Package ‘MCMCprecision’

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Description Estimates the precision of transdimensional Markov chain Monte Carlo (MCMC) output, which is often used for Bayesian analysis of models with different dimensionality (e.g., model selection). Transdimensional MCMC (e.g., reversible jump MCMC) relies on sampling a discrete model-indicator variable to estimate the posterior model probabilities. If only few switches occur between the models, precision may be low and assessment based on the assumption of independent samples misleading. Based on the observed transition matrix of the indicator variable, the method of Heck, Overstall, Gronau, & Wagenmakers (2019, Statistics & Computing, 29, 631-643) <doi:10.1007/s11222-018-9828-0> draws posterior samples of the stationary distribution to (a) assess the uncertainty in the estimated posterior model probabilities and (b) estimate the effective sample size of the MCMC output.
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MCMCprecision-package MCMCprecision: Precision of discrete parameters in transdimensional MCMC

Description

MCMCprecision estimates the precision of the posterior model probabilities in transdimensional Markov chain Monte Carlo methods (e.g., reversible jump MCMC or product-space MCMC). This is useful for applications of transdimensional MCMC such as model selection, mixtures with varying numbers of components, change-point detection, capture-recapture models, phylogenetic trees, variable selection, and for discrete parameters in MCMC output in general.

Details

The main function to assess the estimation uncertainty of discrete MCMC output is stationary. The method is explained in detail in Heck et al. (2019, Statistics & Computing), available in the package by calling: vignette("MCMCprecision")

Author(s)

Daniel W. Heck

References


See Also

Useful links:

• https://github.com/danheck/MCMCprecision
**best_models**  

*Precision for the k Best-Performing Models*

**Description**

Assesses the precision in estimating the ranking of the k best-performing models.

**Usage**

```r
best_models(samples, k, ties.method = "min")
```

**Arguments**

- `samples`: a matrix with posterior samples (one per row) for the model posterior probabilities (one model per column). Can be estimated using `stationary` with the argument `summary = FALSE`.
- `k`: number of best-performing models to be considered
- `ties.method`: a character string specifying how ties are treated, see `rank`

**Examples**

```r
# a sequence of uncorrelated model indices
mult <- rmultinom(1000, 1, c(.05, .6, .15, .12, .08))
idx <- apply(mult, 2, which.max)
z <- letters[idx]
stat <- stationary(z, summary = FALSE)
best_models(stat, 3)
```

**fit_dirichlet**  

*Estimate Parameters of Dirichlet Distribution*

**Description**

C++ implementation of the fixed-point iteration algorithm by Minka (2000).

**Usage**

```r
fit_dirichlet(x, const, maxit = 1e+05, abstol = 0.1)
```

**Arguments**

- `x`: a matrix of Dirichlet samples, one row per observation.
- `const`: constant that is added to avoid problems with zeros in `log(x)`. The default is `const = min(x[x>0])*.01`.
- `maxit`: maximum number of iterations.
- `abstol`: The absolute convergence tolerance: maximum of absolute differences of Dirichlet parameters.
Details

The algorithm is used to estimate the effective sample size based on samples of posterior model probabilities (see `stationary` and `summary.stationary`).

References


See Also

`rdirichlet`

Examples

```r
x <- rdirichlet(100, c(8,1,3,9))
fit_dirichlet(x)
```

---

**rdirichlet**

**Random Sample from Dirichlet Distribution**

**Description**

Random generation from the Dirichlet distribution.

**Usage**

`rdirichlet(n, a)`

**Arguments**

- `n`: number of samples
- `a`: vector or matrix of shape parameters

**See Also**

`fit_dirichlet`

**Examples**

```r
rdirichlet(2, c(1,5,3,8))
```
**rmarkov**

Generate a sample of a discrete-state Markov chain

**Description**

Generates a sequence of discrete states from a discrete-time Markov chain with transition matrix P.

**Usage**

```
rmk <- matrix(c(.30, .50, .20,
                .05, .25, .70,
                .00, .10, .90), 3, 3, byrow=TRUE)
rmk(50, P)
```

**Arguments**

- `n`: length of the generated sequence.
- `P`: transition matrix (rows are normalized to sum to 1).
- `start`: vector with nonnegative values that defines the discrete starting distribution at t=0 (start is normalized to sum to 1). The default is a discrete uniform distribution.

**Examples**

```
P <- matrix(c(.30, .50, .20,
              .05, .25, .70,
              .00, .10, .90), 3, 3, byrow=TRUE)
rmarkov(50, P)
```

**stationary**

Precision of stationary distribution for discrete MCMC variables

**Description**

Transdimensional MCMC methods include a discrete model-indicator variable z with a fixed but unknown stationary distribution with probabilities π (i.e., the model posterior probabilities). The function stationary draws posterior samples to assess the estimation uncertainty of π.

**Usage**

```
stationary(
  z,
  N,
  labels,
  sample = 1000,
  epsilon = "1/M",
  cpu = 1,
  method = "arma",
```


digits = 6,
progress = TRUE,
summary = TRUE
)

Arguments

z MCMC output for the discrete indicator variable with numerical, character, or factor labels (can also be a mcmc.list or a matrix with one MCMC chain per column).

N the observed transitions matrix (if supplied, z is ignored). A quadratic matrix with sampled transition frequencies (N[i,j] = number of switches from z[t]=i to z[t+1]=j).

labels optional: vector of labels for complete set of models (e.g., models not sampled in the chain z). If epsilon=0, this does not affect inferences due to the improper Dirichlet(0,...,0) prior.

cpu number of CPUs used for parallel sampling. Will only speed up computations for large numbers of models (i.e., for large transition matrices).

method how to compute eigenvectors:

- "arma" (default): Uses RcppArmadillo::eig_gen.
- "base": Uses base::eigen, which might be more stable, but also much slower than "arma" for small transition matrices.
- "eigen": Uses package RcppEigen::EigenSolver
- "armas": Uses sparse matrices with RcppArmadillo::eigs_gen, which can be faster for very large number of models if epsilon=0 (might be numerically unstable).
- "iid": Assumes i.i.d. sampling of the model indicator variable z. This is only implemented as a benchmark, because results cannot be trusted if the samples z are correlated (which is usually the case for transdimensional MCMC output)

digits number of digits that are used for checking whether the first eigenvalue is equal to 1 (any difference must be due to low numerical precision)

progress whether to show a progress bar (not functional for cpu>1)

summary whether the output should be summarized. If FALSE, posterior samples of the stationary probabilities are returned.
Details

The method draws independent posterior samples of the transition matrix \( P \) for the discrete-valued indicator variable \( z \) (usually, a sequence of sampled models). For each row of the transition matrix, a Dirichlet(\( \epsilon, \ldots, \epsilon \)) prior is assumed, resulting in a conjugate Dirichlet posterior. For each sample, the eigenvector with eigenvalue 1 is computed and normalized. These (independent) posterior samples can be used to assess the estimation uncertainty in the stationary distribution \( p_i \) of interest (e.g., the model posterior probabilities) and to estimate the effective sample size (see `summary.stationary`).

Value

default: a summary for the posterior distribution of the model posterior probabilities (i.e., the fixed but unknown stationary distribution of \( z \)). If `summary=FALSE`, posterior samples for \( p_i \) are returned.

See Also

`best_models`, `summary.stationary`

Examples

# data-generating transition matrix
P <- matrix(c(0.1, 0.5, 0.4,
              0, 0.5, 0.5,
              0.9, 0.1, 0), ncol = 3, byrow=TRUE)

# input: sequence of sampled models
z <- rmarkov(500, P)
stationary(z)

# input: transition frequencies
N <- transitions(z)
samples <- stationary(N = N, summary = FALSE)

# summaries:
best_models(samples, k = 3)
summary(samples)

---

stationary_mle  

**MLE for stationary distribution of discrete MCMC variables**

Description

Maximum-likelihood estimation of stationary distribution \( \pi \) based on (a) a sampled trajectory \( z \) of a model-indicator variable or (b) a sampled transition count matrix \( N \).

Usage

stationary_mle(z, N, labels, method = "rev", abstol = 1e-05, maxit = 1e+05)
Arguments

z MCMC output for the discrete indicator variable with numerical, character, or factor labels (can also be a `mcmc.list` or a matrix with one MCMC chain per column).

N the observed transitions matrix (if supplied, z is ignored). A quadratic matrix with sampled transition frequencies (\( N[i, j] \) = number of switches from \( z[t] = i \) to \( z[t+1] = j \)).

labels optional: vector of labels for complete set of models (e.g., models not sampled in the chain z). If \( \text{epsilon}=0 \), this does not affect inferences due to the improper Dirichlet(0,..,0) prior.

method Different types of MLEs:

- "iid": Assumes i.i.d. sampling of the model indicator variable z and estimates \( \pi \) as the relative frequencies each model was sampled.
- "rev": Estimate stationary distribution under the constraint that the transition matrix is reversible (i.e., fulfills detailed balance) based on the iterative fixed-point algorithm proposed by Trendelkamp-Schroer et al. (2015)
- "eigen": Computes the first left-eigenvector (normalized to sum to 1) of the sampled transition matrix

abstol absolute convergence tolerance (only for method = "rev")

maxit maximum number of iterations (only for method = "rev")

Details

The estimates are implemented mainly for comparison with the Bayesian sampling approach implemented in `stationary`, which quantify estimation uncertainty (i.e., posterior SD) of the posterior model probability estimates.

Value

a vector with posterior model probability estimates

References


See Also

`stationary`

Examples

```r
P <- matrix(c(.1,.5,.4,
            0,.5,.5,
            .9,.1,0), ncol = 3, byrow=TRUE)
z <- rmarkov(1000, P)
```
summary.stationary

stationary_mle(z)

# input: transition frequency
tab <- transitions(z)
stationary_mle(N = tab)

---

**summary.stationary**  
*Summary for Posterior Model Probabilities*

**Description**

Summary for a sample of posterior model probabilities (`stationary`). Also provides the estimated effective sample size and summaries for all pairwise Bayes factors.

**Usage**

```r
## S3 method for class 'stationary'
summary(object, BF = FALSE, logBF = FALSE, ...)
```

**Arguments**

- `object`  
  posterior samples of the stationary distribution (rows = samples; columns = models).
- `BF`  
  whether to compute summaries for all pairwise Bayes factors.
- `logBF`  
  whether to summarize log Bayes factors instead of Bayes factors.
- `...`  
  passed to `fit_dirichlet` to estimate effective sample size (e.g., `maxit` and `abstol`).

**Details**

Effective sample is estimated by fitting a Dirichlet model to the posterior model probabilities (thereby assuming that samples were drawn from an equivalent multinomial distribution based on independent sampling). More precisely, sample size is estimated by the sum of the Dirichlet parameters \( \sum \alpha[i] \) minus the prior sample size \( \epsilon \times M^2 \) (where \( M \) is the number of sampled models and \( \epsilon \) the prior parameter for each transition frequency).

**Value**

a list with estimates for "pp" = posterior model probabilities, "n.eff" = effective sample size, and "bf" = pairwise Bayes factors (optional)

**See Also**

`stationary`, `fit_dirichlet`
transitions

Examples

```r
P <- matrix(c(.1,.5,.4,
              0,.5,.5,
              .9,.1,0), ncol = 3, byrow=TRUE)
z <- rmarkov(1000, P)
samples <- stationary(z, summary = FALSE)
summary(samples)
```

---

transitions

*Get matrix of observed transition frequencies*

Description

Summarizes a sequence of discrete values by the observed transition frequencies.

Usage

```r
transitions(z, labels, order = 1)
```

Arguments

- `z`: vector of model indices (numerical or character)
- `labels`: fixed labels for models that should be included in transition matrix, e.g., `labels=1:20` or `c("m1","m2",...)`
- `order`: order of the transition table. If `order=1`, a matrix with transition frequencies from `z[t+1]` is returned. If `order=2`, a 3-dimensional array is returned with transition frequencies for `z[t], z[t+1],` and `z[t+2]`.

Value

a square matrix with transition frequencies

Examples

```r
P <- matrix(c(.9,.1,0,
              .1,.6,.3,
              .2,.3,.5), 3, byrow=TRUE)
z <- rmarkov(2000, P)
transitions(z)
transitions(z, order = 2)
```
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