

Package ‘MortCast’

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Author Hana Sevcikova, Nan Li and Patrick Gerland

Maintainer Hana Sevcikova <hanas@uw.edu>

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MortCast-package	<i>MortCast: Estimation and Projection of Age-Specific Mortality Rates</i>
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Description

Age-specific mortality rates are estimated and projected using the Kannisto, Lee-Carter and related methods as described in Sevcikova et al. (2016) <doi:10.1007/978-3-319-26603-9_15>.

Details

The package implements methodology described in Sevcikova et al. (2016) that is related to estimating and predicting age-specific mortality rates. The main functions are:

- `cokannisto`: Extrapolates given mortality rates into higher ages using the Coherent Kannisto method. The original Kannisto method (with sex-independent extrapolation) is available in the function `kannisto`.
- `lileecarter.estimate`: Estimates the coherent Lee-Carter parameters for male and female mortality rates (Li and Lee 2005), i.e. sex-independent parameters a_x and k_t , and the coherent parameter b_x . In addition, it computes the ultimate b_x^u for rotation (Li et al. 2013). The underlying sex-independent estimation is implemented in the function `leecarter.estimate`.
- `mortcast`: Using estimated coherent Lee-Carter parameters and given future sex-specific life expectancies, it projects age-specific mortality rates, while (by default) rotating the b_x parameter as described in Li et al. (2013).

Functions contained in the package can be used to apply Algorithm 2 in Sevcikova et al. (2016) as shown in the Example below.

Other methods for forecasting mortality rates are available:

- `pmd`: pattern of mortality decline
- `mlt`: model life tables
- `logquad`: log-quadratic mortality model
- `mortcast.blend`: combining two different methods

A life table can be constructed using the `life.table` function.

References

- Li, N. and Lee, R. D. (2005). Coherent mortality forecasts for a group of populations: An extension of the Lee-Carter method. *Demography*, 42, 575-594.
- Li, N., Lee, R. D. and Gerland, P. (2013). Extending the Lee-Carter method to model the rotation of age patterns of mortality decline for long-term projections. *Demography*, 50, 2037-2051.
- Sevcikova H., Li N., Kantorova V., Gerland P., Raftery A.E. (2016). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. In: Schoen R. (eds) *Dynamic Demographic Analysis. The Springer Series on Demographic Methods and Population Analysis*, vol 39. Springer, Cham. [Earlier version](#).

Examples

```
# This example applies Algorithm 2 in Sevcikova et al. (2016)
# on data from WPP2017 for China
#
data(mxM, mxF, e0Fproj, e0Mproj, package = "wpp2017")
country <- "China"

# extract observed mortality rates for male and female
mxm <- subset(mxM, name == country)[,4:16]
mxf <- subset(mxF, name == country)[,4:16]
rownames(mxm) <- rownames(mxf) <- c(0,1, seq(5, 100, by=5))

# Step 1: extrapolate from 100+ to 130+ using Coherent Kannisto
mx130 <- cokannisto(mxm, mxf)

# Steps 2-5: estimate coherent Lee-Carter parameters
# (here ax is computed from the last observed period
# and smoothed over ages)
lc.est <- lillecarter.estimate(mx130$male, mx130$female,
                             ax.index = ncol(mx130$male), ax.smooth = TRUE)

# Steps 6-9: project future mortality rates based on future
# life expectancies from WPP2017
e0f <- as.numeric(subset(e0Fproj, name == country)[-(1:2)])
e0m <- as.numeric(subset(e0Mproj, name == country)[-(1:2)])
names(e0f) <- names(e0m) <- colnames(e0Fproj)[-(1:2)]
pred <- mortcast(e0m, e0f, lc.est)

# plot projection for the first and last future time period
plot(pred$female$mx[, "2015-2020"], type="l", log="y",
     ylim=range(pred$female$mx, pred$male$mx), xaxt="n",
     ylab="mx", xlab="Age", main=country, col="red")
axis(1, at=1:nrow(pred$female$mx),
     labels=rownames(pred$female$mx))
lines(pred$male$mx[, "2015-2020"], col="blue")
lines(pred$female$mx[, "2095-2100"], col="red", lty=2)
lines(pred$male$mx[, "2095-2100"], col="blue", lty=2)
legend("topleft", legend=c("male 2015-2020", "female 2015-2020",
                          "male 2095-2100", "female 2095-2100"), bty="n",
     col=rep(c("blue", "red"),2), lty=c(1,1,2,2))
```

 cokannisto

Coherent Kannisto Method

Description

Extrapolate given mortality rates into higher ages using the Coherent Kannisto method as described in Sevcikova et al. (2016).

Usage

```
cokannisto(mxM, mxF, est.ages = seq(80, 95, by = 5),
           proj.ages = seq(100, 130, by = 5))
```

Arguments

<code>mxM</code>	A vector or matrix of male mortality rates. If it is a matrix, rows correspond to age groups with rownames identifying the corresponding age as integers. By default five-years age groups are assigned to rows if rownames are not given.
<code>mxF</code>	A vector or matrix of female mortality rates. Its length or dimension should be the same <code>mxM</code> .
<code>est.ages</code>	A vector of integers identifying the ages to be used for estimation. It should be a subset of rownames of <code>mxM</code> .
<code>proj.ages</code>	A vector of integers identifying the age groups for which mortality rates are to be projected.

Details

The function first estimates the coherent Kannisto parameters by passing mortality rates for age groups `est.ages` into the `cokannisto.estimate` function. The estimated parameters are then passed to the projection function `kannisto.predict` to extrapolate into ages `proj.ages`. Lastly, the input mortality objects are extended by results for the extrapolated ages. If `proj.ages` contains age groups that are included in `mxM` and `mxF`, values for those age groups are overwritten.

Value

A list of two vectors or matrices (for male and female) containing the input mortality objects extended by the extrapolated age groups.

References

Sevcikova H., Li N., Kantorova V., Gerland P., Raftery A.E. (2016). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. In: Schoen R. (eds) Dynamic Demographic Analysis. The Springer Series on Demographic Methods and Population Analysis, vol 39. Springer, Cham

See Also

[cokannisto.estimate](#), [kannisto.predict](#)

Examples

```
data(mxM, mxF, package = "wpp2017")
country <- "South Africa"
mxm <- subset(mxM, name == country)[,-(1:3)]
mxf <- subset(mxF, name == country)[,-(1:3)]
rownames(mxm) <- rownames(mxf) <- c(0,1, seq(5, 100, by=5))
mxnew <- cokannisto(mxm, mxf)
ages <- as.integer(rownames(mxnew$male))
plot(ages, mxnew$male[, "2095-2100"], type="l", log="y",
     xlab="age", ylab="mx", col="blue", main=country)
lines(ages, mxnew$female[, "2095-2100"], col="red")
lines(ages, mxnew$male[, "2010-2015"], lty=2, col="blue")
lines(ages, mxnew$female[, "2010-2015"], lty=2, col="red")
legend("bottomright", legend=c("male 2010-2015", "female 2010-2015",
                               "male 2095-2100", "female 2095-2100"), bty="n",
       col=rep(c("blue", "red"),2), lty=c(2,2,1,1))
```

cokannisto.estimate *Coherent Kannisto Estimation*

Description

Estimate the coherent Kannisto parameters as described in Sevcikova et al. (2016).

Usage

```
cokannisto.estimate(mxM, mxF, ages, fitted = TRUE)
```

Arguments

mxM	A vector of male mortality rates.
mxF	A vector of female mortality rates.
ages	A vector of ages corresponding to mxM and mxF.
fitted	Logical. If TRUE the fitted values and residuals are returned.

Details

Given the Kannisto equation $\text{logit}(m_x) = \log(c) + dx$, the Coherent Kannisto method estimates the d parameter jointly for male and female data, in order to prevent mortality crossovers in higher ages.

Value

List of two lists, one for male and one for female. Each of the two lists contains the following components:

coefficients: named vector with the coherent Kannisto coefficients c and d . The d values are the same in both lists.

fitted.values: the fitted values (not included if `fitted` is FALSE)

residuals: input rates minus the fitted values (not included if `fitted` is FALSE)

References

Sevcikova H., Li N., Kantorova V., Gerland P., Raftery A.E. (2016). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. In: Schoen R. (eds) Dynamic Demographic Analysis. The Springer Series on Demographic Methods and Population Analysis, vol 39. Springer, Cham

See Also

[cokannisto](#), [kannisto.predict](#), [kannisto](#)

Examples

```
data(mxM, mxF, package = "wpp2017")
country <- "Brazil"
mxm <- subset(mxM, name == country)[,"2010-2015"]
mxf <- subset(mxF, name == country)[,"2010-2015"]
cokannisto.estimate(mxm[18:21], mxf[18:21], ages = 18:21)
```

kannisto

Kannisto Method

Description

Extrapolate given mortality rates using the original Kannisto method.

Usage

```
kannisto(mx, est.ages = seq(80, 95, by = 5), proj.ages = seq(100, 130,
  by = 5))
```

Arguments

<code>mx</code>	A vector or matrix of mortality rates. If it is a matrix, rows correspond to age groups with rownames identifying the corresponding age as integers. By default five-years age groups are assigned to rows if rownames are not given.
<code>est.ages</code>	A vector of integers identifying the ages to be used for estimation. It should be a subset of rownames of <code>mx</code> .
<code>proj.ages</code>	A vector of integers identifying the age groups for which mortality rates are to be projected.

Details

The function first estimates the original Kannisto parameters by passing mortality rates for age groups `est.ages` into the `kannisto.estimate` function. The estimated parameters are then passed to the projection function `kannisto.predict` to extrapolate into ages `proj.ages`. Lastly, the input mortality object is extended by results for the extrapolated ages. If `proj.ages` contains age groups that are included in `mx`, values for those age groups are overwritten.

Value

A vector or matrix containing the input mortality object `mx` extended by the extrapolated age groups.

References

Thatcher, A. R., Kannisto, V. and Vaupel, J. W. (1998). The Force of Mortality at Ages 80 to 120, volume 5 of Odense Monographs on Population Aging Series. Odense, Denmark: Odense University Press.

See Also

[kannisto.estimate](#), [kannisto.predict](#), [cokannisto](#)

Examples

```
data(mxM, package = "wpp2017")
mx <- subset(mxM, name == "Burkina Faso")[,-(1:3)]
rownames(mx) <- c(0,1, seq(5, 100, by=5))
mxnew <- kannisto(mx)
ages <- as.integer(rownames(mxnew))
plot(ages, mxnew[, "2095-2100"], type="l", log="y",
     xlab="age", ylab="mx", col="red")
lines(ages, mxnew[, "2010-2015"])
```

kannisto.estimate *Kannisto Estimation*

Description

Estimate the Kannisto parameters (Thatcher et al. 1998).

Usage

```
kannisto.estimate(mx, ages)
```

Arguments

`mx` A vector of mortality rates.
`ages` A vector of ages corresponding to `mx`.

Details

Given the Kannisto equation $\text{logit}(m_x) = \log(c) + dx$, the function estimates the c and d parameters using values of ages as the covariate x .

Value

List with the following components:

`coefficients`: named vector with Kannisto coefficients c and d .

`fitted.values`: the fitted values

`residuals`: input rates minus the fitted values

References

Thatcher, A. R., Kannisto, V. and Vaupel, J. W. (1998). The Force of Mortality at Ages 80 to 120, volume 5 of Odense Monographs on Population Aging Series. Odense, Denmark: Odense University Press.

See Also

[kannisto.predict](#), [kannisto](#), [cokannisto.estimate](#)

Examples

```
data(mxM, package = "wpp2017")
mx <- subset(mxM, name == "Canada")[, "2010-2015"]
kannisto.estimate(mx[18:21], ages = 18:21)
```

kannisto.predict	<i>Kannisto Prediction</i>
------------------	----------------------------

Description

Given estimated Kannisto parameters (coherent or original), it predicts mortality rates for given ages.

Usage

```
kannisto.predict(pars, ages)
```

Arguments

pars	A named vector with Kannisto coefficients c and d (e.g. result of kannisto.estimate or cokannisto.estimate).
ages	A vector of ages to make prediction for.

Details

Given parameters c and d in `pars`, the function uses the Kannisto equation $\text{logit}(m_x) = \log(c) + dx$, to predict mortality rates for age groups x given by `ages`.

Value

Vector of predicted mortality rates.

References

Thatcher, A. R., Kannisto, V. and Vaupel, J. W. (1998). The Force of Mortality at Ages 80 to 120, volume 5 of Odense Monographs on Population Aging Series. Odense, Denmark: Odense University Press.

See Also

[cokannisto](#), [kannisto.estimate](#), [cokannisto.estimate](#)

Examples

```
data(mxM, mxF, package = "wpp2017")
mxm <- subset(mxM, name == "Germany")[, "2010-2015"]
ages <- c(0, 1, seq(5, 130, by=5))

# using original Kannisto parameters
pars <- kannisto.estimate(mxm[18:21], ages = ages[18:21])
mxm.pred <- kannisto.predict(pars$coefficients, ages = ages[22:28])
plot(ages, c(mxm[1:21], mxm.pred), type="l", log="y",
      xlab="age", ylab="mx")
```

```
# Coherent Kannisto
mxf <- subset(mxF, name == "Germany"), "2010-2015"]
copars <- cokannisto.estimate(
  mxm[18:21], mxf[18:21], ages = ages[18:21])
cmxm.pred <- kannisto.predict(copars[["male"]]$coefficients, ages = ages[22:28])
cmxf.pred <- kannisto.predict(copars[["female"]]$coefficients, ages = ages[22:28])
plot(ages, c(mxm[1:21], cmxm.pred), type="l", log="y",
  xlab="age", ylab="mx", col="blue")
lines(ages, c(mxf[1:21], cmxf.pred), col="red")
```

leecarter.estimate *Lee-Carter Estimation*

Description

Estimate Lee-Carter parameters (Lee and Carter 1992).

Usage

```
leecarter.estimate(mx, ax.index = NULL, ax.smooth = FALSE,
  bx.postprocess = TRUE, nx = 5)
```

Arguments

mx	A matrix of age-specific mortality rates where rows correspond to age groups and columns correspond to time periods.
ax.index	A vector of column indices of mx to be used to estimate the a_x parameter. By default all time periods are used.
ax.smooth	Logical allowing to smooth the a_x over ages.
bx.postprocess	Logical determining if numerical anomalies in b_x should be dealt with.
nx	Size of age groups. By default ages are determined by rownames of mx. This argument is only used if mx has no rownames. If nx is 5, the age groups are 0, 1, 5, 10, For nx equals 1, the age groups are 0, 1, 2, 3,

Details

The function estimates parameters of $\log(m_x(t)) = a_x + b_x k(t) + \epsilon_x(t)$ (Lee and Carter 1992). The argument `ax.index` determines which time periods to use to estimate the a_x parameter, while `ax.smooth` controls if the resulting a_x should be smoothed over ages (see Sevcikova et al. 2016 for details).

Value

List with elements `ax`, `bx` and `kt` corresponding to the estimated parameters.

References

Lee, R. D. and Carter, L. (1992). Modeling and forecasting the time series of US mortality. *Journal of the American Statistical Association*, 87, 659-671.

Sevcikova H., Li N., Kantorova V., Gerland P., Raftery A.E. (2016). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. In: Schoen R. (eds) *Dynamic Demographic Analysis*. The Springer Series on Demographic Methods and Population Analysis, vol 39. Springer, Cham

See Also

[mortcast](#), [lileecarter.estimate](#)

Examples

```
data(mxM, package = "wpp2017")
mx <- subset(mxM, name == "Netherlands")[,4:16]
rownames(mx) <- c(0,1, seq(5, 100, by=5))
lc.ax.avg <- leecarter.estimate(mx)
lc.ax.last <- leecarter.estimate(mx, ax.index=ncol(mx))
plot(lc.ax.avg$ax, type="l")
lines(lc.ax.last$ax, col="blue")
```

life.table

Life Table Function

Description

Function for obtaining life table quantities from mortality rates.

Usage

```
life.table(mx, sex = c("male", "female", "total"), abridged = TRUE,
  radix = 1, open.age = 130)
```

Arguments

mx	Vector of age-specific mortality rates nm_x . If <code>abridged</code> is <code>TRUE</code> (default), the elements correspond to $1m_0, 4m_1, 5m_5, 5m_{10}, \dots$. If <code>abridged</code> is <code>FALSE</code> , they correspond to $1m_0, 1m_1, 1m_2, 1m_3, \dots$.
sex	Which sex the mortality rates correspond to.
abridged	Is it an abridged life table (<code>TRUE</code> , default) or not (<code>FALSE</code>). In the former case, the <code>mx</code> vector is interpreted as corresponding to age groups 0, 1-4, 5-9, 10-14, \dots . If <code>FALSE</code> , the <code>mx</code> vector is interpreted as corresponding to one-year age groups, i.e. 0, 1, 2, 3, \dots .
radix	Base of the life table.
open.age	Open age group. If smaller than the last age group of <code>mx</code> , the life table is truncated. It does not have any effect if larger than the last age group.

Details

Computes a life table corresponding to given mortality rates for either 5- or 1-year age groups. The implementation follows Preston et al. (2001), including the choice of a_x (see Table 3.3 on page 48). For compatibility with computations done at the UN, we set a_x for ages 5 and 10 (in the abridged version) to 2.5.

References

Preston, S.H., Heuveline, P., Guillot, M. (2001). Demography: Measuring and Modeling Population Processes. Oxford: Blackwell Publishers Ltd.

Examples

```
data(mxF, e0Fproj, package = "wpp2017")
# get female mortality of Mexico for the current year
country <- "Mexico"
mxf <- subset(mxF, name == country)[,"2010-2015"]
life.table(mxf, sex = "female")
```

lileecarter.estimate *Coherent Lee-Carter Estimation*

Description

Estimate coherent Lee-Carter parameters (Li and Lee 2005).

Usage

```
lileecarter.estimate(mXM, mxF, ...)
```

Arguments

mXM	A matrix of male age-specific mortality rates where rows correspond to age groups and columns correspond to time periods.
mxF	A matrix of female mortality rates of the same shape as mXM.
...	Additional arguments passed to leecarter.estimate .

Details

The coherent Lee-Carter parameters for male and female mortality rates share the same b_x which is the average of the age-specific b_x parameters.

The function in addition computes the ultimate b_x^u as defined in Li et al. (2013) based on the coherent b_x .

Value

List containing elements b_x (coherent b_x parameter), `ultimate.bx` (ultimate b_x^u parameter), and lists `female` and `male`, each with the Lee-Carter parameters.

References

Li, N. and Lee, R. D. (2005). Coherent mortality forecasts for a group of populations: An extension of the Lee-Carter method. *Demography*, 42, 575-594.

Li, N., Lee, R. D. and Gerland, P. (2013). Extending the Lee-Carter method to model the rotation of age patterns of mortality decline for long-term projections. *Demography*, 50, 2037-2051.

Examples

```
data(mxM, mxF, package = "wpp2017")
country <- "Germany"
mxm <- subset(mxM, name == country)[,4:16]
mxf <- subset(mxF, name == country)[,4:16]
rownames(mxm) <- rownames(mxf) <- c(0,1, seq(5, 100, by=5))
lc <- lileecarter.estimate(mxm, mxf)
plot(lc$bx, type="l")
lines(lc$ultimate.bx, lty=2)
```

logquad

Log-Quadratic Mortality Model

Description

Predict age-specific mortality rates using the Log-Quadratic Mortality Model (Wilmoth et al. 2012).

Usage

```
logquad(e0, sex = c("male", "female", "total"), my.coefs = NULL,
        q5ranges = c(1e-04, 0.9), k = 0, keep.lt = FALSE)
```

```
logquadj(e0m, e0f, ...)
```

Arguments

<code>e0</code>	Vector of target life expectancies.
<code>sex</code>	Which sex does the give <code>e0</code> corresponds to.
<code>my.coefs</code>	Data frame with columns “sex”, “age”, “ax”, “bx”, “cx”, “vx”. The “sex” column should contain values “female”, “male” and/or “total”. The “age” column must be sorted so that it assures that rows correspond to ages in increasing order. Any NAs are internally converted to zeros. If not given, the dataset LQcoef is used.

q5ranges	A vector of size two, giving the min and max of 5q0 used in the bisection method.
k	Value of the k parameter.
keep.lt	Logical. If TRUE additional life table columns are kept in the resulting object.
e0m	A time series of target male life expectancy.
e0f	A time series of target female life expectancy.
...	Additional arguments passed to the underlying function.

Details

The LogQuad method in this implementation projects mortality rates using the equation

$$\log(m_x) = a_x + b_x h + c_x h^2 + v_x k$$

where a_x , b_x , c_x and v_x are age-specific coefficients, $h = \log(5q0)$ (i.e. reflects child mortality), and k should be chosen to match 45q15 (adult mortality) or set to 0 (default). The coefficients can be passed as inputs, or taken from the package default dataset `LQcoef` which are taken from <http://www.demog.berkeley.edu/~jrw/LogQuad>.

For the given inputs and values of life expectancy $e0$, the function finds values of h that best match $e0$, using life tables and the bisection method. It returns the corresponding mortality schedule for each value of $e0$.

Function `logquad` is for one sex, while `logquadj` can be used for both sexes.

Value

Function `logquad` returns a list with the following elements: a matrix `mx` with the predicted mortality rates. If `keep.lt` is TRUE, it also contains matrices `sr` (survival rates), and life table quantities `Lx` and `lx`. Function `logquadj` returns a list of objects, one for each sex.

References

Wilmoth, J., Zureick, S., Canudas-Romo, V., Inoue, M., Sawyer, C. (2012). A Flexible Two-Dimensional Mortality Model for Use in Indirect Estimation. *Population studies*, 66(1), 1-28. doi: [10.1080/00324728.2011.611411](https://doi.org/10.1080/00324728.2011.611411)

See Also

[LQcoef](#), [mortcast.blend](#), [mortcast](#), [pmd](#), [mlt](#)

Examples

```
data(e0Mproj, package = "wpp2017")
country <- "Brazil"
# get target e0
e0m <- as.numeric(subset(e0Mproj, name == country)[-(1:2)])
# project into future
pred <- logquad(e0m, sex = "male")
# plot first projection in black and the remaining ones in heat colors
```

```
plot(pred$mx[,1], type = "l", log = "y", ylim = range(pred$mx),
      ylab = "male mx", xlab = "Age", main = country)
for(i in 2:ncol(pred$mx)) lines(pred$mx[,i],
                                col = heat.colors(20)[i])
```

LQcoef

Coefficients for the Log-Quadratic Mortality Model

Description

Data object containing a table of coefficients to be used in the Log-Quadratic Model as implemented in the [logquad](#) function.

Usage

```
data(LQcoef)
```

Format

Data frame containing columns “sex”, “age”, “ax”, “bx”, “cx”, “vx”. Rows correspond to sex and age groups.

Source

<http://www.demog.berkeley.edu/~jrw/LogQuad>

References

Wilmoth, J., Zureick, S., Canudas-Romo, V., Inoue, M., Sawyer, C. (2012). A Flexible Two-Dimensional Mortality Model for Use in Indirect Estimation. *Population studies*, 66(1), 1-28. doi: [10.1080/00324728.2011.611411](https://doi.org/10.1080/00324728.2011.611411)

See Also

[logquad](#)

Examples

```
data(LQcoef)
head(LQcoef)
```

mlt

*Model Life Tables Mortality Patterns***Description**

Predict age-specific mortality rates using Coale-Demeny and UN model life tables.

Usage

```
mlt(e0, sex = c("male", "female"), type = "CD_West")
```

```
mltj(e0m, e0f, ...)
```

Arguments

e0	A time series of target life expectancy.
sex	Either "male" or "female".
type	Type of the model life table. Available options are "CD_East", "CD_North", "CD_South", "CD_West", "UN_Chilean", "UN_Far_Eastern", "UN_General", "UN_Latin_American", "UN_South_Asian".
e0m	A time series of target male life expectancy.
e0f	A time series of target female life expectancy.
...	Additional arguments passed to the underlying function.

Details

Given a level of life expectancy (e0), sex and a type of model life table, the function extracts the corresponding mortality pattern from [MLTlookup](#), while interpolating between neighboring e0 groups. Function `mlt` is for one sex, while `mltj` can be used for both sexes.

Value

Function `mlt` returns a matrix with the predicted mortality rates. Columns correspond to the values in the `e0` vector and rows correspond to age groups. Function `mltj` returns a list of such matrices, one for each sex.

References

<https://population.un.org/wpp/Download/Other/MLT>

Coale, A., P. Demeny, and B. Vaughn. 1983. Regional model life tables and stable populations. 2nd ed. New York: Academic Press.

See Also

[mortcast](#), [mortcast.blend](#), [pmd](#), [MLTlookup](#)

Examples

```

data(e0Fproj, package = "wpp2017")
country <- "Uganda"
# get target e0
e0f <- subset(e0Fproj, name == country)[- (1:2)]
# project into future using life table Coale-Demeny North
mx <- mlt(e0f, sex = "female", type = "CD_North")
# plot first projection in black and the remaining ones in grey
plot(mx[,1], type = "l", log = "y", ylim = range(mx),
      ylab = "female mx", xlab = "Age", main = country)
for(i in 2:ncol(mx)) lines(mx[,i], col = "grey")

```

MLTlookup

*Model Life Tables Lookup***Description**

Lookup table containing values for various model life tables, including Coale-Demeny and UN life tables.

Usage

```
data(MLTlookup)
```

Format

Data frame with the following columns:

type Type of the model life table. Available options are “CD_East”, “CD_North”, “CD_South”, “CD_West”, “UN_Chilean”, “UN_Far_Eastern”, “UN_General”, “UN_Latin_American”, “UN_South_Asian”. For the CD types, see Coale et al. (1983). For the UN types, see <https://population.un.org/wpp/Download/Other/MLT>.

sex Code for distinguishing sexes. 1 is for male, 2 is for female.

age Starting age of an age group. These are 0, 1, 5, 10, ... 130.

e0 Level of life expectancy, starting at 20 and going by steps of 2.5 up to 130.

mx Mortality rates.

lx, Lx, sx Other life table columns.

References

Coale, A., P. Demeny, and B. Vaughn. 1983. Regional model life tables and stable populations. 2nd ed. New York: Academic Press.

<https://population.un.org/wpp/Download/Other/MLT>

See Also[mlt](#)**Examples**

```
data(MLTlookup)
str(MLTlookup)
# CD West life table for male at e0 of 80
subset(MLTlookup, type == "CD_West" & sex == 1 & e0 == 80)
```

mortcast

Coherent Rotated Lee-Carter Prediction

Description

Predict age-specific mortality rates using the coherent rotated Lee-Carter method.

Usage

```
mortcast(e0m, e0f, lc.pars, rotate = TRUE, keep.lt = FALSE,
         constrain.all.ages = FALSE)
```

Arguments

<code>e0m</code>	A time series of future male life expectancy.
<code>e0f</code>	A time series of future female life expectancy.
<code>lc.pars</code>	A list of coherent Lee-Carter parameters with elements <code>bx</code> , <code>ultimate.bx</code> , <code>female</code> and <code>male</code> as returned by <code>lileecarter.estimate</code> . The female and male objects are again lists that should contain a vector <code>ax</code> and optionally a matrix <code>axt</code> if the a_x parameter needs to be defined as time dependent. In such a case, rows are age groups and columns are time periods corresponding to the length of the <code>e0f</code> and <code>e0m</code> vectors.
<code>rotate</code>	If TRUE the rotation method of b_x is used as described in Li et al. (2013).
<code>keep.lt</code>	Logical. If TRUE additional life table columns are kept in the resulting object.
<code>constrain.all.ages</code>	By default the method constrains the male mortality to be above female mortality for old ages if the male life expectancy is below the female life expectancy. Setting this argument to TRUE causes this constraint to be applied to all ages.

Details

This function implements Steps 6-9 of Algorithm 2 in Sevcikova et al. (2016). It uses an abridged life table function to find the level of mortality that corresponds to the given life expectancy.

Value

List with elements `female` and `male`, each of which contains a matrix `mx` with the predicted mortality rates. If `keep.lt` is `TRUE`, it also contains matrices `sr` (survival rates), and life table quantities `Lx` and `lx`.

References

Li, N., Lee, R. D. and Gerland, P. (2013). Extending the Lee-Carter method to model the rotation of age patterns of mortality decline for long-term projections. *Demography*, 50, 2037-2051.

Sevcikova H., Li N., Kantorova V., Gerland P., Raftery A.E. (2016). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. In: Schoen R. (eds) *Dynamic Demographic Analysis*. The Springer Series on Demographic Methods and Population Analysis, vol 39. Springer, Cham

See Also

[rotate.leecarter](#), [leecarter.estimate](#), [lileecarter.estimate](#), [mortcast.blend](#)

Examples

```
data(mxM, mxF, e0Fproj, e0Mproj, package = "wpp2017")
country <- "Brazil"
# estimate parameters from historical mortality data
mxm <- subset(mxM, name == country)[,4:16]
mxf <- subset(mxF, name == country)[,4:16]
rownames(mxm) <- rownames(mxf) <- c(0,1, seq(5, 100, by=5))
lc <- lileecarter.estimate(mxm, mxf)
# project into future
e0f <- as.numeric(subset(e0Fproj, name == country)[-(1:2)])
e0m <- as.numeric(subset(e0Mproj, name == country)[-(1:2)])
pred <- mortcast(e0m, e0f, lc)
# plot first projection in black and the remaining ones in grey
plot(pred$female$mx[,1], type="l", log="y", ylim=range(pred$female$mx),
      ylab="female mx", xlab="Age", main=country)
for(i in 2:ncol(pred$female$mx)) lines(pred$female$mx[,i], col="grey")
```

Description

Predict age-specific mortality rates using a blend of two different methods (Coherent Lee-Carter, Coherent Pattern Mortality Decline, or Model Life Tables). Weights can be applied to fine-tune the blending mix.

Usage

```
mortcast.blend(e0m, e0f, meth1 = "lc", meth2 = "mlt", weights = c(1,
  0.5), apply.kannisto = TRUE, min.age.groups = 28,
  meth1.args = NULL, meth2.args = NULL, kannisto.args = NULL)
```

Arguments

e0m	A time series of future male life expectancy.
e0f	A time series of future female life expectancy.
meth1	Character string giving the name of the first method to blend. It is one of “lc”, “pmd”, “mlt” or “logquad”, corresponding to Coherent Lee-Carter (function mortcast), Pattern Mortality Decline (function copmd), Log-Quadratic model (function logquadj), and Model Life Tables (function mltj), respectively.
meth2	Character string giving the name of the second method to blend. One of the same choices as meth1.
weights	Numeric vector with values between 0 and 1 giving the weight of meth1. If it is a single value, the same weight is applied for all time periods. If it is a vector of size two, it is assumed these are weights for the first and the last time period. Remaining weights will be interpolated. Note that meth2 is weighted by 1 - weights.
apply.kannisto	Logical. If TRUE and if any of the methods results in less than min.age.groups age categories, the coherent Kannisto method (cokannisto) is applied to extend the age groups into old ages.
min.age.groups	Minimum number of age groups. Triggers the application of Kannisto, see above.
meth1.args	List of arguments passed to the function that corresponds to meth1.
meth2.args	List of arguments passed to the function that corresponds to meth2.
kannisto.args	List of arguments passed to the cokannisto function if Kannisto is applied.

Details

The function allows to combine two different methods using given weights. The weights can change over time - by default they are interpolated from the starting weight to the end weight. The projection is done for both sexes, so that coherent methods can be applied.

Value

List with elements `female` and `male`, each of which contains a matrix `mx` with the predicted mortality rates. In addition, it contains elements `meth1res` and `meth2res` which contain the results of the functions corresponding to the two methods. Elements `meth1` and `meth2` contain the names of the methods. A vector `weights` contains the final (possibly interpolated) weights.

See Also

[mortcast](#), [copmd](#), [mltj](#), [logquad](#), [cokannisto](#)

Examples

```

data(mxM, mxF, e0Fproj, e0Mproj, package = "wpp2017")
country <- "Brazil"
# estimate parameters from historical mortality data
mxm <- subset(mxM, name == country)[,4:16]
mxf <- subset(mxF, name == country)[,4:16]
rownames(mxm) <- rownames(mxf) <- c(0,1, seq(5, 100, by=5))
lcest <- lileecarter.estimate(mxm, mxf)
# project into future
e0f <- subset(e0Fproj, name == country)[- (1:2)]
e0m <- subset(e0Mproj, name == country)[- (1:2)]
# Blend LC and MLT
pred1 <- mortcast.blend(e0m, e0f, meth1 = "lc", meth2 = "mlt",
  meth1.args = list(lc.pars = lcest),
  meth2.args = list(type = "CD_North"),
  weights = c(1,0.25))
# Blend PMD and MLT
pred2 <- mortcast.blend(e0m, e0f, meth1 = "pmd", meth2 = "mlt",
  meth1.args = list(mxm0 = mxm[, "2010-2015"],
    mxf0 = mxf[, "2010-2015"]))
# plot projection by time
plotmx <- function(pred, iage, main)
  with(pred, {
    # blended projections
    plot(female$mx[iage,], type="l",
      ylim=range(meth1res$female$mx[iage,],
        meth2res$female$mx[iage,]),
      ylab="female mx", xlab="Time", main=main, col = "red")
    lines(meth1res$female$mx[iage,], lty = 2)
    lines(meth2res$female$mx[iage,], lty = 3)
    legend("topright", legend=c("blend", meth1, meth2),
      lty = 1:3, col = c("red", "black", "black"), bty = "n")
  })
age.group <- 3 # 5-9 years old
par(mfrow=c(1,2))
plotmx(pred1, age.group, "LC-MLT (age 5-9)")
plotmx(pred2, age.group, "PMD-MLT (age 5-9)")

```

Description

Predict age-specific mortality rates using the Pattern of mortality decline (PMD) method (Andreev et al. 2013).

Usage

```
pmd(e0, mx0, sex = c("male", "female"), interp.rho = FALSE,
    kranges = c(0, 25), keep.lt = FALSE, keep.rho = FALSE)

copmd(e0m, e0f, mxm0, mx0, interp.rho = FALSE, keep.rho = FALSE, ...)
```

Arguments

<code>e0</code>	A vector of target life expectancy, one element for each predicted time point.
<code>mx0</code>	A vector with starting age-specific mortality rates.
<code>sex</code>	Either "male" or "female".
<code>interp.rho</code>	Logical controlling if the ρ coefficients should be interpolated (TRUE) or if the raw (binned) version should be used (FALSE), as stored in the dataset PMDrho .
<code>kranges</code>	A vector of size two, giving the min and max of the k parameter which is estimated to match the target <code>e0</code> using the bisection method.
<code>keep.lt</code>	Logical. If TRUE additional life table columns are kept in the resulting object.
<code>keep.rho</code>	Logical. If TRUE the ρ coefficients are included in the resulting object.
<code>e0m</code>	A time series of target male life expectancy.
<code>e0f</code>	A time series of target female life expectancy.
<code>mxm0</code>	A vector with starting age-specific male mortality rates.
<code>mx0</code>	A vector with starting age-specific female mortality rates.
<code>...</code>	Additional arguments passed to the underlying function. For <code>copmd</code> , in addition to <code>kranges</code> and <code>keep.lt</code> , it can be <code>sexratio.adjust</code> which is a logical controlling if a sex-ratio adjustment should be applied to prevent crossovers between male and female <code>mx</code> . In such a case it uses coefficients from the PMDadjcoef dataset. However, if the argument <code>adjust.with.mx0</code> is set to TRUE (in addition to <code>sexratio.adjust</code>), the adjustment is done using the female mortality rates as the lower constraint for male mortality rates. If the argument <code>adjust.sr.if.needed</code> is set to TRUE, a sex-ratio adjustment is performed dynamically, using the sex ratio in the previous time point. In such a case, an adjustment in time t is applied only if there was a drop of sex ratio below one at time $t-1$.

Details

These functions implements the PMD method introduced in Andreev et al. (2013). It assumes that the future decline in age-specific mortality will follow a certain pattern with the increase in life expectancy at birth (`e0`):

$$\log mx(t) = \log mx(t-1) - k(t)\rho_x(t)$$

Here, $\rho_x(t)$ is the age-specific pattern of mortality decline between $t-1$ and t . Such patterns for each sex and various levels of `e0` are stored in the dataset [PMDrho](#). The `pmd` function can be instructed to interpolate between neighboring levels of `e0` by setting the argument `interp.rho` to TRUE. The k parameter is estimated to match the `e0` level using the bisection method.

Function `pmd` evaluates the method for a single sex, while `copmd` does it coherently for both sexes. In the latter case, the same ρ_x (namely the average over sex-specific ρ_x) is used for both, male and female.

Value

Function `pmd` returns a list with the following elements: a matrix `mx` with the predicted mortality rates. If `keep.lt` is TRUE, it also contains matrices `sr` (survival rates), and life table quantities `Lx` and `lx`. If `keep.rho` is TRUE, it contains a matrix `rho` where columns correspond to the values in the `e0` vector and rows correspond to age groups.

Function `copmd` returns a list with one element for each sex (male and female) where each of them is a list as described above. In addition if `keep.rho` is TRUE, element `rho.sex` gives the sex-dependent (i.e. not averaged) ρ_x coefficient.

References

Andreev, K., Gu, D., Gerland, P. (2013). Age Patterns of Mortality Improvement by Level of Life Expectancy at Birth with Applications to Mortality Projections. Paper presented at the Annual Meeting of the Population Association of America, New Orleans, LA. <http://paa2013.princeton.edu/papers/132554>.

Gu, D., Pelletier, F., Sawyer, C. (2017). Projecting Age-sex-specific Mortality: A Comparison of the Modified Lee-Carter and Pattern of Mortality Decline Methods, UN Population Division, Technical Paper No. 6. New York: United Nations. https://population.un.org/wpp/Publications/Files/WPP2017_TechnicalPaperNo6.pdf

See Also

[mortcast](#), [mortcast.blend](#), [PMDrho](#)

Examples

```
data(mxF, e0Fproj, package = "wpp2017")
country <- "Hungary"
# get initial mortality for the current year
mxf <- subset(mxF, name == country)[,"2010-2015"]
names(mxf) <- c(0,1, seq(5, 100, by=5))
# get target e0
e0f <- subset(e0Fproj, name == country)[- (1:2)]
# project into future
pred <- pmd(e0f, mxf, sex = "female")
# plot first projection in black and the remaining ones in grey
plot(pred$mx[,1], type = "l", log = "y", ylim = range(pred$mx),
      ylab = "female mx", xlab = "Age", main = country)
for(i in 2:ncol(pred$mx)) lines(pred$mx[,i], col = "grey")
```

PMDadjcoef

Coefficients for Sex Ratio Adjustments in the PMD Method

Description

Data object containing a table of coefficients to be used to adjust the sex ratio in the coherent Pattern Mortality Decline method as implemented in the `copmd` function. To invoke the adjustment, argument `sexratio.adjust` should be set to TRUE.

Usage

```
data(PMDadjcoef)
```

Format

Data frame containing columns “age”, “intercept”, “lmxf”, “e0f”, “e0f2”, and “gap”. Rows correspond to age groups. The values are estimates of the following regression

$$\log_{10} mx^M = \beta_0 + \beta_1 \log_{10} mx^F + \beta_2 e_0^F + \beta_3 (e_0^F)^2 + \beta_4 (e_0^F - e_0^M)$$

The order of the columns starting with intercept corresponds to the order of the coefficients in the above equation.

Source

The coefficients were estimated and provided by Danan Gu, UN Population Division.

References

Gu, D., Pelletier, F. and Sawyer, C. (2017). Projecting Age-sex-specific Mortality: A Comparison of the Modified Lee-Carter and Pattern of Mortality Decline Methods, UN Population Division, Technical Paper No. 6. New York: United Nations. https://population.un.org/wpp/Publications/Files/WPP2017_TechnicalPaperNo6.pdf

See Also

`copmd`

Examples

```
data(PMDadjcoef)
PMDadjcoef
```

PMDrho

Pattern Mortality Decline Lookup Tables

Description

Data object containing two tables with ρ coefficients for the Pattern Mortality Decline method as implemented in the `pmd` function.

Usage

```
data(PMDrho)
```

Format

Using `data(PMDrho)` loads two objects into memory: `RhoFemales` and `RhoMales`. They both are data frames with 22 rows corresponding to age groups, and 17 columns corresponding to different levels of life expectancy in 5-years intervals (from 50 to 135). The names of the columns reflect the middle of the respective interval.

References

Andreev, K. Gu, D., Gerland, P. (2013). Age Patterns of Mortality Improvement by Level of Life Expectancy at Birth with Applications to Mortality Projections. Paper presented at the Annual Meeting of the Population Association of America, New Orleans, LA. <http://paa2013.princeton.edu/papers/132554>.

Gu, D., Pelletier, F. and Sawyer, C. (2017). Projecting Age-sex-specific Mortality: A Comparison of the Modified Lee-Carter and Pattern of Mortality Decline Methods, UN Population Division, Technical Paper No. 6. New York: United Nations. https://population.un.org/wpp/Publications/Files/WPP2017_TechnicalPaperNo6.pdf

See Also

[pmd](#)

Examples

```
data(PMDrho)
head(RhoFemales)
head(RhoMales)

# plot a few male patterns
e0lev <- colnames(RhoMales)[c(1, 5, 9, 13, 17)]
plot(RhoMales[, e0lev[1]], type="l", log="y", ylim=range(RhoMales[,e0lev]),
     ylab="male rho", xlab="Age")
for(i in 2:length(e0lev)) lines(RhoMales[,e0lev[i]], lty = i)
legend("bottomleft", legend = e0lev, lty = 1:length(e0lev), bty= "n")
```

rotate.leecarter	<i>Rotated Lee-Carter</i>
------------------	---------------------------

Description

Rotate the Lee-Carter parameter b_x over time to reach an ultimate b_x^u , as described in Li et al. (2013).

Usage

```
rotate.leecarter(bx, ultimate.bx, e0, e0l = 80, e0u = 102, p = 0.5)

ultimate.bx(bx)
```

Arguments

bx	A vector of the Lee-Carter b_x parameter (from e.g. lileecarter.estimate or leecarter.estimate).
ultimate.bx	A vector of the ultimate b_x^u parameter as defined in Li, Lee, Gerland (2013) (obtained using lileecarter.estimate or ultimate.bx).
e0	A time series of life expectancies.
e0l	Level of life expectancy at which the rotation starts.
e0u	Level of life expectancy at which the rotation finishes.
p	Exponent of the smooth function.

Value

Function `rotate.leecarter` returns a matrix of rotated $B_x(t)$ where rows correspond to age groups and columns correspond to time periods (given by the vector `e0`).

Function `ultimate.bx` returns a vector of the ultimate b_x^u .

References

Li, N., Lee, R. D. and Gerland, P. (2013). Extending the Lee-Carter method to model the rotation of age patterns of mortality decline for long-term projections. *Demography*, 50, 2037-2051.

Examples

```
data(mxF, mxM, e0Fproj, e0Mproj, package = "wpp2017")
country <- "Japan"
mxm <- subset(mxM, name == country)[,4:16]
mxf <- subset(mxF, name == country)[,4:16]
e0f <- as.numeric(subset(e0Fproj, name == country)[-(1:2)])
e0m <- as.numeric(subset(e0Mproj, name == country)[-(1:2)])
rownames(mxm) <- rownames(mxf) <- c(0,1, seq(5, 100, by=5))
lc <- lileecarter.estimate(mxm, mxf)
rotlc <- rotate.leecarter(lc$bx, lc$ultimate.bx, (e0f + e0m)/2)
```

```
plot(lc$bx, type="l")  
lines(lc$ultimate.bx, col="red")  
for(i in 1:ncol(rotlc)) lines(rotlc[,i], col="grey")
```

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