

# Package ‘bayesPop’

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MortCast

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**Description** Generating population projections for all countries of the world using several probabilistic components, such as total fertility rate and life expectancy.

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bayesPop-package	<i>Probabilistic Population Projection</i>
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## Description

The package allows to generate population projections for all countries of the world using several probabilistic components, such as total fertility rate (TFR) and life expectancy. Generating subnational projections is also supported.

## Details

Package: bayesPop  
 Type: Package  
 Version: 8.1-1  
 Date: 2019-10-25  
 URL: <https://bayespop.csss.washington.edu>

The main function is called `pop.predict`. It uses trajectories of TFR from the **bayesTFR** package and life expectancy from the **bayesLife** package and for each trajectory it computes a population projection using the Cohort component method. It results in probabilistic age and sex specific projections. Various plotting functions are available for results visualization (`pop.trajectories.plot`, `pop.pyramid`, `pop.trajectories.pyramid`, `pop.map`), as well as a `summary` function. Aggregations can be derived using `pop.aggregate`. An `expression language` is available to obtain the distribution of various population quantities.

Subnational projections can be generated using `pop.predict.subnat`. Function `pop.aggregate.subnat` aggregates such projections.

## Author(s)

Hana Sevcikova, Adrian Raftery, Thomas Buettner

Maintainer: Hana Sevcikova <hanas@uw.edu>

## References

H. Sevcikova, A. E. Raftery (2016). bayesPop: Probabilistic Population Projections. *Journal of Statistical Software*, 75(5), 1-29. doi:10.18637/jss.v075.i05

A. E. Raftery, N. Li, H. Sevcikova, P. Gerland, G. K. Heilig (2012). Bayesian probabilistic population projections for all countries. *Proceedings of the National Academy of Sciences* 109:13915-13921.

P. Gerland, A. E. Raftery, H. Sevcikova, N. Li, D. Gu, T. Spoorenberg, L. Alkema, B. K. Fosdick, J. L. Chunn, N. Lalic, G. Bay, T. Buettner, G. K. Heilig, J. Wilmoth (2014). World Population Stabilization Unlikely This Century. *Science* 346:234-237.

H. Sevcikova, N. Li, V. Kantorova, P. Gerland and A. E. Raftery (2015). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. arXiv:1503.05215. <http://arxiv.org/abs/1503.05215>

## See Also

[bayesTFR](#), [bayesLife](#)

## Examples

```
## Not run:
sim.dir <- tempfile()
# Generates population projection for one country
country <- "Netherlands"
pred <- pop.predict(countries=country, output.dir=sim.dir)
summary(pred, country)
pop.trajectories.plot(pred, country)
dev.off()
pop.trajectories.plot(pred, country, sum.over.ages=TRUE)
pop.pyramid(pred, country)
pop.pyramid(pred, country, year=2100, age=1:26)
unlink(sim.dir, recursive=TRUE)

## End(Not run)

# Here are commands needed to run probabilistic projections
# from scratch, i.e. including TFR and life expectancy.
# Note that running the first four commands
# (i.e. predicting TFR and life expectancy) can take
# LONG time (up to several days; see below for possible speed-up).
# For a toy simulation, set the number of iterations (iter)
# to a small number.
## Not run:
sim.dir.tfr <- "directory/for/TFR"
sim.dir.e0 <- "directory/for/e0"
sim.dir.pop <- "directory/for/pop"

# Estimate TFR parameters (speed-up by including parallel=TRUE)
run.tfr.mcmc(iter="auto", output.dir=sim.dir.tfr, seed=1)
```

```

# Predict TFR (if iter above < 4000, reduce burnin and nr.traj accordingly)
tfr.predict(sim.dir=sim.dir.tfr, nr.traj=2000, burnin=2000)

# Estimate e0 parameters (females) (speed-up by including parallel=TRUE)
# Can be run independently of the two commands above
run.e0.mcmc(sex="F", iter="auto", output.dir=sim.dir.e0, seed=1)

# Predict female and male e0
# (if iter above < 22000, reduce burnin and nr.traj accordingly)
e0.predict(sim.dir=sim.dir.e0, nr.traj=2000, burnin=20000)

# Population prediction
pred <- pop.predict(output.dir=sim.dir.pop, verbose=TRUE,
  inputs = list(tfr.sim.dir=sim.dir.tfr,
    e0F.sim.dir=sim.dir.e0, e0M.sim.dir="joint_"))
pop.trajectories.plot(pred, "Madagascar", nr.traj=50, sum.over.ages=TRUE)
pop.trajectories.table(pred, "Madagascar")

## End(Not run)

```

---

age.specific.migration

*Reconstruction of Sex- and Age-specific Migration*

---

## Description

Reconstructs the sex- and age-specific net migration datasets out of the total net migration using a residual method.

## Usage

```

age.specific.migration(wpp.year = 2019, years = seq(1955, 2100, by = 5),
  countries = NULL, smooth = TRUE, rescale = TRUE, ages.to.zero = 18:21,
  write.to.disk = FALSE, directory = getwd(), file.prefix = "migration",
  depratio = wpp.year == 2015, verbose = TRUE)

```

## Arguments

wpp.year	Integer determining which <b>wpp</b> package should be used to get the necessary data from. That package is required to have a dataset on total net migration (called <a href="#">migration</a> ). Currently, only packages <a href="#">wpp2015</a> , <a href="#">wpp2017</a> and <a href="#">wpp2019</a> contain it (see <a href="#">Details</a> ).
years	Array of years that the reconstruction should be made for. This should be a subset of years for which the total net migration is available.
countries	Numerical country codes to do the reconstruction for. By default it is performed on all countries included in the <a href="#">migration</a> dataset where aggregations are excluded.

<code>smooth</code>	Logical controlling if smoothing of the reconstructed curves is required. Due to rounding issues the residual method often yields unrealistic zig-zags on migration curves by age. Smoothing usually improves their look.
<code>rescale</code>	Logical controlling if the resulting migration should be rescaled to match the total migration.
<code>ages.to.zero</code>	Indices of age groups where migration should be set to zero. Default is 85 and older.
<code>write.to.disk</code>	If TRUE results are written to disk.
<code>directory</code>	Directory where to write the results if <code>write.to.disk</code> is TRUE.
<code>file.prefix</code>	If <code>write.to.disk</code> is TRUE results are written into two text files with this prefix, a letter “M” and “F” determining the sex, and concluded by the “.txt” suffix. By default “migrationM.txt” and “migrationF.txt”.
<code>depratio</code>	If it is TRUE it will use an internal dataset on migration dependency ratios to adjust the first three age groups. It can also be a name of a binary file containing such dataset.
<code>verbose</code>	Logical controlling the amount of output messages.

### Details

Unlike in [wpp2012](#), for the three latest releases of the WPP, the **wpp2015**, **wpp2017**, and **wpp2019**, the UN Population Division did not publish the sex- and age-specific net migration counts, only the totals. However, since the sex- and age-schedules are needed for population projections, this function attempts to reconstruct those missing datasets. It uses the published population projections by age and sex, fertility and mortality projections from the **wpp** package. It computes the population projection without migration and sets the residual to the published population projection as the net migration. By default such numbers are then scaled so that the sum over sexes and ages corresponds to the total migration count.

If `smooth` is TRUE a smoothing procedure is performed over ages where necessary. Also, for simplicity, we set migration of old ages to zero (default is 85+). Both is done before the scaling. If it is desired to obtain raw residuals without any additional processing, set `smooth=FALSE`, `rescale=FALSE`, `ages.to.zero=c()`.

### Value

List of two data frames (male and female), each having the same structure as [migrationM](#).

### Warning

Due to rounding issues and slight differences in the methodology, this function does not reproduce the unpublished UN datasets exactly. It is only an approximation! Especially, the first age groups might be more off than other ages.

### Note

The function is called automatically from [pop.predict](#) if no migration inputs is given. Thus, only users that need sex- and age-specific migration for other purposes will need to call this function explicitly.

**Author(s)**

Hana Sevcikova

**See Also**

[pop.predict](#), [migration](#) [migrationM](#)

**Examples**

```
## Not run:  
asmig <- age.specific.migration()  
head(asmig$male)  
head(asmig$female)  
## End(Not run)
```

---

`get.countries.table`    *Accessing Country Information*

---

**Description**

The function returns a data frame containing codes and names of all countries used in the prediction.

**Usage**

```
## S3 method for class 'bayesPop.prediction'  
get.countries.table(object, ...)
```

**Arguments**

<code>object</code>	Object of class <a href="#">bayesPop.prediction</a> .
<code>...</code>	Not used.

**Value**

Data frame with columns code and name.

**Author(s)**

Hana Sevcikova

---

get.pop.prediction      *Accessing Prediction Object*

---

### Description

Function `get.pop.prediction` retrieves results of a prediction from disk and creates an object of class `bayesPop.prediction`. Function `has.pop.prediction` checks an existence of such results.

### Usage

```
get.pop.prediction(sim.dir, aggregation = NULL, write.to.cache = TRUE)
```

```
has.pop.prediction(sim.dir)
```

```
pop.cleanup.cache(pop.pred)
```

### Arguments

<code>sim.dir</code>	Directory where the prediction is stored. It should correspond to the value of the <code>output.dir</code> argument used in the <code>pop.predict</code> function.
<code>aggregation</code>	If given, the prediction object is considered to be an aggregation and both arguments are passed to <code>get.pop.aggregation</code> .
<code>write.to.cache</code>	Logical controlling if other functions are allowed to write the cache of this prediction object (see Details).
<code>pop.pred</code>	Object of class <code>bayesPop.prediction</code> .

### Details

The `pop.predict` function stores resulting trajectories into a directory called `output.dir/prediction`. Here the argument `sim.dir` should correspond to `output.dir` (i.e. without the “prediction” part).

In addition to retrieving prediction results, the `get.pop.prediction` function also looks for a file called ‘`cache.rda`’ and loads it into an environment called `cache`. If it does not exist, it creates an empty cache environment. See `pop.map` - Section Performance and Caching. The environment can be cleaned up using the `pop.cleanup.cache` function which also deletes the ‘`cache.rda`’ file on disk. If `write.to.cache` is `FALSE`, other functions are not allowed to manipulate the ‘`cache.rda`’ file.

### Value

Function `has.pop.prediction` returns a logical indicating if a prediction exists.

Function `get.pop.prediction` returns an object of class `bayesPop.prediction`.

### Author(s)

Hana Sevcikova

**See Also**

[bayesPop.prediction](#), [get.pop.aggregation](#)

**Examples**

```
sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir)
summary(pred)
```

---

LifeTableMx

*Life Table Functions*

---

**Description**

Functions for obtaining life table quantities.

**Usage**

```
LifeTableMx(mx, sex = c("Male", "Female", "Total"), include01 = TRUE,
radix = 1, open.age = 130)
```

```
LifeTableMxCol(mx, colname = c("Lx", "lx", "qx", "mx", "dx", "Tx", "sx", "ex", "ax"), ...)
```

**Arguments**

<code>mx</code>	Vector of age-specific mortality rates $nm_x$ . The elements correspond to $1m_0$ , $4m_1$ , $5m_5$ , $5m_{10}$ , ... It can have no more than 28 elements which corresponds to age up to 130. In the <code>LifeTableMxCol</code> function, this argument can be a two-dimensional matrix with first dimension being the age.
<code>sex</code>	For which sex is the life table.
<code>include01</code>	Logical. If it is <code>FALSE</code> the first two age groups (0-1 and 1-4) are collapsed to one age group (0-4).
<code>radix</code>	Base of the life table.
<code>open.age</code>	Open age group. If smaller than the last age group of $mx_m$ , the life table is truncated.
<code>colname</code>	Name of the column of the life table that should be returned.
<code>...</code>	Arguments passed to underlying functions. Argument <code>age05</code> is a logical vector of size three, specifying if the age groups 0-1, 1-4 and 0-5 should be included. Default value of <code>c(FALSE, FALSE, TRUE)</code> includes the 0-5 age group only.

**Details**

Function `LifeTableMx` returns a life table for one set of mortality rates. Function `LifeTableMxCol` returns one column of the life table for (possibly) multiple sets of mortality rates.

**Value**

Function `LifeTableMx` returns a data frame with the following elements:

<code>age</code>	Age groups
<code>mx</code>	<code>mx</code> , the input vector of mortality rates.
<code>qx</code>	<code>nqx</code> , probability of dying between ages $x$ and $x+n$ .
<code>lx</code>	<code>lx</code> , number left alive at age $x$ .
<code>dx</code>	<code>ndx</code> , cohort deaths between ages $x$ and $x+n$ .
<code>Lx</code>	<code>nLx</code> , person-years lived between ages $x$ and $x+n$ .
<code>sx</code>	<code>sx</code> , survival rate at age $x$ .
<code>Tx</code>	<code>Tx</code> , person-years lived above age $x$ .
<code>ex</code>	<code>e0x</code> , expectation of life at age $x$ .
<code>ax</code>	<code>nax</code> , average person-years lived in the interval by those dying in the interval.

Function `LifeTableMxCol` returns one given column of the life table, possibly as a matrix (if `mx` is a matrix).

**Author(s)**

Hana Sevcikova, Thomas Buettner, Nan Li, Patrick Gerland

**References**

Preston, P., Heuveline, P., Guillot, M. (2001): Demography. Blackwell Publishing Ltd.

**See Also**

[pop.expressions](#) for examples on retrieving some life table quantities.

**Examples**

```
## Not run:
sim.dir <- tempfile()
pred <- pop.predict(countries="Ecuador", output.dir=sim.dir, wpp.year=2015,
  present.year=2015, keep.vital.events=TRUE, fixed.mx=TRUE, fixed.pasfr=TRUE)
# get male mortality rates from 2020 for age groups 0-1, 1-4, 5-9, ...
mxm <- pop.byage.table(pred, expression="MEC_M{age.index01(27)}", year=2020)[,1]
print(LifeTableMx(mxm), digits=3)
# female LT with first two age categories collapsed
mxf <- pop.byage.table(pred, expression="MEC_F{age.index01(27)}", year=2020)[,1]
print(LifeTableMx(mxf, sex="Female", include01=FALSE), digits=3)
unlink(sim.dir, recursive=TRUE)
## End(Not run)
```

---

mac.expression	<i>Expression Generator</i>
----------------	-----------------------------

---

## Description

Help functions to easily generate commonly used expressions.

## Usage

```
mac.expression(country)
```

## Arguments

country            Country code as defined for [expressions](#).

## Details

mac.expression generates an expression for the mean age of childbearing of the given country. Note that [pop.predict](#) has to be run with `keep.vital.events=TRUE` for this to work.

## Value

mac.expression returns a character string corresponding to the formula  $(17.5 * R_c(15 - 19) + 22.5 * R_c(20 = 24) + \dots + 47.5 * R_c(45 - 49))/100$  where  $R_c(x)$  denotes the country-specific percent age-specific fertility for the age group  $x$ .

## See Also

[pop.expressions](#)

## Examples

```
## Not run:
sim.dir <- tempfile()
# Run pop.predict with storing vital events
pred <- pop.predict(countries=c("Germany", "France"), nr.traj=3,
                   keep.vital.events=TRUE, output.dir=sim.dir)
# plot the mean age of childbearing
pop.trajectories.plot(pred, expression=mac.expression("FR"), cex.main = 0.7)
unlink(sim.dir, recursive=TRUE)
## End(Not run)
```

---

MLTbx

*Dataset on Lee-Carter bx for Modeled Countries*

---

### Description

Dataset with values of the Lee-Carter bx parameter for countries where mortality was obtained using model life tables.

### Usage

```
data(MLTbx)
```

### Format

A data frame with nine rows and 28 columns. Each row corresponds to one mortality age pattern as defined in the [vwBaseYear](#) dataset. Each column corresponds to an age group, starting with 0-1, 1-4, 5-9, 10-14, ... up to 125-129, 130+.

### Details

These values are used for countries for which the column `AgeMortalityType` in [vwBaseYear](#) is equal to “Model life tables”. In such a case a row is selected that corresponds to the corresponding value of the column `AgeMortalityPattern` (also in [vwBaseYear](#)). These values are then used instead of estimating the Lee-Carter  $b_x$  from the country’s historical data.

### Source

Data provided by the United Nations Population Division.

### See Also

[vwBaseYear](#)

### Examples

```
data(MLTbx)
str(MLTbx)
```

---

 pop.aggregate

 Aggregation of Population Projections
 

---

### Description

Aggregation of existing countries' population projections into projections of given regions, and accessing such aggregations.

### Usage

```
pop.aggregate(pop.pred, regions,
              input.type = c("country", "region"), name = input.type,
              inputs = list(e0F.sim.dir = NULL, e0M.sim.dir = "joint_", tfr.sim.dir = NULL),
              my.location.file = NULL, verbose = FALSE, ...)
```

```
get.pop.aggregation(sim.dir = NULL, pop.pred = NULL, name = NULL,
                   write.to.cache = TRUE)
```

```
pop.aggregate.subnat(pop.pred, regions, locations, ..., verbose = FALSE)
```

### Arguments

- |            |  |
|------------|--|
| pop.pred   | Object of class <a href="#">bayesPop.prediction</a> containing country-specific population projections.  |
| regions    | Vector of numerical codes of regions. It should correspond to values in the column "country_code" in the <a href="#">UNlocations</a> dataset or in <code>my.location.file</code> (see below). For <code>pop.aggregate.subnat</code> it is a numerical code of a country over which subregions are aggregated.  |
| input.type | There are two methods for aggregating projections depending on the type of inputs, "country"- and "region"-based, see Details.   |
| name       | Name of the aggregation. It becomes a part of a directory name where aggregation results are stored.   |
| inputs     | This argument is only used when the "region"-based method is selected. It is a list of inputs of probabilistic components of the projection: <ul style="list-style-type: none"> <li><b>e0F.sim.dir</b> Simulation directory with projections of female life expectancy (generated using <a href="#">bayesLife</a>). It must contain projections for the given regions (see functions <a href="#">run.e0.mcmc.extra</a>, <a href="#">e0.predict.extra</a>). If it is not given, the same e0 directory is taken which was used for generating the <code>pop.pred</code> object, in which case the e0 projections are re-loaded from disk.</li> <li><b>e0M.sim.dir</b> Simulation directory with projections of male life expectancy. By default (value NULL or "joint_") the function assumes a joint female-male projections of life expectancy and thus tries to load the male projections from the female projection object created using the <code>e0F.sim.dir</code> argument.</li> </ul> |

<b>tfr.sim.dir</b>	Simulation directory with projections of total fertility rate (generated using <a href="#">bayesTFR</a> ). It must contain projections for the given regions (see functions <a href="#">run.tfr.mcmc.extra</a> , <a href="#">tfr.predict.extra</a> ). If it is not given, the same TFR directory is taken which was used for generating the <code>pop.pred</code> object, in which case the TFR projections are re-loaded from disk.
<code>my.location.file</code>	User-defined location file that can contain other aggregation groups than the default UN location file. It should have the same structure as the <a href="#">UNlocations</a> dataset, see below.
<code>verbose</code>	Logical switching log messages on and off.
<code>sim.dir</code>	Simulation directory where aggregation is stored. It is the same directory used for creating the <code>pop.pred</code> object. Alternatively, <code>pop.pred</code> can be used. Either <code>sim.dir</code> or <code>pop.pred</code> must be given.
<code>write.to.cache</code>	Logical controlling if functions operating on this object are allowed to write into its cache (see Details of <a href="#">get.pop.prediction</a> ).
<code>locations</code>	Name of a tab-delimited file that contains definitions of the sub-regions. It should be the same file as used for the <code>locations</code> argument in <a href="#">pop.predict.subnat</a> .
<code>...</code>	Additional arguments. For a country-type aggregation, it can be logical use <code>kannisto</code> which determines if the Kannisto method should be used for old ages when aggregating mortality rates. A logical argument <code>keep.vital.events</code> determines if vital events should be computed for aggregations. Argument <code>adjust</code> determines if country-level population numbers should be adjusted to the WPP values.

## Details

Function `pop.aggregate` triggers an aggregations over countries while function `pop.aggregate.subnat` is used for aggregation over sub-regions to a country. The following details refer to the use of `pop.aggregate`. For sub-national aggregation see Example in [pop.predict.subnat](#).

The dataset [UNlocations](#) or `my.location.file` is used to determine countries to be aggregated, in particular the field “location\_type” of the entries with “country\_code” given in the `regions` argument. One can aggregate over the following location types: Type 0 means aggregating all countries of the world (or in the file), type 2 is aggregating over continents, type 3 is aggregating over regions within continents, and any other integer (except 4) corresponds to user-defined aggregations. Note that type 4 is reserved as a location type of countries and thus, all aggregations are performed over entries of this type. For type 2, countries are matched using the “area\_code” column; for type 3 the matching is done using the “reg\_code” column of the [UNlocations](#) dataset. E.g., if `regions=908` (Europe) which has location type 2 in the default [UNlocations](#) dataset, all countries are aggregated for which values of 908 are found in the “area\_code” column. If the location type is other than 0, 2, 3 and 4, there must be a column in the file called “agcode\_x” with  $x$  being the location type. This column is then used to match the countries to be aggregated.

Consider the following example. Say we want to pair four countries (Germany [DE], France [FR], Netherlands [NL], Italy [IT]) in two different ways, so we have two overlapping groupings, each of which has two groups (A,B):

1. group A = (DE, FR), group B = (NL, IT)

2. group A = (DE, NL), group B = (FR, IT)

Then, my.location.file should have the following entries:

country_code	name	location_type	agcode_98	agcode_99
1001	grouping1_groupA	98	-1	-1
1002	grouping1_groupB	98	-1	-1
1003	grouping2_groupA	99	-1	-1
1004	grouping2_groupB	99	-1	-1
276	Germany	4	1001	1003
250	France	4	1001	1004
258	Netherlands	4	1002	1003
380	Italy	4	1002	1004
1005	all	0	-1	-1

The “country\_code” of the groups is user-specific, but it must be unique within the file. Values of “country\_code” for countries must match those in the prediction object. To run the aggregation for the four groups above we set `regions=1001:1004`. Having “location\_type” being 98 and 99, it is expected the file to have columns “agcode\_98” and “agcode\_99” containing assignments to each of the two groupings. Values in this columns corresponding to groups are not used and thus can have any value. For aggregating over all four countries, set `regions=1005` which has “location\_type” equal 0 and thus, it is aggregated over all entries with “location\_type” equals 4.

There are two methods available for generating aggregations of population projection:

**Country-based Method** Aggregations are created by summing trajectories over countries of the given region.

**Region-based Method** The aggregation is generated using the same algorithm as population projections for single countries (function `pop.predict`), but it operates on aggregated input components. These are created as follows. Here  $c$  denotes countries over which we aggregate a region  $R$ ,  $s \in \{m, f\}$ ,  $a$ , and  $t$  denote sex, age category and time, respectively.  $t = P$  denotes the present year of the prediction.  $N_{s,a,t}^c$  and  $M_{s,a,t}^c$ , respectively, denotes the historical population count and the Bayesian predictive median of population, respectively, of sex  $s$ , in age category  $a$  at time  $t$  for country  $c$  (refer to the links in parentheses for description of the data):

**Initial sex and age-specific population (popM, popF):**  $N_{s,a,t=P}^R = \sum_c N_{s,a,t=P}^c$

**Sex and age-specific death rates (mxM, mxF):**  $mx_{s,a,t}^R = \frac{\sum_c (mx_{s,a,t}^c \cdot N_{s,a,t}^c)}{\sum_c N_{s,a,t}^c}$

**Sex ratio at birth (srb):**  $SRB_t^R = \frac{\sum_c M_{s=m,a=1,t}^c}{\sum_c M_{s=f,a=1,t}^c}$

**Percentage age-specific fertility rate (pasfr):**  $PASFR_{a,t}^R = \frac{\sum_c (PASFR_{a,t}^c \cdot M_{s=f,a,t}^c)}{\sum_c M_{s=f,a,t}^c}$

**Migration code and start year (mig.type):** Aggregated migration code is the code of maximum counts over aggregated countries weighted by  $N_{t=P}^c$ . Migration start year is the maximum of start years over aggregated countries.

**Sex and age-specific migration (migM, migF):**  $mig_{s,a,t}^R = \sum_c mig_{s,a,t}^c$

**Probabilistic projection of life expectancy:** We assume an aggregation of life expectancy for the given regions was generated prior to this call, using the `run.e0.mcmc.extra` and `e0.predict.extra` functions of the `bayesLife` package.

**Probabilistic projection of total fertility rate:** We assume an aggregation of total fertility for the given regions was generated prior to this call, using the `run.tfr.mcmc.extra` and `tfr.predict.extra` functions of the **bayesTFR** package.

Results of the aggregations are stored in the same top directory as the `pop.pred` object, in a subdirectory called `'aggregations_name'`. They can be accessed using the function `get.pop.aggregation`. Note that multiple runs of this function with the same name will overwrite previous aggregations results of the same name.

### Value

Object of class `bayesPop.prediction` containing the aggregated results. In addition it contains elements `aggregation.method` giving the input `type` used, and `aggregated.countries` which is a list of countries aggregated for each region.

### Author(s)

Hana Sevcikova, Adrian Raftery

### References

H. Sevcikova, A. E. Raftery (2016). `bayesPop`: Probabilistic Population Projections. *Journal of Statistical Software*, 75(5), 1-29. doi:10.18637/jss.v075.i05

### See Also

`pop.predict`, `tfr.predict.extra`, `e0.predict.extra`

### Examples

```
## Not run:
sim.dir <- tempfile()
pred <- pop.predict(countries=c(528,218,450), output.dir=sim.dir)
aggr <- pop.aggregate(pred, 900) # aggregating World (i.e. all countries available in pred)
pop.trajectories.plot(aggr, 900, sum.over.ages=TRUE)
# countries over which we aggregated:
subset(UNlocations, country_code %in% aggr$aggregated.countries[["900"]])
unlink(sim.dir, recursive=TRUE)
## End(Not run)
```

---

pop.cohorts.plot

*Extracting and Plotting Cohort Data*

---

### Description

Extracts and plots population counts or results of expressions by cohorts.

**Usage**

```
cohorts(pop.pred, country = NULL, expression = NULL, pi = c(80, 95))

pop.cohorts.plot(pop.pred, country = NULL, expression = NULL, cohorts = NULL,
  cohort.data = NULL, pi = c(80, 95), dev.ncol = 5, show.legend = TRUE,
  legend.pos = "bottomleft", ann = par("ann"), add = FALSE, xlab = "", ylab = "",
  main = NULL, xlim = NULL, ylim = NULL, col = "red", ...)
```

**Arguments**

pop.pred	Object of class <a href="#">bayesPop.prediction</a> .
country	Name or numerical code of a country. If it is not given, expression must be specified.
expression	Expression defining the population measure to be plotted. For syntax see <a href="#">pop.expressions</a> . It must be country-specific, i.e. "XXX" is not allowed, and it must contain curly braces, i.e. be age specific.
pi	Probability interval. It can be a single number or an array.
cohorts	Years of the cohorts to be plotted. By default, 10 future cohorts (starting from the last observed one) are used. It can be a single number or an array.
cohort.data	List with the cohort data obtained via the cohorts function. If it is not given, function cohorts is called internally, but by passing this argument the processing is faster.
dev.ncol	Number of column for the graphics device.
show.legend	Logical controlling whether the legend should be drawn.
legend.pos	Position of the legend passed to the <a href="#">legend</a> function.
ann, xlab, ylab, main, xlim, ylim, col, ...	Graphical parameters passed to the plot function.
add	Logical specifying if the plot should be added to an existing graphics.

**Details**

pop.cohorts.plot plots all cohorts passed in the cohorts argument on the same scale of the *y*-axis.

**Value**

Function cohorts returns a list where each element corresponds to one cohort. Each cohort element is a matrix with columns corresponding to years and rows corresponding to the median (first row) and quantiles of the given probability intervals.

**Author(s)**

Hana Sevcikova

**See Also**

[pop.trajectories.plot](#), [pop.byage.plot](#), [pop.expressions](#)

## Examples

```
sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir)
# Population cohorts
pop.cohorts.plot(pred, "Netherlands")
# plot specific cohorts using expression (must contain {})
pop.cohorts.plot(pred, expression="P528{}", cohorts=c(1960, 1980, 2000, 2020))
# the same as
cohort.data <- cohorts(pred, expression="P528{}")
pop.cohorts.plot(pred, cohort.data=cohort.data, cohorts=c(1960, 1980, 2000, 2020))
```

---

pop.expressions

*Expressions as used in Population Output Functions*

---

## Description

Documentation of expressions supported by functions [pop.trajectories.plot](#), [pop.trajectories.plotAll](#), [pop.trajectories.table](#), [pop.byage.plot](#), [pop.byage.table](#), [cohorts](#), [pop.cohorts.plot](#), [pop.map](#), [pop.map.gvis](#), [write.pop.projection.summary](#), [get.pop.ex](#), [get.pop.exba](#).

## Details

The functions above accept an argument `expression` which should define a population measure, i.e. a quantity that can be computed from population projections, observed population data or vital events. Such an expression is a collection of *basic components* connected via usual arithmetic operators, such as `+`, `-`, `*`, `/`, `^`, `%`, `%%`, and combined using parentheses. In addition, standard R functions or predefined functions (see below) can be used within expressions.

A **basic component** is a character string constituted of four parts, two of which are optional. They must be in the following order:

1. Measure identification. One of the following upper-case characters:
  - ‘P’ - population,
  - ‘D’ - deaths,
  - ‘B’ - births,
  - ‘S’ - survival ratio,
  - ‘F’ - fertility rate,
  - ‘R’ - percent age-specific fertility,
  - ‘M’ - mortality rate,
  - ‘Q’ - probability of dying,
  - ‘E’ - life expectancy,
  - ‘G’ - net migration.

All but the ‘P’ and ‘G’ indicators are available only if the [pop.predict](#) function was run with `keep.vital.events=TRUE`.

2. Country part. One of the following:

- Numerical country code (as used in [UNlocations](#), see [http://en.wikipedia.org/wiki/ISO\\_3166-1\\_numeric](http://en.wikipedia.org/wiki/ISO_3166-1_numeric)),
  - two- or three-character ISO 3166 code, see [http://en.wikipedia.org/wiki/ISO\\_3166-1\\_alpha-2](http://en.wikipedia.org/wiki/ISO_3166-1_alpha-2), [http://en.wikipedia.org/wiki/ISO\\_3166-1\\_alpha-3](http://en.wikipedia.org/wiki/ISO_3166-1_alpha-3),
  - characters “XXX” which serves as a wildcard for a country code.
3. Sex part (optional): The country part can be followed by either “\_F” (for female) or “\_M” (for male).
  4. Age part (optional): If used, the basic component is concluded by an age index given as an array. Such array is embraced by either brackets (“[” and “]”) or curly braces (“{” and “}”). The former invokes a summation of counts over given ages, the latter is used when no summation is desired. Note that if this part is missing, counts are automatically summed over all ages. To use all ages without summing, empty curly braces can be used. Age index one corresponds to age 0-4, index two corresponds to age 5-9 etc. Indicators ‘S’, ‘M’, ‘Q’ and ‘E’ allow an index -1 which corresponds to age 0-1 and an index 0 which corresponds to age 1-4. Use the pre-defined functions `age.index01(...)` and `age.index05(...)` (see below) to define the right indices.

Not all combinations of the four parts above make sense. For example, ‘F’ and ‘R’ can be only combined with female sex, ‘B’, ‘F’ and ‘R’ can be only combined with a subset of the age groups, namely child-bearing ages (indices 4 to 10). Or, there is no point in summing the life table based indicators (M, Q, E, S) over sexes, i.e. using it without the sex part, or over multiple age groups, i.e. using brackets.

Examples of basic components are “P276”, “D50\_F[4:10]”, “PXXX{14:27}”, “SCZE\_M{ }”, “QIE\_M[-1]”.

When the expression is evaluated on a prediction object, each basic component is substituted by an array of four dimensions (using the `get.pop` function):

1. Country dimension: Equals to one if a specific country code is given, or it equals the number of countries in the prediction object if a wildcard is used.
2. Age dimension: Equals to one if the third component above is missing or the age is defined within square brackets. If the age is defined within curly braces, this dimension corresponds to the length of the age array.
3. Time dimension: Depending on the time context of the expression, this dimension corresponds to either the number of projection periods or the number of observation periods.
4. Trajectory dimension: Corresponds to the number of trajectories in the prediction object, or one if the component is evaluated on observed data.

Depending on the context from which the expression is called, the trajectory dimension of the result of the expression can be reduced by computing given quantiles, and if only one country is evaluated, the first dimension is removed. In addition, with an exception of functions `pop.byage.plot`, `pop.byage.table`, `cohorts`, and `pop.cohorts.plot`, the expression should be constructed in a way that the age dimension is eliminated. This can be done for example by using brackets to define age, by using the `apply` function or one of the pre-defined functions described below. When using within `pop.byage.plot`, `pop.byage.table`, `cohorts`, or `pop.cohorts.plot`, the expression MUST include curly braces.

While `get.pop` can be used to obtain results of a basic component, functions `get.pop.ex` and `get.pop.exba` evaluate whole expressions.

## Pre-defined functions

The following functions can be used within an expression:

- `gmedian(f, cat)`  
It gives a median for grouped data with frequencies `f` and categories `cat`. This function is to be used in combination with `apply` or `pop.apply` (see below) along the age dimension. For example,  
“`apply(P380{ }, c(1,3,4), gmedian, cats=seq(0, by=5, length=28))`”  
is an expression for median age in Italy. (See `pop.apply` below for a simplified version.)
- `gmean(f, cat)`  
Works like `gmedian` but gives the grouped mean.
- `age.func(data, fun="*")`  
This function applies `fun` to data and the corresponding age (the middle point of each age category). The default case would multiply data by the corresponding age. As `gmedian`, it is to be used in combination with `apply` or `pop.apply`.
- `drop.age(data)`  
Drops the age dimension of the data. For example, if two basic components are combined where one is used within the `apply` function, the other will need to change its dimension in order to have conformable arrays. For example,  
“`apply(age.func(P752{ }), c(1,3,4), sum) / drop.age(P752)`”  
is an expression for the average age in Sweden. (See `pop.apply` below for a simplified version.)
- `pop.apply(data, fun, ..., split.along=c("None", "age", "traj", "country"))`  
By default applies function `fun` to the age dimension of data and converts the result into the same format as returned by a basic component. This allows combining the `apply` function with other basic components without having to modify their dimensions. For example,  
“`pop.apply(age.func(P752{ }), fun=sum) / P752`” gives the average age in Sweden, or  
“`pop.apply(P380{ }, gmedian, cats=seq(0, by=5, length=28))`” gives the median age of Italy. If `slice.along` is not ‘None’, it can be used as an `apply` function where the data is sliced along one axis.
- `pop.combine(data1, data2, fun, ..., split.along=c("age", "traj", "country"))`  
Can be used if two basic components should be combined that result in different shapes. It tries to put data into the right format and calls `pop.apply`. For example,  
“`pop.combine(PIND{ }, PIND, '/')`” give population by age per total population in India, or  
“`pop.combine(BFR - DFR, GFR, '+', split.along='traj')`” gives births minus deaths plus net migration in France. Here, `pop.combine` is necessary, because ‘GFR’ is a deterministic component and thus, has only one trajectory, whereas births and deaths are probabilistic.
- `age.index01(end)`  
Can be used with indicators ‘S’, ‘M’, ‘Q’ and ‘E’ only. It returns an array of age group indices that include ages 0-1 and 1-4 and exclude 0-4. The last age index is `end`.
- `age.index05(end)`  
Returns an array of age group indices starting with group 0-4, 5-9 until the age group corresponding to index `end`.

There is also a help function available that generates an expression for the mean age of childbearing, see [mac.expression](#).

**Note**

The expression parser is simple and far from being perfect. We recommend to leave spaces around the basic components.

**Author(s)**

Hana Sevcikova, Adrian Raftery

**References**

H. Sevcikova, A. E. Raftery (2016). bayesPop: Probabilistic Population Projections. *Journal of Statistical Software*, 75(5), 1-29. doi:10.18637/jss.v075.i05

**See Also**

[mac.expression](#), [get.pop](#), [pop.trajectories.plot](#), [pop.map](#), [write.pop.projection.summary](#).

**Examples**

```
sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir, write.to.cache=FALSE)

# median age of women in child-bearing ages in Netherlands and all countries - trajectories
pop.trajectories.plot(pred, nr.traj=0,
  expression="pop.apply(P528_F{4:10}, gmedian, cats= seq(15, by=5, length=8))")
## Not run:
pop.trajectories.plotAll(pred, nr.traj=0,
  expression="pop.apply(PXXX_F{4:10}, gmedian, cats= seq(15, by=5, length=8))")

## End(Not run)
# mean age of women in child-bearing ages in Netherlands - table
pop.trajectories.table(pred,
  expression="pop.apply(age.func(P528_F{4:10}), fun=sum) / P528_F[4:10]")
# - gives the same results as with "pop.apply(P528_F{4:10}, gmean, cats=seq(15, by=5, length=8))"
# - for the mean age of childbearing, see ?mac.expression

# migration per capita by age
pop.byage.plot(pred, expression="GNL{} / PNL{}", year=2000)

## Not run:
# potential support ratio - map (with the two countries
#   contained in pred object)
pop.map(pred, expression="PXXX[5:13] / PXXX[14:27]")
## End(Not run)

# proportion of 0-4 years old to whole population - export to an ASCII file
dir <- tempfile()
write.pop.projection.summary(pred, expression="PXXX[1] / PXXX", output.dir=dir)
unlink(dir)

## Not run:
# These are vital events only available if keep.vital.events=TRUE in pop.predict, e.g.
```

```
# sim.dir.tmp <- tempfile()
# pred <- pop.predict(countries="Netherlands", nr.traj=3,
#                    keep.vital.events=TRUE, output.dir=sim.dir.tmp)
# log female mortality rate by age for Netherlands in 2050, including 0-1 and 1-4 age groups
pop.byage.plot(pred, expression="log(MNL_F{age.index01(27)})", year=2050)

# trajectories of male 1q0 and table of 5q0 for Netherlands
pop.trajectories.plot(pred, expression="QNLD_M[-1]")
pop.trajectories.table(pred, expression="QNLD_M[1]")
# unlink(sim.dir.tmp)
## End(Not run)
```

---

pop.map

*World Map of Population Measures*


---

### Description

Generates a world map of various population measures for a given quantile and a projection or observed period.

### Usage

```
pop.map(pred, sex = c("both", "male", "female"), age = "all", expression = NULL, ...)

get.pop.map.parameters(pred, expression = NULL, sex = c("both", "male", "female"),
  age = "all", range = NULL, nr.cats = 50, same.scale = TRUE, quantile = 0.5, ...)

pop.map.gvis(pred, ...)
```

### Arguments

pred	Object of class <a href="#">bayesPop.prediction</a> .
sex	One of “both” (default), “male” or “female”. By default the male and female counts are summed up. This argument is only used if expression is NULL.
age	Either a character string “all” (default) or an integer vector of age indices. Value 1 corresponds to age 0-4, value 2 corresponds to age 5-9 etc. Last age group 130+ corresponds to index 27. This argument is only used if expression is NULL.
expression	Expression defining the population measure to be plotted. For syntax see <a href="#">pop.expressions</a> . The country components of the expression should be given as “XXX”.
range	Range of the population measure to be displayed. It is of the form <code>c(min, max)</code> .
nr.cats	Number of color categories.
same.scale	Logical controlling if maps for all years of this prediction object should be on the same color scale.

quantile	Quantile for which the map should be generated. It must be equal to one of the values in <code>dimnames(pred\$quantiles[[2]])</code> , i.e. 0, 0.025, 0.05, 0.1, 0.2, 0.25, 0.3, 0.4, 0.5, 0.6, 0.7, 0.75, 0.8, 0.9, 0.95, 0.975, 1. Value 0.5 corresponds to the median.
...	Additional arguments passed to the underlying functions. In <code>pop.map</code> , these are <code>quantile</code> , <code>year</code> , <code>projection.index</code> , <code>device</code> , <code>main</code> , and <code>device.args</code> (see <code>tfr.map</code> ). In <code>pop.map.gvis</code> , these are all arguments that can be passed to <code>tfr.map.gvis</code> . In addition, the first two functions accept arguments passed to the <code>mapCountryData</code> function of the <b>rworldmap</b> package.

### Details

`pop.map` creates a single map for the given time period and quantile. If the package **fields** is installed, a color bar legend at the bottom of the map is created.

Function `get.pop.map.parameters` can be used in combination with `pop.map`. It sets breakpoints for the color scheme.

Function `pop.map.gvis` creates an interactive map using the **googleVis** package and opens it in an internet browser. It also generates a table of the mapped values that can be sorted by columns interactively in the browser.

### Value

`get.pop.map.parameters` returns a list with elements:

<code>pred</code>	The object of class <code>bayesPop.prediction</code> used in the function.
<code>quantile</code>	Value of the argument <code>quantile</code> .
<code>catMethod</code>	If the argument <code>same.scale</code> is <code>TRUE</code> , this element contains breakpoints for categorization. Otherwise, it is <code>NULL</code> .
<code>numCats</code>	Number of categories.
<code>coulourPalette</code>	Subset of the rainbow palette, starting from dark blue and ending at red.
...	Additional arguments passed to the function.

### Performance and Caching

If the `expression` argument or a non-standard combination of sex and age is used, quantiles are computed on the fly. In such a case, trajectory files for all countries have to be loaded from disk, which can be quite time expensive. Therefore a simple caching mechanism was added to the prediction object which allows re-using data from previously used expressions. The prediction object points to an environment called `cache` which is a collection of data arrays that are results of evaluating expressions. The space-trimmed expressions are the names of the cache entries. Every time a map function is called, it is checked if the corresponding expression is contained in the cache. If it is not the case, the quantiles are computed on the fly, otherwise the existing values are taken.

When computing on the fly, the function tries to process it in parallel if possible, using the package **parallel**. In such a case, the computation is split into  $n$  nodes where  $n$  is either the number of cores detected automatically (default), or the value of `getOption("cl.cores")`. Use `options(cl.cores=n)` to modify the default. If a sequential processing is desired, set `cl.cores` to 1.

The cache data are also stored on disk, namely in the simulation directory of the prediction object. By default, every update of the cache in memory is also updated on the disk. Thus, data expression results can be re-used in multiple R sessions. Function `pop.cleanup.cache` deletes the content of the cache. This behaviour can be turned off by setting the argument `write.to.cache=FALSE` in the `get.pop.prediction` function. We use this settings in the examples throughout this manual whenever the example data from the installation directory is used, in order to prevent writing into the installation directory.

### Author(s)

Hana Sevcikova

### See Also

[tfr.map](#)

### Examples

```
## Not run:
#####
# This example only makes sense if there is a simulation
# for all countries. Below, only two countries are included,
# so the map is useless.
#####
sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir=sim.dir, write.to.cache=FALSE)
# Uses heat colors with seven categories by default
pop.map(pred, sex="female", age=4:10)
# Female population in child-bearing age as a proportion of totals
pop.map(pred, expression="PXXX_F[4:10] / PXXX")
# The same with more colors
params <- get.pop.map.parameters(pred, expression="PXXX_F[4:10] / PXXX")
do.call("pop.map", params)
# Another projection year on the same color scale
do.call("pop.map", c(list(year=2043), params))
# Potential support ratio using googleVis
pop.map.gvis(pred, expression="PXXX[5:13] / PXXX[14:27]")
## End(Not run)
```

---

pop.predict

*Probabilistic Population Projection*

---

### Description

The function generates trajectories of probabilistic population projection for all countries for which input data is available, or any subset of them.

## Usage

```
pop.predict(end.year = 2100, start.year = 1950, present.year = 2020,
  wpp.year = 2019, countries = NULL,
  output.dir = file.path(getwd(), "bayesPop.output"),
  inputs = list(popM=NULL, popF=NULL, mxM=NULL, mxF=NULL, srb=NULL,
    pasfr=NULL, patterns=NULL, migM=NULL, migF=NULL,
    e0F.file=NULL, e0M.file=NULL, tfr.file=NULL,
    e0F.sim.dir=NULL, e0M.sim.dir=NULL, tfr.sim.dir=NULL,
    migMtraj = NULL, migFtraj = NULL),
  nr.traj = 1000, keep.vital.events = FALSE,
  fixed.mx = FALSE, fixed.pasfr = FALSE,
  lc.for.hiv = TRUE, lc.for.all = TRUE,
  my.locations.file = NULL, replace.output = FALSE,
  verbose = TRUE, ...)
```

## Arguments

end.year	End year of the projection.
start.year	First year of the historical data.
present.year	Year for which initial population data is to be used.
wpp.year	Year for which WPP data is used. The functions loads a package called <b>wpp<math>x</math></b> where $x$ is the wpp.year and uses the various datasets as default if the corresponding inputs element is missing (see below).
countries	Array of country codes or country names for which a projection is generated. If it is NULL, all available countries are used. If it is NA and there is an existing projection in output.dir and replace.output=FALSE, then a projection is performed for all countries that are not included in the existing projection. Names of countries are matched to those in the <a href="#">UNlocations</a> dataset (or in the dataset loaded from my.locations.file if used).
output.dir	Output directory of the projection. If there is an existing projection in output.dir and replace.output=TRUE, everything in the directory will be deleted.
inputs	A list of file names where input data is stored. It contains the following elements (Unless otherwise noted, these are tab delimited ASCII files; Names of default datasets from the corresponding <b>wpp</b> package which are used if the corresponding element is NULL are shown in brackets): <ul style="list-style-type: none"> <li><b>popM, popF</b> Initial male/female age-specific population (at time present.year) [<a href="#">popM</a>, <a href="#">popF</a>].</li> <li><b>mxM, mxF</b> Historical data and (optionally) projections of male/female age-specific death rates [<a href="#">mxM</a>, <a href="#">mxF</a>] (see also argument fixed.mx).</li> <li><b>srb</b> Projection of sex ratio at birth. [<a href="#">sexRatio</a>]</li> <li><b>pasfr</b> Historical data and (optionally) projections of percentage age-specific fertility rate [<a href="#">percentASFR</a>] (see also argument fixed.pasfr).</li> <li><b>patterns, mig.type</b> Migration type and base year of the migration. In addition, this dataset gives information on country's specifics regarding mortality and fertility age patterns as defined in [<a href="#">vwBaseYear</a>]. patterns and mig.type have the same meaning and can be used interchangeably.</li> </ul>

- migM, migF** Projection of male/female age-specific migration as net counts on the same scale as initial population [[migrationM](#), [migrationF](#)]. If not available, the migration schedules are reconstructed from total migration counts derived from [migration](#) using the [age.specific.migration](#) function.
- e0F.file** Comma-delimited CSV file with results of female life expectancy (generated using [bayesLife](#), function [convert.e0.trajectories](#), file “ascii\_trajectories.csv”). Required columns are “LocID”, “Year”, “Trajectory”, and “e0”. If this element is not NULL, the argument [e0F.sim.dir](#) is ignored. If both [e0F.file](#) and [e0F.sim.dir](#) are NULL, data from the corresponding **wpp** package is taken, namely the median projections as one trajectory and the low and high variants (if available) as second and third trajectory.
- e0M.file** Comma-delimited CSV file containing results of male life expectancy (generated using [bayesLife](#), function [convert.e0.trajectories](#), file “ascii\_trajectories.csv”). Required columns are “LocID”, “Year”, “Trajectory”, and “e0”. If this element is not NULL, the argument [e0M.sim.dir](#) is ignored. As in the female case, if both [e0M.file](#) and [e0M.sim.dir](#) are NULL, data from the corresponding **wpp** package is taken.
- tfr.file** Comma-delimited CSV file with results of total fertility rate (generated using [bayesTFR](#), function [convert.tfr.trajectories](#), file “ascii\_trajectories.csv”). Required columns are “LocID”, “Year”, “Trajectory”, and “TF”. If this element is not NULL, the argument [tfr.sim.dir](#) is ignored. If both [tfr.file](#) and [tfr.sim.dir](#) are NULL, data from the corresponding **wpp** package is taken (median and the low and high variants as three trajectories). Alternatively, this argument can be the keyword “median\_” in which case only the wpp median is taken.
- e0F.sim.dir** Simulation directory with results of female life expectancy (generated using [bayesLife](#)). It is only used if [e0F.file](#) is NULL.
- e0M.sim.dir** Simulation directory with results of male life expectancy (generated using [bayesLife](#)). Alternatively, it can be the string “joint\_”, in which case it is assumed that the male life expectancy was projected jointly from the female life expectancy (see [joint.male.predict](#)) and thus contained in the [e0F.sim.dir](#) directory. The argument is only used if [e0M.file](#) is NULL.
- tfr.sim.dir** Simulation directory with results of total fertility rate (generated using [bayesTFR](#)). It is only used if [tfr.file](#) is NULL.
- migMtraj, migFtraj** Comma-delimited CSV file with male/female age-specific migration trajectories. If present, it replaces deterministic projections given by the [migM](#) and [migF](#) items. It has a similar format as e.g. [e0M.file](#) with columns “LocID”, “Year”, “Trajectory”, “Age” and “Migration”. The “Age” column must have values “0-4”, “5-9”, “10-14”, ..., “95-99”, “100+”.
- nr.traj** Number of trajectories to be generated. If this number is smaller than the number of available trajectories of the probabilistic components (TFR, life expectancy and migration), the trajectories are equidistantly thinned. If all of those components contain less trajectories than [nr.traj](#), the value is adjusted to the maximum of available trajectories of the components. For those that have less trajectories than the adjusted number, the available trajectories are re-sampled, so that all components have the same number of trajectories.

keep.vital.events	Logical. If TRUE age- and sex-specific vital events of births and deaths as well as other objects are stored in the prediction object, see Details.
fixed.mx	Logical. If TRUE, it is assumed the dataset of death rates (mxM and mxF) include data for projection years and they are then used instead of the life expectancy.
fixed.pasfr	Logical. If TRUE, it is assumed the dataset on percent age-specific fertility rate (percentASFR) include data for projection years and they are then used instead of computing it on the fly.
lc.for.hiv	Logical controlling if the modified Lee-Carter method should be used for projection of mortality rates for countries with HIV epidemics. If FALSE, the function <code>hiv.mortmod</code> from the <code>HIV.LifeTables</code> package is used.
lc.for.all	Logical controlling if the modified Lee-Carter method should be used for projection of mortality rates for all countries. If FALSE, the corresponding method is determined by the columns “AgeMortProjMethod1” and “AgeMortProjMethod2” of the <code>vwBaseYear</code> dataset.
my.locations.file	Name of a tab-delimited ascii file with a set of all locations for which a projection is generated. Use this argument if you are projecting for a country/region that is not included in the standard <code>UNlocations</code> dataset. It must have the same structure.
replace.output	Logical. If TRUE, everything in the directory <code>output.dir</code> is deleted prior to the prediction.
verbose	Logical controlling the amount of output messages.
...	Additional arguments passed to the underlying function. These can be <code>parallel</code> and <code>nr.nodes</code> for parallel processing and the number of nodes, respectively, as well as further arguments passed for creating a parallel cluster.

## Details

The population projection is computed using the Cohort Component method and is based on an algorithm used by the United Nation Population Division (see also Sevcikova et al (2015) in the References below). For each country, one projection is calculated for each trajectory of male and female life expectancy, TFR and possibly migration. This results in a set of trajectories of population projection which forms its posterior distribution. The trajectories of life expectancy and TFR can be given either in its binary form generated by the packages `bayesLife` and `bayesTFR`, respectively (as directories `e0M.sim.dir`, `e0F.sim.dir`, `tfr.sim.dir` of the `inputs` argument), or they can be given as ASCII tables in csv format, see above. The number of trajectories for male and female life expectancy must match, as does for male and female migration.

The projection is generated sequentially country by country. Results are stored in a sub-directory of `output.dir` called ‘prediction’. There is one binary file per country, called ‘`totpop_countryx.rda`’, where  $x$  is the country code. It contains six objects: `totp`, `totpf`, `totpm` (trajectories of total population, age-specific female and age-specific male, respectively), `totp.hch`, `totpf.hch`, `totpm.hch` (the UN half-child variant for total population, age-specific female and age-specific male, respectively). Optionally, if `keep.vital.events` is TRUE, there is an additional file per country, called ‘`vital_events_countryx.rda`’, containing the following objects: `btm`, `btf` (trajectories for births by age of mothers for male and female child, respectively), `deathsm`, `deathsf` (trajectories for

age-specific male and female deaths, respectively), `asfert` (trajectories of age-specific fertility), `mxm`, `mxm` (trajectories of male and female age-specific mortality rates), `migm`, `migf` (if used, these are trajectories of male and female age-specific migration), `btm.hch`, `btf.hch`, `deathsm.hch`, `deathsf.hch`, `asfert.hch`, `mxm.hch`, `mxm.hch` (the UN half-child variant for age- and sex-specific births, deaths, fertility rates and mortality rates). An object of class `bayesPop.prediction` is stored in the same directory in a file `'prediction.rda'`. It is updated every time a country projection is finished.

See [pop.trajectories](#) for extracting trajectories.

To access a previously stored prediction object, use [get.pop.prediction](#).

## Value

Object of class `bayesPop.prediction` with the following elements:

<code>base.directory</code>	Full path to the base directory output.dir.
<code>output.directory</code>	Sub-directory relative to <code>base.directory</code> with the projections.
<code>nr.traj</code>	The actual number of trajectories of the projections.
<code>quantiles</code>	Three-dimensional array of projection quantiles (countries x number of quantiles x projection periods). The second dimension corresponds to the following quantiles: 0.025, 0.05, 0.1, 0.25, 0.5, 0.75, 0.9, 0.95, 0.975.
<code>traj.mean.sd</code>	Three-dimensional array of projection mean and standard deviation (countries x 2 x projection periods). First and second matrix of the second dimension, respectively, is the mean and standard deviation, respectively.
<code>quantilesM</code> , <code>quantilesF</code>	Quantiles of male and female projection, respectively. Same structure as <code>quantiles</code> .
<code>traj.mean.sdM</code> , <code>traj.mean.sdF</code>	Same as <code>traj.mean.sd</code> corresponding to male and female projection, respectively.
<code>quantilesMage</code> , <code>quantilesFage</code>	Four-dimensional array of age-specific quantiles of male and female projection, respectively (countries x age groups x number of quantiles x projection periods). The same quantiles are used as in <code>quantiles</code> .
<code>quantilesPropMage</code> , <code>quantilesPropFage</code>	Array of age-specific quantiles of male and female projection, respectively, divided by the total population. The dimensions are the same as in <code>quantilesMage</code> .
<code>estim.years</code>	Vector of time for which historical data was used in the projections.
<code>proj.years</code>	Vector of projection time periods starting with the present period.
<code>wpp.year</code>	The wpp year used.
<code>inputs</code>	List of input data used for the projection.
<code>function.inputs</code>	Content of the <code>inputs</code> argument passed to the function.
<code>countries</code>	Matrix of countries for which projection exists. It contains two columns: code, name.
<code>ages</code>	Vector of age groups.

`cache` This component is added by `get.pop.prediction` and modified and used by `pop.map` and `write.pop.projection.summary`. It is an environment for caching and re-using results of expressions.

`write.to.cache` Logical determining if cache should be modified.

`is.aggregation` Logical determining if this object is a result of `pop.predict` or `pop.aggregate`.

### Author(s)

Hana Sevcikova, Thomas Buettner, based on code of Nan Li and helpful comments from Patrick Gerland

### References

H. Sevcikova, A. E. Raftery (2016). `bayesPop`: Probabilistic Population Projections. *Journal of Statistical Software*, 75(5), 1-29. doi:10.18637/jss.v075.i05

A. E. Raftery, N. Li, H. Sevcikova, P. Gerland, G. K. Heilig (2012). Bayesian probabilistic population projections for all countries. *Proceedings of the National Academy of Sciences* 109:13915-13921.

P. Gerland, A. E. Raftery, H. Sevcikova, N. Li, D. Gu, T. Spoorenberg, L. Alkema, B. K. Fosdick, J. L. Chunn, N. Lalic, G. Bay, T. Buettner, G. K. Heilig, J. Wilmoth (2014). World Population Stabilization Unlikely This Century. *Science* 346:234-237.

H. Sevcikova, N. Li, V. Kantorova, P. Gerland and A. E. Raftery (2015). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. arXiv:1503.05215. <http://arxiv.org/abs/1503.05215>

### See Also

[pop.trajectories.plot](#), [pop.pyramid](#), [pop.trajectories](#), [get.pop.prediction](#), [age.specific.migration](#)

### Examples

```
## Not run:
sim.dir <- tempfile()
# Countries can be given as a combination of numerical codes and names
pred <- pop.predict(countries=c("Netherlands", 218, "Madagascar"), nr.traj=3,
  output.dir=sim.dir)
pop.trajectories.plot(pred, "Ecuador", sum.over.ages=TRUE)
unlink(sim.dir, recursive=TRUE)

## End(Not run)
```

---

pop.predict.subnat      *Subnational Probabilistic Population Projection*

---

## Description

Generates trajectories of probabilistic population projection for subregions of a given country.

## Usage

```
pop.predict.subnat(end.year = 2060, start.year = 1950, present.year = 2020,
  wpp.year = 2019, output.dir = file.path(getwd(), "bayesPop.output"),
  locations = NULL, default.country = NULL,
  inputs = list(
    popM = NULL, popF = NULL,
    mxM = NULL, mxF = NULL, srb = NULL,
    pasfr = NULL, patterns = NULL,
    migM = NULL, migF = NULL,
    e0F.file = NULL, e0M.file = NULL, tfr.file = NULL,
    e0F.sim.dir = NULL, e0M.sim.dir = NULL, tfr.sim.dir = NULL,
    migMtraj = NULL, migFtraj = NULL
  ),
  nr.traj = 1000, keep.vital.events = FALSE,
  fixed.mx = FALSE, fixed.pasfr = FALSE,
  replace.output = FALSE, verbose = TRUE)
```

## Arguments

end.year	End year of the projection.
start.year	First year of the historical data on mortality rates. It determines the length of the historical time series used in the Lee-Carter estimation.
present.year	Year for which initial population data is to be used.
wpp.year	Year for which WPP data is used. The function loads a package called <b>wppx</b> where <i>x</i> is the wpp.year and uses its data (corresponding to the default.country) as default datasets if region-specific alternatives are not given (see more details below).
output.dir	Output directory of the projection.
locations	Name of a tab-delimited file that contains definitions of the subregions. It has a similar structure as <a href="#">UNlocations</a> , with mandatory columns reg_code (unique identifier of the subregions) and name (name of the subregions). Optionally, location_type should be set to 4 for subregions to be processed. Column country_code can be included with the numerical code of the corresponding country. A row with location_type of 0 determines the country that the subregions belong to and is used for extracting default "national" datasets if the argument default.country is missing. In such a case, the code of the default country is taken from its column country_code. This is a mandatory argument.

default.country	Numerical code of a country to which the subregions belong to. It is used for extracting default datasets from the <b>wpp</b> package if some region-specific input datasets are missing. Alternatively, it can be also included in the <code>locations</code> file, see above. In either case, the code must exist in the <code>UNlocations</code> dataset.
inputs	<p>A list of file names where input data is stored. Unless otherwise noted, these are tab delimited ASCII files with a mandatory column <code>reg_code</code> giving the numerical identifier of the subregions. If an element of this list is NULL, usually a default dataset corresponding to <code>default.country</code> is extracted from the <b>wpp</b> package. Names of these default datasets are shown in brackets. This list contains the following elements:</p> <p><b>popM, popF</b> Initial male/female age-specific population (at time <code>present.year</code>). Mandatory items, no defaults. Must contain columns <code>reg_code</code> and <code>age</code> and be of the same structure as <code>popM</code> from <b>wpp</b>.</p> <p><b>mxM, mxF</b> Historical data and (optionally) projections of male/female age-specific death rates [<code>mxM, mxF</code>] (see also argument <code>fixed.mx</code>).</p> <p><b>srb</b> Projection of sex ratio at birth. [<code>sexRatio</code>]</p> <p><b>pasfr</b> Historical data and (optionally) projections of percentage age-specific fertility rate [<code>percentASFR</code>] (see also argument <code>fixed.pasfr</code>).</p> <p><b>patterns</b> Information on region's specifics regarding migration type, base year of the migration, mortality and fertility age patterns as defined in [<code>vwBaseYear</code>]. In addition, it can contain columns defining migration shares between the subregions, see Details below.</p> <p><b>migM, migF</b> Projection of male/female age-specific migration as net counts on the same scale as initial population. It should have the same format as <code>migrationM</code>. If not available, the migration schedules are constructed from total migration counts of the <code>default.country</code> derived from <code>migration</code> using Rogers Castro for age distribution. Migration shares between subregions (including sex-specific shares) can be given in the <code>patterns</code> file, see above and Details below.</p> <p><b>e0F.file</b> Comma-delimited CSV file with projected female life expectancy. It has the same structure as the file "<code>ascii_trajectories.csv</code>" generated using <code>bayesLife::convert.e0.trajectories</code> (which currently works for country-level results only). Required columns are "<code>LocID</code>", "<code>Year</code>", "<code>Trajectory</code>", and "<code>e0</code>". If <code>e0F.file</code> is NULL, data from the corresponding <b>wpp</b> package (for <code>default.country</code>) is taken, namely the median projections as one trajectory and the low and high variants (if available) as second and third trajectory. Alternatively, this element can be the keyword "<code>median_</code>" in which case only the median is taken.</p> <p><b>e0M.file</b> Comma-delimited CSV file containing projections of male life expectancy of the same format as <code>e0F.file</code>. As in the female case, if <code>e0M.file</code> is NULL, data for <code>default.country</code> from the corresponding <b>wpp</b> package is taken.</p> <p><b>tfr.file</b> Comma-delimited CSV file with results of total fertility rate (generated using <code>bayesTFR</code>, function <code>convert.tfr.trajectories</code>, file "<code>ascii_trajectories.csv</code>"). Required columns are "<code>LocID</code>", "<code>Year</code>", "<code>Trajectory</code>", and "<code>TF</code>". If this element is not NULL, the argument <code>tfr.sim.dir</code> is ignored. If both <code>tfr.file</code></p>

and `tfr.sim.dir` are NULL, data for `default.country` from the corresponding **wpp** package is taken (median and the low and high variants as three trajectories). Alternatively, this argument can be the keyword “`median_`” in which case only the wpp median is taken.

**e0F.sim.dir** Simulation directory with results of female life expectancy. Since **bayesLife** does not support subnational projections yet, this element should not be used. Instead use `e0F.file` if region-specific e0 projections are available. Alternatively, it can be set to the keyword “`median_`” which has the same effect as when `e0F.file` is “`median_`”.

**e0M.sim.dir** This is analogous to `e0F.sim.dir`, here for male life expectancy. Use `e0M.file` instead of this item.

**tfr.sim.dir** Simulation directory with projections of total fertility rate (generated using `bayesTFR: :tfr.predict.subnat`). It is only used if `tfr.file` is NULL.

**migMtraj, migFtraj** Comma-delimited CSV file with male/female age-specific migration trajectories. If present, it replaces deterministic projections given by the `migM` and `migF` items. It has a similar format as e.g. `e0M.file` with columns “`LocID`”, “`Year`”, “`Trajectory`”, “`Age`” and “`Migration`”. The “`Age`” column must have values “`0-4`”, “`5-9`”, “`10-14`”, ..., “`95-99`”, “`100+`”.

`nr.traj`, `keep.vital.events`, `fixed.mx`, `fixed.pasfr`, `replace.output`, `verbose`

These arguments have the same meaning as in [pop.predict](#).

## Details

Population projection for subnational units (regions) is performed by applying the cohort component method to subnational datasets on projected fertility (TFR), mortality and net migration, starting from given sex- and age-specific population counts. The only required inputs are the initial sex- and age-specific population counts in each region (`popM` and `popF` elements of the `inputs` argument) and a file with a set of locations (argument `locations`). If no other input datasets are given, those datasets are replaced by the corresponding "national" values, taken from the corresponding **wpp** package. The argument `default.country` determines the country for those default "national" values. The default country can be also included in the `locations` file as a record with `location.type` being set to 0.

The TFR component can be given as a set of trajectories generated using the `tfr.predict.subnat` function of the **bayesTFR** package (`tfr.sim.dir` element). Alternatively, trajectories can be given in an ASCII file (`tfr.file`). Having a set of subnational TFR trajectories, the cohort component method is applied to each of them to yield a distribution of future subnational population.

Net migration can either be given as disaggregated sex- and age-specific datasets `migM` and `migF`. Alternatively, it can be given as shares between regions as columns in the `patterns` dataset. These are: `inmigrationM_share`, `inmigrationF_share`, `outmigrationM_share`, `outmigrationF_share`. The sex specification and/or direction specification (in/out) can be omitted, e.g. it can be simply `migration_share`. The function extracts the values of net migration projection on the national level and distributes it to regions according to the given shares. For positive (national) values, it uses the in-migration shares; for negative values it uses the out-migration shares. If the in/out prefix is omitted in the column names, the given migration shares are used for both, positive and negative net migration projection. By default, if no `migM` and `migF` neither region-specific shares are given, the distribution between regions is proportional to the size of population. The age-specific schedules follow by default the

Rogers-Castro age schedules. Note that when handling migration using shares as described here, it only affects the distribution of international migration into regions. It does not take into account between-region migration.

The package contains example datasets for Canada. Use these as templates for your own data. See Example below.

### Value

Object of class `bayesPop.prediction` containing the subnational projections. Note that this object can be use in the various **bayesPop** functions exactly the same way as an object with national projections. However, the meaning of the argument `country` in many of these functions (e.g. in `pop.trajectories.plot`) changes to an identification of the region (either as a numerical code or name as defined in the `locations` file).

### Acknowledgment

We are grateful to Patrice Dion from Statistics Canada for providing us with example data. Note that the example datasets included in the package are not official STATCAN data - they only serve the purpose of illustration and templates. Data for the time period 2015-2020 has been imputed by the author.

### Author(s)

Hana Sevcikova

### See Also

[pop.predict](#), [tfr.predict.subnat](#), [pop.aggregate.subnat](#)

### Examples

```
## Not run:
# Subnational projections for Canada
#####
data.dir <- file.path(find.package("bayesPop"), "extdata")

# Use national data for tfr and e0
###
sim.dir <- tempfile()
pred <- pop.predict.subnat(output.dir = sim.dir,
  locations = file.path(data.dir, "CANlocations.txt"),
  inputs = list(popM = file.path(data.dir, "CANpopM.txt"),
    popF = file.path(data.dir, "CANpopF.txt"),
    tfr.file = "median_"
  ),
  verbose = TRUE)
pop.trajectories.plot(pred, "Alberta", sum.over.ages = TRUE)
unlink(sim.dir, recursive=TRUE)

# Use subnational TFR simulation
###
```

```

# Subnational TFR projections for Canada (from ?tfr.predict.subnat)
my.subtfr.file <- file.path(find.package("bayesTFR"), 'extdata', 'subnational_tfr_template.txt')
tfr.nat.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
tfr.reg.dir <- tempfile()
tfr.preds <- tfr.predict.subnat(124, my.tfr.file = my.subtfr.file,
  sim.dir = tfr.nat.dir, output.dir = tfr.reg.dir, start.year = 2013)

# Pop projections
sim.dir <- tempfile()
pred <- pop.predict.subnat(output.dir = sim.dir,
  locations = file.path(data.dir, "CANlocations.txt"),
  inputs = list(popM = file.path(data.dir, "CANpopM.txt"),
    popF = file.path(data.dir, "CANpopF.txt"),
    patterns = file.path(data.dir, "CANpatterns.txt"),
    tfr.sim.dir = file.path(tfr.reg.dir, "subnat", "c124")
  ),
  verbose = TRUE)
pop.trajectories.plot(pred, "Alberta", sum.over.ages = TRUE)
pop.pyramid(pred, "Manitoba", year = 2050)
get.countries.table(pred)

# Aggregate to country level
aggr <- pop.aggregate.subnat(pred, regions = 124,
  locations = file.path(data.dir, "CANlocations.txt"))
pop.trajectories.plot(aggr, "Canada", sum.over.ages = TRUE)

unlink(sim.dir, recursive = TRUE)
unlink(tfr.reg.dir, recursive = TRUE)

## End(Not run)

```

---

pop.pyramid

*Probabilistic Population Pyramid*


---

## Description

Functions for plotting probabilistic population pyramid. `pop.pyramid` creates a classic pyramid using rectangles; `pop.trajectories.pyramid` creates one or more pyramids using vertical lines (possibly derived from population trajectories). They can be used to view a prediction object created with this package, or any user-defined sex- and age-specific dataset. For the latter, function `get.bPop.pyramid` should be used to translate user-defined data into a `bayesPop.pyramid` object.

## Usage

```

## S3 method for class 'bayesPop.prediction'
pop.pyramid(pop.object, country, year = NULL,
  indicator = c("P", "B", "D"), pi = c(80, 95),
  proportion = FALSE, age = 1:21, plot = TRUE, pop.max = NULL, ...)

```

```

## S3 method for class 'bayesPop.pyramid'
pop.pyramid(pop.object, main = NULL, show.legend = TRUE,
  pyr1.par = list(border="black", col=NA, density=NULL, height=0.9),
  pyr2.par = list(density = -1, height = 0.3),
  col.pi = NULL, ann = par("ann"), axes = TRUE, grid = TRUE,
  cex.main = 0.9, cex.sub = 1, cex = 1, cex.axis = 1, ...)

pop.pyramidAll(pop.pred, year = NULL,
  output.dir = file.path(getwd(), "pop.pyramid"),
  output.type = "png", one.file = FALSE, verbose = FALSE, ...)

## S3 method for class 'bayesPop.prediction'
pop.trajectories.pyramid(pop.object, country, year = NULL,
  indicator = c("P", "B", "D"), pi = c(80, 95), nr.traj = NULL,
  proportion = FALSE, age = 1:21, plot = TRUE, pop.max = NULL, ...)

## S3 method for class 'bayesPop.pyramid'
pop.trajectories.pyramid(pop.object, main = NULL, show.legend = TRUE,
  col = rainbow, col.traj = "#00000020", lwd = 2, ann = par("ann"), axes = TRUE,
  grid = TRUE, cex.main = 0.9, cex.sub = 1, cex = 1, cex.axis = 1, ...)

pop.trajectories.pyramidAll(pop.pred, year = NULL,
  output.dir = file.path(getwd(), "pop.traj.pyramid"),
  output.type = "png", one.file = FALSE, verbose = FALSE, ...)

## S3 method for class 'bayesPop.pyramid'
plot(x, ...)

## S3 method for class 'bayesPop.prediction'
get.bPop.pyramid(data, country, year = NULL,
  indicator = c("P", "B", "D"), pi = c(80, 95),
  proportion = FALSE, age = 1:21, nr.traj = 0, sort.pi=TRUE, pop.max = NULL, ...)

## S3 method for class 'data.frame'
get.bPop.pyramid(data, main.label = NULL, legend = "observed",
  is.proportion = FALSE, ages = NULL, pop.max = NULL,
  LRmain = c("Male", "Female"), LRcolnames = c("male", "female"), CI = NULL, ...)

## S3 method for class 'matrix'
get.bPop.pyramid(data, ...)

## S3 method for class 'list'
get.bPop.pyramid(data, main.label = NULL, legend = NULL, CI = NULL, ...)

```

## Arguments

`pop.object` Object of class [bayesPop.prediction](#) or `bayesPop.pyramid` (see Value section).

pop.pred	Object of class <a href="#">bayesPop.prediction</a> .
x	Object of class <a href="#">bayesPop.pyramid</a> .
data	Data frame, matrix, list or object of class <a href="#">bayesPop.prediction</a> . For data frame and matrix, it must have columns defined by <code>LRcolnames</code> (“male” and “female” by default). The row names will determine the age labels. For lists, it can be a collection of such data frames. The names of the list elements are used for legend, unless <code>legend</code> is given.
country	Name or numerical code of a country.
year	Year within the projection or estimation period to be plotted. Default is the start year of the prediction. It can also be a vector of years. <code>pop.pyramid</code> draws the first two, <code>pop.trajectories.pyramid</code> draws all of them. In the functions <code>pop.pyramidAll</code> and <code>pop.trajectories.pyramidAll</code> , the <code>year</code> argument can be a list of years, in which case the pyramids are created for all elements in the list.
indicator	One of the characters “P” (population), “B” (births), “D” (deaths) determining the pyramid indicator.
pi	Probability interval. It can be a single number or an array.
proportion	Logical. If TRUE the pyramid contains the distribution of rates of age-specific counts and population totals.
age	Integer vector of age indices. Value 1 corresponds to age 0-4, value 2 corresponds to age 5-9 etc. Last available age group is 130+ which corresponds to index 27. The purpose of this argument here is mainly to control the height of the pyramid.
plot	If FALSE, nothing is plotted. It can be used to retrieve the pyramid object without drawing it.
main	Titel of the plot. By default it is the country name and projection year if known.
show.legend	Logical controlling if the plot legend is drawn.
pyr1.par, pyr2.par	List of graphical parameters (color, border, density and height) for drawing the pyramid rectangles, for the first and second pyramid, respectively (see Details). The height component should be a number between 0 (corresponds to a line) and 1 (for non-overlapping rectangles). If density is NULL, the rectangles are transparent, see the argument <code>density</code> in <a href="#">rect</a> .
col.pi	Vector of colors for drawing the probability boxes. If it is given, it must be of the same length as <code>pi</code> .
ann	Logical controlling if any annotation (main and legend) is plotted.
axes	Logical controlling if axes are plotted.
grid	Logical controlling if grid lines are plotted.
cex.main, cex.sub, cex, cex.axis	Magnification to be used for the title, secondary titles on the right and left panels, legend and axes, respectively.
output.dir	Directory into which resulting graphs are stored.

output.type	Type of the resulting files. It can be “png”, “pdf”, “jpeg”, “bmp”, “tiff”, or “postscript”.
one.file	Logical. If TRUE the output is put into one single file, by default a PDF.
verbose	Logical switching log messages on and off.
nr.traj	Number of trajectories to be plotted. If NULL, all trajectories are plotted, otherwise they are thinned evenly.
col	Colors generating function. It is called with an argument giving the number of pyramids to be plotted. Each color is then used for one pyramid, including its confidence intervals.
col.traj	Color used for trajectories. If more than one pyramid is drawn with its trajectories, this can be a vector of the size of number of pyramids.
lwd	Line width for the pyramids.
sort.pi	Logical controlling if the probability intervals are sorted in decreasing order. This has an effect on the order in which they are plotted and thus on overlapping of pyramid boxes. By default the largest intervals are plotted first.
main.label	Optional argument for the main title.
legend	Legend to be used. In case of multiple pyramids, this can be a vector for each of them. If not given and data is a list, names of the list elements are taken as legend.
is.proportion	Either logical, indicating if the values in data are proportions, or NA in which case the proportions are computed.
ages	Vector of age labels. It must be of the same length as the number of rows of data. If it is not given, the age labels are considered to be the row names of data.
pop.max	Maximum value to be drawn in the pyramid. If it is not given, max(data) is taken.
LRmain	Vector of character strings giving the secondary titles for the left and right panel, respectively.
LRcolnames	Vector of character strings giving the column names of data to be used for the left and right panel of the pyramid, respectively.
CI	Confidence intervals. It should be of the same format as the bayesPop.pyramid\$CI object, see below.
...	Arguments passed to the underlying functions.

## Details

The `pop.pyramid` function generates one or two population pyramids in one plot. The first (main) one is usually the median of a future year prediction, but it can also be the current year or any population estimates. The second one serves the purpose of comparing two pyramids with one another and is drawn on top of the main pyramid. For example, one can use it to compare a future prediction with the present, or two different time points in the past, or two different geographies. The main pyramid can have confidence intervals associated with it, which are also plotted. If `pop.pyramid` is called on a `bayesPop.prediction` object, the main and secondary pyramid, respectively, is generated from data of a time period given by the first and second element, respectively, of the `year` argument. In

such a case, confidence intervals only of the first year are shown. Thus, it makes sense to set the first year to be a prediction year and the second year to an observed time period. If `pop.pyramid` is called on a `bayesPop.pyramid` object, data in the first and second element, respectively, of the `bayesPop.pyramid$pyramid` list are used, and only the first element of `bayesPop.pyramid$CI` is used.

Pyramids generated via the `pop.trajectories.pyramid` function have different appearance and therefore more than two pyramids can be put into one figure. Furthermore, confidence intervals of more than one pyramid can be shown. Thus, all elements of `bayesPop.pyramid$pyramid` and `bayesPop.pyramid$CI` are plotted. In addition, single trajectories given in `bayesPop.pyramid$trajectories` can be shown by setting the argument `nr.traj` larger than 0.

Both, `pop.pyramid` and `pop.trajectories.pyramid` (if called with a `bayesPop.prediction` object) use data from one country. Functions `pop.pyramidAll` and `pop.trajectories.pyramidAll` create such pyramids for all countries for which a projection is available and for all years given by the `year` argument which should be a list. In this case, one pyramid figure (possibly containing multiple pyramids) is created for each country and each element of the `year` list.

The core of these functions operates on a `bayesPop.pyramid` object which is automatically created when called with a `bayesPop.prediction` object. If used with a user-defined data set, one has to convert such data into `bayesPop.pyramid` using the function `get.bPop.pyramid` (see an example below). In such a case, one can simply use the `plot` function which then calls `pop.pyramid`.

## Value

`pop.pyramid`, `pop.trajectories.pyramid` and `get.bPop.pyramid` return an object of class `bayesPop.pyramid` which is a list with the following components:

<code>label</code>	Label used for the main titel.
<code>pyramid</code>	List of pyramid data, one element per pyramid. Each component is a data frame with at least two columns, containing data for the left and right panels of the pyramid. Their names must correspond to <code>LRcolnames</code> (see below). There is one row per age group and the row names are used for labeling the y-axis. Names of the list elements are used in the legend.
<code>CI</code>	List of lists of confidence intervals with one element per pyramid. The order corresponds to the order in the <code>pyramid</code> component and it is <code>NULL</code> if the corresponding pyramid does not have confidence intervals. Each element is a list with one element per probability interval whose names are the values of the intervals. Each element is again a list with components <code>low</code> and <code>high</code> which have the same structure as <code>pyramid</code> and contain the lower and upper bounds of the corresponding interval.
<code>trajectories</code>	List of lists of trajectories with one element per pyramid. As in the case of <code>CI</code> , it is ordered the same way as the <code>pyramid</code> component and is <code>NULL</code> if the corresponding pyramid does not have any trajectories to be shown. Each element is again a list with two components, one for the left part and one for the right part of the pyramid. Their names correspond to <code>LRcolnames</code> and each of them is a matrix of size number of age categories x number of trajectories. This is only used by the <code>pop.trajectories.pyramid</code> function.
<code>is.proportion</code>	Logical indicating if values in the various data frames in this object are proportions or raw values.

pop.max	Maximum value for the x-axis.
LRmain	Vector of character strings determining the titles for the left and right panels, respectively.
LRcolnames	Vector of character strings determining the column names in pyramid, CI and trajectories used to plot data into the left and right panel, respectively.

**Author(s)**

Hana Sevcikova, Adrian Raftery, using feedback from Sam Clark and the bayesPop group at the University of Washington.

**See Also**

[pop.trajectories.plot](#), [bayesPop.prediction](#), [summary.bayesPop.prediction](#)

**Examples**

```
# pyramids for bayesPop prediction objects
#####
sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir)
pop.pyramid(pred, "Netherlands", c(2045, 2010))
dev.new()
pop.trajectories.pyramid(pred, "Netherlands", c(2045, 2010, 1960), age=1:25, proportion=TRUE)
# using manual manipulation of the data: e.g. show only the prob. intervals
pred.pyr <- get.bPop.pyramid(pred, country="Ecuador", year=2090, age=1:27)
pred.pyr$pyramid <- NULL
plot(pred.pyr)

# pyramids for user-defined data
#####
# this example dataset contains population estimates for the Washington state and King county
# (Seattle area) in 2011
data <- read.table(file.path(find.package("bayesPop"), "ex-data", "popestimates_WAKing.txt"),
  header=TRUE, row.names=1)
# extract data for two pyramids and put it into the right format
head(data)
WA <- data[,c("WA.male", "WA.female")]; colnames(WA) <- c("male", "female")
King <- data[,c("King.male", "King.female")]; colnames(King) <- c("male", "female")
# create and plot a bayesPop.pyramid object
pyramid <- get.bPop.pyramid(list(WA, King), legend=c("Washington", "King"))
plot(pyramid, main="Population in 2011", pyr2.par=list(height=0.7, col="violet", border="violet"))
# show data as proportions
pyramid.prop <- get.bPop.pyramid(list(WA, King), is.proportion=NA, legend=c("Washington", "King"))
pop.pyramid(pyramid.prop, main="Population in 2011 (proportions)",
  pyr1.par=list(col="lightgreen", border="lightgreen", density=30),
  pyr2.par=list(col="darkred", border="darkred", density=50))
```

## Description

Obtain projection trajectories of population and vital events/rates. `get.pop` allows to access trajectories using a basic component of an expression. `get.pop.ex` and `get.pop.exba` returns results of an expression defined “by time” and “by age”, respectively. `get.trajectory.indices` creates a link to the probabilistic components of the projection by providing indices to the trajectories of TFR, e0 and migration. `extract.trajectories.eq` returns trajectories (of population or expression) and their indices that are closest to given values or a quantile. Similarly, functions `extract.trajectories.ge` and `extract.trajectories.le` return trajectories and their indices that are greater equal and less equal, respectively, to the given values or a quantile.

## Usage

```
pop.trajectories(pop.pred, country, sex = c("both", "male", "female"),
  age = "all", ...)
```

```
get.pop(object, pop.pred, aggregation = NULL, observed = FALSE, ...)
```

```
get.pop.ex(expression, pop.pred, observed = FALSE, ...)
```

```
get.pop.exba(expression, pop.pred, observed = FALSE, ...)
```

```
get.trajectory.indices(pop.pred, country,
  what = c("TFR", "e0M", "e0F", "migM", "migF"))
```

```
extract.trajectories.eq(pop.pred, country = NULL, expression = NULL,
  quant = 0.5, values = NULL, nr.traj = 1, ...)
```

```
extract.trajectories.ge(pop.pred, country = NULL, expression = NULL,
  quant = 0.5, values = NULL, all = TRUE, ...)
```

```
extract.trajectories.le(pop.pred, country = NULL, expression = NULL,
  quant = 0.5, values = NULL, all = TRUE, ...)
```

## Arguments

pop.pred	Object of class <a href="#">bayesPop.prediction</a> .
country	Name or numerical code of a country.
sex	One of “both” (default), “male” or “female”. By default the male and female projections are summed up.
age	Either a character string “all” (default) or an integer vector of age indices. Value 1 corresponds to age 0-4, value 2 corresponds to age 5-9 etc. Last age group 130+ corresponds to index 27. Results is summed over the given age categories.

object	Character string giving a basic component of an expression (see <a href="#">pop.expressions</a> ).
aggregation	If the basic component is to be evaluated on an aggregated prediction object, this argument gives the name of the aggregation (corresponds argument name in <a href="#">pop.aggregate</a> ). By default, the function searches for available aggregations and gives priority to the one called “country”.
observed	Logical. Determines if the evaluation uses observed data (TRUE) or predictions (FALSE).
expression	Expression defining the trajectories measure. For syntax see <a href="#">pop.expressions</a> . It must be define by age (i.e. contain curly braces) if used in <code>get.pop.exba</code> , and the opposite applies to <code>get.pop.ex</code> .
what	A character string that defines to which component should the indices link to. Allowable options are “TFR”, “e0M” (male life expectancy), “e0F” (female life expectancy), “migM” (male migration), “migF” (female migration).
quant	Quantile used to select the closest trajectories to.
values	Vector of values used to select the closest trajectories to. If it is not of length 1, it has to be of the same length as the number of projected time periods. If it is not given, quant is used.
nr. traj	Number of trajectories to return. This argument can be passed to any of the functions that contains . . . .
all	Logical indicating if the corresponding condition should apply to all time periods of a trajectory. If it is FALSE, a trajectory is extracted if the condition is fulfilled in at least one time period.
. . .	Additional argument passed to the underlying functions. In case of <code>get.pop</code> , <code>get.pop.ex</code> and <code>get.pop.exba</code> , this is only used for <code>observed=FALSE</code> . It can be either <code>nr. traj</code> giving the number of trajectories or logical <code>typical. trajectory</code> .

## Details

Function `pop.trajectories` returns an array of population trajectories for given sex and age.

Function `get.pop` evaluates a basic component of an expression and results in a four-dimensional array. Internally, this function is used for evaluation after an expression is decomposed into basic components. It can be useful for example for debugging purposes, to obtain results from parts of an expression. In addition, while `pop.trajectories` works only for population counts, `get.pop` can be used for obtaining trajectories of vital events and rates. Note that if `object` contains the wildcard “XXX”, the function only works on observed data, i.e. `observed` must be TRUE.

Functions `get.pop.ex` and `get.pop.exba` evaluate a whole expression and the dimensions of the resulting array is collapsed depending on the specific expression. Use `get.pop.ex` if the expected result of the expression does not contain the age dimension, i.e. it uses no brackets or square brackets. If it is not the case, i.e. the expression is defined using curly braces in order to include the age dimension, the `get.pop.exba` function is to be used. Argument `nr. traj` can be used to restrict the number of trajectories returned.

Function `get.trajectory.indices` returns an array of indices that link back to the given probabilistic component. It is of the same length as number of trajectories in the prediction object. For example, an array of `c(10, 15, 20)` (for a prediction with three trajectories) obtained with `what="TFR"` means that the 1st, 2nd and 3rd population trajectory, respectively, were generated with the 10th,

15th and 20th TFR trajectory, respectively. If the input TFR and e0 were generated using `bayesTFR` and `bayesLife`, functions `get.tfr.trajectories` and `get.e0.trajectories` can be used to extract the corresponding TFR and e0 trajectories.

Function `extract.trajectories.eq` can be used to select a given number of trajectories of any population quantity, including vital events, that are close to either specific values or to a given quantile. For example the default setting with `quant=0.5` and `nr.traj=1` returns the one trajectory that is “closest” to the median projection. As a measure of “closeness” the sum of absolute differences (across all time periods) is used.

Similarly, function `extract.trajectories.ge` (`extract.trajectories.le`) selects all trajectories that are greater (less) equal to the specific values or a given quantile. The argument `all` specifies, if the greater/less condition should be valid for all time periods of the selected trajectories or at least one time period.

## Value

Function `pop.trajectories` returns a two-dimensional array (time x trajectory).

Function `get.pop` returns an array of four dimensions (country x age x time x trajectory). See [pop.expressions](#) for more details.

Functions `get.pop.ex` and `get.pop.exba` return an array of trajectories. Its dimensions depend on the expression and whether it is evaluated on observed data or projections.

Function `get.trajectory.indices` returns a 1-d array of indices. If the given component is deterministic, it returns NULL.

Functions `extract.trajectories.eq`, `extract.trajectories.ge`, `extract.trajectories.le` return a list with two components. `trajectories`: 2-d array of trajectories; `index`: indices of the selected trajectories relative to the whole set of available trajectories.

## Author(s)

Hana Sevcikova

## See Also

[pop.expressions](#)

## Examples

```
sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir, write.to.cache=FALSE)

# observed female of Netherlands by age; 1x21x14x1 array
popFNL <- get.pop("PNL_F{}", pred, observed=TRUE)

# observed migration for all countries in the prediction object,
# here 2 countries; 2x1x14x1 array
migAll <- get.pop("GXXX", pred, observed=TRUE)

# projection population for Ecuador with 3 trajectories;
# 1x1x18x3 array
popEcu <- get.pop("P218", pred, observed=FALSE)
```

```

# the above is equivalent to
popEcu2 <- pop.trajectories(pred, "Ecuador")

# Expression "PNL_F{} / PNL_M{}" evaluated on projections
# is internally replaced by
FtoM <- get.pop("PNL_F{}", pred) / get.pop("PNL_M{}", pred)
# should return the same result as
FtoMa <- get.pop.exba("PNL_F{} / PNL_M{}", pred)

# the same expression by time (summed over ages)
FtoMt <- get.pop.ex("PNL_F / PNL_M", pred)

# the example simulation was generated with 3 TFR trajectories ...
get.trajectory.indices(pred, "Netherlands", what="TFR")
# ... and 1 e0 trajectory
get.trajectory.indices(pred, "Netherlands", what="e0M")

# The three trajectories of the population ratio of Ecuador to Netherlands
get.pop.ex("PEC/PNL", pred)
# Returns the trajectory closest to the upper 80% bound, including the corresponding index
extract.trajectories.eq(pred, expression="PEC/PNL", quant=0.9)
# Returns the median trajectory and the high variant, including the corresponding index
extract.trajectories.ge(pred, expression="PEC/PNL", quant=0.45)

```

---

pop.trajectories.plot *Output of Probabilistic Population Projection*

---

## Description

The functions plot and tabulate the distribution of population projection for a given country, or for all countries, including the median and given probability intervals.

## Usage

```

pop.trajectories.plot(pop.pred, country = NULL, expression = NULL, pi = c(80, 95),
  sex = c("both", "male", "female"), age = "all", sum.over.ages = FALSE,
  half.child.variant = FALSE, nr.traj = NULL, typical.trajectory = FALSE,
  main = NULL, dev.ncol = 5, lwd = c(2, 2, 2, 2, 1),
  col = c("black", "red", "red", "blue", "#00000020"), show.legend = TRUE,
  ann = par("ann"), ...)

```

```

pop.trajectories.plotAll(pop.pred,
  output.dir=file.path(getwd(), "pop.trajectories"),
  output.type="png", expression = NULL, verbose=FALSE, ...)

```

```

pop.trajectories.table(pop.pred, country = NULL, expression = NULL, pi = c(80, 95),
  sex = c("both", "male", "female"), age = "all", half.child.variant = FALSE, ...)

```

```

pop.byage.plot(pop.pred, country = NULL, year = NULL, expression = NULL,
  pi = c(80, 95), sex = c("both", "male", "female"),
  half.child.variant = FALSE, nr.traj = NULL, typical.trajectory=FALSE,
  xlim = NULL, ylim = NULL, xlab = "", ylab = "Population projection",
  main = NULL, lwd = c(2,2,2,1), col = c("red", "red", "blue", "#00000020"),
  show.legend = TRUE, add = FALSE, ann = par("ann"), type = "l", pch = NA,
  pt.cex = 1, ...)

pop.byage.plotAll(pop.pred,
  output.dir=file.path(getwd(), "pop.byage"),
  output.type="png", expression = NULL, verbose=FALSE, ...)

pop.byage.table(pop.pred, country = NULL, year = NULL, expression = NULL,
  pi = c(80, 95), sex = c("both", "male", "female"),
  half.child.variant = FALSE)

```

### Arguments

pop.pred	Object of class <a href="#">bayesPop.prediction</a> .
country	Name or numerical code of a country.
expression	Expression defining the population measure to be plotted. For syntax see <a href="#">pop.expressions</a> . For <code>pop.trajectories.plot</code> , <code>pop.trajectories.table</code> , <code>pop.byage.plot</code> and <code>pop.byage.table</code> the basic components of the expression must be country-specific. For <code>pop.trajectories.plotAll</code> and <code>pop.byage.plotAll</code> the country part should be given as "XXX". In addition, expressions passed into <code>pop.byage.plot</code> and <code>pop.byage.table</code> must contain curly braces (i.e. be age specific).
pi	Probability interval. It can be a single number or an array.
sex	One of "both" (default), "male" or "female". By default the male and female projections are summed up.
age	Either a character string "all" (default) or an integer vector of age indices. Value 1 corresponds to age 0-4, value 2 corresponds to age 5-9 etc. Last age group 130+ corresponds to index 27. It can also be the string "psr" (potential support ratio), in which case a ratio of population of ages 20-64 to 65+ (indices 5:13 to 14:27) is shown.
sum.over.ages	Logical. If TRUE, the values are summed up over given age groups. Otherwise there is a separate plot for each age group.
half.child.variant	Logical. If TRUE the United Nations "+/-0.5 child" variant computed with fertility $\pm 0.5 \times$ TFR median and the median of life expectancy is shown.
nr.traj	Number of trajectories to be plotted. If NULL, all trajectories are plotted, otherwise they are thinned evenly.
typical.trajectory	Logical. If TRUE one trajectory is shown that has the smallest distance to the median.
xlim, ylim, xlab, ylab, main, ann, pt.cex	Graphical parameters passed to the plot function.

<code>dev.ncol</code>	Number of column for the graphics device if <code>sum.over.ages</code> is FALSE. If the number of age groups is smaller than <code>dev.ncol</code> , the number of columns is automatically decreased.
<code>lwd, col</code>	For the first three functions it is a vector of five elements giving the line width and color for: 1. observed data, 2. median, 3. quantiles, 4. half-child variant, 5. trajectories. For functions that show results by age it is a vector of four elements - as above without the first item (observed data).
<code>type, pch</code>	Currently works for plotting by age only. It is a vector of four elements giving the plot type and point type for: 1. median, 2. quantiles, 3. half-child variant, 4. trajectories. The last element of the array is recycled.
<code>show.legend</code>	Logical controlling whether the legend should be drawn.
<code>...</code>	Additional graphical arguments. Functions <code>pop.trajectories.plotAll</code> and <code>pop.byage.plotAll</code> accept also any arguments of <code>pop.trajectories.plot</code> and <code>pop.byage.plot</code> , respectively, except <code>country</code> .
<code>output.dir</code>	Directory into which resulting graphs are stored.
<code>output.type</code>	Type of the resulting files. It can be “png”, “pdf”, “jpeg”, “bmp”, “tiff”, or “postscript”.
<code>verbose</code>	Logical switching log messages on and off.
<code>year</code>	Any year within the time period to be outputted.
<code>add</code>	Logical specifying if the plot should be added to an existing graphics.

## Details

`pop.trajectories.plot` plots trajectories of population projection by time for a given country. `pop.trajectories.table` gives the same output as a table. `pop.trajectories.plotAll` creates a set of graphs (one per country) that are stored in `output.dir`. The projections can be visualized separately for each sex and age groups, or summed up over both sexes and/or given age groups. This is controlled by the arguments `sex`, `age` and `sum.over.ages`.

`pop.byage.plot` and `pop.byage.table` plots/tabulate the posterior distribution by age for a given country and time period. `pop.byage.plotAll` creates such plots for all countries.

The median and given probability intervals are computed using all available trajectories. Thus, `nr.traj` does not influence those values - it is used only to control the number of trajectories plotted.

If plotting results of an expression and the function fails, to debug obtain values of that expression using the functions `get.pop.ex` (for `pop.trajectories.plot`) and `get.pop.exba` (for `pop.byage.plot`).

## Author(s)

Hana Sevcikova

## See Also

[bayesPop.prediction](#), [summary.bayesPop.prediction](#), [pop.pyramid](#), [pop.expressions](#), [get.pop](#)

**Examples**

```

sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir)
pop.trajectories.plot(pred, country="Ecuador", pi=c(80, 95))
pop.trajectories.table(pred, country="Ecuador", pi=c(80, 95))
# female population of Ecuador in child bearing ages (by time)
pop.trajectories.plot(pred, expression="PEC_F[4:10]")
# Population by age in Netherands for two different years
pop.byage.plot(pred, country="Netherlands", year=2050)
pop.byage.plot(pred, expression="PNL{}", year=2000)

```

project.pasfr

*Projections of Percent Age-Specific Fertily Rate***Description**

The projections of percent age-specific fertility rate (PASFR) is normally computed within the [pop.predict](#) function for each trajectory. This function allows to project PASFR outside of population projections for the median total fertility and export it.

**Usage**

```

project.pasfr(inputs = NULL, present.year = 2020, end.year = 2100,
             wpp.year = 2019, digits = 2, out.file.name = "percentASFR.txt")

project.pasfr.traj(inputs = NULL, countries = NULL, nr.traj = NULL,
                  present.year = 2020, end.year = 2100, wpp.year = 2019,
                  digits = 2, out.file.name = "percentASFRtraj.txt")

```

**Arguments**

inputs	List of input data (file names) with the same meaning as in <a href="#">pop.predict</a> . The relevant items here are: either <code>tfr.file</code> or <code>tfr.sim.dir</code> (TFR projections), <code>pasfr</code> (PASFR for observed time periods), <code>srb</code> (observed and predicted sex ratio at birth), <code>patterns</code> (PASFR patterns). All entries are optional. By default the data is taken from the corresponding <code>wpp</code> package.
present.year	Year of the last observed data point.
end.year	End year of the projection.
wpp.year	Year for which WPP data is used if one of the <code>inputs</code> components is left out.
digits	Number of decimal places in the results.
out.file.name	Name of the resulting file. If <code>NULL</code> nothing is written.
countries	Vector of numerical country codes. By default the function is applied to all countries.
nr.traj	Number of trajectories on which the function should be applied. By default all trajectories are taken. Otherwise they are thinned appropriately.

### Details

For `project.pasfr`, the median total fertility rate is derived from the inputs and PASFR corresponding to this median is projected using the method from Sevcikova et al (2015).

For `project.pasfr.traj`, the PASFR is projected for single trajectories of TFR.

### Value

Returns invisible data frame with the projected PASFR.

### Author(s)

Igor Ribeiro, Hana Sevcikova

### References

H. Sevcikova, N. Li, V. Kantorova, P. Gerland and A. E. Raftery (2015). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. arXiv:1503.05215. <http://arxiv.org/abs/1503.05215>

### See Also

[pop.predict](#)

### Examples

```
## Not run:
inputs <- list(tfr.sim.dir=file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output"))
pasfr <- project.pasfr(inputs, out.file.name=NULL)
head(pasfr)

pasfr.traj <- project.pasfr.traj(inputs, out.file.name=NULL)
head(pasfr.traj)

## End(Not run)
```

---

summary.bayesPop.prediction

*Summary of Probabilistic Population Projection*

---

### Description

Summary of an object `bayesPop.prediction` created using the `pop.predict` function. The summary contains the mean, standard deviation and several commonly used quantiles of the simulated trajectories.

**Usage**

```
## S3 method for class 'bayesPop.prediction'
summary(object, country = NULL,
        sex = c("both", "male", "female"), compact = TRUE, ...)
```

**Arguments**

object	Object of class <a href="#">bayesPop.prediction</a> .
country	Country name or code. If it is NULL, only meta information included.
sex	One of “both” (default), “male”, or “female”. If it is not “both”, the summary is given for sex-specific trajectories.
compact	Logical switching between a smaller and larger number of displayed quantiles.
...	A list of further arguments.

**Author(s)**

Hana Sevcikova

**See Also**

[bayesPop.prediction](#)

**Examples**

```
sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir)
summary(pred, "Netherlands")
```

---

vwBaseYear	<i>Datasets on Migration Base Year and Type, and Mortality and Fertility Age Patterns</i>
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---

**Description**

Datasets giving information on the baseyear and type of migration for each country. The 2012, 2015, 2017 and 2019 datasets also give information on country’s specifics regarding mortality and fertility age patterns.

**Usage**

```
data(vwBaseYear2019)
data(vwBaseYear2017)
data(vwBaseYear2015)
data(vwBaseYear2012)
data(vwBaseYear2010)
```

**Format**

A data frame containing the following variables:

`country_code` Numerical Location Code (3-digit codes following ISO 3166-1 numeric standard)  
- see [http://en.wikipedia.org/wiki/ISO\\_3166-1\\_numeric](http://en.wikipedia.org/wiki/ISO_3166-1_numeric).

`country` Country name. Not used by the package.

`isSmall` UN internal code. Not used by the package.

`ProjFirstYear` The base year of migration.

`MigCode` Type of migration. Zero means migration is evenly distributed over each time interval.  
Code 9 means migration is captured at the end of each interval.

`WPPAIDS` Dummy indicating if the country has generalized HIV/AIDS epidemics.

`AgeMortalityType` Type of mortality age pattern. Only relevant for countries with the entry “Model life tables”. In such a case, the  $b_x$  Lee-Carter parameter is not estimated from historical data. Instead is taken from the dataset `MLTbx` using a pattern given in the `AgeMortalityPattern` column.

`AgeMortalityPattern` If `AgeMortalityType` is equal to “Model life tables”, this value determines which  $b_x$  is selected from the `MLTbx` dataset. It must correspond to one of the rownames of `MLTbx`, e.g. “CD East”, “CD West”, “UN Latin American”.

`AgeMortProjMethod1` Method for projecting age-specific mortality rates. It is one of “LC” (modified Lee-Carter, uses function `mortcast`), “PMD” (pattern mortality decline, uses function `copmd`), “MLT” (model life tables, uses function `mlt`), “LogQuad” (log quadratic method, uses function `logquad`), or “HIVmortmod” (HIV model life tables as implemented in the **HIV.LifeTables** package which can be installed from the **PPgP/HIV.LifeTables** GitHub repo).

`AgeMortProjMethod2` If the mortality rates are to be projected via a blend of two methods (see `mortcast.blend`), this column determines the second method. The options are the same as in the column `AgeMortProjMethod1`.

`AgeMortProjPattern` If one of the `AgeMortProjMethodX` columns contains the “MLT” method, this column determines the type of the life table (see the argument `type` in the `mlt` function).

`AgeMortProjMethodWeights` If the mortality rates are to be projected via a blend of two methods, this column determines the weights in the first and the last year of the projection, respectively. It should be given as an R vector, e.g. “c(1, 0.5)” (see the argument `weights` in `mortcast.blend`).

`AgeMortProjAdjSR` Code determining how the “PMD” method should be adjusted if it’s used. 0 means no adjustment, 1 means the argument `sexratio.adjust` in `copmd` is set to TRUE, and code 3 means that the argument `adjust.sr.if.needed` in `copmd` is set to TRUE.

`LatestAgeMortalityPattern` Indicator  $n$  for how many latest time periods should be used to compute the  $a_x$  Lee-Carter parameter. If  $n$  is zero, all time periods are used. If  $n$  is one, only the latest time period is used. If  $n$  is negative, the latest  $n$  time periods are excluded.

`SmoothLatestAgeMortalityPattern` If `LatestAgeMortalityPattern` is 1, this column indicates if the  $a_x$  should be smoothed.

`PasfrNorm` Type of norm for computing age-specific fertility pattern to which the country belongs to. Currently only “GlobalNorm” is used.

`PasfrGlobalNorm`, `PasfrFarEastAsianNorm`, `PasfrSouthAsianNorm` Dummies indicating which country to include to compute the specific norms.

## Details

There is one record for each country. See Sevcikova et al (2015) on how information from the various columns is used for projections.

## Source

Data provided by the United Nations Population Division.

## References

H. Sevcikova, N. Li, V. Kantorova, P. Gerland and A. E. Raftery (2015). Age-Specific Mortality and Fertility Rates for Probabilistic Population Projections. arXiv:1503.05215. <http://arxiv.org/abs/1503.05215>

## Examples

```
data(vwBaseYear2019)
str(vwBaseYear2019)
```

---

```
write.pop.projection.summary
```

*Writing Projection Summary Files*

---

## Description

The function creates ASCII files containing projection summaries, such as the median, the lower and upper bound of the 80 and 95% probability intervals, respectively.

## Usage

```
write.pop.projection.summary(pop.pred, what = NULL, expression = NULL,
  output.dir = NULL, ...)
```

## Arguments

pop.pred	Object of class <a href="#">bayesPop.prediction</a> .
what	A character vector specifying what kind of projection to write. Total population is specified by “pop”. Vital events are specified by “births”, “deaths”, “sr” (survival rate), “fertility” and “pfertility” (percent fertility). Each of these strings can (some must) have a suffix “sex” and/or “age” if sex- and/or age-specific measure is desired. For example, “popage”, “birthssexage”, “deaths”, “deaths-sex”, are all valid values. Note that for survival, only “srsexage” is allowed. For percent fertility, only “pfertilityage” is allowed. Suffix “sex” cannot be used in combination with “fertility”. Moreover, “fertility” (without age) corresponds to the total fertility rate. If the argument is NULL, all valid combinations are used. The argument is not used if expression is given. Note that vital events can be only used if the prediction object contains vital events, i.e. if it was generated with the <code>keep.vital.events</code> argument being TRUE (see <a href="#">pop.predict</a> ).

expression	Expression defining the measure to be written. If it is not NULL, argument what is ignored. For expression syntax see <a href="#">pop.expressions</a> . The country components of the expression should be given as “XXX”.
output.dir	Directory in which the resulting files will be stored. If NULL pop.pred\$output.directory is used.
...	These are arguments used if expression is given: file.suffix defines the file suffix; expression.label (defaults to the expression) is put as the first line in the resulting file; logical include.observed determines if observed data should be included; integer digits defines the number of decimal places in the resulting file.

### Details

There is one file created per value of what, or expression, called ‘projection\_summary\_’*suffix*‘.csv’, where *suffix* is either what or, if an expression is given, the value of file.suffix. It is a comma-separated table with the following columns:

- “country\_name”: country name
- “country\_code”: country code
- “variant”: name of the variant, such as “median”, “lower 80”, “upper 80”, “lower 95”, “upper 95”
- period1: e.g. “2005-2010”: Given population measure for the first time period
- period2: e.g. “2010-2015”: Given population measure for the second time period
- ... further time period columns

If expression is given, expression.label (by default the full expression) is written as the first line of the file starting with #. The file contains one line per country, and possibly sex and age.

### Note

If the expression argument is used, the same applies as for [pop.map](#) in terms of Performance and Caching.

### Author(s)

Hana Sevcikova

### See Also

[pop.predict](#), [pop.map](#), [pop.expressions](#)

### Examples

```
outdir <- tempfile()
dir.create(outdir)
sim.dir <- file.path(find.package("bayesPop"), "ex-data", "Pop")
pred <- get.pop.prediction(sim.dir=sim.dir, write.to.cache=FALSE)

# proportion of 65+ years old to the whole population
```

```
write.pop.projection.summary(pred, expression="PXXX[14:27] / PXXX", file.suffix="age65plus",
  output.dir=outdir, include.observed=TRUE, digits=2)

# various measures
write.pop.projection.summary(pred, what=c("pop", "popsexage", "popsex"),
  output.dir=outdir)

unlink(outdir, recursive=TRUE)
```

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