Package ‘bamp’

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Type Package

Title Bayesian Age-Period-Cohort Modeling and Prediction

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Description Bayesian Age-Period-Cohort Modeling and Prediction using effi-
cient Markov Chain Monte Carlo Methods. This is the R version of the previous BAMP soft-
ware as described in Volker Schmid and Leon-
hard Held (2007) <DOI:10.18637/jss.v021.i08> Bayesian Age-Period-Cohort Modeling and Pre-
diction - BAMP, Journal of Statistical Software 21:8. This package includes checks of conver-
gence using Gelman’s R.

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Depends R (>= 3.5.0)

Imports coda, graphics, parallel, stats, abind

LazyData true

RoxygenNote 7.1.2

VignetteBuilder knitr, R.rsp

Suggests knitr, rmarkdown, R.rsp

URL https://volkerschmid.github.io/bamp/

BugReports https://github.com/volkerschmid/bamp/issues

NeedsCompilation yes

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R topics documented:

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Description

Class for (Bayesian) age-period-cohort objects

Usage

apc()

Details

bamp will return an object of class apc. Available functions are

- plot.apc plots main effects
- print.apc print summary of model and effects
- effects.apc extract effects (mean, median and quantiles)

Value

apc class
Example dataset for APC model

Description
A dataset containing case counts and population numbers in eight age groups for ten years. Each age group consists of five years.

Usage
data(apc)

Format
- population: matrix of population data
- cases: matrix of case counts
- cov_p: covariate for period
- cov_c: covariate for cohort

Simulate from age-period-cohort model

Description
This functions simulates a data set of cases on the Lexis diagram from given age, period and cohort effects. Population numbers have to be given; can be one number for all age group/period combinations.

Usage
apcSimulate(intercept, age, period, cohort, periods_per_agegroup, population)

Arguments
- intercept: Intercept
- age: Vector of effect for age groups
- period: Vector of effects for periods
- cohort: Vector of effect for cohorts
- periods_per_agegroup: Periods per age group
- population: Population number. Either a matrix or a scalar.
Value

List with number of cases (matrix) and population numbers (matrix).

See Also

vignette("simulation", package = "bamp")

Examples

age=sqrt(seq(5,0,length=10)); age<-1-age-mean(age)
period=15:1; period[8:15]<-8:15; period<-period/6; period<-period-mean(period)
periods_per_agegroup=5; number_of_cohorts<- periods_per_agegroup*(10-1)+15
cohort<-rep(0,60); cohort[1:10]<-1:10; cohort[41:60]<- -(1:20)/2; cohort<-cohort/10;
cohort<-cohort-mean(cohort)
simdata<-apcSimulate(-5, age, period, cohort, periods_per_agegroup, 1e6)
par(mfrow=c(3,1))
plot(age, type="l")
plot(period, type="l")
plot(cohort, type="l")
## Not run:
simmod <- bamp(cases = simdata$cases, population = simdata$population, age = "rw1",
period = "rw1", cohort = "rw1", periods_per_agegroup =periods_per_agegroup)
plot(simmod)
## End(Not run)

bamp

Bayesian Age-Period-Cohort Modeling and Prediction (bamp)

Description

Bayesian Age-Period-Cohort Modeling for the analyze of incidence or mortality data on the Lexis diagram. For each pixel in the Lexis diagram (that is for a specific age group and specific period) data must be available on the number of persons under risk (population number) and the number of disease cases (typically cancer incidence or mortality). A hierarchical model is assumed with a binomial model in the first-stage. As smoothing priors for the age, period and cohort parameters random walks of first and second order (RW1 or RW2) available. Deviance information criterion and effective number of parameters is computed for model comparison. Note that there is a non-identifiability in the likelihood of the APC-model, see e.g. Clayton and Schifflers (1987, DOI:10.1002/sim.4780060406), which indices some problems in interpreting the latent effects. Only for RW1 model, the parameters are (weakly) identifiable. Period and age groups do not need to be on the same grid, for example periods can be in one year intervals and age groups in five year intervals.
Additionally to the model described in Knorr-Held and Rainer (2001, DOI:10.1093/biostatistics/2.1.109), bamp can handle

• AP and AC models,
• models with and without global heterogeneity parameter (overdispersion),
• models with additional age, period and/or cohort heterogeneity,
• additional covariates.

Usage

```r
bamp(
cases, population, age, period, cohort, overdisp = FALSE, period_covariate = NULL, cohort_covariate = NULL, periods_per_agegroup, mcmc.options = list(number_of_iterations = 1e+05, burn_in = 50000, step = 50, tuning = 500), hyperpar = list(age = c(1, 0.5), period = c(1, 5e-04), cohort = c(1, 5e-04), overdisp = c(1, 0.05)), dic = TRUE, parallel = TRUE, verbose = FALSE)
```

Arguments

cases number of cases
population population number
age prior for age groups ("rw1", "rw2", "rw1+het", "rw2+het", " ")
period prior for periods ("rw1", "rw2", "rw1+het", "rw2+het", " ")
cohort prior for cohorts ("rw1", "rw2", "rw1+het", "rw2+het", " ")
overdisp logical, add overdispersion to model
period_covariate covariate for period
cohort_covariate covariate for cohort
periods_per_agegroup periods per age group
mcmc.options list of options for MCMC.

• number_of_iterations: number of iterations per chain.
• burn_in: number of iterations used as burnin at the beginning of the algorithm, these iterations will be removed.
• step: Step size, for example default is 50, so only every 50th iterations will be stored.
checkConvergence

• tuning: number of iterations for automatic tuning. Depending on the model, the MCMC algorithm will tune certain parameters for more efficient MCMC chains. After tuning, the algorithm is restarted.

hyperpar

list of hyper parameters. The hyper prior for the precision (inverse variance) in the random walk priors is a Gamma distribution with parameters \( a \) and \( b \); expected value is \( a/b \), variance is \( a/b^2 \). Weak hyper parameters are suggested, defaults are \( a = 1, b = 0.5 \) for age, \( a = 1, b = 0.0005 \) for period and cohort effects and \( a = 1, b = 0.05 \) for overdispersion (if added). It is recommended to choose the hyper priors depending on the model, in particular on the order of the random walk.

dic

logical. If true, DIC will be computed

parallel

logical, should computation be done in parallel. This uses the parallel package, which does not allow parallel computing under Windows.

verbose

verbose mode

Details

This function returns an \texttt{apc} object. Only samples from the posterior are computed, point estimates and credible intervals will be computed in \texttt{effects.apc}, \texttt{print.apc} and \texttt{plot.apc}. \texttt{predict_apc} can be used for prediction of the future rates and number of cases and for a retrospective prediction for model checking.

See Also

\vignette{modeling, package = "bamp"}

Examples

```r
## Not run:
data(apc)
model <- bamp(cases, population, age="rw1", period="rw1", cohort="rw1", periods_per_agegroup = 5)
## End(Not run)
```

checkConvergence \hspace{1cm} Check \texttt{apc} object, whether MCMC has converged

Description

This function uses Gelman and Rubin’s \( R \) to check convergence for all main parameters. All parameters should have \( R < 1.1 \). \texttt{bamp} runs at least four MCMC chains by default (more if parallel is more than four).

Usage

```r
checkConvergence(x, info = FALSE, level = 2, auto = FALSE)
```
coh

Arguments

<table>
<thead>
<tr>
<th>x</th>
<th>An apc object</th>
</tr>
</thead>
<tbody>
<tr>
<td>info</td>
<td>logical; print more information</td>
</tr>
<tr>
<td>level</td>
<td>level of check; 1 uses point point estimation, 2 uses upper C.I.</td>
</tr>
<tr>
<td>auto</td>
<td>logical; should be TRUE if called automatically from bamp #'</td>
</tr>
</tbody>
</table>

Value

logical; TRUE if check is fine.

Examples

```r
## Not run:
data(apc)
model <- bamp(cases, population, age="rw1", period="rw1", cohort="rw1", periods_per_agegroup = 5)
checkConvergence(model)

## End(Not run)
```

coh Compute cohort index from age and period index

Description

Compute cohort index from age and period index

Usage

coh(agegroup, period, noa, periods_per_agegroup)

Arguments

<table>
<thead>
<tr>
<th>agegroup</th>
<th>age group index</th>
</tr>
</thead>
<tbody>
<tr>
<td>period</td>
<td>period index</td>
</tr>
<tr>
<td>noa</td>
<td>number of age groups in total</td>
</tr>
<tr>
<td>periods_per_agegroup</td>
<td>periods per age group</td>
</tr>
</tbody>
</table>

Value

cohort index

Examples

```r
# last agegroup in first period equals first cohort
coh(10, 1, 10, 5)

# first agegroup in last period equals last cohort
coh(1, 8, 10, 5)
```
effects.apc

**Effects from Fitted APC Model**

**Description**

Effects from Fitted APC Model

**Usage**

```r
## S3 method for class 'apc'
effects(object, mean = FALSE, quantiles = 0.5, update = FALSE, ...)
```

**Arguments**

- `object`: an apc object
- `mean`: logical. If TRUE, mean effects are computed
- `quantiles`: Scalar or vector of quantiles to compute (only if mean=FALSE)
- `update`: logical. If TRUE, the apc object including the effects is returned
- `...`: Additional arguments will be ignored

**Value**

List of age, period, cohort effects or apc object including effects (if update=TRUE)

**Examples**

```r
## Not run:
data(apc)
model <- bamp(cases, population, age="rw1", period="rw1", cohort="rw1", periods_per_agegroup = 5)
effects(model)
## End(Not run)
```

---

plot.apc

**Plot apc object**

**Description**

Plot apc object

**Usage**

```r
## S3 method for class 'apc'
plot(x, quantiles = c(0.05, 0.5, 0.95), ...)
```
predict_apc

Arguments

x          apc object
quantiles  quantiles to plot. Default: c(0.05, 0.5, 0.95) is median and 90% credible interval.
...        Additional arguments will be ignored

Details

Plot of age, period and cohort effects from apc objects. If covariates have been used for period/cohort, a second plot with covariate, absolute effect and relative effect is created. Absolute effect is relative effect times covariate.

Value

plot

Examples

## Not run:
data(apc)
model <- bamp(cases, population, age="rw1", period="rw1", cohort="rw1", periods_per_agegroup = 5)
plot(model)
## End(Not run)

predict_apc

Prediction for age-period-cohort models

Description

Prediction of rates and, if possible, cases from the Bayesian age-period-cohort model using the prior assumptions (random walks) of the model and the estimated variance of the random walk. For example, random walk of first order (rw1) for period effect predicts constant effects for future periods plus noise.

Usage

predict_apc(
  object,
  periods = 0,
  population = NULL,
  quantiles = c(0.05, 0.5, 0.95),
  update = FALSE
)
Arguments

- **object**: apc object
- **periods**: number of periods to predict
- **population**: matrix of (predicted) population, if NULL, population data from original bamp call will be used
- **quantiles**: vector of quantiles to compute
- **update**: boolean. If TRUE, object will be returned with results added to the object

Details

This function will return predicted rates for future periods. For this, future period and cohort effects will be predicted. Further age group effects will not be predicted. The rates are random samples from the predictive distribution; number of samples is equal to number of MCMC iterations. Quantiles will be provided for convenience, but all samples are available. If population numbers are given, number of cases will also be predicted. Number of cases will not only be predicted for future periods, but also for the time periods where data are available; this can be used for model assessment.

Value

list with quantiles of predicted probabilities (pr), predicted cases (cases) and predicted cases per period (cases_period) and a list samples with MCMC samples of pr, cases and cases_period. If update=TRUE, the apc object will be returned with this list (predicted) added.

See Also

vignette("prediction", package = "bamp")

Examples

```r
## Not run:
data(apc)
model <- bamp(cases, population, age="rw1", period="rw1", cohort="rw1", periods_per_agegroup = 5)
pred <- predict_apc(model, periods=1)
plot(pred$pr[2,11,], main="Predicted rate per agegroup", ylab="p")

## End(Not run)
```

Description

Print apc objects
print.apc

Usage

## S3 method for class 'apc'
print(x, ...)

Arguments

x apc object
...
additional arguments will be ignored

Value

print

Examples

## Not run:
data(apc)
model <- bamp(cases, population, age="rw1", period="rw1", cohort="rw1", periods_per_agegroup = 5)
print(model)

## End(Not run)
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