Package ‘DemoDecomp’

August 14, 2018

Type Package
Title Decompose Demographic Functions
Version 1.0.1
Date 2018-08-13
Author Tim Riffe
Maintainer Tim Riffe <tim.riffe@gmail.com>
License GPL-3
LazyLoad yes
RoxygenNote 6.1.0
Suggests covr
RdMacros Rdpack
Imports Rdpack
Depends R (>= 2.10)

BugReports https://github.com/timriffe/DemoDecomp/issues
Encoding UTF-8
NeedsCompilation no
Repository CRAN
Date/Publication 2018-08-14 09:30:09 UTC

R topics documented:

  Compare ................................................................. 2
  horiuchi ............................................................... 2
  LTabr ................................................................. 5
Comparison decomposition results by age and cause

A matrix containing the contributions to the difference in life expectancy at birth between 2002 US males and England and Wales males. Ages (in rows) are in abridged categories, 0-85, and there are six causes, including other, in columns. The sum of the matrix is the difference in life expectancy at birth between the two populations. Values are based on symmetrical stepwise replacement from young to old ages only. This is just to make sure implementation is close.

Usage

Compare

Format

A matrix with 19 rows and 6 columns

Source


---

Numerical Approximation of Continuous Decomposition

Description

This is an exact R implementation of the decomposition code in Matlab offered by the authors in the supplementary material given here: <http://www.demog.berkeley.edu/~jrw/Papers/decomp.suppl.pdf>. The difference between DecompContinuous() and this function is that DecompContinuousOrig takes rates1 and rates2 as single vectors, rather than as matrices, and output is also returned as a vector. This difference makes the function more flexible, but may add a step when writing the function to be decomposed. See examples.
Usage
horiuchi(func, pars1, pars2, N, ...)

Arguments

func A function specified by the user. This must be able to take the vectors rates1 or rates2 as its argument, and to return the value of the function, y, when evaluated for these rates. It may also have additional arguments, not to be decomposed.

pars1 vector of covariates to be passed on as arguments to func(). Covariates can be in any order, as long as func() knows what to do with them. pars1 is for time 1 (or population 1).

pars2 is the same as pars2 but for time/population 2.

N The number of intervals to integrate over.

... optional parameters to pass on to func(). These are not decomposed.

Details
The decomposition works by assuming a linear change in all covariates between time 1 and time 2 (or population 1 and population 2). At each small time step approaching time 2 (the size of which is the inverse of N) each covariate is moved forward along its linear trajectory. One at a time, each covariate (of which there are ages*variables of) is switched out twice, once for its value at 1/(2N) forward and once for its value at 1/(2N) backward in time. The difference between func() evaluated with these two rate matrices is the change in y attributable to that particular covariate and that particular time step. Summing over all N time steps, we get the contribution to the difference of each covariate, effectmat. The sum of effectmat should come very close to func(rates2)−func(rates1). The error decreases with larger N, but there is not much point in having an N higher than 100, and 20 is usually sufficient. This ought to be able to handle a very wide variety of functions.

If pars1 are observations from 2005 and pars2 are observations from 2006 an N of 20 would imply a delta of 1/20 of a year for each integration step. Higher N provides finer results (a lower total residual), but takes longer to compute. In general, there are decreasing returns to higher N. sum(effectmat) ought to approximate func(rates2)−func(rates1).

Value
returns effectmat, a matrix of the variable effects that is organized in the same way as pars1 and pars2.

References
Examples

data(rates1)
data(rates2)

# we need rates1 and rates2 as vectors
rates1 <- c(rates1)
rates2 <- c(rates2)
# look at the function:
R0vec
# 2 things to point out:
# 1) it has an argument pfem, proportion female of births (1/(1+SRB)),
# that must be specified, but that we don’t care about decomposing
# 2) x is a single vector. The the inside of the function needs to
# either refer to parts of it by indexing, as done here, or else
# re-assign x to various objects. In this case x[1:1] is Lx and
# x[(1+1):(2*1)] is Fx...
A <- horiuchi(func = R0vec,
    pars1 = rates1,
    pars2 = rates2,
    N = 10,
    pfem = .4886)
# the output, A, is also a single vector. Each element corresponds
# to the effect of changes in that particular covariate toward the
# overall change in the function value. sum(A) should be close to
# original difference
(check1 <- R0vec(rates2) - R0vec(rates1))
(check2 <- sum(A))

# This package does not supply default plotting functions, but one
# strategy might be the following:

# reorder A into a matrix (sideways):
A <- t(matrix(A,ncol=2))
# call barplot() twice, once for positive values and again for
# negative values
Apos <- A * .5 * (sign(A) + 1)
Aneg <- A * .5 * abs(sign(A) - 1)
## Not run:
barplot(Apos,
    width = rep(1, length(A) / 2),
    space = 0,
    ylim = range(A),
    main = "A fake decomposition of R0",
    col=c("yellow","green"),
    axisnames = FALSE,
    xlim=c(0, 90),
    ylab = "contrib to change in R0",
    cex.axis = .7)
barplot(Aneg,
LTabr

width = rep(1, length(A) / 2),
add = TRUE,
space = 0,
col = c("yellow", "green"),
axes = FALSE, axisnames = FALSE)
segments(seq(from=0, to=90, by=10), 0, seq(from=0, to=90, by=10), -0.27, lty=2, col="grey")
text(seq(from=0, to=90, by=10), -0.27, seq(from=0, to=90, by=10), pos=1, xpd=T)
legend("bottomright", fill=c("yellow", "green"), legend=c("contrib from change in Lx",
"contrib from change in Fx"), title="age specific contrib of changes in Fx and Lx", bg="white")

## End(Not run)

LTabr

an abridged lifetable based on $M(x)$

Description

Implements the abridged lifetable formulas given in the supplementary material to Andreev et. al. (2012). An entire lifetable is calculated, but only life expectancy at birth is returned.

Usage

LTabr(Mx, Age = c(0, 1, cumsum(rep(5, length(Mx) - 2))), radix = 1e+05)

Arguments

Mx numeric vector of abridged mortality rates.
Age integer, abridged age lower bounds.
radix numeric. Can be anything positive.

Details

Chiang’s $a(x)$ is assumed in the following way: $a(0) = 0.07 + 1.7 * M(0)$, $a(1) = 1.6$, $a(\omega) = \frac{1}{M(\omega)}$, and all others are assumed at mid interval. The last age is assumed open. Everything else is pretty standard.

Value

numeric life expectancy at birth

References

### Mxc1

**Year 2002 death rates by cause for US males in abridged age classes**

**Description**

A matrix containing death rates for six causes (one of which is other) for abridged age classes 0-85. Ages are labelled in rows, and causes in column names.

**Usage**

Mxc1

**Format**

A matrix with 19 rows and 6 columns

**Source**


### Mxc2

**Year 2002 death rates by cause for England and Wales males in abridged age classes**

**Description**

A matrix containing death rates for six causes (one of which is other) for abridged age classes 0-85. Ages are labelled in rows, and causes in column names.

**Usage**

Mxc2

**Format**

A matrix with 19 rows and 6 columns

**Source**

### Mxc2e0abr

**get life expectancy at birth from an (abridged)age-cause matrix**

**Description**

Given a matrix with abridged ages in rows and causes of death in columns, then calculate life expectancy at birth using `LTabr()`.

**Usage**

```
Mxc2e0abr(Mxc)
```

**Arguments**

- **Mxc**: numeric matrix

**Details**

This assumes that the marginal row sums give all-cause mortality rates. Give an other category if you need to top-up to all-cause mortality. Do not include all-cause mortality itself!

**Value**

numeric life expectancy at birth

---

### Mxc2e0abrvec

**get life expectancy at birth from the vec of an age-cause matrix**

**Description**

Given a vector with abridged ages stacked within causes of death, assign its dimensions, take the age marginal sums using `Mxc2e0abr`, then calculate life expectancy at birth using `LTabr()`.

**Usage**

```
Mxc2e0abrvec(Mxcvec, dims, trans = FALSE)
```

**Arguments**

- **Mxcvec**: numeric vector, `c(Mxc)`
- **dims**: integer vector of length two, `c(nrow(Mxc), ncol(Mxc))`
- **trans**: do we need to transpose in order to arrive back to an age-cause matrix?
Details

This assumes that the marginal row sums give all-cause mortality rates. Give an other category if you need to top-up to all-cause mortality. Do not include all-cause mortality itself! length(Mxcvec) must equal prod(dim(Mxc)). This function is meant to be fed to a generic decomposition function, such as stepwise_replacement(), or DecompContinuousOrig().

Value

numeric life expectancy at birth

R0vec

R0vec Calculates net reproduction, R0, according to a given set of rates Lx,fx and a fixed proportion female of births, pfem.

Description

This function is only provided for the examples of horiuchi(). It calculates the sum of the row products of rates multiplied by pfem.

Usage

R0vec(x, pfem = 0.4886)

Arguments

- **x**: a single vector containing Lx followed by Fx or vice versa. Here, Lx is the survival function integrated within each age interval and with a lifetable radix of 1. Fx is the fertility function, calculated as births/person years of exposure. Fx should simply contain zeros in ages with no fertility, OR, all vectors should be limited to reproductive ages. Both Lx and Fx should for this function be of the same length.
- **pfem**: the proportion female of births. Something like .49, .48, or (1/(2.05)). This can either be specified as a single number, or it may be allowed to vary by age. For the later case, be sure to specify a value for each age (length(x)/2 values). Default .4886.

Details

The main feature that functions need to have when specified for horiuchi() or stepwise_replacement() is that the rates must all go into a (potentially long) vector, probably consisting in your rate vectors one after the other. Really the decomposition function does not care how things are arranged in the vector- the components of change vector that is returned from horiuchi() will be arranged in exactly the same way as its input rate vectors, so as long as you know how to sort it out, and your function can extract what it needs from the vectors, then it can be specified in any way. For this particular example function, R0vec(), x must be specified with either Lx followed by Fx or vice versa. It would also be possible to redefine the function to place pfem in with the rates vector, x, which would allow this item to be decomposed too. Here it is specified separately in order to demonstrate passing on parameters to the function within horiuchi().
Value

the value of R0 for the given set of rates and proportion female of births.

Examples

data(rates1)
# take vec:
x <- c(rates1)
R0vec(x)

rates1

Fake data generated for example.

Description

The first column lx is a discrete survival function for time point 1. The second column fx are age specific fertility rates.

Usage

rates1

Format

An object of class matrix with 101 rows and 2 columns.

Examples

## Not run:
data(rates1)
data(rates2)
# nothing fancy
# compare lx
plot(rates1[,1],type='l',col="blue")
lines(rates2[,1],col="green")
# compare fx
plot(rates1[,2],type='l',col="blue")
lines(rates2[,2],col="green")

## End(Not run)
rates2

 Fake data generated for example.

Description

The first column $l_x$ is a discrete survival function for time point 2. The second column $f_x$ are age specific fertility rates.

Usage

rates2

Format

An object of class matrix with 101 rows and 2 columns.

Examples

```r
## Not run:
data(rates1)
data(rates2)
# nothing fancy
# compare lx
plot(rates1[,1], type='l', col="blue")
lines(rates2[,1], col="green")
# compare fx
plot(rates1[,2], type='l', col="blue")
lines(rates2[,2], col="green")

## End(Not run)
```

stepwise_replacement

implementation of the decomposition algorithm of stepwise replacement

Description

This implements the algorithm described in Andreev et al (2002), with defaults set to approximate their recommendations for replacement ordering and result averaging.

Usage

```r
stepwise_replacement(func, pars1, pars2, symmetrical = TRUE,
direction = "up", ...)
```
Arguments

- **func**
  A function specified by the user. This must be able to take the vectors `pars1` or `pars2` as its argument, and to return the value of the function, `y`, when evaluated for these rates. It may also have additional arguments, not to be decomposed.

- **pars1**
  Vector of covariates to be passed on as arguments to `func()`. Covariates can be in any order, as long as `func()` knows what to do with them. `pars1` is for time 1 (or population 1).

- **pars2**
  Is the same as `pars1` but for time/population 2.

- **symmetrical**
  Logical. Default `TRUE` as recommended by authors. Shall we average the results of replacing 1 with 2 and 2 with 1?

- **direction**
  Character. One of "up", "down", or "both". Default "up", as recommended by authors.

- **...**
  Optional parameters to pass on to `func()`.

Details

The `symmetrical` argument toggles whether or not we replace `pars1` with `pars2` (FALSE), or take the arithmetic average or replacement in both directions. `direction` refers to whether we go from the bottom up or top down, or take the arithmetic average of these when replacing vector elements. Although the total difference will always sum correctly, the calculated contribution from individual components can vary greatly depending on the order in general. Defaults are set to symmetrically replace from the bottom up, per the authors’ suggestion.

Value

A matrix of the variable effects that is organized in the same way as `pars1` and `pars2`.

References


Examples

data(Mxc1)
data(Mxc2)
# we'll want to pass in these dimensions
dims <- dim(Mxc1)
# we need parameters in vec form
Mxc1v <- c(Mxc1)
Mxc2v <- c(Mxc2)
B <- stepwise_replacement(func = Mxc2e0abrvec,
pars1 = Mxc1v, pars2 = Mxc2v, dims = dims,
# authors' recommendations:
symmetrical = TRUE, direction = "up")
dim(B) <- dims
# the output, B, is also a single vector. Each element corresponds
# to the effect of changes in that particular covariate toward the
# overall change in the function value. sum(B) should equal the
# original difference
(check1 <- Mxc2e@abr(Mxc2) - Mxc2e@abr(Mxc1))
(check2 <- sum(B))

# This package does not supply default plotting functions, but one
# strategy might be the following:
## Not run:
Age <- c(0, 1, seq(5, 85, by = 5))
matplot(Age, B, type = 'l',
xlab = "Age", ylab = "Contrib to diff in e(0)", col = 1:6)
legend("bottomleft",lty=1:5,col=1:6,
    legend = c("Neoplasms","Circulatory","Respiratory",
        "Digestive","Acc/viol","Other"))

## End(Not run)
Index

*Topic datasets
  Compare, 2
  Mxc1, 6
  Mxc2, 6
  rates1, 9
  rates2, 10

Compare, 2

horiuchi, 2

LTabr, 5

Mxc1, 6
Mxc2, 6
Mxc2e0abr, 7
Mxc2e0abrvec, 7

R0vec, 8
rates1, 9
rates2, 10

stepwise_replacement, 10