# Introduction to MortalityLaws

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# 1 Package structure

MortalityLaws in an R package which exploits the available optimization methods to provide tools for fitting and analyzing a wide range of complex mortality models. The package can be used to construct full and abridged life tables given various input indices and to download data from Human Mortality Database as well. The main functions in the package are: MortalityLaw, LifeTable, LawTable, convertFx and ReadHMD. The package also provides support functions like availableLaws, availableLF and availableHMD that return information about the mortality laws implemented in the package, the loss functions used in optimization processes, and HMD data availability. Generic functions like predict, plot, summary, fitted, residuals are created for MortalityLaws objects. Small data set for testing purposes ahmd is saved in the package.

All functions are documented in the standard way, which means that once you load the package using library(MortalityLaws) you can just type, for example, ?MortalityLaw to see the help file.

# 2 Downloading HMD Data

Download data form Human Mortality Database (2016) using the ReadHMD function:

library(MortalityLaws)

Here we download all the registered death counts Dx in Sweden from 1751 until 2014. In the same way one can download the following records: birth records births, exposure-to-risk Ex, deaths by Lexis triangles lexis, population size population, death-rates mx, life tables for females LT\_f, life tables for males LT\_m, life tables both sexes combined LT\_t, life expectancy at birth e0, cohort death-rates mxc and cohort exposures Exc for over 40 countries and regions in different formats.

# 3 Model fitting and diagnosis

Once we have data from HMD or other sources we can start analyzing it. For example, let's fit a Heligman-Pollard (1980) model under a Poisson setting which is already implemented as one of the standard models in the package. We have to use the MortalityLaw function in this regard.

```
<- 1950
vear
         <- 0:100
ages
         <- ahmd$Dx[paste(ages), paste(year)]
deaths
exposure <- ahmd$Ex[paste(ages), paste(year)]</pre>
fit <- MortalityLaw(x</pre>
                         = ages,
                                    # vector with death counts
                     Dx = deaths,
                     Ex = exposure, # vector containing exposures
                     law = "HP",
                     opt.method = "LF2")
# inspect the output object
ls(fit)
                                             "df"
## [1] "coefficients"
                          "deviance"
                                                                "fitted.values"
## [5] "goodness.of.fit" "info"
                                             "input"
                                                                 "opt.diagnosis"
```

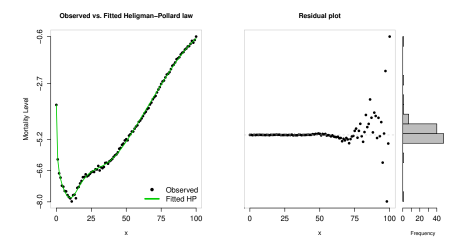
A summary can be obtained using the summary function:

## [9] "residuals"

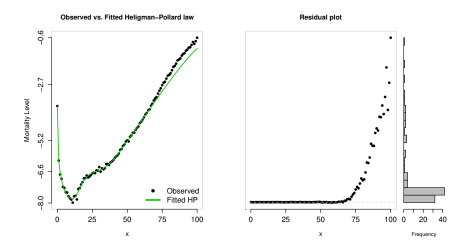
```
summary(fit)
## Heligman-Pollard model: q[x]/p[x] = A^{(x + B)^{2}} + D \exp[-E \log(x/F)^{2}] + G H^{x}
## Fitted values: mx
##
## Call: MortalityLaw(x = ages, Dx = deaths, Ex = exposure, law = "HP",
##
       opt.method = "LF2")
##
## Deviance Residuals:
      Min. 1st Qu. Median
                            Mean 3rd Qu.
                                               Max.
## -0.0312 -0.0002 0.0000 0.0009 0.0003 0.0462
##
## Parameters:
                                                 F_{-}
                 В
                          C
                                  D
                                          \boldsymbol{E}
   0.0022 0.0146 0.1229 0.0009 2.7565 29.0080 0.0000 1.1141
## Residual standard error: 0.07118 on 93 degrees of freedom
```

The standard plot helps us to investigate visually the goodness of fit.

```
plot(fit)
```



A model can be fitted using a subset of the data only by specifying in fit.this.x age range to be covered:



The gray area on the plot showing the fitted value indicates the age range used in fitting the model.

# 4 Mortality laws in the package

In **R** one can check the availability of the implemented models using availableLaws:

```
availableLaws()
```

See table 1.

# 5 Loss function in the package

A parametric model is fitted by optimizing a loss function e.g. a likelihood function or a function that minimizes errors. In MortalityLaws 8 such functions are implemented and can be used to better capture different portions of a mortality curve. Check availableLF for more details.

```
availableLF()
##
## Loss functions available in the package:
##
##
   LOSS FUNCTION
                                                   CODE
## L = -[Dx * log(mu) - mu*Ex]
                                                   poissonL
   L = -[Dx * log(1 - exp(-mu)) - (Ex - Dx)*mu]
                                                   binomialL
## L = [1 - mu/ov]^2
                                                   LF1
##
   L = log[mu/ov]^2
                                                   LF2
   L = [(ov - mu)^2]/ov
                                                   LF3
   L = [ov - mu]^2
                                                   LF4
   L = [ov - mu] * log[ov/mu]
                                                   LF5
   L = abs(ov - mu)
                                                   LF6
##
## LEGEND:
## Dx: Death counts
## Ex: Population exposed to risk
## mu: Estimated value
## ov: Observed value
```

# 6 Custom mortality laws

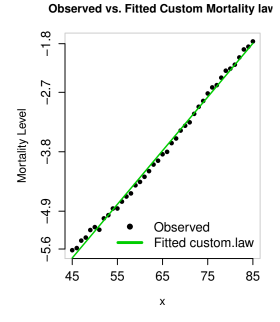
Now let's fit a mortality law that is not defined in the package, say a reparametrize version of Gompertz in terms of modal age at death (Missov et al. 2015),

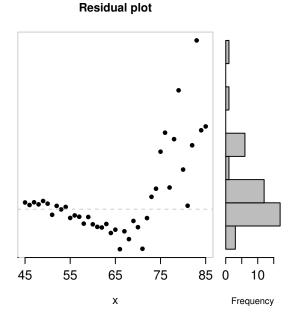
$$\mu_x = \beta e^{\beta(x-M)}. (1)$$

We have to define a function containing the desired hazard function and then using the custom.law argument it can be used in the MortalityLaw function.

```
# Here we define a function for our new model and provide start parameters
my_gompertz <- function(x, par = c(b = 0.13, M = 45)){
   hx <- with(as.list(par), b*exp(b*(x - M)) )</pre>
```

```
# return everything inside this function
  return(as.list(environment()))
}
# Select data
       <- 1950
year
        <- 45:85
ages
deaths <- ahmd$Dx[paste(ages), paste(year)]</pre>
exposure <- ahmd$Ex[paste(ages), paste(year)]</pre>
# Use 'custom.law' argument to instruct the MortalityLaw function how to behave
my_model <- MortalityLaw(x = ages,</pre>
                         Dx = deaths,
                         Ex = exposure,
                         custom.law = my_gompertz)
summary(my_model)
## Custom Mortality Law
## Fitted values: mx
##
## Call: MortalityLaw(x = ages, Dx = deaths, Ex = exposure, custom.law = my_gompertz)
##
## Deviance Residuals:
## Min. 1st Qu. Median Mean 3rd Qu.
                                             {\it Max} .
## -0.0029 -0.0011 0.0002 0.0009 0.0015 0.0122
##
## Parameters:
##
        Ъ
## 0.0993 35.8021
##
## Residual standard error: 0.02097 on 39 degrees of freedom
plot(my_model)
```





### 7 Full life tables

Using LifeTable function one can build full or abridged life table with various input choices like: death counts and mid-interval population estimates (Dx, Ex) or age-specific death rates (mx) or death probabilities (qx) or survivorship curve (1x) or a distribution of deaths (dx). If one of these options are specified, the other can be ignored.

```
# Life table for year of 1900
y <- 1900
x <- as.numeric(rownames(ahmd$mx))
Dx <- ahmd$Dx[, paste(y)]
Ex <- ahmd$Ex[, paste(y)]

LT1 <- LifeTable(x, Dx = Dx, Ex = Ex)
LT2 <- LifeTable(x, mx = LT1$lt$mx)
LT3 <- LifeTable(x, qx = LT1$lt$qx)
LT4 <- LifeTable(x, lx = LT1$lt$lx)
LT5 <- LifeTable(x, dx = LT1$lt$dx)</pre>
```

```
##
## Number of life tables: 1
## Dimension: 111 x 10
## Age intervals: [0,1) [1,2) [2,3) ... [108,109) [109,110) [110,+)
##
##
        x.int
                                  ax
                                         lx
                                               dx
                                                     Lx
                                                             Tx
                      mx
                              qx
                                                                    ex
##
        [0,1)
                0 0.1513 0.1404 0.49 1e+05 14042 92802 4808384 48.08
##
        [1,2)
                1 0.0479 0.0468 0.5 85958 4023 83931 4715582 54.86
##
        [2,3)
                2 0.0191 0.0189 0.5 81935 1552 81157 4631651 56.53
                3 0.013 0.0129 0.5 80383 1041 79861 4550494 56.61
##
        [3,4)
                                             740 78971 4470633 56.35
##
                4 0.0094 0.0093 0.5 79342
        [4,5)
##
        [5,6)
                5 0.0067 0.0067 0.5 78602
                                              523 78340 4391661 55.87
##
         <NA> ...
                     . . .
                             . . .
    [108,109) 108
                    6.75 0.9988 0.15
                                          0
                                                0
                                                      0
                                                              0 0.15
    [109,110) 109
                    6.75 0.9988 0.15
                                          0
                                                0
                                                      0
                                                              0 0.15
##
      [110,+) 110
                    6.75 0.9988 0.15
                                                              0 0.15
ls(LT1)
## [1] "call"
                      "lt"
                                      "process_date"
```

# 8 Abridged life tables

```
# Example
x \leftarrow c(0, 1, seq(5, 110, by = 5))
mx \leftarrow c(.053, .005, .001, .0012, .0018, .002, .003, .004,
       .004, .005, .006, .0093, .0129, .019, .031, .049,
       .084, .129, .180, .2354, .3085, .390, .478, .551)
lt <- LifeTable(x, mx = mx, sex = "female")</pre>
lt
##
## Abridged Life Table
## Number of life tables: 1
## Dimension: 24 x 10
## Age intervals: [0,1) [1,5) [5,10) ... ... [100,105) [105,110) [110,+)
##
##
        x.int
                                         lx
                                              dx
                                                      Lx
                                                              Tx
                              qx
                                   ax
##
        [0,1)
                   0.053 0.0516 0.2 1e+05 5162 95878 6485467 64.85
##
        [1,5)
                1 0.005 0.0198 1.44 94838 1878 374547 6389590 67.37
##
       [5,10)
                5 0.001 0.005 2.5 92960 464 463640 6015042 64.71
##
      [10, 15)
               10 0.0012 0.006 2.5 92496 553 461098 5551402 60.02
##
               15 0.0018 0.009 2.5 91943 824 457653 5090304 55.36
      [15,20)
##
      [20, 25)
               20 0.002
                           0.01 2.5 91119 907 453326 4632651 50.84
         <NA>
##
```

```
##
    [100,105) 100
                    0.39 0.8577 1.73
                                        407
                                             349
                                                     896
                                                            1014 2.49
##
    [105,110) 105 0.478 0.9084 1.59
                                         58
                                              53
                                                     110
                                                             119
                                                                  2.05
                                               5
##
      [110,+) 110 0.551 0.9364 1.59
                                          5
                                                                 1.59
```

#### 9 Citation

```
citation(package = "MortalityLaws")
## Warning in citation(package = "MortalityLaws"): no date field in
## DESCRIPTION file of package 'MortalityLaws'
## Warning in citation(package = "MortalityLaws"): could not determine year
## for 'MortalityLaws' from package DESCRIPTION file
##
## To cite package 'MortalityLaws' in publications use:
##
##
     Marius D. Pascariu (NA). MortalityLaws: Parametric Mortality
##
    Models, Life Tables and HMD. R package version 1.8.0.
     https://github.com/mpascariu/MortalityLaws
##
##
## A BibTeX entry for LaTeX users is
##
     @Manual{,
##
       title = {MortalityLaws: Parametric Mortality Models, Life Tables and HMD},
##
##
       author = {Marius D. Pascariu},
       note = {R package version 1.8.0},
##
##
       url = {https://qithub.com/mpascariu/MortalityLaws},
##
```

### References

Heligman, Larry, and John H Pollard. 1980. "The Age Pattern of Mortality." *Journal of the Institute of Actuaries* 107 (01): 49–80.

Missov, Trifon I, Adam Lenart, Laszlo Nemeth, Vladimir Canudas-Romo, and James W Vaupel. 2015. "The Gompertz Force of Mortality in Terms of the Modal Age at Death." *Demographic Research* 32: 1031–48.

University of California Berkeley, USA, and Max Planck Institute for Demographic Research, Germany. 2016. "Human Mortality Database." http://www.mortality.org/.

Table 1: Parametric functions build in the Mortality Laws package  $\,$ 

Mortality Laws	Predictor	Code
Gompertz	$\mu(x) = Ae^{Bx}$	gompertz
Gompertz	$\mu(x) = \frac{1}{\sigma} exp\left\{\frac{x-M}{\sigma}\right\}$	gompertz0
Inverse-Gompertz	$\mu(x) = \frac{1}{\sigma} exp\left\{\frac{x-M}{\sigma}\right\} / \left(exp\left\{e^{\frac{-(x-M)}{\sigma}}\right\} - 1\right)$	invgompertz
Makeham	$\mu(x) = Ae^{Bx} + C$	'makeham'
Makeham	$\mu(x) = \frac{1}{\sigma} exp\left\{\frac{x-M}{\sigma}\right\} + C$	makeham0
Opperman	$\mu(x) = \frac{A}{\sqrt{x}} + B + C\sqrt[3]{x}$	opperman
Thiele	$\mu(x) = A_1 e^{-B_1 x} + A_2 e^{-\frac{1}{2}B_2(x-C)^2} + A_3 e^{B_3 x}$	thiele
Wittstein & Bumstead	$q(x) = \frac{1}{B}A^{-(Bx)^N} + A^{-(M-x)^N}$	wittstein
Perks	$\mu(x) = (A + BC^x)/(BC^{-x} + 1 + DC^x)$	perks
Weibull	$\mu(x) = \frac{1}{\sigma} \left(\frac{x}{M}\right)^{\frac{M}{\sigma} - 1}$	weibull
Inverse-Weibull	$\mu(x) = \frac{1}{\sigma} \left(\frac{x}{M}\right)^{-\frac{M}{\sigma} - 1} / \left(exp\left\{\left(\frac{x}{M}\right)^{-\frac{M}{\sigma}}\right\} - 1\right)$	invweibull
Van der Maen	$\mu(x) = A + Bx + Cx^2 + I/(N-x)$	vandermaen
Van der Maen	$\mu(x) = A + Bx + I/(N - x)$	vandermaen2
Quadratic	$\mu(x) = A + Bx + Cx^2$	quadratic
Beard	$\mu(x) = Ae^{B^x}/(1 + KAe^{B^x})$	beard
Makeham-Beard	$\mu(x) = Ae^{B^x}/(1 + KAe^{B^x}) + C$	makehambeard
Gamma-Gompertz	$\mu(x) = Ae^{B^x}/[1 + \frac{AG}{B}(e^{B^x} - 1)]$	ggompertz
Siler	$\mu(x) = A_1 e^{-B_1 x} + A_2 + A_3 e^{B_3 x}$	siler
Heligman - Pollard	$q(x)/p(x) = A^{(x+B)^C} + De^{-E(\ln x - \ln F)^2} + GH^x$	HP
Heligman - Pollard	$q(x) = A^{(x+B)^C} + De^{-E(\ln x - \ln F)^2} + \frac{GH^x}{1+GH^x}$	HP2
Heligman - Pollard	$q(x) = A^{(x+B)^C} + De^{-E(\ln x - \ln F)^2} + \frac{GH^x}{1 + KGH^x}$	HP3
Heligman - Pollard	$q(x) = A^{(x+B)^C} + De^{-E(\ln x - \ln F)^2} + \frac{GH^{x^K}}{1 + GH^{x^K}}$	HP3
Rogers-Planck	$q(x) = A_0 + A_1 e^{-Ax} + A_2 e^{\left\{B(x-U) - e^{-C(x-U)}\right\}} + A_3 e^{Dx}$	rogersplanck
Martinelle	$\mu(x) = (Ae^{Bx} + C)/(1 + De^{Bx}) + Ke^{Bx}$	martinelle
Carriere	$S(x) = \psi_1 S_1(x) + \psi_2 S_2(x) + \psi_3 S_3(x)$	carriere1
Carriere	$S(x) = \psi_1 S_1(x) + \psi_4 S_4(x) + \psi_3 S_3(x)$	carriere2
Kostaki	$q(x)/p(x) = A^{(x+B)^C} + De^{-E_i(\ln x - \ln F)^2} + GH^x$	kostaki
Kannisto	$\mu(x) = Ae^{Bx}/(1 + Ae^{Bx}) + C$	kannisto