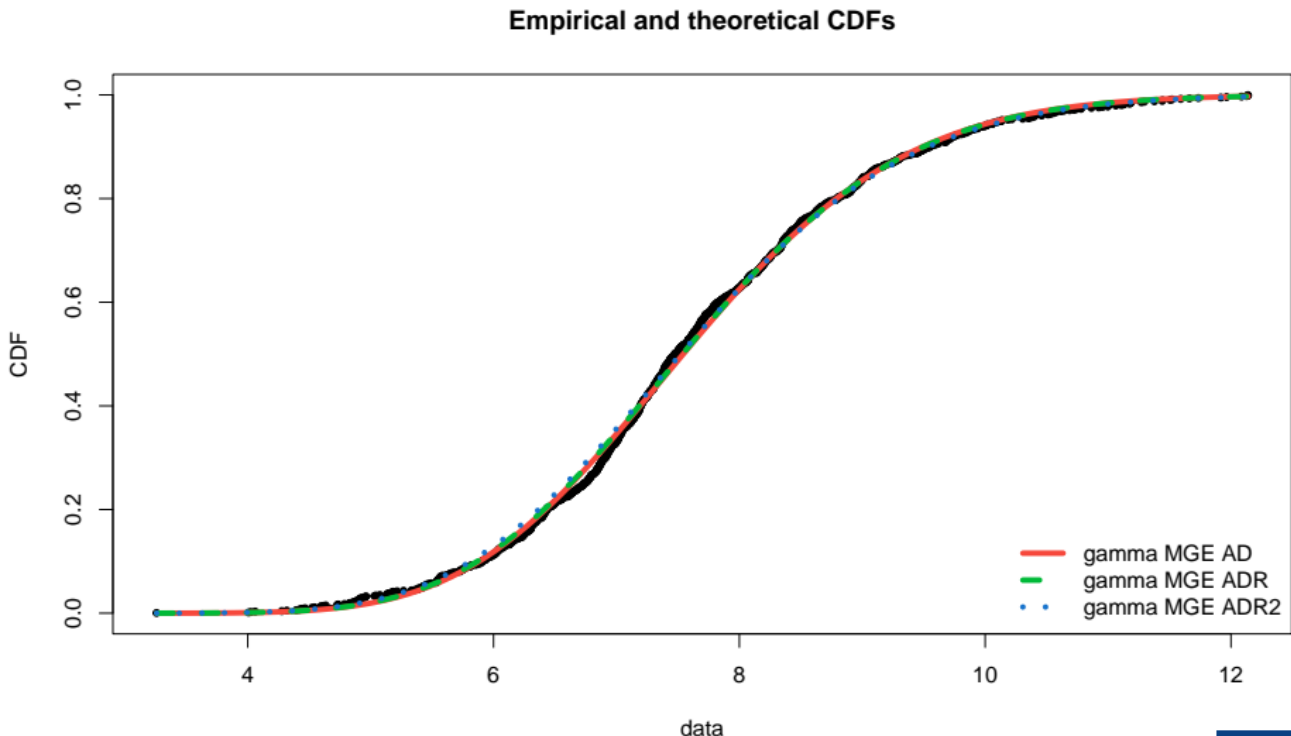
```
fit.gamma.mge.adr <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "gamma", method = "mge", gof = "ADR")
fit.gamma.mge.adr2 <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "gamma", method = "mge", gof = "AD2R")
gof.mge.legend2 <- c("gamma MGЕ AD", "gamma MGЕ ADR", "gamma MGЕ AD2R")
```

```
gofstat(list(fit.gamma.mge.ad, fit.gamma.mge.adr, fit.gamma.mge.adr2),  
        fitnames = gof.mge.legend2)  
  
## Goodness-of-fit statistics  
##                                     gamma MGE AD gamma MGE ADR  
## Kolmogorov-Smirnov statistic      0.02841676   0.0309970  
## Cramer-von Mises statistic       0.14533386   0.1553754  
## Anderson-Darling statistic       0.96063188   0.9799768  
##                                     gamma MGE ADR2  
## Kolmogorov-Smirnov statistic      0.03980158  
## Cramer-von Mises statistic       0.24760685  
## Anderson-Darling statistic       1.38154307  
##  
## Goodness-of-fit criteria  
##                                     gamma MGE AD gamma MGE ADR  
## Akaike's Information Criterion    3908.736     3908.223  
## Bayesian Information Criterion   3918.740     3918.228  
##                                     gamma MGE ADR2  
## Akaike's Information Criterion    3908.049  
## Bayesian Information Criterion   3918.054
```

```
denscomp(list(fit.gamma.mge.ad, fit.gamma.mge.adr, fit.gamma.mge.adr2),  
        legendtext = gof.mge.legend2, fitlwd = 3)
```

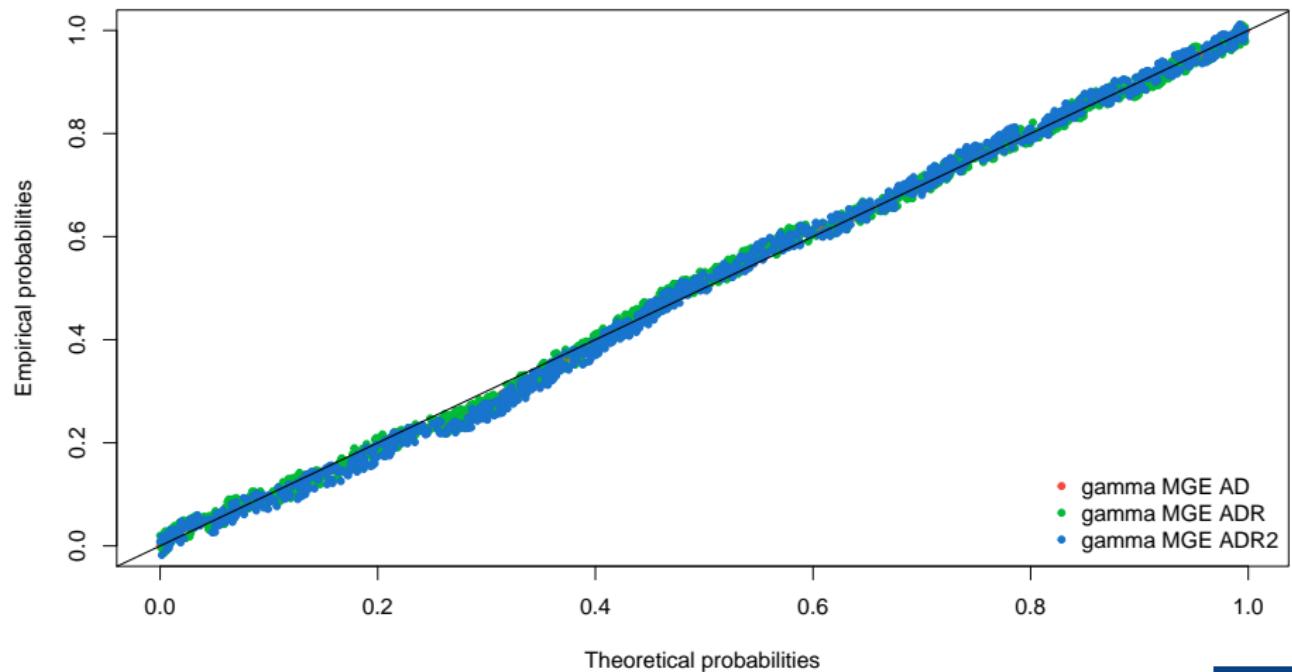


```
cdfcomp(list(fit.gamma.mge.ad, fit.gamma.mge.adr, fit.gamma.mge.adr2),  
        legendtext = gof.mge.legend2, fitlwd = 4, datapch = 20)
```



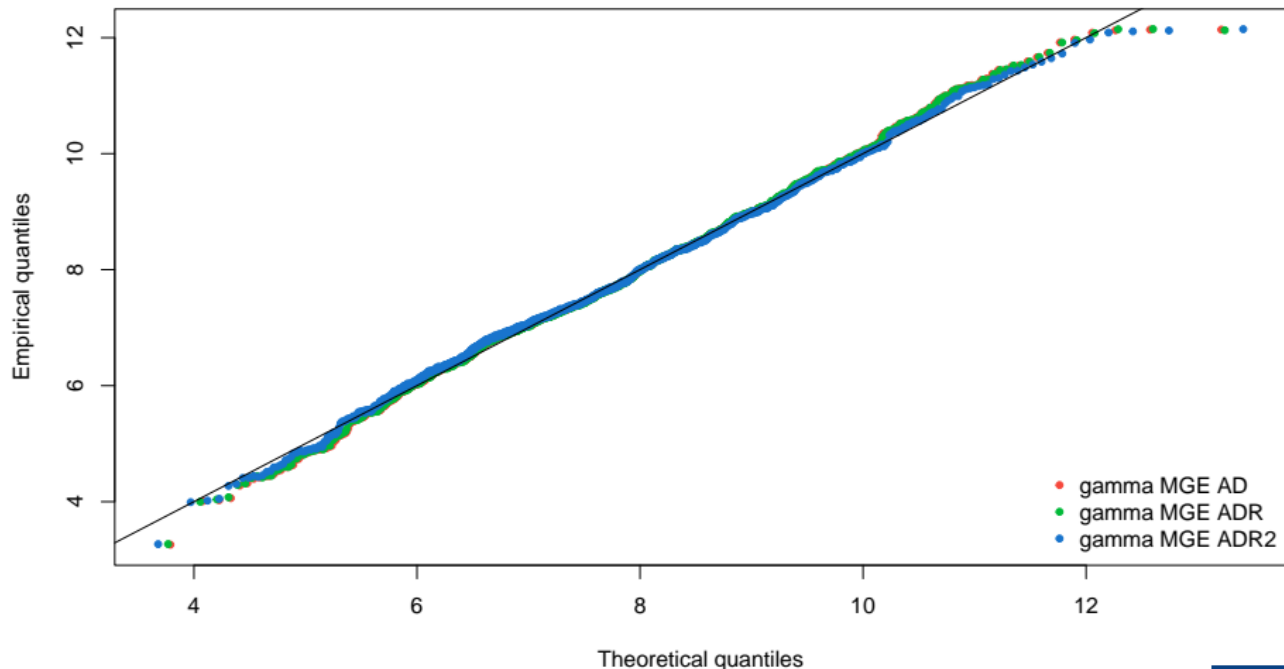
```
ppcomp(list(fit.gamma.mge.ad, fit.gamma.mge.adr, fit.gamma.mge.adr2),  
       legendtext = gof.mge.legend2, fitpch = 20)
```

P-P plot



```
qqcomp(list(fit.gamma.mge.ad, fit.gamma.mge.adr, fit.gamma.mge.adr2),  
       legendtext = gof.mge.legend2, fitpch = 20)
```

Q-Q plot



✖ Quantile matching

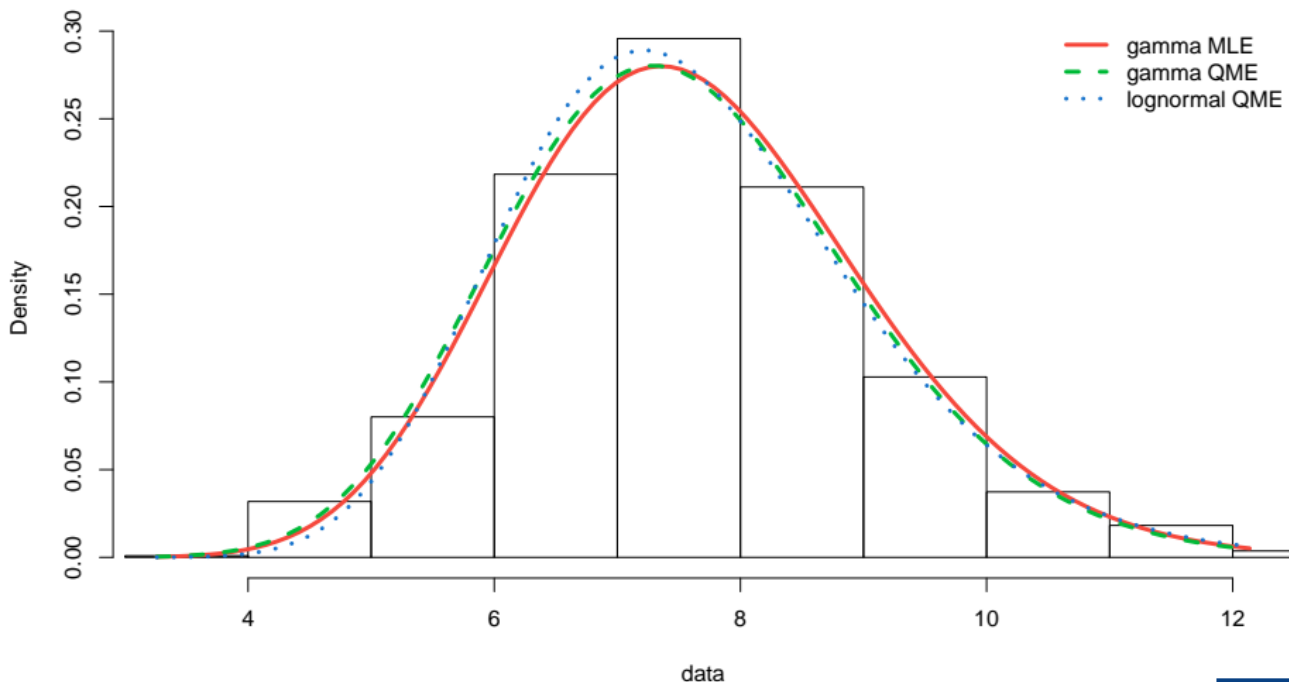
- Quantile matching is easily implemented in R:
 - set `method="qme"` in the call to `fitdist`;
 - and add an argument `probs` defining the probabilities for which the quantile matching is performed.
- The number of quantiles to match must be the same as the number of parameters to estimate.
- The quantile matching is carried out numerically, by minimising the sum of squared differences between observed and theoretical quantiles.

For example:

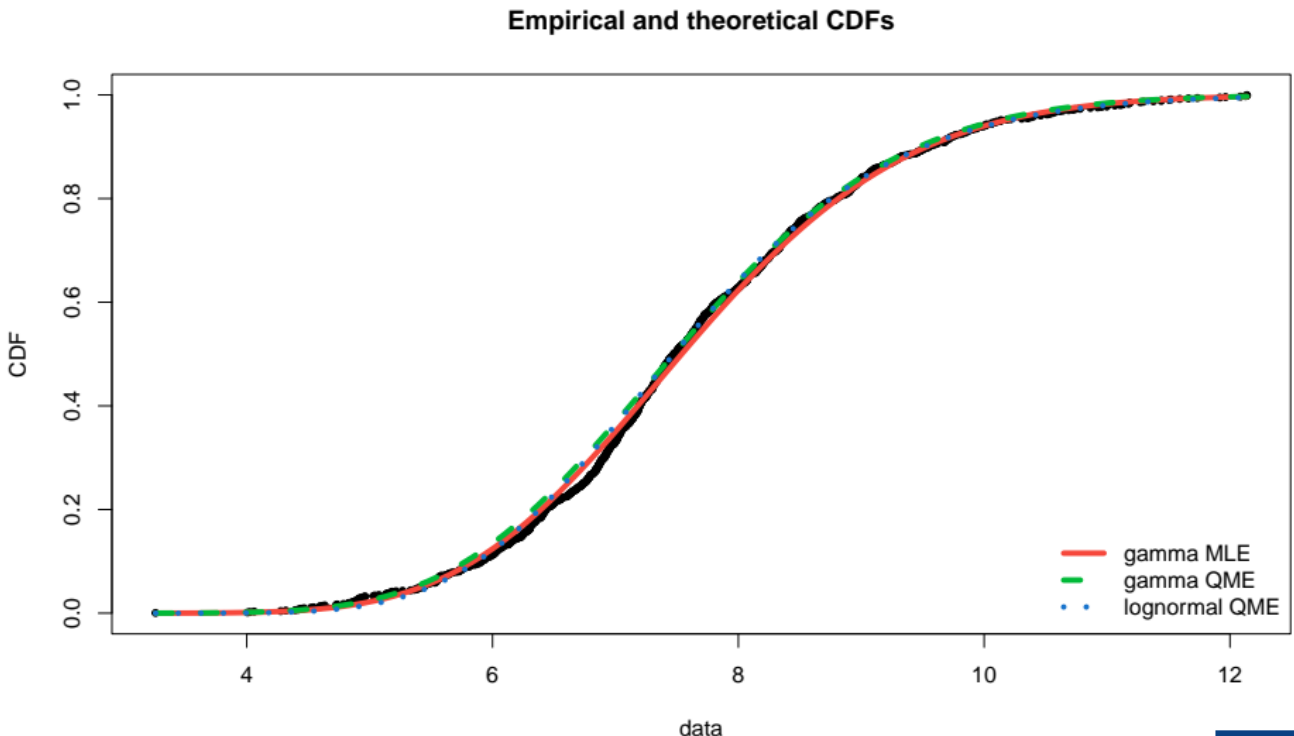
```
fit.gamma.qme <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "gamma", method = "qme", probs = c(0.5, 0.75))
fit.lnorm.qme <- fitdist(log(SUVA$dailyallow[SUVA$dailyallow > 0]), "lnorm", method = "qme", probs = c(0.5, 0.75))
gof.qme.legend <- c("gamma MLE", "gamma QME", "lognormal QME")
```

```
denscomp(list(fit.gamma.mle, fit.gamma.qme, fit.lnorm.qme), legendtext = go  
fitlwd = 3)
```

Histogram and theoretical densities

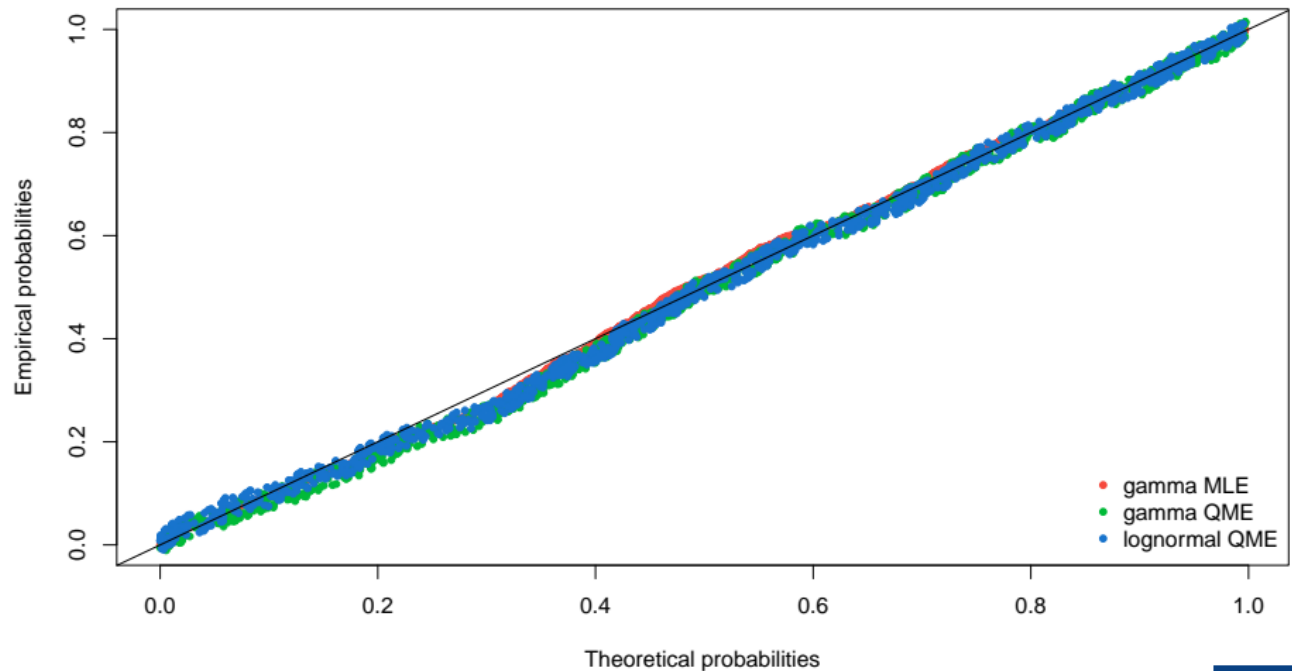


```
cdfcomp(list(fit.gamma.mle, fit.gamma.qme, fit.lnorm.qme), legendtext = gof  
fitlwd = 4, datapch = 20)
```



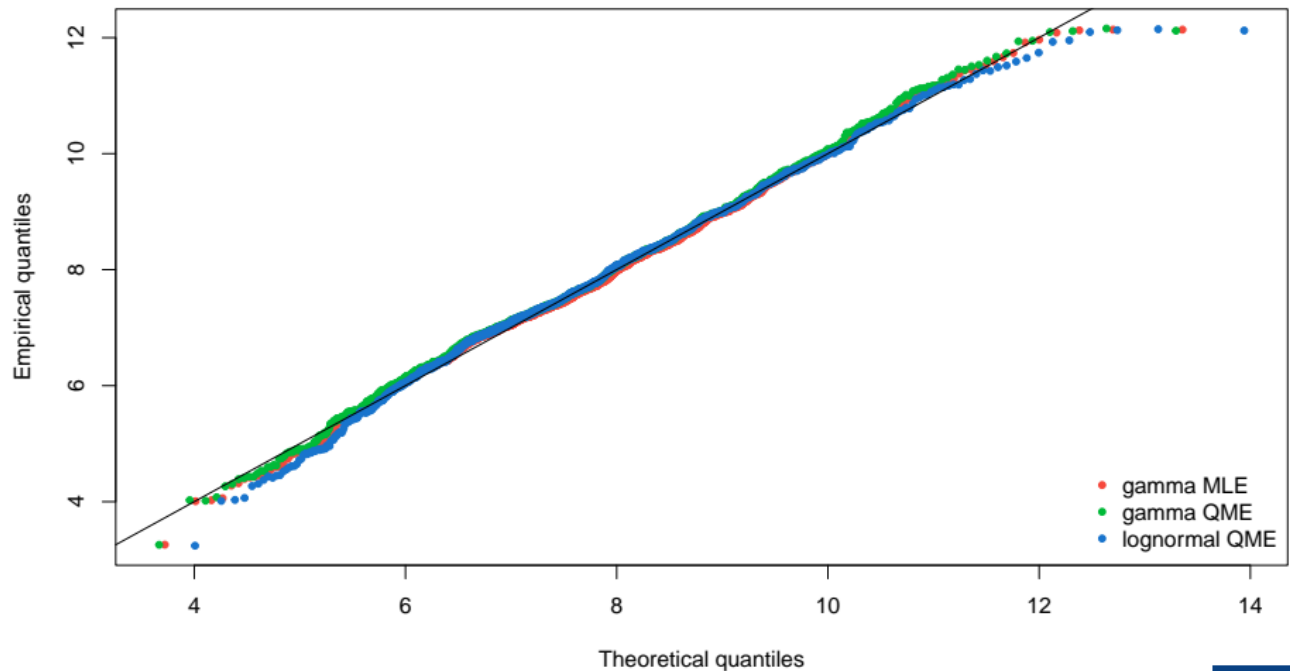
```
ppcomp(list(fit.gamma.mle, fit.gamma.qme, fit.lnorm.qme), legendtext = gof.  
       fitpch = 20)
```

P-P plot



```
qqcomp(list(fit.gamma.mle, fit.gamma.qme, fit.lnorm.qme), legendtext = gof.  
       fitpch = 20)
```

Q-Q plot



6 ✖ Other advanced topics

- ✖ Alternative methods for estimation
- ✖ Zero-inflated severity model $X = IB$

✖ Zero-inflated severity model $X = IB$

In this approach $X = IB$, where

- I is an indicator of claim with

$$\Pr [I = 1] = q \text{ and } \Pr [I = 0] = 1 - q$$

- B is the {claim amount given $I = 1$ } (given a claim occurs).

This allows us to avoid a large probability mass at 0 for rare losses.

- Note that this is a good approach for modifying continuous distributions, which are generally used for severity.
- In the case of discrete distributions—usually used for frequency, modifications of the usual distributions (in the form of “zero-truncated” or “zero-modified”) are well known, and readily available in the package `actuar`. This is discussed in Module 2.
- In practice a frequency and severity model would be chosen at the same time, and the way zero claims are dealt with should be determined in a consistent way, e.g.:
 - “frequency” models strictly positive claims, and “severity” is a strictly positive continuous distribution;
 - “frequency” models insurable events (which may lead to claims of 0), and “severity” includes a mass at 0 (such as in this section);
 - etc...

✖ Resulting distribution

As a consequence,

$$\begin{aligned}\Pr[X \leq x] &= \Pr[X \leq x | I = 0] \Pr[I = 0] \\ &\quad + \Pr[X \leq x | I = 1] \Pr[I = 1] \\ &= 1 - q + q \Pr[B \leq x]\end{aligned}$$

and

$$\begin{aligned}M_X(t) &= E[e^{tX} | I = 0] \Pr[I = 0] \\ &\quad + E[e^{tX} | I = 1] \Pr[I = 1] \\ &= 1 - q + qE[e^{tB}].\end{aligned}$$

✖ Mean and Variance

The **mean** can be determined using

$$E[X] = E[E[X|I]] = E[X|I=1]\Pr[I=1] = qE(B),$$

after noting that $E[X|I=0] = 0$.

The **variance** can be determined using

$$\begin{aligned} \text{Var}(X) &= \text{Var}(E[X|I]) + E[\text{Var}(X|I)] \\ &= [E(B)]^2 \text{Var}(I) + q\text{Var}(B) \\ &= q(1-q)(E[B])^2 + q\text{Var}(B) \end{aligned}$$

after noting that

$$\begin{aligned} E[X|I] &= I \cdot E[B], \text{ and that} \\ \text{Var}(X|I) &= I^2 \cdot \text{Var}(B), \end{aligned}$$

which are both random variables (functions of I).

✖ Special case: $X = Ib$

- Fixed claim: $B = b$ with probability 1.
- The individual claim random variable becomes

$$X = Ib = \begin{cases} b, & \text{w.p. } q \\ 0, & \text{w.p. } 1 - q \end{cases} .$$

- Mean: $E[X] = bq$
- Variance: $Var(X) = b^2 Var(I) = b^2 q(1 - q)$

This is nothing more than a scaled Bernoulli... (and if you add them, the sum becomes a scaled Binomial)

✖ Example: Bicycle Theft (get B out of X)

- Insurance policy against bicycle theft (insured amount is 400)
- Only half is paid if bicycle is not locked.
- Assume: $\Pr[X = 400] = 0.05$ and $\Pr[X = 200] = 0.15$.
- Probability of a claim: $q = \Pr[I = 1] = 0.20$ [law of total probability]
- The pmf of B is computed in the following way

$$\begin{aligned}\Pr[B = 400] &= \Pr[X = 400 | I = 1] = \frac{\Pr[X=400 \cap I=1]}{\Pr[I=1]} \\ &= \frac{0.05}{0.20} = 0.25\end{aligned}$$

✖ Example

In an insurance portfolio, there are 15 insured:

- ten of the insured persons have 0.1 probability of making a claim, and
- the other 5 have a 0.2 probability of making a claim.

All claims are independent and follow an exponential distribution with mean $1/\lambda$. What is the mgf of the aggregate claims distribution?

Let X_i be the total amount of claims incurred from the i th person, and B_i denote the amount of claim, if there is one. Then $X_i = I_i B_i$.

If $q_i = \Pr(I_i = 1)$ then $q_1 = \dots = q_{10} = 0.1$ and $q_{11} = \dots = q_{15} = 0.2$.

The mgf of X_i :

$$\begin{aligned} M_{X_i}(t) &= E[e^{tX_i} | I_i = 0] \Pr(I_i = 0) + E[e^{tX_i} | I_i = 1] \Pr(I_i = 1) \\ &= 1 - q_i + E[e^{tB_i}] q_i = 1 - q_i + \frac{\lambda}{\lambda-t} q_i \end{aligned}$$

Since the aggregate claims are $S = X_1 + \dots + X_{15}$ the mgf of S is

$$\begin{aligned} M_S(t) &= \prod_{i=1}^{15} E[e^{tX_i}] \\ &= \left(1 - 0.1 + 0.1 \frac{\lambda}{\lambda-t}\right)^{10} \left(1 - 0.2 + 0.2 \frac{\lambda}{\lambda-t}\right)^5 \end{aligned}$$

- 1 Introduction
- 2 Data analysis and descriptive statistics (MW 3.1)
- 3 Selected parametric claims size distributions (MW 3.2)
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- 6 ✕ Other advanced topics
- 7 Calculating within layers for claim sizes (MW 3.4)

7 Calculating within layers for claim sizes (MW 3.4)

- Usual policy transformations
- Reinsurance
- Proportional reinsurance
- Nonproportional reinsurance
- Stop loss premiums
- Leverage effect of claims inflation

Deductible and Policy Limit

One way to control the cost (and variability) of individual claim losses is to introduce deductibles and policy limits.

- **Deductible d :** the insurer starts paying claim amounts above the deductible d
- **Limit M :** the insurer pays up to the limit M .

If we denote the damage random variable by D , then if a claim occurs the insurer is liable for

$$X = \min [\max (D - d, 0), M].$$

7 Calculating within layers for claim sizes (MW 3.4)

- Usual policy transformations
- Reinsurance
- Proportional reinsurance
- Nonproportional reinsurance
- Stop loss premiums
- Leverage effect of claims inflation

Reinsurance

Reinsurance is a **risk transfer** from an insurer (the direct writer) to a reinsurer:

- in other words, some of the (random) risk faced by the insurer is “transferred” to the reinsurer (that means the reinsurer will cover that risk), in exchange of a (deterministic) premium (which will obviously generally be higher than the expected value of the risk that was transferred)
- the risk that the insurer keeps is called the retention

There are different types of reinsurance:

- proportional
 - quota share: the proportion is the same for all risks
 - surplus: the proportion can vary from risk to risk
- nonproportional
 - (individual) excess of loss: on each individual loss (X_i)
 - stop loss: on the aggregate loss (S)
- cheap (reinsurance premium is the expected value), or non cheap (reinsurance premium is loaded)
- Alternative Risk Transfers (“ART”), where usually the idea is to transfer the risk to different / deeper pockets. For instance, for transfers to the financial markets:
 - Catastrophe bonds (“CAT bonds”)
 - Longevity bonds
 - Pandemic bonds

7 Calculating within layers for claim sizes (MW 3.4)

- Usual policy transformations
- Reinsurance
- **Proportional reinsurance**
- Nonproportional reinsurance
- Stop loss premiums
- Leverage effect of claims inflation

Proportional reinsurance

The **retained proportion** α defines who pays what:

- the insurer pays $Y = \alpha X$
- the reinsurer pays $Z = (1 - \alpha)X$

This is nothing else but a change of scale and we have

$$\mu_Y = \alpha \mu_X, \quad \sigma_Y^2 = \alpha^2 \sigma_X^2, \quad \gamma_Y = \gamma_X.$$

In some cases it suffices to adapt the scale parameter. Example:

- If X is exponential with parameter β

$$\Pr[Y \leq y] = \Pr[\alpha X \leq y] = \Pr[X \leq y/\alpha] = 1 - e^{-\beta y/\alpha}$$

and thus Y is exponential with parameter β/α .

7 Calculating within layers for claim sizes (MW 3.4)

- Usual policy transformations
- Reinsurance
- Proportional reinsurance
- Nonproportional reinsurance**
- Stop loss premiums
- Leverage effect of claims inflation

Nonproportional reinsurance

Basic arrangements:

- the reinsurer pays the excess over a **retention (excess point) d**
 - the insurer pays $Y = \min(X, d)$
 - the reinsurer pays $Z = (X - d)_+$
 - $E[(X - d)_+]$ is called **stop-loss premium**.
- the reinsurer limits his payments to an amount M . In that case
 - the insurer pays $Y = \min(X, d) + (X - M - d)_+$
 - the reinsurer pays $Z = \min\{(X - d)_+, M\}$

Example

Consider a life insurance company with 16,000 1-year term life insurance policies. The associated insured amounts are:

<i>Benefit (10000's)</i>	<i># policies</i>
1	8000
2	3500
3	2500
5	1500
10	500

The probability of death (q) for each of the 16,000 lives is 0.02. This company has an EoL reinsurance contract with retention limit 30,000 at a cost of 0.025 per dollar of coverage.

What is the approximate probability (using CLT) that the total cost will exceed 8,250,000?

The portfolio of retained business is given by

<i>k</i>	<i>retained benefit b_k (10000's)</i>	<i># policies n_k</i>
1	1	8000
2	2	3500
3	3	4500

Now

$$\begin{aligned}
 E[S] &= \sum_{k=1}^3 n_k E[X_k] = \sum_{k=1}^3 n_k b_k q_k \\
 &= 8000 \cdot 1 \cdot 0.02 + 3500 \cdot 2 \cdot 0.02 + 4500 \cdot 3 \cdot 0.02 \\
 &= 570, \quad \text{and}
 \end{aligned}$$

$$\begin{aligned}
 \text{Var}[S] &= \sum_{k=1}^3 n_k \text{Var}[X_k] = \sum_{k=1}^3 n_k b_k^2 q_k (1 - q_k) \\
 &= 8000 \cdot 1^2 \cdot 0.02 \cdot 0.98 + 3500 \cdot 2^2 \cdot 0.02 \cdot 0.98 \\
 &\quad + 4500 \cdot 3^2 \cdot 0.02 \cdot 0.98 \\
 &= 1225
 \end{aligned}$$

The reinsurance cost is

$$[(5 - 3) \cdot 1500 + (10 - 3) \cdot 500] \cdot 0.025 = 162.5.$$

Thus, the desired probability becomes

$$\begin{aligned}\Pr[S + 162.5 > 825] &= \Pr\left[\frac{S - E[S]}{\sqrt{\text{Var}(S)}} > \frac{662.5 - E[S]}{\sqrt{\text{Var}(S)}}\right] \\ &\approx \Pr\left[Z > \frac{662.5 - 570}{\sqrt{1225}}\right] \\ &= \Pr[Z > 2.643] = 0.0041.\end{aligned}$$

Discussion:

- Without reinsurance, exp/var is $700/2587.20$ so the associated probability of shortfall is $\approx \Pr[Z > 2.458]$, which is higher even though it is not cheap reinsurance.
- However, there is lower expected gain:
 - With reinsurance the expected gain is

$$P - 570 - 162.5 = P - 732.5$$

- Without reinsurance it is

$$P - 700,$$

which is higher.

✖ The actuar coverage function

- The package `actuar` allows for direct specification of the pdf of a modified random variable after possible left-truncation and right-censoring.
- Given the pdf or cdf of the original loss D , `coverage` returns a function object to compute the pdf or cdf of the modified random variable after one or several of the following modifications:
 - ordinary deductible d ;
 - franchise deductible d ;
 - limit u ;
 - coinsurance α ;
 - inflation r .
- The vignette on loss modeling features of `actuar` provides precise definitions, and this document summarises all the formulas.

Assume that insurance payments are

$$X = \begin{cases} D - d, & d \leq D \leq u \\ u - d, & D \geq u \end{cases}$$

with mixed distribution

$$f_X(x) = \begin{cases} 0, & x = 0 \\ \frac{f_D(y+d)}{1-F_D(d)}, & 0 < x < u - d \\ \frac{1-F_D(u)}{1-F_D(d)}, & x = u - d \\ 0 & x > u - d \end{cases}$$

as seen before. Note however that the u is expressed on the raw variable D , not the payment, so that the maximum payment is $u - d$.

If D is gamma, $d = 2$, and $u = 20$, one can get $f_X(x)$ in R as follows:

```
f <- coverage(pdf = dgamma, cdf = pgamma, deductible = 2, limit = 20)
```

The function can be then used to fit distributions to data.

As an example we will use the previously generated data `xcens`. Note it needs to be shifted by d down, because what we have is the left-truncated and right-censored D , not the insurance payments as per the formulation above.

```
fit.gamma.xcens2 <- fitdistr(xcens$left - 2, f, start = list(shape = mean(xcens$left),  
rate = mean(xcens$left)/var(xcens$left)))  
fit.gamma.xcens2  
##      shape          rate  
## 2.03990496 0.20142871  
## (0.10516140) (0.01009694)  
fit.tgamma.xcens # our previous fit with fitdist  
## Fitting of the distribution ' tgamma ' on censored data by maximum likelihood  
## Parameters:  
## Fixed parameters:  
c(fit.gamma.xcens2$loglik, fit.tgamma.xcens$loglik)  
## [1] -5341.551 -5340.151
```

Note that this is using MASS::fitdistr rather than the (arguably more flexible and possibly advanced) fitdistrplus::fitdist. This approach works as well, but does not seem as precise in this particular instance.

A useful identity

Note that

$$\min(X, c) = X - (X - c)_+$$

and thus

$$E[\min(X, c)] = E[X] - E[(X - c)_+].$$

The amount $E[(X - c)_+] = \Pr[X > c]e(c)$

- is commonly called “stop loss premium” with retention c .
- is identical to the expected payoff of a call with strike price c , and thus results from financial mathematics can sometimes be directly used (and vice versa).

7 Calculating within layers for claim sizes (MW 3.4)

- Usual policy transformations
- Reinsurance
- Proportional reinsurance
- Nonproportional reinsurance
- Stop loss premiums
- Leverage effect of claims inflation

Stop loss premiums

Let

$$E[(X - d)_+] = P_d.$$

Then we have (for positive rv's)

$$P_d = \begin{cases} \int_d^\infty [1 - F_X(x)] dx & \text{if } X \text{ is continuous} \\ \sum_d^\infty [1 - F_X(x)] & \text{if } X \text{ is discrete} \end{cases}$$

Example

Calculate P_d if X is Exponential with mean $1/\beta$.

✚ Recursive formulas in the discrete case

First moment: - if d is an integer

$$P_{d+1} = P_d - [1 - F_X(d)] \text{ with } P_0 = E[X]$$

- if d is not an integer

$$P_d = P_{\lfloor d \rfloor} - (d - \lfloor d \rfloor)[1 - F_X(\lfloor d \rfloor)].$$

Second moment $P_d^2 = E[(X - d)_+^2]$:

$$P_{d+1}^2 = P_d^2 - 2P_d + [1 - F_X(d)] \text{ with } P_0^2 = E[X^2].$$

Note that $\lfloor x \rfloor$ is the integer part of x (e.g. $\lfloor 2.5 \rfloor = 2$).

Numerical example

For the distribution F_{1+2+3} derived in Module 2 we have $E[S] = 4 = 128/32$ and $E[S^2] = 19.5 = 624/32$ and thus

d	$f_{1+2+3}(d)$	$F_{1+2+3}(d)$	P_d	P_d^2	$\text{Var}((X - d)_+)$
0	1/32	1/32	128/32	624/32	3.500
1	2/32	3/32	97/32	399/32	3.280
2	4/32	7/32	68/32	234/32	2.797
3	6/32	13/32	43/32	123/32	2.038
4	6/32	19/32	24/32	56/32	1.188
5	6/32	25/32	11/32	21/32	0.538
6	4/32	29/32	4/32	6/32	0.172
7	2/32	31/32	1/32	1/32	0.030
8	1/32	32/32	0	0	0.000

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Leverage effect of claims inflation

Choose a fixed deductible $d > 0$ and assume that the claim at time 0 is given by Y_0 . Assume that there is a deterministic inflation index $i > 0$ such that the claim at time 1 can be represented by $Y_1 = (1 + i)Y_0$. We have

$$E[(Y_1 - d)_+] \geq (1 + i)E[(Y_0 - d)_+].$$

When tax brackets are not adapted, this leads to the so-called “bracket creep”...

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