# Package 'bayesTFR'

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Author Hana Sevcikova (hanas@uw.edu), Leontine Alkema (alkema@nus.edu.sg), Adrian Raftery (raftery@uw.edu) with code contribution from Bailey Fosdick (bfosdick@uw.edu) and Patrick Gerland (gerland@un.org)
Maintainer Hana Sevcikova <hanas@uw.edu></hanas@uw.edu>
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# Description

Collection of functions for making probabilistic projections of total fertility rate (TFR) for all countries of the world, using a Bayesian hierarchical model (BHM) and the United Nations demographic time series.

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#### **Details**

Package: bayesTFR
Version: 4.0-5
Date: 2014-02-07
License: GPL (>= 2)

URL: http://bayespop.csss.washington.edu

The projection follows a method developed by Alkema et al. (2011) and Raftery et al (2013). It uses historical data provided by the United Nations (included in the package) to simulate a posterior distribution of total fertility rates for all countries in the world simultaneously.

The estimation is split into two parts:

- 1. BHM for fertility in a transition phase (Phase II), as described in Alkema et al. (2011),
- 2. BHM for fertility in a post-transition phase (Phase III), as described in Raftery et al (2013).

The second part is optional and can be replaced by a simple AR(1) process.

The main functions of the package are:

- run.tfr.mcmc: Evokes running a Markov Chain Monte Carlo (MCMC) simulation for TFR in Phase II using one or more chains, possibly in parallel. It results in a posterior sample of the mcmc parameters. Existing simulation runs can be resumed using continue.tfr.mcmc.
- run.tfr3.mcmc: Starts MCMCs for TFR in Phase III. Existing simulation runs can be resumed using continue.tfr3.mcmc.
- tfr.predict: Using the posterior parameter samples it derives posterior trajectories of the total fertility rate for all countries.
- run.tfr.mcmc.extra: Runs MCMC for extra countries or regions, i.e. for countries not included in the Bayesian hierarchical model. It can be also used for aggregations.
- tfr.predict.extra: Generates predictions for extra countries or aggregated regions.

The order of the functions above roughly corresponds to a typical workflow when using the package: 1. run a Phase II MCMC simulation, 2. run a Phase III MCMC simulation (optional but recommended), 3. generate predictions, 4. analyze results (using the functions below). If there are countries that were not included in steps 1.-3., or if there are aggregated regions for which a prediction is desired, one proceeds with the two functions at the bottom of the list above, followed by the analyzing functions below.

A number of functions analyzing results are included in the package:

- tfr.trajectories.plot: Shows the posterior trajectories for a given country, including their median and given probability intervals.
- tfr.trajectories.table: Shows the posterior trajectories for a given country in a tabular form.
- tfr.map: Shows a TFR world map for a given projection period.
- DLcurve.plot: Shows the posterior curves of the double logistic function used in the simulation, including their median and given probability intervals.

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• tfr.partraces.plot and tfr.partraces.cs.plot: Plot the Phase II MCMC traces of country-independent parameters and country-specific parameters, respectively. tfr3.partraces.plot and tfr3.partraces.cs.plot do the same for Phase III MCMCs.

- tfr.pardensity.plot and tfr.pardensity.cs.plot: Plot the posterior density of the Phase II MCMCs for country-independent parameters and country-specific parameters, respectively. tfr3.pardensity.plot and tfr3.pardensity.cs.plot do the same for Phase III MCMCs.
- summary.bayesTFR.mcmc.set: Summary function for the MCMC results.
- summary.bayesTFR.prediction: Summary function for the prediction results.

For MCMC diagnostics, functions coda.list.mcmc and coda.list.mcmc3 create an object of type "mcmc.list" that can be used with the **coda** package. Furthermore, function tfr.diagnose and tfr3.diagnose analyze the MCMCs using the Raftery diagnostics implemented in the **coda** package and gives information about parameters that did not converge.

Existing simulation results can be accessed using the get.tfr.mcmc (Phase II) and get.tfr3.mcmc (Phase III) function. An existing prediction can be accessed via get.tfr.prediction. Existing convergence diagnostics can be accessed using the get.tfr.convergence, get.tfr.convergence.all, get.tfr3.convergence and get.tfr3.convergence.all functions.

The historical TFR data are taken from one of the packages wpp2008, wpp2010 or wpp2012, depending on users settings.

#### Note

There is a directory ex-data shipped with the package which contains results from an example simulation, containing one chain with 60 iterations. The Example section below shows how these results were created. These data are used in Example sections throughout the manual. The user can either reproduce the data in her/his local directory, or use the ones from the package.

# Author(s)

Maintainer: Hana Sevcikova <hanas@uw.edu>

#### References

Hana Sevcikova, Leontine Alkema, Adrian E. Raftery (2011). bayesTFR: An R Package for Probabilistic Projections of the Total Fertility Rate. Journal of Statistical Software, 43(1), 1-29. http://www.jstatsoft.org/v43/i01/.

Raftery, A.E., Alkema, L. and Gerland, P. (2013). Bayesian Population Projections for the United Nations. Statistical Science, in press.

L. Alkema, A. E. Raftery, P. Gerland, S. J. Clark, F. Pelletier, Buettner, T., Heilig, G.K. (2011). Probabilistic Projections of the Total Fertility Rate for All Countries. Demography, Vol. 48, 815-839. (A Working Paper version at http://www.csss.washington.edu/Papers [nr. 97])

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# **Examples**

```
## Not run:
# This command produces output data such as in the directory ex-data
sim.dir <- tempfile()
# Phase II MCMCs
m <- run.tfr.mcmc(nr.chains=1, iter=60, output.dir=sim.dir, seed=1, verbose=TRUE)
# Phase III MCMCs (not included in the package)
m3 <- run.tfr3.mcmc(sim.dir=sim.dir, nr.chains=2, iter=100, thin=1, seed=1, verbose=TRUE)
# Prediction
pred <- tfr.predict(m, burnin=30, burnin3=50, verbose=TRUE)
summary(pred, country='Ghana')
unlink(sim.dir, recursive=TRUE)
## End(Not run)</pre>
```

bayesTFR.mcmc

MCMC Simulation Object

### **Description**

MCMC simulation object bayesTFR.mcmc containing information about one MCMC chain, either from Phase II or Phase III simulation. A set of such objects belonging to the same simulation together with a bayesTFR.mcmc.meta object constitute a bayesTFR.mcmc.set object.

## **Details**

An object bayesTFR.mcmc points to a place on disk (element output.dir) where MCMC results from all iterations are stored. They can be retrieved to the memory using get.tfr.mcmc(...) (Phase II) or get.tfr3.mcmc(...) (Phase III), and tfr.mcmc(...).

The object is in standard cases not to be manipulated by itself, but rather as part of a bayesTFR.mcmc.set object.

#### Value

A bayesTFR.mcmc object contains parameters of the Bayesian hierarchical model, more specifically, their values from the last iteration. If it is a **Phase II** object these parameters are:

psi, chi, a\_sd, b\_sd, const\_sd, S\_sd, sigma0, mean\_eps\_tau, sd\_eps\_tau, Triangle4 - non-country specific parameters, containing one value each.

alpha, delta - non-country specific parameters, containing three values each.

U\_c, d\_c, Triangle\_c4 - country-specific parameters (1d array).

gamma\_ci - country-specific parameter with three values for each country, i.e. an  $n \times 3$  matrix where n is the number of countries.

**Phase III** MCMC objects contain single-value parameters mu, rho, sigma.mu, sigma.rho, sigma.eps and *n*-size vectors mu.c and rho.c.

Furthermore, the object (independent of Phase) contains components:

iter Total number of iterations the simulation was started with.

bayesTFR.mcmc.meta

finished.iter Number of iterations that were finished. Results from the last finished iteration

are stored in the parameters above.

length Length of the MCMC stored on disk. It differs from finished.iter only if

thin is larger than one.

thin Thinning interval used when simulating the MCMCs.

id Identifier of this chain.

output.dir Subdirectory (relative to output.dir in the bayesTFR.mcmc.meta object) where

results of this chain are stored.

This is a placeholder for keeping whole parameter traces in the memory. If the

processing operates in a low memory mode, it will be 0. It can be filled in using the function <code>get.tfr.mcmc(..., low.memory=FALSE)</code>. In such a case, traces is a  $I \times J$  array where I is the MCMC length and J is the number of

parameters.

traces.burnin Burnin used to retrieve the traces, i.e. how many stored iterations are missing

from the beginning in the traces array comparing to the 'raw' traces on the

disk.

rng. state State of the random number generator at the end of the last finished interation.

compression.type

Type of compression of the underlying files.

meta Object of class bayesTFR.mcmc.meta used for simulation of this chain.

# Author(s)

Hana Sevcikova

# See Also

```
run.tfr.mcmc, get.tfr.mcmc, run.tfr3.mcmc, get.tfr3.mcmc, bayesTFR.mcmc.set, bayesTFR.mcmc.meta
```

# **Examples**

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
# loads traces from one chain
m <- get.tfr.mcmc(sim.dir, low.memory=FALSE, burnin=35, chain.ids=1)
# should have 25 rows, since 60 iterations in total minus 35 burnin
dim(tfr.mcmc(m, 1)$traces)
summary(m, chain.id=1)</pre>
```

bayesTFR.mcmc.meta

MCMC Simulation Meta Object

# **Description**

Simulation meta object bayesTFR.mcmc.meta used by all chains of the same MCMC simulation. It contains information that is common to all chains. It is part of a bayesTFR.mcmc.set object.

#### **Details**

The object is in standard cases not to be manipulated by itself, but rather as part of a bayesTFR.mcmc.set object.

# Value

A bayesTFR.mcmc.meta object contains various components that correspond to the input arguments of the run.tfr.mcmc and run.tfr3.mcmc functions. Furthermore, it contains components:

nr.chains Number of MCMC chains.

phase Value 2 or 3, depending which Phase the object belongs to.

output.dir Directory for storing simulation output.

#### Value - Phase II

Furthermore, Phase II meta objects contain components:

**tfr\_matrix\_all** A  $q \times n$  matrix with the United Nations TFR estimates. q is number of years (see T\_end below), n is number of countries (see nr\_countries below). The first  $n_e$  columns correspond to countries included in the MCMC estimation (see nr\_countries\_estimation below), where  $n_e <= n$ .

tfr\_matrix Like tfr\_matrix\_all, but it has NA values before and after country"s fertility transition

tfr\_matrix\_observed Like tfr\_matrix, but it has NA values for years where no historical data is available.

nr\_countries Number of countries included in the tfr matrices.

**nr\_countries\_estimation** Number of countries included in the MCMC estimation. It must be smaller or equal to nr\_countries.

tau\_c Estimated start year of the fertility decline for each country (as a row index within the tfr matrices). -1 means that the decline started before start.year.

id\_Tistau Index of countries for which present.year is equal to tau\_c.

id\_DL Index of countries for which the projection is made using the double logistic function, i.e. high fertility countries.

id\_early Index of countries with early decline, i.e. countries for which tau\_c=-1.

**id\_notearly** Index of countries with not early decline.

lambda\_c Start period of the recovery phase for each country (as an index of the tfr\_matrix).

**start\_c** Maximum of tau\_c and 1 for each country. Thus, it is a row index of the tfr\_matrix where the fertility decline starts.

**proposal\_cov\_gammas\_cii** Proposal covariance matrices of  $\gamma_{ci}$  for each country.

**T\_end** Number of years for which United Nations historical data are available (i.e. number of rows of tfr\_matrix).

**T\_end\_c** Like **T\_end** but country specific.

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```
regions List of arrays of length nr_countries. These are:

name - Region name for each country.

code - Region code for each country.

area_name - Area name for each country.

area_code - Area code for each country.

country_name - Array of country names.

country_code - Array of country codes.

Any country indices in the bayesTFR.mcmc.meta object are derived from this component.
```

#### Value - Phase III

Phase III meta objects contain additional components:

id\_phase3 Indices of countries included in the Phase III estimation. It is relative to the order of countries in the region object in the parent meta object.

**nr.countries** Number of countries included in the estimation.

parent Link to the Phase II meta object.

# Author(s)

Hana Sevcikova, Leontine Alkema

# See Also

```
run.tfr.mcmc, get.tfr.mcmc, run.tfr3.mcmc, get.tfr3.mcmc
```

# **Examples**

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
m <- get.tfr.mcmc(sim.dir)
summary(m, meta.only = TRUE)
names(m$meta)</pre>
```

 ${\tt coda.list.mcmc}$ 

Convertion to coda's Objects

# Description

The functions convert MCMC traces (simulated using run.tfr.mcmc and run.tfr3.mcmc) into objects that can be used with the **coda** package.

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# Usage

```
coda.list.mcmc(mcmc = NULL, country = NULL, chain.ids = NULL,
    sim.dir = file.path(getwd(), "bayesTFR.output"),
    par.names = tfr.parameter.names(),
    par.names.cs = tfr.parameter.names.cs(),
    rm.const.pars = FALSE, burnin = 0,
    low.memory = FALSE, ...)

coda.list.mcmc3(mcmc = NULL, country = NULL, chain.ids = NULL,
    sim.dir = file.path(getwd(), "bayesTFR.output"),
    par.names = tfr3.parameter.names(),
    par.names.cs = tfr3.parameter.names.cs(),
    burnin = 0, low.memory = FALSE, ...)

## S3 method for class 'bayesTFR.mcmc'
coda.mcmc(mcmc, country = NULL,
    par.names = NULL, par.names.cs = NULL,
    burnin = 0, thin = 1, ...)
```

Thinning interval.

# **Arguments**

thin

. . .

мсмс	In coda.mcmc, it is an object of class bayesTFR.mcmc. In coda.list.mcmc and coda.list.mcmc3, it is either a list of bayesTFR.mcmc objects, or an object of class bayesTFR.mcmc.set or in case of coda.list.mcmc it can be bayesTFR.prediction. If it is NULL, the MCMCs are loaded from sim.dir. Either mcmc or sim.dir must be given.
country	Country name or code. It is used in connection with the par.names.cs argument (see below).
chain.ids	Vector of chain identifiers. By default, all chains available in the mcmc.list object are included.
sim.dir	Directory with the MCMC simulation results. Only used if mcmc.list is NULL.
par.names	Names of country-independent parameters to be included. In coda.mcmc the default names are tfr.parameter.names() if the mcmc object is an MCMC of phase II or tfr3.parameter.names() if the MCMC is of phase III.
par.names.cs	Names of country-specific parameters to be included. The argument country is used to filter out traces that correspond to a specific country. If country is not given, for each parameter, traces for all countries are included. In coda.mcmc the default names are tfr.parameter.names.cs() if the mcmc object is an MCMC of phase II or tfr3.parameter.names.cs() if the MCMC is of phase III.
rm.const.pars	Logical indicating if parameters with constant values should be removed.
burnin	Burnin indicating how many iterations should be removed from the beginning of each chain.
low.memory	Logical indicating if the function should run in a memory-efficient mode.

Additional arguments passed to the  ${\bf coda}$ 's  ${\bf mcmc}$  function.

# **Details**

Function coda.list.mcmc is for accessing all chains of phase II MCMCs; Function coda.list.mcmc3 is for accessing all chains of phase III MCMCs.

#### Value

The function coda.list.mcmc and coda.list.mcmc3 return an object of class "mcmc.list". The function coda.mcmc returns an object of class "mcmc", both defined in the **coda** package.

# Author(s)

Hana Sevcikova

```
convert.tfr.trajectories
```

Converting TFR Trajectories into ACSII Files

# **Description**

Converts TFR trajectories stored in a binary format into two CSV files of a UN-specific format.

# Usage

```
convert.tfr.trajectories(dir = file.path(getwd(), 'bayesTFR.output'),
    n = 1000, output.dir = NULL, verbose = FALSE)
```

# **Arguments**

dir	Directory containing the prediction object. It should correspond to the output.dir argument of the tfr.predict function.
n	Number of trajectories to be stored. It can be either a single number or the word "all" in which case all trajectories are stored.
output.dir	Directory in which the resulting files will be stored. If NULL the same directory is used as for the prediction.
verbose	Logical switching log messages on and off.

# **Details**

The function creates two files. One is called "ascii\_trajectories.csv", it is a comma-separated table with the following columns:

- "LocID" country code
- "Period" prediction interval, e.g. 2015-2020
- "Year" middle year of the prediction interval
- "Trajectory" identifier of the trajectory

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• "TF"total fertility rate

The second file is called "ascii\_trajectories\_wide.csv", it is also a comma-separated table and it contains the same information as above but in a 'transposed' format. I.e. the data for one country are ordered in columns, thus, there is one column per country. The country columns are ordered alphabetically.

If n is smaller than the total number of trajectories, the trajectories are selected using equal spacing.

#### Note

This function is automatically called from the tfr.predict function, therefore in standard cases it will not be needed to call it directly. However, it can be useful for example, if different number of trajectories are to be converted, without having to re-run the prediction.

# Author(s)

Hana Sevcikova

#### See Also

```
write.projection.summary,tfr.predict
```

# **Examples**

country.names

Accessing Country Names

# **Description**

The function returns country names for countries given either by their codes or by index.

# Usage

```
country.names(meta, countries = NULL, index = FALSE)
```

# **Arguments**

meta Object of class bayesTFR.mcmc.meta, bayesTFR.mcmc.set, bayesTFR.mcmc,

or bayesTFR.prediction.

countries Vector of country codes or indices. If it is not given, names of all countries are

returned.

index Logical indicating if the argument countries is an index.

#### **Details**

The function considers countries that are included in the simulations and predictions. If the argument countries is not given, all countries are returned in the same order as they are stored in the meta object.

#### Value

Vector of country names.

# Author(s)

Hana Sevcikova

## See Also

```
get.country.object
```

# **Examples**

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
m <- get.tfr.mcmc(sim.dir)
country.names(m)
# these two calls should give the same answer
country.names(m, c(800, 120))
country.names(m, c(15, 20), index=TRUE)</pre>
```

DLcurve.plot

Plotting Posterior Distribution of the Double Logistic Function

# **Description**

The functions plot the posterior distribution of the double logistic function used in the simulation of Phase II, including their median and given probability intervals.

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# Usage

```
DLcurve.plot(mcmc.list, country, burnin = NULL, pi = 80, tfr.max = 10,
    nr.curves = NULL, predictive.distr = FALSE, ylim = NULL,
    xlab = 'TFR (reversed)', ylab = 'TFR decrement', main = NULL,
    show.legend = TRUE, ...)
DLcurve.plot.all(mcmc.list = NULL, sim.dir = NULL,
    output.dir = file.path(getwd(), 'DLcurves'),
    output.type = "png", burnin = NULL, verbose = FALSE, ...)
```

# **Arguments**

mcmc.list	List of bayesTFR.mcmc objects, an object of class bayesTFR.mcmc.set or of class bayesTFR.prediction. In case of DLcurve.plot.all if it si NULL, it is loaded from sim.dir.
country	Name or numerical code of a country.
burnin	Number of iterations to be discarded from the beginning of parameter traces.
pi	Probability interval. It can be a single number or an array.
tfr.max	Maximum TFR to be shown in the plot.
nr.curves	Number of curves to be plotted. If NULL, all curves are plotted.
predictive.dist	er en
	Logical. If TRUE, an error term is added to each trajectory.
ylim, xlab, yla	ab, main
	Graphical parameters passed to the plot function.
show.legend	Logical determining if the legend should be shown.
• • •	Additional graphical parameters. For DL curve.plot.all, $\dots$ contains also arguments pi, tfr.max and nr.curves.
sim.dir	Directory with the simulation results. Only relevant, if mcmc.list is NULL.
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".
verbose	Logical switching log messages on and off.

# **Details**

DLcurve.plot plots double logistic curves for the given country. DLcurve.plot.all creates such plots for all countries and stores them in output.dir. Parameters inputting the double logistic function are either thinned traces created by the tfr.predict function (if mcmc.list is an object of class bayesTFR.prediction), or they are selected by equal spacing from the MCMC traces. In the former case, burnin is set automatically; in the latter case, burnin defaults to 0. If nr.curves is smaller than 2000, the median and probability intervals are computed on a sample of 2000 equally spaced data points, otherwise on all plotted curves.

# Author(s)

Hana Sevcikova, Leontine Alkema

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# **Examples**

```
## Not run:
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
mcmc.set <- get.tfr.mcmc(sim.dir=sim.dir)
DLcurve.plot(country="Burkina Faso", mcmc.set, burnin=15)
## End(Not run)</pre>
```

get.country.object

Accessing Country Information

# **Description**

Function get.country.object returns an object containing country name, code and index. Functions get.countries.table return a data frame containing codes and names of all countries.

# Usage

```
get.country.object(country, meta = NULL, country.table = NULL, index = FALSE)
## S3 method for class 'bayesTFR.mcmc.set'
get.countries.table(object, ...)
## S3 method for class 'bayesTFR.prediction'
get.countries.table(object, ...)
```

# **Arguments**

country Country name, code or index. If it is an index, the argument index must be set

to TRUE.

meta Object of class bayesTFR.mcmc.meta. If it is not given, the argument country.table

must be given.

country.table A table containing columns "name" and "code" from which the country info can

be extracted. Only relevant, if meta is NULL.

index Logical determining if the argument country is an index.

object Object of class bayesTFR.mcmc.set or bayesTFR.prediction.

... Not used.

## **Details**

Given partial information about a country (i.e. having either name or code or index), the function get.country.object returns an object containing all three pieces of information. Only countries are considered that are included in the simulations and predictions. Country index is an internal index used in various components of a bayesTFR.mcmc.meta object.

get.cov.gammas 15

# Value

Function get.country.object returns a list with components:

name Country name code Country code index Country index

Function get.countries.table return a data frame with columns code and name.

## Author(s)

Hana Sevcikova

#### See Also

```
country.names
```

# **Examples**

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
m <- get.tfr.mcmc(sim.dir)
# all four calls should give the same answer
get.country.object('China', m$meta)
get.country.object(156, m$meta)
get.country.object(56, m$meta, index=TRUE)
get.country.object(156, NULL, country.table=get.countries.table(m))</pre>
```

get.cov.gammas

Covariance Matrices of Gamma Parameters

# **Description**

From a given MCMC, obtain a covariance matrix of the  $\gamma_{ci}$  (i=1,2,3) parameters for each country c.

# Usage

```
get.cov.gammas(mcmc.set = NULL, sim.dir = NULL, burnin = 200, chain.id = 1)
```

# **Arguments**

mcmc.set	Object of class bayesTFR.mcmc.set. If it is NULL, the sim.dir is used to load existing simulation results.
sim.dir	Directory with existing MCMC simulation results. It is only used if ${\tt mcmc.set}$ is NULL.
burnin	Number of burn-in iterations to be discarded from the beginning of the chain.
chain.id	Identifier of the MCMC to be used. By default the first chain is used.

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#### **Details**

In order to speed-up MCMC convergence, the package contains default values of gamma covariance that were obtained from a previous run (they can be loaded using data(proposal\_cov\_gammas\_cii)). However, this function allows to obtain new values and overwrite the default values by passing the resulting object to the run.tfr.mcmc function as the proposal\_cov\_gammas argument.

## Value

A list with components:

values An array of size  $nr_countries \times 3 \times 3$  containing values of the covariance matrix

of  $\gamma_{ci}$  (i = 1, 2, 3) for each country c.

country\_codes A vector of size nr\_countries. A covariance matrix values[i,,] corresponds

to a country with the code country\_codes[i].

# Author(s)

Leontine Alkema, Hana Sevcikova

#### See Also

```
run.tfr.mcmc
```

# **Examples**

```
## Not run:
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
cov.gammas <- get.cov.gammas(sim.dir=sim.dir, burnin=0)
m <- run.tfr.mcmc(nr.chains=1, iter=10, proposal_cov_gammas=cov.gammas, verbose=TRUE)
## End(Not run)</pre>
```

# **Description**

The function loads objects of class bayesTFR.convergence from disk.

# Usage

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```
get.tfr3.convergence.all(sim.dir = file.path(getwd(), "bayesTFR.output"))
```

# **Arguments**

sim.dir Simulation directory used for computing the diagnostics.

thin Thinning interval used with this diagnostics.
burnin Burnin used for computing the diagnostics.

#### **Details**

Function get.tfr.convergence loads an object of class bayesTFR.convergence for the specific thin and burnin generated for Phase II MCMCs. Function get.tfr.convergence.all loads all Phase II bayesTFR.convergence objects available for sim.dir. Functions get.tfr3.convergence and get.tfr3.convergence.all do the same thing for Phase III MCMCs.

# Value

```
get.tfr.convergence and get.tfr3.convergence return an object of class bayesTFR.convergence; get.tfr.convergence.all and get.tfr3.convergence.all return a list of objects of class bayesTFR.convergence.
```

#### Author(s)

Hana Sevcikova

#### See Also

bayesTFR.convergence, summary.bayesTFR.convergence.

get.tfr.mcmc Accessing MCMC Results

# **Description**

The function get.tfr.mcmc retrieves results of an MCMC simulation of Phase II and creates an object of class bayesTFR.mcmc.set. Function has.tfr.mcmc checks the existence of such results. Functions get.tfr3.mcmc and has.tfr3.mcmc do the same for Phase III MCMCs. Function tfr.mcmc extracts a single chain and tfr.mcmc.list extracts several or all chains from the simulation results.

# Usage

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```
get.tfr3.mcmc(sim.dir = file.path(getwd(), "bayesTFR.output"), ...)
has.tfr3.mcmc(sim.dir)

tfr.mcmc(mcmc.set, chain.id)

tfr.mcmc.list(mcmc.set, chain.ids=NULL)
```

# Arguments

sim.dir Directory where the simulation results are stored. chain.ids Chain identifiers in case only specific chains should be included in the resulting object. By default, all available chains are included. If FALSE full MCMC traces are loaded into memory. low.memory burnin Burnin used for loading traces. Only relevant, if low.memory=FALSE. Logical switching log messages on and off. verbose Chain identifier. chain.id mcmc.set Object of class bayesTFR.mcmc.set. Arguments passed to get.tfr.mcmc.

#### **Details**

Function get.tfr.mcmc is an accessor of results generated using run.tfr.mcmc and continue.tfr.mcmc. Function get.tfr3.mcmc can be used to access results generated using run.tfr3.mcmc and continue.tfr3.mcmc.

# Value

get.tfr.mcmc and get.tfr3.mcmc return an object of class bayesTFR.mcmc.set. has.tfr.mcmc and has.tfr3.mcmc return a logical value. tfr.mcmc returns an object of class bayesTFR.mcmc, and tfr.mcmc.list returns a list of bayesTFR.mcmc objects.

#### Author(s)

Hana Sevcikova

#### See Also

```
bayesTFR.mcmc.set
```

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
m <- get.tfr.mcmc(sim.dir)
summary(m)

# summary of the single chains
for(mc in tfr.mcmc.list(m)) print(summary(mc))</pre>
```

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```
get.tfr.parameter.traces
```

Accessing MCMC Parameter Traces

# Description

Functions for accessing traces of the MCMC parameters, either country-independent or country-specific. Functions get.tfr.parameter.traces and get.tfr.parameter.traces.cs access Phase II MCMCs; Functions get.tfr3.parameter.traces and get.tfr3.parameter.traces.cs access Phase III MCMCs.

## Usage

```
get.tfr.parameter.traces(mcmc.list, par.names = tfr.parameter.names(),
    burnin = 0, thinning.index = NULL, thin = NULL)

get.tfr.parameter.traces.cs(mcmc.list, country.obj,
    par.names=tfr.parameter.names.cs(),
    burnin=0, thinning.index=NULL, thin=NULL)

get.tfr3.parameter.traces(mcmc.list, par.names = tfr3.parameter.names(), ...)

get.tfr3.parameter.traces.cs(mcmc.list, country.obj,
    par.names=tfr3.parameter.names.cs(), ...)
```

# **Arguments**

mcmc.list	List of bayesTFR.mcmc objects.
country.obj	Country object list (see get.country.object).
par.names	Names of country-independent parameters (in case of get.tfr.parameter.traces) or country-specific parameters (in case of get.tfr.parameter.traces.cs) to be included.
burnin	Burnin indicating how many iterations should be removed from the beginning of each chain.
thinning.index	Index of the traces for thinning. If it is NULL, thin is used. thinning.index does not include burnin. For example, if there are two MCMC chains of length 1000, burnin=200 and we want a sample of length 400, then the value should be thinning.index=seq(1,1600, length=400).
thin	Alternative to thinning.index. In the above example it would be thin=4.
•••	Arguments passed to underlying functions (i.e. to get.tfr.parameter.traces or get.tfr.parameter.traces.cs).

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#### Value

All functions return a matrix with columns being the parameters and rows being the MCMC values, attached to one another in case of multiple chains. get.tfr.parameter.traces and get.tfr3.parameter.traces return country-independent parameters, get.tfr.parameter.traces.cs and get.tfr3.parameter.traces.cs return country-specific parameters.

## Author(s)

Hana Sevcikova

#### See Also

coda.list.mcmc for another way of retrieving parameter traces.

# **Examples**

get.tfr.prediction

Accessing a Prediction Object

## **Description**

Function get.tfr.prediction retrieves results of a prediction and creates an object of class bayesTFR.prediction. Function has.tfr.prediction checks an existence of such results.

# Usage

```
get.tfr.prediction(mcmc = NULL, sim.dir = NULL, mcmc.dir = NULL)
has.tfr.prediction(mcmc = NULL, sim.dir = NULL)
```

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#### **Arguments**

mcmc Object of class bayesTFR.mcmc.set used to make the prediction. It must corre-

spond to a Phase II MCMC. If it is NULL, the prediction is loaded from directory

given by sim.dir.

sim.dir Directory where the prediction is stored. It should correspond to the value of the

output.dir argument used in the tfr.predict function. Only relevant if mcmc

is NULL.

mcmc.dir Optional argument to be used only in a special case when the mcmc object con-

tained in the prediction object was estimated in different directory than in the one to which it points to (for example due to moving or renaming the original directory). The argument causes that the mcmc is redirected to the given

directory.

# **Details**

If mcmc is not NULL, the search directory is set to mcmc\$meta\$output.dir. This approach assumes that the prediction was stored in the same directory as the MCMC simulation, i.e. the output.dir argument of the tfr.predict function was set to NULL. If it is not the case, the argument mcmc.dir should be used.

# Value

Function has.tfr.prediction returns a logical indicating if a prediction exists for the given mcmc. Function get.tfr.prediction returns an object of class bayesTFR.prediction.

## Author(s)

Hana Sevcikova

# See Also

```
bayesTFR.prediction, tfr.predict, summary.bayesTFR.prediction
```

## **Examples**

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
pred <- get.tfr.prediction(sim.dir=sim.dir)
summary(pred, country="Canada")</pre>
```

# **Description**

Function for accessing TFR trajectories.

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# Usage

```
get.tfr.trajectories(tfr.pred, country)
```

# **Arguments**

tfr.pred Object of class bayesTFR.prediction.
country Name or numerical code of a country.

# **Details**

The function loads TFR trajectories for the given country from disk, offsets it if needed (see tfr.median.shift) and returns it as a matrix.

# Value

Array of size number of projection periods (including the present year) times the number of trajectories. The row names correspond to the mid-years of the prediction periods.

# Author(s)

Hana Sevcikova

#### See Also

```
bayesTFR.prediction, get.tfr.prediction, tfr.trajectories.table, tfr.median.shift
```

# **Examples**

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
pred <- get.tfr.prediction(sim.dir=sim.dir)
get.tfr.trajectories(pred, "Germany")</pre>
```

```
get.thinned.tfr.mcmc Creating and Accessing Thinned MCMCs
```

# Description

The function get.thinned.tfr.mcmc accesses a thinned and burned version of the given Phase II MCMC set. create.thinned.tfr.mcmc creates such a set.

# Usage

```
get.thinned.tfr.mcmc(mcmc.set, thin = 1, burnin = 0)
create.thinned.tfr.mcmc(mcmc.set, thin = 1, burnin = 0,
    output.dir = NULL, verbose = TRUE)
```

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# Arguments

mcmc.set	Object of class bayesTFR.mcmc.set of Phase II.
thin, burnin	Thinning interval and burnin used for creating or identifying the thinned object.
output.dir	Output directory. It is only used if the output goes to a non-standard directory.
verbose	Logical switching log messages on and off.

# **Details**

The function create.thinned.tfr.mcmc is called from tfr.predict and thus, the resulting object contains exactly the same MCMCs used for generating projections. In addition, it can be also called from tfr.diagnose if desired, so that the projection process can re-use such a set that lead to a convergence.

The thinning is done as follows: The given burnin is removed from the beginning of each chain in the original MCMC set. Then each chain is thinned by thin using equal spacing and all chains are collapsed into one single chain per parameter. They are stored in the main simulation directory under the name 'thinned\_mcmc\_t\_b' where *t* is the value of thin and *b* the value of burnin.

#### Value

Both functions return an object of class bayesTFR.mcmc.set. get.thinned.tfr.mcmc returns NULL if such object does not exist.

## Author(s)

Hana Sevcikova

# See Also

```
bayesTFR.mcmc.set, tfr.predict, tfr.diagnose
```

```
## Not run:
sim.dir <- tempfile()
m <- run.tfr.mcmc(nr.chains=2, iter=30, seed=1, output.dir=sim.dir, verbose=TRUE)
tfr.predict(m, burnin=15, use.tfr3=FALSE) # creates thinned MCMCs
mb <- get.thinned.tfr.mcmc(m, thin=1, burnin=15)
summary(mb, meta.only=TRUE) # length 30 = 2chains x (30-15)iters.
unlink(sim.dir, recursive=TRUE)
## End(Not run)</pre>
```

24 get.total.iterations

```
get.total.iterations Total Number of Iterations
```

# **Description**

Function get.total.iterations gives the total number of iterations of MCMCs summed over chains whith burnin being subtracted from each chain. Function get.stored.mcmc.length gives the total length of the MCMCs stored on disk minus those iterations that correspond to burnin. Result of the latter will be different from the former only if the MCMCs were run with value of thin larger than one.

# Usage

```
get.total.iterations(mcmc.list, burnin = 0)
get.stored.mcmc.length(mcmc.list, burnin = 0)
```

#### **Arguments**

mcmc.list List of bayesTFR.mcmc objects.

burnin Number of iterations to be subtracted from each chain.

#### Value

A single number.

#### Author(s)

Hana Sevcikova

## See Also

bayesTFR.mcmc

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
mcmc.set <- get.tfr.mcmc(sim.dir=sim.dir)
get.total.iterations(mcmc.set$mcmc.list) # 60=1chain x 60iters
get.total.iterations(mcmc.set$mcmc.list, burnin=20) # 40=1x(60-20)

## Not run:
sim.dir <- tempfile()
m <- run.tfr.mcmc(iter=10, nr.chains=2, output.dir=sim.dir, thin=5, verbose=TRUE)
get.total.iterations(m$mcmc.list) # 20=2x10
get.stored.mcmc.length(m$mcmc.list) # 6=2x3
unlink(sim.dir, recursive=TRUE)

## End(Not run)</pre>
```

include 25

include

**Inclusion Codes** 

# Description

Data sets containing codes that determine which countries are to be included into a simulation or/and projections.

## Usage

```
data(include_2008)
data(include_2010)
data(include_2012)
```

#### **Format**

Data frames containing one record per country or region. It has the following variables:

```
country Name of country or region. Not used.
```

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

**include\_code** Entries for which include\_code=2 are included in MCMC simulations (i.e. estimation of the model parameters). Entries for which include\_code is 1 or 2 are included in the prediction.

#### **Details**

In a simulation, an include\_\* dataset is selected that corresponds to the given wpp.year passed to the function run.tfr.mcmc. It is merged with a tfr dataset from the corresponding wpp package using the country\_code column. Thus, the country entries in this dataset should correspond to entries in the tfr dataset.

The package contains also a dataset called 'my\_tfr\_template' (in 'extdata' directory) which is a template for user-specified TFR time series. It has the same structure as the tfr dataset, except that most of the columns are optional. The only required column is country\_code (see description of the argument my.tfr.file in run.tfr.mcmc).

#### Source

Data provided by the United Nations Population Division.

```
data(include_2012)
head(include_2012)
```

run.tfr.mcmc

Running Markov Chain Monte Carlo for Parameters of Total Fertility Rate in Phase II

# Description

Runs (or continues running) MCMCs for simulating the total fertility rate of all countries of the world (phase II), using a Bayesian hierarchical model.

# Usage

```
run.tfr.mcmc(nr.chains = 3, iter = 62000,
   output.dir = file.path(getwd(), "bayesTFR.output"),
    thin = 1, replace.output = FALSE,
   start.year = 1750, present.year = 2010, wpp.year = 2012,
   my.tfr.file = NULL, buffer.size = 100,
   U.c.low = 5.5, U.up = 8.8, U.width = 3,
   mean.eps.tau0 = -0.25, sd.eps.tau0 = 0.4, nu.tau0 = 2,
   Triangle_c4.low = 1, Triangle_c4.up = 2.5,
   Triangle_c4.trans.width = 2,
   Triangle4.0 = 0.3, delta4.0 = 0.8, nu4 = 2,
   S.low = 3.5, S.up = 6.5, S.width = 0.5,
   a.low = 0, a.up = 0.2, a.width = 0.02,
   b.low = a.low, b.up = a.up, b.width = 0.02,
    sigma0.low = 0.01, sigma0.up = 0.6, sigma0.width = 0.1,
    sigma0.min = 0.001,
   const.low = 0.8, const.up = 2, const.width = 0.3,
   d.low = 0.05, d.up = 0.5, d.trans.width = 1,
   chi0 = -1.5, psi0 = 0.6, nu.psi0 = 2,
   alpha0.p = c(-1, 0.5, 1.5), delta0 = 1, nu.delta0 = 2,
   dl.p1 = 9, dl.p2 = 9,
   S.ini = NULL, a.ini = NULL, b.ini = NULL, sigma0.ini = NULL,
   Triangle_c4.ini = NULL, const.ini = NULL, gamma.ini = 1,
   proposal_cov_gammas = NULL,
    seed = NULL, parallel = FALSE, nr.nodes = nr.chains,
    save.all.parameters = FALSE, compression.type = 'None'
   auto.conf = list(max.loops=5, iter=62000, iter.incr=10000,
       nr.chains=3, thin=80, burnin=2000),
   verbose = FALSE, verbose.iter = 10, ...)
continue.tfr.mcmc(iter, chain.ids=NULL,
   output.dir=file.path(getwd(), "bayesTFR.output"),
   parallel = FALSE, nr.nodes = NULL, auto.conf = NULL,
   verbose=FALSE, verbose.iter = 10, ...)
```

# **Arguments**

nr.chains

Number of MCMC chains to run.

iter Number of iterations to run in each chain. In addition to a single value, it can have the value 'auto' in which case the function runs for the number of iterations given in the auto. conf list (see below), then checks if the MCMCs converged (using the auto.conf settings). If it did not converge, the procedure is repeated until convergence is reached or the number of repetition exceeded auto.conf\$max.loops. output.dir Directory which the simulation output should be written into. Thinning interval between consecutive observations to be stored on disk. thin replace.output If TRUE, existing outputs in output dir will be replaced by results of this simulation. Start year for using historical data. start.year End year for using historical data. present.year Year for which WPP data is used. The functions loads a package called wppx wpp.year where x is the wpp. year and uses the tfr\* datasets. my.tfr.file File name containing user-specified TFR time series for one or more countries. See Details below. buffer.size Buffer size (in number of iterations) for keeping data in the memory. The smaller the buffer. size the more often will the process access the hard disk and thus, the slower the run. On the other hand, the smaller the buffer. size the less data will be lost in case of failure. Lower and upper bound of the parameter  $U_c$ , the start level of the fertility tran-U.c.low, U.up sition. The lower bound is set for each country as the maximum of U.c.low and the minimum of historical TFR for that country. U.width Width for slice sampling used when updating the  $U_c$  parameter. mean.eps.tau0, sd.eps.tau0 Mean and standard deviation of the prior distribution of  $m_{\tau}$  which is the mean of the distortion terms  $\epsilon_{c,\tau}$  in start periods  $\tau_c$ . The shape parameter of the prior Gamma distribution of  $1/s_{\tau}^2$  is nu.tau0/2.  $s_{\tau}$ nu.tau0 is standard deviation of the distortion terms in start periods  $\tau_c$ . Triangle\_c4.low, Triangle\_c4.up Lower and upper bound of the  $\Delta_{c4}$  parameter. Triangle\_c4.trans.width Width for slice sampling used when updating the logit-transformed  $\Delta_{c4}$  parameter. Triangle4.0, delta4.0 Mean and standard deviation of the prior distribution of the  $\Delta_4$  parameter which is the hierarchical mean of the logit-transformed  $\Delta_{c4}$ . The shape parameter of the prior Gamma distribution of  $1/\delta_4^2$  is nu4/2.  $\delta_4$  is nu4 standard deviation of the logit-transformed  $\Delta_{c4}$ . Lower and upper bound of the uniform prior distribution of the S parameter S.low, S.up which is the TFR at which the distortion has maximum variance. S.width Width for slice sampling used when updating the S parameter. a.low, a.up Lower and upper bound of the uniform prior distribution of the a parameter

which is a coefficient for linear decrease of the TFR for TFR larger than S.

a.width	Width for slice sampling used when updating the <i>a</i> parameter.
b.low, b.up	Lower and upper bound of the uniform prior distribution of the $b$ parameter which is a coefficient for linear decrease of the TFR for TFR smaller than $S$ .
b.width	Width for slice sampling used when updating the $b$ parameter.
sigma0.low, sig	gma0.up
	Lower and upper bound of the uniform prior distribution of the $\sigma_0$ parameter. $\sigma_0^2$ is the maximum variance of the distortions at TFR equals $S$ .
sigma0.width	Width for slice sampling used when updating the $\sigma_0$ parameter.
sigma0.min	Minimum value that $\sigma_0$ can take.
const.low, cons	st.up
	Lower and upper bound of the uniform prior distribution of the $c$ parameter which is the multiplier of standard deviation of the distortions before 1975.
const.width	Width for slice sampling used when updating the $c$ parameter.
d.low, d.up	Lower and upper bound of the parameter $d_c$ , the maximum annual decrement for country $c$ . (Note that in Alkema et al. this parameter is a five-years decrement.)
d.trans.width	Width for slice sampling used when updating the logit-transformed $d_{c}$ parameter.
chi0, psi0	Prior mean and standard deviation of the $\chi$ parameter which is the hierarchical mean of logit-transformed maximum decline parameter $d_c$ .
nu.psi0	The shape parameter of the prior Gamma distribution of $1/\psi^2$ is nu.psi0/2. $\psi$ is the standard devation of logit-transformed maximum decline parameter $d_c$ .
alpha0.p	Vector of prior means of the $\alpha_i$ parameters, $i=1,2,3.$ $\alpha_i$ is the hierarchical mean of $\gamma_{ci}$ .
delta0	Prior standard deviation of the $\alpha_i$ parameters. It is a single value, i.e. the same standard deviation is used for all $i$ .
nu.delta0	The shape parameter of the prior Gamma distribution of $1/\delta_i^2$ is nu.delta0/2. $\delta_i$ is the standard deviation of $\gamma_{ci}$ .
dl.p1, dl.p2	Values of the parameters $p_1$ and $p_2$ of the double logistic function.
S.ini	Initial value for the $S$ parameter. It can be a single value or an array of the size nr.chains. By default, if nr.chains is 1, it is the middle point of the interval [S.low, S.up]. Otherwise, it is equally spaced distributed between S.low and S.up.
a.ini	Initial value for the $a$ parameter. It can be a single value or an array of the size nr.chains. By default, if nr.chains is 1, it is the middle point of the interval [a.low, a.up]. Otherwise, it is equally spaced distributed between a.low and a.up.
b.ini	Initial value for the $b$ parameter. It can be a single value or an array of the size nr.chains. By default, if nr.chains is 1, it is the middle point of the interval [b.low, b.up]. Otherwise, it is equally spaced distributed between b.low and b.up.
sigma0.ini	Initial value for the $\sigma_0$ parameter. It can be a single value or an array of the size nr.chains. By default, if nr.chains is 1, it is the middle point of the interval [sigma0.low, sigma0.up]. Otherwise, it is equally spaced distributed between sigma0.low and sigma0.up.

Triangle\_c4.ini

Initial value for the  $\Delta_{c4}$  parameter. It can be a single value or an array of the size nr.chains. By default, if nr.chains is 1, it is the middle point of the interval [Triangle\_c4.low, Triangle\_c4.up]. Otherwise, it is equally spaced distributed between Triangle\_c4.low and Triangle\_c4.up.

const.ini

Initial value for the c parameter. It can be a single value or an array of the size nr.chains. By default, if nr.chains is 1, it is the middle point of the interval [const.low, const.up]. Otherwise, it is equally spaced distributed between const.low and const.up.

gamma.ini Initial value for the  $\gamma_c$  parameter. The same value is used for all countries.

proposal\_cov\_gammas

Proposal for the gamma covariance matrices for each country. It should be a list with two values: values and country\_codes. The structure corresponds to the object returned by the function get.cov.gammas. By default the covariance matrices are obtained using data(proposal\_cov\_gammas\_cii). This argument overwrite the defaults for countries contained the argument.

seed

Seed of the random number generator. If NULL no seed is set. It can be used to generate reproducible results.

parallel

Logical determining if the simulation should run multiple chains in parallel. If it is TRUE, the package **snowFT** is required.

nr.nodes

Relevant only if parallel is TRUE. It gives the number of nodes for running the simulation in parallel. By default it equals to the number of chains.

save.all.parameters

If TRUE, the distortion terms  $\epsilon_{c,t}$  for all t are stored on disk, otherwise not.

compression.type

One of 'None', 'gz', 'xz', 'bz', determining type of a compression of the MCMC files.

auto.conf

List containing a configuration for an 'automatic' run (see description of argument iter). Item iter gives the number of iterations in the first chunk of the MCMC simulation; item iter.incr gives the number of iterations in the following chunks; nr.chains gives the number of chains in all chunks of the MCMC simulation; items thin and burnin are used in the convergence diagnostics following each chunk; max.loops controls the maximum number of chunks. All items must be integer values. This argument is only used if the function argument iter is set to 'auto'.

verbose Logical switching log messages on and off.

verbose.iter Integer determining how often (in number of iterations) log messages are outputted during the estimation.

Additional parameters to be passed to the function performParallel, if parallel is TRUE.

chain.ids Array of chain identifiers that should be resumed. If it is NULL, all existing chains in output.dir are resumed.

#### **Details**

The function run.tfr.mcmc creates an object of class bayesTFR.mcmc.meta and stores it in output.dir. It launches nr.chains MCMCs, either sequentially or in parallel. Parameter traces of each chain are stored as (possibly compressed) ASCII files in a subdirectory of output.dir, called mcx where x is the identifier of that chain. There is one file per parameter, named after the parameter with the suffix ".txt", possibly followed by a compression suffix if compression.type is given. Country-specific parameters  $(U,d,\gamma)$  have the suffix \_cy where y is the country code. In addition to the trace files, each mcx directory contains the object bayesTFR.mcmc in binary format. All chain-specific files are written into disk after the first, last and each buffer.size-th iteration.

Using the function continue.tfr.mcmc one can continue simulating an existing MCMCs by iter iterations for either all or selected chains.

The function loads observed data (further denoted as WPP dataset) from the tfr and tfr\_supplemental datasets in a wppx package where x is the wpp.year. It is then merged with the include dataset that corresponds to the same wpp.year. The argument my.tfr.file can be used to overwrite those default data. Such a file can include a subset of countries contained in the WPP dataset, as well as a set of new countries. In the former case, the function replaces the corresponding country data from the WPP dataset by values in this file. Only columns are replaced that match column names of the WPP dataset, and in addition, columns 'last.observed' and 'include\_code' are used, if present. Countries are merged with WPP using the column 'country\_code'. In addition, in order the countries to be included in the simulation, in both cases (whether they are included in the WPP dataset or not), they must be contained in the UN table of locations (UNlocations). In addition, their corresponding include\_code must be set to 2. If the column 'include\_code' is present in my.tfr.file, its value overwrites the default include code, unless is -1.

For simulation of the hyperparameters of the Bayesian hierarchical model, all countries are used that are included in the WPP dataset, possibly complemented by the my.tfr.file, that have include\_code equal to 2. The hyperparameters are used to simulate country-specific parameters, which is done for all countries with include\_code equal 1 or 2. The following values of include\_code in my.tfr.file are recognized: -1 (do not overwrite the default include code), 0 (ignore), 1 (include in prediction but not estimation), 2 (include in both, estimation and prediction). Thus, the set of countries included in the estimation and prediction can be fully user-specific.

Optionally, my.tfr.file can contain a column called last.observed containing the year of the last observation for each country. In such a case, the code would ignore any data after that time point. Furthermore, the function tfr.predict fills in the missing values using the median of the BHM procedure (stored in tfr\_matrix\_reconstructed of the bayesTFR.prediction object). For last.observed values that are below a middle year of a time interval  $[t_i, t_{i+1}]$  (computed as  $t_i+3$ ) the last valid data point is the time interval  $[t_{i-1}, t_i]$ , whereas for values larger equal a middle year, the data point in  $[t_i, t_{i+1}]$  is valid.

The package contains a dataset called 'my\_tfr\_template' (in 'extdata' directory) which is a template for user-specified my.tfr.file.

## Value

An object of class bayesTFR.mcmc.set which is a list with two components:

meta An object of class bayesTFR.mcmc.meta.

mcmc.list A list of objects of class bayesTFR.mcmc, one for each MCMC.

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# Author(s)

Hana Sevcikova, Leontine Alkema

#### References

L. Alkema, A. E. Raftery, P. Gerland, S. J. Clark, F. Pelletier, Buettner, T., Heilig, G.K. (2011). Probabilistic Projections of the Total Fertility Rate for All Countries. Demography, Vol. 48, 815-839. (A Working Paper version at http://www.csss.washington.edu/Papers [nr. 97])

# See Also

```
get.tfr.mcmc, summary.bayesTFR.mcmc.set.
```

# **Examples**

```
## Not run:
m <- run.tfr.mcmc(nr.chains=1, iter=5, verbose=TRUE)
summary(m)
m <- continue.tfr.mcmc(iter=5, verbose=TRUE)
summary(m)
## End(Not run)</pre>
```

run.tfr.mcmc.extra

Run MCMC for Extra Countries, Areas or Regions

# Description

Run MCMC for extra countries, areas or regions. It uses the posterior distribution of model hyper-parameters from an existing simulation to generate country-specific parameters.

# Usage

```
run.tfr.mcmc.extra(sim.dir = file.path(getwd(), "bayesTFR.output"),
    countries = NULL, my.tfr.file = NULL,
    iter = NULL, thin = 1, burnin = 2000,
    parallel = FALSE, nr.nodes = NULL, verbose = FALSE, verbose.iter = 100, ...)
```

# **Arguments**

sim.dir	Directory with an existing simulation.
countries	Vector of country codes. These include codes of areas and regions (see column country_code in UNlocations).
my.tfr.file	File name containing user-specified TFR time series for countries for which the simulation should run (see Details below).

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iter	Number of iterations to be used for sampling from the posterior distribution of the hyperparameters. By default, the number of iterations used in the existing simulation is taken.
thin	Thinning interval for sampling from the posterior distribution of the hyperparameters.
burnin	Number of iterations discarded before sampling from the posterior distribution of the hyperparameters. It is also used when computing proposal of gamma covariance matrices (see get.cov.gammas).
parallel	Logical determining if the simulation should run multiple chains in parallel.
nr.nodes	Relevant only if parallel is TRUE. It gives the number of nodes for running the simulation in parallel. By default it equals to the number of chains contained in the existing simulation.
verbose	Logical switching log messages on and off.
verbose.iter	Integer determining how often (in number of iterations) log messages are outputted during the estimation.
	Additional parameters to be passed to the function performParallel, if parallel is TRUE.

#### **Details**

The function can be used to make predictions for countries, areas or regions (further denoted as 'countries') that were not included in the MCMC estimation (invoked by run.tfr.mcmc). It creates MCMC traces for country-specific parameters. The purpose of this function is to have country-specific parameters available in order to be able to generate projections for additional countries or their aggregations, without having to re-run the often time-expensive MCMC simulation.

The set of countries to be considered by this function can be given either by their codes, using the argument countries, in which case the countries must be included in the UN WPP tfr dataset. Or, it can be given by a user-specific TFR file, using the argument my.tfr.file. The function considers a union of both arguments. The function will ignore all countries that were used in the existing MCMC simulation for estimating the hyperparameters. Countries that already own country-specific parameters (e.g. because they were included in my.tfr.file passed to run.tfr.mcmc) get their parameters recomputed. Note that all countries must be included in the UNlocations dataset, but unlike in run.tfr.mcmc, their include\_code is ignored.

# Value

An object of class bayesTFR.mcmc.set.

#### Note

If there is an existing projection for the directory sim.dir, use tfr.predict.extra to obtain projections for the extra countries used in this function.

# Author(s)

Hana Sevcikova, Leontine Alkema

#### See Also

```
run.tfr.mcmc,tfr.predict.extra
```

# **Examples**

```
## Not run:
sim.dir <- tempfile()
m <- run.tfr.mcmc(nr.chains=1, iter=20, output.dir=sim.dir, verbose=TRUE)
m <- run.tfr.mcmc.extra(sim.dir=sim.dir, countries=c(908,924), burnin=10, verbose=TRUE)
summary(m, country=924)
pred <- tfr.predict(m, burnin=10, use.tfr3=FALSE, verbose=TRUE)
summary(pred, country=908)
unlink(sim.dir, recursive=TRUE)
## End(Not run)</pre>
```

run.tfr3.mcmc

Running Markov Chain Monte Carlo for Parameters of Total Fertility Rate in Phase III

# **Description**

Runs (or continues running) MCMCs for simulating phase III total fertility rate, using a Bayesian hierarchical version of an AR(1) model.

# Usage

```
run.tfr3.mcmc(sim.dir, nr.chains = 3, iter = 50000, thin = 10,
    replace.output = FALSE, my.tfr.file = NULL, buffer.size = 100,
    use.extra.countries = FALSE,
   mu.prior.range = c(0, 2.1), rho.prior.range = c(0, 1 - .Machine double.xmin),
   sigma.mu.prior.range = c(1e-05, 0.318), sigma.rho.prior.range = c(1e-05, 0.289),
    sigma.eps.prior.range = c(1e-05, 0.5),
    mu.ini = NULL, mu.ini.range = mu.prior.range,
    rho.ini = NULL, rho.ini.range = rho.prior.range,
    sigma.mu.ini = NULL, sigma.mu.ini.range = sigma.mu.prior.range,
    sigma.rho.ini = NULL, sigma.rho.ini.range = sigma.rho.prior.range,
    sigma.eps.ini = NULL, sigma.eps.ini.range = sigma.eps.prior.range,
   seed = NULL, parallel = FALSE, nr.nodes = nr.chains, compression.type = "None",
   auto.conf = list(max.loops = 5, iter = 50000, iter.incr = 20000, nr.chains = 3,
                    thin = 60, burnin = 10000),
    verbose = FALSE, verbose.iter = 1000, ...)
continue.tfr3.mcmc(sim.dir, iter, chain.ids=NULL,
    parallel = FALSE, nr.nodes = NULL, auto.conf = NULL,
    verbose=FALSE, verbose.iter = 1000, ...)
```

# **Arguments**

sim.dir Directory with an existing simulation of phase II TFR (see run.tfr.mcmc).

nr. chains Number of MCMC chains to run.

iter Number of iterations to run in each chain. In addition to a single value, it can

have the value 'auto' in which case the function runs for the number of iterations given in the auto.conf list (see below), then checks if the MCMCs converged (using the auto.conf settings). If it did not converge, the procedure is repeated until convergence is reached or the number of repetition exceeded

auto.conf\$max.loops.

thin Thinning interval between consecutive observations to be stored on disk.

replace.output If TRUE, previously stored results of a phase III simulation will be overwritten.

my.tfr.file File name containing user-specified TFR time series for one or more countries.

See description of this argument in run.tfr.mcmc.

buffer.size Buffer size (in number of iterations) for keeping data in the memory.

use.extra.countries

By default, only countries are used in the MCMCs that were assigned for estimation (i.e. their 'include\_code' is 2 in the include) dataset and are in phase III at present time (argument present.year in run.tfr.mcmc). If this argument is TRUE, countries that were added using run.tfr.mcmc.extra and are in phase III are also included

TRUE, countries that were added using run.tfr.mcmc.extra and are in phase

III are also included.
mu.prior.range, rho.prior.range, sigma.mu.prior.range, sigma.rho.prior.range, sigma.eps.prior.range

Min and max for the prior (uniform) distribution of these paraemters.

mu.ini, rho.ini, sigma.mu.ini, sigma.rho.ini, sigma.eps.ini

Initial value(s) of the parameters. It can be a single value or an array of the size nr.chains. By default, if nr.chains is 1, it is the middle point of the corresponding range. Otherwise, it is uniformly randomly distributed within the range.

mu.ini.range, rho.ini.range, sigma.mu.ini.range, sigma.rho.ini.range, sigma.eps.ini.range
Min and max for the initial values.

will and max for the initial values.

seed Seed of the random number generator. If NULL no seed is set.

parallel Logical determining if the simulation should run multiple chains in parallel. If

it is TRUE, the package **snowFT** is required.

nr.nodes Relevant only if parallel is TRUE. It gives the number of nodes for running the

simulation in parallel.

compression.type

One of 'None', 'gz', 'xz', 'bz', determining type of a compression of the MCMC

auto.conf

List containing a configuration for an 'automatic' run (see description of argument iter). Item iter gives the number of iterations in the first chunk of the MCMC simulation; item iter.incr gives the number of iterations in the following chunks; nr.chains gives the number of chains in all chunks of the MCMC simulation; items thin and burnin are used in the convergence diagnostics following each chunk; max.loops controls the maximum number of chunks. All items must be integer values. This argument is only used if the function argument iter is set to 'auto'.

verbose	Logical switching log messages on and off.
verbose.iter	Integer determining how often (in number of iterations) messages are outputted during the estimation.
•••	Additional parameters to be passed to the function performParallel, if parallel is TRUE.
chain.ids	Array of chain identifiers that should be resumed. If it is NULL, all existing chains are resumed.

# **Details**

The MCMCs are stored in sim.dir in a subdirectory called "phaseIII". It has exactly the same structure as phase II MCMCs described in run.tfr.mcmc.

# Value

An object of class bayesTFR.mcmc.set which is a list with two components:

meta An object of class bayesTFR.mcmc.meta.

mcmc.list A list of objects of class bayesTFR.mcmc, one for each MCMC.

# Author(s)

Hana Sevcikova

# References

Raftery, A.E., Alkema, L. and Gerland, P. (2013). Bayesian Population Projections for the United Nations. Statistical Science, in press.

## See Also

```
run.tfr.mcmc, get.tfr3.mcmc
```

```
## Not run:
sim.dir <- tempfile()
# Runs Phase II MCMCs (must be run before Phase III)
m <- run.tfr.mcmc(nr.chains=1, iter=5, output.dir=sim.dir, verbose=TRUE)
# Runs Phase III MCMCs
m3 <- run.tfr3.mcmc(sim.dir=sim.dir, nr.chains=2, iter=50, thin=1, verbose=TRUE)
m3 <- continue.tfr3.mcmc(sim.dir=sim.dir, iter=10, verbose=TRUE)
summary(m3, burnin=10)
unlink(sim.dir, recursive=TRUE)
## End(Not run)</pre>
```

```
summary.bayesTFR.convergence
```

Summary of a TFR Convergence Object

# **Description**

Summary of an object of class bayesTFR. convergence created using the tfr.diagnose or tfr3.diagnose functions. It gives an overview about parameters that did not converge.

# Usage

```
## S3 method for class 'bayesTFR.convergence'
summary(object, expand = FALSE, ...)
```

# **Arguments**

object Object of class bayesTFR.convergence.

expand By default, the function does not show parameters for each country for which

there was no convergence, if the status is 'red'. This argument can switch that

option on.

... Not used.

# Author(s)

Hana Sevcikova

# See Also

```
tfr.diagnose, tfr3.diagnose
```

```
summary.bayesTFR.mcmc.set
```

Summary Statistics for TFR Markov Chain Monte Carlo Chains

# Description

Summary of an object bayesTFR.mcmc.set or bayesTFR.mcmc, computed via run.tfr.mcmc or run.tfr3.mcmc. It can be obtained either for all countries or for a specific country, and either for all parameters or for specific parameters. The function uses the summary.mcmc function of the **coda** package.

# Usage

```
## S3 method for class 'bayesTFR.mcmc.set'
summary(object, country = NULL, chain.id = NULL,
    par.names = NULL, par.names.cs = NULL, meta.only = FALSE,
    thin = 1, burnin = 0, ...)

## S3 method for class 'bayesTFR.mcmc'
summary(object, country = NULL, par.names = NULL, par.names.cs = NULL,
    thin = 1, burnin = 0, ...)
```

## **Arguments**

object	Object of class bayesTFR.mcmc.set or bayesTFR.mcmc.
country	Country name or code if a country-specific summary is desired. By default, summary for all countries is generated.
chain.id	Identifiers of MCMC chains. By default, all chains are considered.
par.names	Country independent parameters to be included in the summary. If the underlying object is an MCMC of phase II, the default names are given by tfr.parameter.names(), if it is phase III the names are tfr3.parameter.names().
par.names.cs	Country-specific parameters to be included in the summary. If the underlying object is an MCMC of phase II, the default names are given by tfr.parameter.names.cs(), if it is phase III the names are tfr3.parameter.names.cs().
meta.only	If it is TRUE, only meta information of the simulation is included.
thin	Thinning interval. Only used if larger than the thin argument used in run.tfr.mcmc or run.tfr3.mcmc.
burnin	Number of iterations to be discarded from the beginning of each chain before computing the summary.
	Additional arguments passed to the summary.mcmc function of the <b>coda</b> package.

# Author(s)

Hana Sevcikova

## See Also

```
bayesTFR.mcmc.set, summary.mcmc
```

# **Examples**

```
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
m <- get.tfr.mcmc(sim.dir)
summary(m, country="Czech Republic", burnin=15)</pre>
```

```
summary.bayesTFR.prediction
```

Summary of a Prediction of the Total Fertility Rate

# **Description**

Country-specific summary of an object of class bayesTFR.prediction, created using the function tfr.predict. The summary contains the mean, standard deviation and several commonly used quantiles of the simulated trajectories.

# Usage

```
## S3 method for class 'bayesTFR.prediction'
summary(object, country=NULL, compact = TRUE, ...)
```

## **Arguments**

object Object of class bayesTFR.prediction.

country Country name or code. If it is NULL, only prediction parameters are included.

compact Logical switching between a smaller and larger number of displayed quantiles.

. . . A list of further arguments.

## Author(s)

Hana Sevcikova

## See Also

```
bayesTFR.prediction
```

## **Examples**

39 tfr.diagnose

tfr.diagnose Conve	rgence Diagnostics of TFR Markov Chain Monte Carlo

**Description** 

Functions tfr.diagnose and tfr3.diagnose run convergence diagnostics of existing TFR MCMCs for phase II and phase III, respectively, using the raftery diag function from the coda package. has.mcmc.converged checks if the existing diagnostics converged.

#### Usage

```
tfr.diagnose(sim.dir, thin = 80, burnin = 2000, express = FALSE,
    country.sampling.prop = NULL, keep.thin.mcmc=FALSE, verbose = TRUE)
tfr3.diagnose(sim.dir, thin = 60, burnin = 10000, express = TRUE,
    country.sampling.prop = NULL, verbose = TRUE, ...)
has.mcmc.converged(diag)
```

## **Arguments**

sim.dir Directory with the MCMC simulation results.

Thinning interval. thin

burnin Number of iterations to be discarded from the beginning of the parameter traces.

express Logical. If TRUE, the convergence diagnostics is run only on the country-independent

parameters. If FALSE, the country-specific parameters are included in the diag-

nostics. The number of countries can be controlled by country. sampling.prop.

country.sampling.prop

Proportion of countries that are included in the diagnostics. If it is NULL and express=FALSE, all countries are included. Setting here a number between 0 and 1, one can limit the number of countries which are then randomly sampled. Note that for long MCMCs, this argument may significantly influence the run-

time of this function.

keep.thin.mcmc Logical. If TRUE the thinned traces used for computing the diagnostics are

stored on disk (see create.thinned.tfr.mcmc). It is only available for phase

II MCMCs.

verbose Logical switching log messages on and off. Object of class bayesTFR. convergence. diag

Not used. . . .

#### **Details**

The diagnose functions invoke the tfr.raftery.diag (or tfr3.raftery.diag) function separately for country-independent parameters and for country-specific parameters. It results in two possible states: red, i.e. it did not converge, and green, i.e. it converged. The resulting object from 40 tfr.diagnose

tfr.diagnose is stored in

'{sim.dir}/diagnostics/bayesTFR.convergence\_{thin}\_{burnin}.rda' and can be accessed using the function get.tfr.convergence. Function tfr3.diagnose stores its result into '{sim.dir}/phaseIII/diagnostics/bayesTFR.convergence\_{thin}\_{burnin}.rda' which can be accessed via get.tfr3.convergence.

#### Value

has.mcmc.converged returns a logical value determining if there is convergence or not.

tfr.diagnose and tfr3.diagnose return an object of class bayesTFR.convergence with components:

result Table containing all not-converged parameters. Its columns include 'Total itera-

tions needed' and 'Remaining iterations'.

lresult.country.independent

Number of rows in result that correspond to country-independent paramters.

These rows are groupped at the beginning of the table.

country.independent

Result of tfr.raftery.diag processed on country-independent parameters.

country.specific

Result of tfr.raftery.diag processed on country-specific parameters.

iter.needed Number of additional iterations suggested in order to achieve convergence.

iter.total Total number of iterations of the original unthinned set of chains.

use.nr.traj Suggestion for number of trajectories in generating predictions.

burnin Burnin used.

thin Thinning interval used.

status Vector of character strings containing the result status. Possible values: 'green',

'red'.

mcmc.set Object of class bayesTFR.mcmc.set that corresponds to the original set of MCMCs

on which the diagnostics was run.

thin.mcmc If keep.thin.mcmc is TRUE, it is an object of class bayesTFR.mcmc.set that

corresponds to the thinned mcmc set on which the diagnostics was run, other-

wise NULL.

express Value of the input argument express.

nr.countries Vector with elements used - number of countries used in this diagnostics, and

total - number of countries that this mcmc.set object was estimated on.

#### Author(s)

Hana Sevcikova, Leontine Alkema, Adrian Raftery

# See Also

tfr.raftery.diag, raftery.diag, summary.bayesTFR.convergence, get.tfr.convergence, create.thinned.tfr.mcmc

tfr.dl.coverage 41

tfr.dl.coverage	Goodness of Fit of the Double Logistic Function	
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# **Description**

The function computes coverage, i.e. the ratio of observed data fitted within the given probability intervals of the predictive posterior distribution of the double logistic function, as well as the root mean square error and mean absolute error of the simulation.

## Usage

```
tfr.dl.coverage(sim.dir, pi = c(80, 90, 95), burnin = 2000, verbose = TRUE)
```

# Arguments

sim.dir	Directory with the MCMC simulation results. If a prediction and its corresponding thinned MCMCs are available in the simulation directory, those are taken for assessing the goodness of fit.
pi	Probability interval. It can be a single number or an array.
burnin	Burnin. Only relevant if sim. dir does not contain thinned chains.
verbose	Logical switching log messages on and off.

# Value

time.rmse

country.rmse

List with the following components:

total.coverage	Vector of the coverage, one element per probability interval. For each pi, it is the ratio of the number of observed data points that fall within the probability interval of the posterior distribution over the total number of data points, i.e. TFR for all countries and historical time periods.
time.coverage	Matrix corresponding to the coverage computed per time period. (Rows correspond to probability intervals, columns correspond to time.) It is derived like total.coverage except that both, the nominator and denominator, contain only data points belonging to the corresponding time period.
country.coverag	ge
	Matrix corresponding to the coverage computed per country. (Rows correspond to probability intervals, columns correspond to countries.) It is derived like total.coverage except that both, the nominator and denominator, contain only data points belonging to the corresponding country.
total.rmse	Root mean square error as $\sqrt{(1/n\sum(x-m)^2)}$ where $x$ are observed data points, $m$ is the mean of the posterior distribution and $n$ is the number of data points. Here the sum is taken over all countries and historical time periods.

Like total.rmse except that each time period is considered separately. Like total.rmse except that each country is considered separately. 42 tfr.map

total.mae	Mean absolute error as $1/n \sum  x-m $ where $x$ are observed data points, $m$ is the median of the posterior distribution and $n$ is the number of data points. Here the sum is taken over all countries and historical time periods.
time.mae	Like total.mae except that each time period is considered separately.
country.mae	Like total.mae except that each country is considered separately.
pred.cdf	$T \times C$ matrix (with $T$ being the number of time periods and $C$ being the number of countries), containing the predictive CDF of the observation, i.e. the quantile of each data point within the predictive posterior distribution.
n	$0$ -1 $T \times C$ matrix indicating if the corresponding data point was included in the goodness of fit computation. Zeros indicate missing historical values.

## Note

To see the fit visually per country, use DLcurve.plot(..., predictive.distr=TRUE,...).

# Author(s)

Hana Sevcikova

## See Also

```
DLcurve.plot
```

# **Examples**

```
## Not run:
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
tfr <- get.tfr.mcmc(sim.dir)
# Note that this simulation is a toy example and thus has not converged.
gof <- tfr.dl.coverage(sim.dir)
gof$time.coverage
DLcurve.plot(tfr, country=608, predictive.distr=TRUE, pi=c(80, 90, 95))
## End(Not run)</pre>
```

tfr.map

TFR World Map

# Description

Generates a world map of the total fertility rate for given projection period and quantile. In addition, country specific Phase II MCMC parameters can be projected into the world map.

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## Usage

```
tfr.map(pred, quantile = 0.5,
    year = NULL, par.name = NULL, adjusted = FALSE,
    projection.index = 1, device = "dev.new", main = NULL,
    resolution=c("coarse","low","less islands","li","high"),
    device.args = NULL, data.args = NULL, ...)

tfr.map.all(pred, output.dir, output.type = "png",
    tfr.range = NULL, nr.cats = 50, same.scale = TRUE,
    quantile = 0.5, file.prefix='TFRwrldmap_', ...)

get.tfr.map.parameters(pred, tfr.range = NULL,
    nr.cats = 50, same.scale = TRUE, quantile = 0.5, ...)

tfr.map.gvis(pred, year = NULL, quantile = 0.5, pi = 80,
    par.name = NULL, adjusted = FALSE, ...)
```

#### **Arguments**

pred	Object of class bayesTFR.prediction.
------	--------------------------------------

quantile Quantile for which the map should be generated. It must be equal to one of the

values in dimnames(pred $\quantiles[[2]]$ ), 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.

year Year to be plotted. It can be a year within a projection period or a year within an

estimation period. In the latter case, the observed data are plotted. If not given,

projection.index determines the projection year.

par. name Name of a country-specific parameter to be plotted. If NULL, the TFR is plotted.

Allowed values are any of those returned by tfr.parameter.names.cs.extended()

and 'lambda' (see Details).

adjusted Logical indicating if the measure to be plotted is based on adjusted TFRs.

projection.index

Index of the projection to be displayed. It is only relevant if year is NULL. projection.index=1 means the present year, projection.index=2 means the

first projection period after present year, etc..

device Device for displaying the map. It is passed to the mapDevice function of the

rworldmap package. If it is equal to 'dev.cur', the current device is used. Oth-

erwise, it can be 'dev.new', 'png', 'pdf' etc.

main Title for the map. If it is NULL, a default title is constructed from the projection

year and quantile.

resolution Map resolution as implemented in getMap. High resolution requires the **rworldx**-

tra package.

device.args List of additional arguments to be passed to the mapDevice function of the

rworldmap package.

data.args List of additional arguments to be passed to the underlying data retrieving func-

tion.

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output.dir Directory into which resulting maps are stored.

output.type Type of the resulting files. It can be "png", "pdf", "jpeg", "bmp", "tiff", or

"postscript".

tfr.range Range of the total fertility rate to be displayed. It is of the form c(tfr.min,

tfr.max). By default, the whole range is considered. Note that countries with

values outside of the given range will appear white.

nr.cats Number of color categories.

same.scale Logical controlling if maps for all projection years of this prediction object

should be on the same color scale.

file.prefix Prefix for file names.

.. Arguments passed to the mapCountryData function of the **rworldmap** pack-

age. In case of tfr.map.gvis these are passed to the underlying data retrieving

function (the same as data.args).

pi Probability interval to be shown when a country is selected in an interactive map.

The corresponding quantiles must be available (see argument quantile above).

#### **Details**

tfr.map creates a single map for a given projection period and quantile using the **rworldmap** package. tfr.map.all generates a sequence of such maps, namely one for each projection period. If the package **fields** is installed, a color bar legend at the botom of the map is created.

Function get.tfr.map.parameters can be used in combination with tfr.map. (Note that get.tfr.map.parameters is called from inside of tfr.map.all.) It sets breakpoints for the color scheme using quantiles of a fitted gamma distribution.

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

By default, both tfr.map and tfr.map.gvis produce maps of TFRs. Alternatively, the functions can be used to plot country-specific Phase II MCMC parameters into a world map. They are given by the argument par.name. In addition to the MCMC parameters, if par.name='lambda', the period of the end of TFR decline (i.e. start of Phase III) is computed for each country and projected into the map. In such a case, we recommend to adjust the color scale in tfr.map e.g. using the arguments catMethod='pretty' and numCats=20 (see mapCountryData).

## Value

get.tfr.map.parameters returns a list with elements:

pred The object of class bayesTFR. prediction used in the function.

quantile Value of the argument quantile.

catMethod If the argument same scale is TRUE, this element contains breakpoints for cat-

egorization. It is generated from a fitted gamma distribution. Otherwise, it is

NULL.

numCats Number of categories.

coulourPalette Subset of the rainbow palette, starting from dark blue and ending at red.

... Additional arguments passed to the function.

tfr.median.set 45

## Author(s)

Hana Sevcikova, Patrick Gerland, Adrian Raftery

## **Examples**

```
## Not run:
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
pred <- get.tfr.prediction(sim.dir=sim.dir)
# Uses heat colors and seven categories by default
tfr.map(pred)
# Uses more colors with more suitable categorization
params <- get.tfr.map.parameters(pred)
do.call("tfr.map", params)
# Another projection year on the same scale
do.call("tfr.map", c(list(year=2043), params))
# Using Google Vizualization tool
tfr.map.gvis(pred)
## End(Not run)</pre>
```

tfr.median.set

Editing Medians of the Projection

# **Description**

These functions are to be used by expert analysts. They allow to change the projection medians either to specific values or shift the medians by a given constant, or by a specific adjusting procedure.

# Usage

```
tfr.median.set(sim.dir, country, values, years = NULL)

tfr.median.shift(sim.dir, country, reset = FALSE, shift = 0,
    from = NULL, to = NULL)

tfr.median.adjust(sim.dir, countries, factor1 = 2/3, factor2 = 1/3, forceAR1 = FALSE)

tfr.median.reset(sim.dir, countries)
```

# **Arguments**

sim.dir Directory containing the prediction object.

country Name or numerical code of a country.

countries Vector of country names or codes.

values Array of the new median values.

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	years	Numeric vector giving years which values correspond to. Ideally it should be of the same length as values. If it is NULL, values are set starting from the first prediction period. If values correspond to consecutive years, only the first year might be given here. A year $t$ represents a prediction period $[t_i, t_{i+1}]$ if $t_i < t \le t_{i+1}$ .
	reset	Logical. If TRUE medians in a range of from and to are reset to their original values.
	shift	Constant by which the medians should be offset. It is not used if reset is TRUE.
	from	Year from which the offset/reset should start. By default, it starts at the first prediction period.
	to	Year until which the offset/reset should be done. By default, it is set to the last prediction period.
factor1, factor2		
		Adjusting factors for the first and second projection period, respectively (see below).
	forceAR1	Logical. If TRUE, the given countries are forced to enter Phase III (i.e. the $AR(1)$ process) in the first projection period.

## **Details**

The function tfr.median.set can be used to set the medians of the given country to specific values. Function tfr.median.shift can be used to offset the medians by a specific constant, or to reset the medians to their original BHM values. Function tfr.median.adjust runs the prediction procedure for the given countries with an additional decrement in the model in the first two projection periods. In the first projection period it is computed as factor1\*S where S is a difference between observed decrement and the expected decrement (by the double logistic function) in the last observed period. In the second projection period, in the formula factor1 is replaced by factor2. If forceAR1 is set to TRUE, we recommend to set factor1 and factor2 to 0. The function then calls tfr.median.set in order to store the new median for each country. Function tfr.median.reset resets medians of the given countries to the original values.

In all four functions, if a median is modified, the corresponding offset is stored in the prediction object (element median.shift) and the updated prediction object is written back to disk. All functions in the package that use trajectories and trajectory statistics use the median.shift values to offset the results correspondingly.

#### Value

All three functions return an updated object of class bayesTFR.prediction.

#### Author(s)

Hana Sevcikova, Leontine Alkema

tfr.parameter.names 47

tfr.parameter.names Accessing Parameter Names

# **Description**

Functions for accessing names of the MCMC parameters, either country-independent or country-specific.

#### Usage

```
tfr.parameter.names(trans = NULL)
tfr.parameter.names.cs(country.code = NULL, trans = NULL)
tfr.parameter.names.extended()
tfr.parameter.names.cs.extended(country.code = NULL)

tfr3.parameter.names()
tfr3.parameter.names.cs(country.code = NULL)
```

#### **Arguments**

trans It can be NULL or logical. If TRUE, names of the transformable parameters (i.e.

'alpha' and 'delta' in case of country-independent parameters, or 'gamma' in case of country-specific parameters) are replaced by the names of the transformed parameters (i.e. 'alphat', 'deltat', or 'gammat'). If trans=FALSE, there is no such replacement. If trans=NULL, all parameter names, including the

transformable parameters are returned.

country.code Country code. If it is given, the country-specific parameter names contain the

postfix ' $\_cx$ ' where x is the country.code.

## Value

tfr.parameter.names returns names of the country-independent Phase II parameters.

tfr.parameter.names.cs returns names of the country-specific Phase II parameters.

tfr.parameter.names.extended returns names of all country-independent Phase II parameters, including the transformed parameters. Parameters 'alpha', 'delta', 'alphat', and 'deltat' are in their extended format with the postfix '\_1', '\_2' and '\_3'.

tfr.parameter.names.cs.extended returns names of all country-specific Phase II parameters, including the transformed parameters. Parameters 'gamma' and 'gammat' are in their extended format with the postfix '\_1', '\_2' and '\_3'.

tfr3.parameter.names returns names of the country-independent Phase III parameters.

tfr3.parameter.names.cs returns names of the country-specific Phase III parameters.

#### Author(s)

Hana Sevcikova

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## **Examples**

```
tfr.parameter.names()
tfr.parameter.names.extended()
tfr.parameter.names.cs()
tfr.parameter.names.cs.extended()
tfr3.parameter.names()
tfr3.parameter.names.cs()
```

tfr.pardensity.plot Plotting MCMC Parameter Density

# Description

Functions for plotting density of the posterior distribution of the MCMC parameters.

## Usage

```
tfr.pardensity.plot(mcmc.list = NULL,
   sim.dir = file.path(getwd(), "bayesTFR.output"),
   chain.ids = NULL, par.names = tfr.parameter.names(trans = TRUE),
   burnin = NULL, dev.ncol=5, low.memory = TRUE, ...)
tfr.pardensity.cs.plot(country, mcmc.list=NULL,
    sim.dir=file.path(getwd(), "bayesTFR.output"),
   chain.ids=NULL, par.names=tfr.parameter.names.cs(trans=TRUE),
   burnin=NULL, dev.ncol=3, low.memory=TRUE, ...)
tfr3.pardensity.plot(mcmc.list = NULL,
    sim.dir = file.path(getwd(), "bayesTFR.output"),
   chain.ids = NULL, par.names = tfr3.parameter.names(),
   burnin = NULL, dev.ncol=3, low.memory = TRUE, ...)
tfr3.pardensity.cs.plot(country, mcmc.list=NULL,
    sim.dir=file.path(getwd(), "bayesTFR.output"),
    chain.ids=NULL, par.names=tfr3.parameter.names.cs(),
   burnin=NULL, dev.ncol=2, low.memory=TRUE, ...)
```

## Arguments

country	Name or numerical code of a country.
mcmc.list	List of bayesTFR.mcmc objects, or an object of class bayesTFR.mcmc.set or of class bayesTFR.prediction (allowed only for Phase II MCMCs). If it is NULL, the parameter values are loaded from sim.dir.
sim.dir	Directory with the MCMC simulation results. It is only used if mcmc.list is NULL.

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chain.ids	List of MCMC identifiers to be plotted. If it is NULL, all chains found in mcmc.list or sim.dir are plotted.
par.names	Names of parameters for which density should be plotted. By default all (possibly transformed) country-independent parameters are plotted if used within tfr.pardensity.plot and tfr3.pardensity.plot, or country-specific parameters are plotted if used within tfr.pardensity.cs.plot and tfr3.pardensity.cs.plot.
burnin	Number of iterations to be discarded from the beginning of each chain.
dev.ncol	Number of column for the graphics device. If the number of parameters is smaller than dev.ncol, the number of columns is automatically decreased.
low.memory	Logical indicating if the processing should run in a low-memory mode. If it is FALSE, traces of all available parameters are loaded into memory. Otherwise, parameters are loaded as they are needed and are not kept in the memory.
	Further arguments passed to the density function.

#### Details

The functions plot the density of the posterior distribution either for country-independent parameters (tfr.pardensity.plot for phase II MCMCs and tfr3.pardensity.plot for phase III MCMCs) or for country-specific parameters (tfr.pardensity.cs.plot for phase II and tfr3.pardensity.cs.plot for phase III), one graph per parameter. One can restrict it to specific chains by setting the chain.ids argument and to specific parameters by setting the par.names argument.

If mcmc.list is an object of class bayesTFR.prediction (which is allowed in tfr.pardensity.plot and tfr.pardensity.cs.plot only) and if this object contains thinned traces, they are used instead of the full chains. In such a case, burnin and chain.ids cannot be modified - their value is set to the one used when the thinned traces were created, namely when running tfr.predict. In a situation with long MCMC chains, this approach can significantly speed-up creation of the density plots.

## Author(s)

Hana Sevcikova

## See Also

```
tfr.partraces.plot
```

#### **Examples**

```
## Not run:
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
tfr.pardensity.plot(sim.dir=sim.dir)
tfr.pardensity.cs.plot(country="Ireland", sim.dir=sim.dir, bw=0.2)
## End(Not run)</pre>
```

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tfr.partraces.plot Plotting MCMC Parameter Traces

## **Description**

Functions for plotting the MCMC parameter traces.

# Usage

```
tfr.partraces.plot(mcmc.list = NULL,
   sim.dir = file.path(getwd(), "bayesTFR.output"), chain.ids = NULL,
   par.names = tfr.parameter.names(trans = TRUE),
   nr.points = NULL, dev.ncol=5, low.memory = TRUE, ...)
tfr.partraces.cs.plot(country, mcmc.list = NULL,
   sim.dir = file.path(getwd(), "bayesTFR.output"), chain.ids = NULL,
   par.names = tfr.parameter.names.cs(trans = TRUE),
   nr.points = NULL, dev.ncol=3, low.memory = TRUE, ...)
tfr3.partraces.plot(mcmc.list = NULL,
   sim.dir = file.path(getwd(), "bayesTFR.output"), chain.ids = NULL,
   par.names = tfr3.parameter.names(),
   nr.points = NULL, dev.ncol=3, low.memory = TRUE, ...)
tfr3.partraces.cs.plot(country, mcmc.list = NULL,
   sim.dir = file.path(getwd(), "bayesTFR.output"), chain.ids = NULL,
   par.names = tfr3.parameter.names.cs(),
   nr.points = NULL, dev.ncol=2, low.memory = TRUE, ...)
```

# **Arguments**

country	Name or numerical code of a country.
mcmc.list	List of bayesTFR.mcmc objects, or an object of class bayesTFR.mcmc.set or of class bayesTFR.prediction (allowed only for Phase II MCMCs). If it is NULL, the traces are loaded from sim.dir.
sim.dir	Directory with the MCMC simulation results. It is only used if mcmc.list is NULL.
chain.ids	List of MCMC identifiers to be plotted. If it is NULL, all chains found in mcmc.list or sim.dir are plotted.
par.names	Names of parameters for which traces should be plotted. By default all (possibly transformed) country-independent parameters are plotted if used within tfr.partraces.plot and tfr3.partraces.plot, or country-specific parameters are plotted if used within tfr.partraces.cs.plot and tfr3.partraces.cs.plot.
nr.points	Number of points to be plotted. If NULL, all points are plotted, otherwise the traces are thinned evenly.

dev.ncol	Number of column for the graphics device. If the number of parameters is smaller than dev.ncol, the number of columns is automatically decreased.
low.memory	Logical indicating if the processing should run in a low-memory mode. If it is FALSE, traces of all available parameters are loaded into memory. Otherwise, parameters are loaded as they are needed and are not kept in the memory.
	Additional graphical arguments.

## **Details**

The functions plot MCMC traces either for country-independent parameters (tfr.partraces.plot for phase II MCMCs and tfr3.partraces.plot for phase III MCMCs) or for country-specific parameters (tfr.partraces.cs.plot for phase II MCMCs and tfr3.partraces.cs.plot for phase III MCMCs), one graph per parameter. One can restrict it to specific chains by setting the chain.ids argument, and to specific parameters by setting the par.names argument.

## Author(s)

Hana Sevcikova

#### See Also

coda.list.mcmc for retrieving raw values of the traces.

## **Examples**

```
## Not run:
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
tfr.partraces.plot(sim.dir=sim.dir)
tfr.partraces.cs.plot(country="Netherlands", sim.dir=sim.dir)
## End(Not run)</pre>
```

tfr.predict

Generating Posterior Trajectories of the Total Fertility Rate

## **Description**

Using the posterior parameter samples simulated by run.tfr.mcmc (and possibly run.tfr3.mcmc) the function generates posterior trajectories for the total fertility rate for all countries of the world.

# Usage

```
tfr.predict(mcmc.set = NULL, end.year = 2100,
    sim.dir = file.path(getwd(), "bayesTFR.output"),
    replace.output = FALSE, start.year = NULL,
    nr.traj = NULL, thin = NULL, burnin = 2000,
    use.diagnostics = FALSE, use.tfr3 = TRUE, burnin3 = 10000,
    mu = 2.1, rho = 0.8859, sigmaAR1 = 0.1016,
```

```
use.correlation = FALSE, save.as.ascii = 1000, output.dir = NULL,
low.memory = TRUE, seed = NULL, verbose = TRUE, ...)
```

## **Arguments**

mcmc.set Object of class bayesTFR.mcmc.set corresponding Phase II MCMCs. If it is

NULL, the object is loaded from the directory given by sim.dir.

end. year End year of the prediction.

sim.dir Directory with the MCMC simulation results. It should equal to the output.dir

argument in run.tfr.mcmc.

replace.output Logical. If TRUE, existing predictions in output.dir will be replaced by results

of this run.

start year Start year of the prediction. By default the prediction is started at the next time

period after present. year set in the estimation step. If start. year is smaller than the default, projections for countries and time periods that have data avail-

able after start. year are set to those data.

nr.traj Number of trajectories to be generated. If NULL, the argument thin is taken to

determine the number of trajectories. If both are NULL, the number of trajectories

corresponds to the size of the parameter sample.

thin Thinning interval used for determining the number of trajectories. Only relevant,

if nr.traj is NULL.

burnin Number of iterations to be discarded from the beginning of the parameter traces.

use.diagnostics

Logical determining if an existing convergence diagnostics for phase II MCMCs should be used for choosing the values of thin and burnin. In such a case, arguments nr.traj, thin and burnin are ignored. The 'best' values are chosen from results of running the tfr.diagnose function. Only diagnostics can be used that suggest a convergence of the underlying MCMCs. If there are more than one such objects, the one is chosen whose recommendation for the number

of trajectories is larger and closest to 2000.

use.tfr3 Logical determining if phase III should be predicted via MCMCs (simulated via

run.tfr3.mcmc) or a classic AR(1) process. If TRUE but no phase III MCMCs were simulated, a warning is given and the prediction switches automatically to

a classic AR(1) process.

burnin3 Burnin used for phase III MCMCs. Only relevant if use.tfr3 is TRUE.

save.as.ascii Either a number determining how many trajectories should be converted into an

ASCII file, or "all" in which case all trajectories are converted. It should be set

to 0, if no conversion is desired.

output.dir Directory into which the resulting prediction object and the trajectories are stored.

If it is NULL, it is set to either sim.dir, or to output.dir of mcmc.set\$meta if

mcmc.set is given.

low.memory Logical indicating if the prediction should run in a low-memory mode. If it is

FALSE, the whole traces of all parameters, including the burnin, are loaded into memory. Otherwise, burnins are discarded and parameters are loaded as they

are needed and are not kept in the memory.

mu Long-term mean  $\mu$  in the AR(1) projection model. Only used if use.tfr3 is

FALSE.

rho Autoregressive parameter  $\rho$  in AR(1) projection model. If it is NULL it is esti-

mated from the data. Only used if use.tfr3 is FALSE.

sigmaAR1 Standard deviation s of AR(1) distortion terms in short-term projections. If it

is NULL it is estimated from the data. It can be a single value or a vector giving the standard deviations for single projections. If the vector is shorter than the number of projections simulated via the AR(1) process, the last value is repeated for the remaining projections. In case of a single value (default), the standard deviation is kept constant over all AR(1) projections. Only used if use.tfr3 is

FALSE.

use.correlation

Logical. If TRUE the model errors are sampled jointly for all countries (Fosdick

and Raftery, 2012).

seed Seed of the random number generator. If NULL no seed is set. It can be used to

generate reproducible projections.

verbose Logical switching log messages on and off.

... Further arguments passed to the underlying functions.

#### **Details**

The trajectories are generated using a distribution of country-specific decline curves (Alkema et al 2011) and either a classic AR(1) process or a country-specific AR(1) process (Raftery et al 2013). Phase II parameter samples simulated using run.tfr.mcmc are used from all chains, from which the given burnin was discarded. They are evenly thinned to match nr.traj or using the thin argument. Such thinned parameter traces, collapsed into one chain, if they do not already exist, are stored on disk into the sub-directory '{thinned\_mcmc\_t\_b' where t is the value of thin and b the value of burnin (see create.thinned.tfr.mcmc).

If Phase III is projected using a BHM (i.e. if use.tfr3 is TRUE), parameter samples simulated via run.tfr3.mcmc are used from which burnin (given by burnin3) is discarded and the chains are evenly thinned in a way that the total size corresponds to the final size of the Phase II parameter samples. Countries for which there are no simulated country-specific Phase III parameters (e.g. because their TFR is still in Phase II or it is an aggregated region) use samples of the "world" AR(1) parameters.

The projection is run for all missing values before the present year, if any. Medians over the trajectories are used as imputed values and the trajectories are discarded. The process then continues by projecting the future values where all generated trajectories are kept.

The resulting prediction object is saved into '{output.dir}/predictions'. Trajectories for all countries are saved into the same directory in a binary format, one file per country. At the end of the projection, if save.as.ascii is larger than 0, the function converts the given number of trajectories into a CSV file of a UN-specific format. They are selected by equal spacing (see function convert.tfr.trajectories for more details on the conversion). In addition, two summary files are created: one in a user-friendly format, the other using a UN-specific coding of the variants and time (see write.projection.summary for more details).

#### Value

Object of class bayesTFR.prediction which is a list containing components:

quantiles A  $n \times q \times p$  array of quantile values computed on all trajectories. n is the

number of countries, q is the number of quantile bounds and p is the number of

projections.

traj.mean.sd  $A n \times 2 \times p$  array holding the mean of all trajectories in the first column and the

standard deviation in the second column. n and p are the number of countries

and number of projections, respectively.

nr. traj Number of trajectories.

trf\_matrix\_reconstructed

Matrix containing imputed TFR values on spots where the original TFR matrix has missing values, i.e. between the last observed data point and the present

year.

output.directory

Directory where trajectories corresponding to this prediction are stored.

nr.projections Number of projections.

burnin, thin, burnin3, thin3

Burnin and thin used for this prediction for Phase II and Phase III, respectively.

end. year The end year of this prediction.

mu, rho, sigma\_t, sigmaAR1

Parameters of the AR(1) process. sigma\_t is a vector of actual values of the

standard deviation s used for each projection.

na.index Index of trajectories for which at least one country got NA values.

mcmc.set Object of class bayesTFR.mcmc.set used for this prediction, i.e. the burned,

thinned, and collapsed MCMC chain.

#### Author(s)

Hana Sevcikova, Leontine Alkema, Bailey Fosdick

#### References

L. Alkema, A. E. Raftery, P. Gerland, S. J. Clark, F. Pelletier, Buettner, T., Heilig, G.K. (2011). Probabilistic Projections of the Total Fertility Rate for All Countries. Demography, Vol. 48, 815-839. (A Working Paper version at http://www.csss.washington.edu/Papers [nr. 97])

Raftery, A.E., Alkema, L. and Gerland, P. (2013). Bayesian Population Projections for the United Nations. Statistical Science, in press.

Fosdick, B., Raftery, A.E. (2012). Regional Probabilistic Fertility Forecasting by Modeling Between-Country Correlations. Working paper, CSSS, University of Washington. http://www.csss.washington.edu/Papers [nr. 126]

#### See Also

run.tfr.mcmc, run.tfr3.mcmc, create.thinned.tfr.mcmc, convert.tfr.trajectories, write.projection.summarget.tfr.prediction, summary.bayesTFR.prediction

tfr.predict.extra 55

## **Examples**

# Description

Using the posterior parameter samples the function generates posterior trajectories of the total fertility rate for given countries or regions. It is intended to be used after running run.tfr.mcmc.extra, but it can be also used for purposes of testing specific settings on one or a few countries.

## Usage

```
tfr.predict.extra(sim.dir = file.path(getwd(), 'bayesTFR.output'),
    prediction.dir = sim.dir, countries = NULL,
    save.as.ascii = 1000, verbose = TRUE)
```

## **Arguments**

sim.dir Directory with the MCMC simulation results.

prediction.dir Directory where the prediction object and the trajectories are stored.

Vector of country codes for which the prediction should be made. If it is NULL, the prediction is run for all countries that are included in the MCMC object but for which no prediction was generated.

Save.as.ascii Either a number determining how many trajectories should be converted into an ascii file, or "all" in which case all trajectories are converted. It should be set to 0, if no converions is desired. Note that the convertion is done on all countries. verbose Logical switching log messages on and off.

#### **Details**

In order to use this function, a prediction object must exist, i.e. the function tfr.predict must have been processed prior to using this function.

Trajectories for given countries or regions are generated and stored in binary format along with other countries (in prediction\_dir). The existing prediction object is updated and stored in the same directory. If save.as.ascii is larger than zero, trajectories of ALL countries are converted to an ascii format.

56 tfr.raftery.diag

## Value

Updated object of class bayesTFR.prediction.

#### Author(s)

Hana Sevcikova

#### See Also

```
tfr.predict
```

tfr.raftery.diag

Raftery Diagnostics for Parameters of the Total Fertility Rate

# **Description**

The functions compute the Raftery diagnostics for each parameter of MCMCs of phase II (tfr.raftery.diag) and phase III (tfr.raftery.diag), taking median over all chains.

## Usage

```
tfr.raftery.diag(mcmc = NULL,
    sim.dir = file.path(getwd(), "bayesTFR.output"),
    burnin = 0, country = NULL,
    par.names = tfr.parameter.names(trans = TRUE),
    par.names.cs = tfr.parameter.names.cs(trans=TRUE),
    country.sampling.prop = 1, verbose=TRUE, ...)

tfr3.raftery.diag(mcmc = NULL,
    sim.dir = file.path(getwd(), "bayesTFR.output"),
    burnin = 0, country = NULL,
    par.names = tfr3.parameter.names(),
    par.names.cs = tfr3.parameter.names.cs(),
    country.sampling.prop = 1, verbose=TRUE, ...)
```

# **Arguments**

mcmc	A bayesTFR.mcmc or bayesTFR.mcmc.set object.
sim.dir	Directory with the MCMC simulation results. Only used if mcmc is NULL.
burnin	Burnin.
country	Name or code of a country. If it is given, country-specific parameters are reduced to parameters of that country.
par.names	Names of country-independent parameters for which the Raftery diagnostics should be computed.
par.names.cs	Names of country-specific parameters for which the Raftery diagnostics should be computed.

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country.sampling.prop

Proportion of countries that are included in the diagnostics. It should be between 0 and 1. If it is smaller than 1, the countries are randomly sampled. It is only

relevant if par.names.cs is not NULL.

verbose Logical switching log messages on and off.

... Additional arguments passed to the coda.list.mcmc function.

#### **Details**

The Raftery diagnostics is computed for each parameter, using **coda**'s raftery diag with r=0.0125, q=0.025 and q=0.975. Values of N and burnin are taken as the median over chains. For each country-specific parameter, the maximum over all included countries of such medians is taken.

#### Value

List with the components:

Nmedian 2-d array of N values (processed as described in Details) with two rows: first

corresponding to q=0.025, second corresponding to q=0.975. Each column cor-

responds to one parameter.

burnin 2-d array of the same structure as Nmedian, containing the burnin values (pro-

cessed as described in Details).

not.converged.parameters

List with two elements, each of which is a data frame containing columns "parameter.name", "chain.id", and "N". These are parameters for which the computed value of Raftery diagnostics N is larger than the total number of finished iterations summed over all chains. The first element of the list corresponds to

q=0.025, second corresponds to q=0.975.

not.converged.inchain.parameters

List of the same structure as not.converged.parameters. The parameters included are those for which the computed value of Raftery diagnostics N is larger than the number of finished iterations in the corresponding chain.

N.country.indep

Data frame containing columns "parameter.name", "chain.id", "N0.025", and "N0.975". Each row gives N computed with the two different q for each country-independent parameter and chain.

N.country.spec The same as N.country.indep, but here the country-specific parameters are considered.

Nmedian.country.spec

2-d array of N values for country-specific parameters containing medians over

chains.

thin.ind List with elements '0.025', '0.975' and median. The first two elements are

matrices with one row per chain and one column per parameter. They contain values of thin that makes the MCMC independent, for q=0.025 and q=0.975, respectively. The median element is of the same structure as Nmedian, contain-

ing medians ove rows in the two matrices in this list.

nr.countries Vector with elements used (number of countries used in this diagnostics) and

total (number of countries that this mcmc object was estimated on).

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## Author(s)

Hana Sevcikova, Adrian Raftery

#### See Also

```
raftery.diag
```

tfr.trajectories.plot Output of Posterior Distribution of TFR Trajectories

## **Description**

The functions plot/tabulate the posterior distribution of TFR trajectories for a given country, or for all countries, including their median and given probability intervals.

#### Usage

```
tfr.trajectories.plot(tfr.pred, country, pi = c(80, 95),
    half.child.variant = TRUE, nr.traj = NULL,
    adjusted.only = TRUE, typical.trajectory = FALSE,
    mark.estimation.points = FALSE,
    xlim = NULL, ylim = NULL, type = 'b', xlab = 'Year', ylab = 'TFR',
    main = NULL, lwd = c(2, 2, 2, 2, 2, 1),
    col=c('black', 'green', 'red', 'red', 'blue', 'gray'),
    show.legend = TRUE, add = FALSE, ...)

tfr.trajectories.plot.all(tfr.pred,
    output.dir = file.path(getwd(), 'TFRtrajectories'),
    output.type = "png", main = NULL, verbose = FALSE, ...)

tfr.trajectories.table(tfr.pred, country, pi = c(80, 95),
    half.child.variant = TRUE)
```

## **Arguments**

tfr.pred Object of class bayesTFR.prediction. country Name or numerical code of a country. Probability interval. It can be a single number or an array. рi half.child.variant If TRUE the United Nations variant of "+/-0.5 child" (relative to the median) is shown. nr.traj Number of trajectories to be plotted. If NULL, all trajectories are plotted, otherwise they are thinned evenly. adjusted.only Logical. By default, if the projection median is adjusted using e.g. tfr.median.set, the function plots the adjusted median. If adjusted.only=FALSE the original (non-adjusted) median is plotted as well.

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typical.trajectory

Logical. If TRUE one trajectory is shown that has the smallest distance to the median.

mark.estimation.points

Logical. If TRUE, points that were not used in the estimation (phase I) are shown in lighter color than points in phase II and III.

xlim, ylim, type, xlab, ylab

Graphical parameters passed to the plot function.

lwd, col Vector of six elements giving the line width and color for: 1. observed data,

2. imputed missing data, 3. median, 4. quantiles, 5. half-child variant, 6.

trajectories.

show.legend Logical controlling whether the legend should be drawn.

add Logical controlling whether the trajectories should be plotted into a new graphic

device (FALSE) or into an existing device (TRUE). One can use this argument to

plot trajectories from multiple countries into one graphics.

... Additional graphical parameters. In addition, for tfr.trajectories.plot.all,

... contains any of the arguments of tfr.trajectories.plot.

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".

main Main title for the plot(s). In tfr.trajectories.plot.all any occurence of

the string "XXX" is replaced by the country name.

verbose Logical switching log messages on and off.

#### **Details**

tfr.trajectories.plot plots posterior distribution of TFR trajectories for a given country. tfr.trajectories.table gives the same output as a table. tfr.trajectories.plot.all creates a set of graphs (one per country) that are stored in output.dir.

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.

#### Author(s)

Hana Sevcikova, Leontine Alkema

#### See Also

```
bayesTFR.prediction
```

# **Examples**

```
## Not run:
sim.dir <- file.path(find.package("bayesTFR"), "ex-data", "bayesTFR.output")
pred <- get.tfr.prediction(sim.dir)
tfr.trajectories.plot(pred, country="Burkina Faso", pi=c(80, 95))</pre>
```

UN\_time

```
tfr.trajectories.table(pred, country="Burkina Faso", pi=c(80, 95))
## End(Not run)
```

UN\_time

Dataset with UN-specific Time Coding

# Description

Dataset used by the UN for coding time. It is an TAB-separated ASCII file called "UN\_time.txt".

## Usage

```
data(UN_time)
```

## **Format**

A data frame with 1034 observations on the following 4 variables.

TimeID Time identifier.

TLabel Label of the time, with minimum values of 1950 and 1950-1955, and maximum values of 2399, 2400 and 2400-2405.

TDate Equal to TLabel if it is a single year, or the starting year of TLabel, if it is an interval.

Tinterval Length of the time interval, or zero, if it is a single year.

## **Details**

For 5-year period data, fertility rates are defined from 1 July of year (t) to 1 July of year (t+5), with 1 January of year (t+3) as exact mid-date. This means for example that data for 2000-2005, refer to the period between 2000.5 and 2005.5, with 2003.0 as exact mid-point.

## Source

Data provided by the United Nations Population Division

# **Examples**

```
data(UN_time)
str(UN_time)
```

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UN\_variants

Dataset with UN-specific Coding of Variants

## **Description**

Dataset used by the UN for coding variants. It also includes variants for the lower and upper bounds of the 80 and 95% probability intervals, respectively, resulting from the Bayesian hierarchical model. The dataset is stored in a TAB-separated ASCII file called "UN\_variants.txt".

#### **Usage**

```
data(UN_variants)
```

## **Format**

A data frame with 23 observations on the following 5 variables.

RevID Revision identifier.

VarID Identifier of the variant.

Vshort Short name of the variant.

VName Full name of the variant.

VariantDomain Domain of the variant.

#### Source

Data provided by the United Nations Population Division

## **Examples**

```
data(UN_variants)
str(UN_variants)
```

```
write.projection.summary
```

Writing Projection Summary Files

# Description

The function creates two files containing projection summaries, such as the median, the lower and upper bound of the 80 and 90% probability intervals, respectively, the +/- 0.5 child variant and the constant variant. One file is in a user-friendly format, whereas the other is in a UN-specific format with internal coding of the time and the variants. In addition, a file containing some of the model parameters is created.

## Usage

```
write.projection.summary(dir = file.path(getwd(), "bayesTFR.output"),
    output.dir = NULL, revision = NULL, adjusted = FALSE)
```

## **Arguments**

dir	Directory containing the prediction object. It should correspond to the output.dir argument of the tfr.predict function.
output.dir	Directory in which the resulting file will be stored. If NULL the same directory is used as for the prediction.
revision	UN WPP revision number. If NULL it is determined from the corresponding WPP year: WPP 2008 corresponds to revision 13, every subsequent WPP increases the revision number by one. Used as a constant in the second file only.
adjusted	Logical. By default the function writes summary using the original BHM projections. If the projection medians are adjusted (using e.g. tfr.median.set), setting this argument to TRUE causes writing the adjusted projections.

#### **Details**

The first file that the function creates is called 'projection\_summary\_user\_friendly.csv' (or 'projection\_summary\_user\_friendly\_adjusted.csv' if adjusted=TRUE), 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", "-0.5child", "+0.5child", "constant"
- period1: e.g. "2005-2010": TFR for the first time period
- period2: e.g. "2010-2015": TFR for the second time period
- ... further columns with TFR projections

The second file, called 'projection\_summary.csv' (or 'projection\_summary\_adjusted.csv' if adjusted=TRUE), also comma-separated table, contains the same information as above in a UN-specific format:

- "RevID": revision number, passed to the function as an argument
- "VarID": variant identifier, extracted from the UN\_variants dataset
- "LocID": country code
- "TimeID": time identifier, extracted from the UN\_time dataset
- "TFR": the total fertility rate for this variant, location and time period

The third comma-separated file, called 'projection\_summary\_parameters.csv' contains the following columns:

- "country\_name": country name
- "country\_code": country code

- "TF\_time\_start\_decline": start period of TFR decline
- "TF\_max": TFR at the onset of the fertitility transition (median of the  $\mathcal{U}_c$  parameter)
- "TF\_max\_decrement": maximum decrement of TFR decline (median of the  $d_c$  parameter)
- "TF\_end\_level": median of the end level of the fertility transition ( $\Delta_{c4}$  parameter)
- "TF\_end\_level\_low": 2.5 percentile of the  $\Delta_{c4}$  distribution
- "TF\_end\_level\_high": 97.5 percentile of the  $\Delta_{c4}$  distribution
- "TF\_time\_end\_decline": period of the end decline, measured using the prediction median

Note that this file is not created if adjusted=TRUE.

## Note

This function is automatically called from the tfr.predict function, therefore in standard cases it will not be needed to call it directly.

# Author(s)

Hana Sevcikova

## See Also

```
convert.tfr.trajectories, tfr.predict
```

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