

Package ‘dsp’

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Type Package

Title Dynamic Shrinkage Process and Change Point Detection

Version 1.5.0

Description Provides efficient Markov chain Monte Carlo (MCMC) algorithms for dynamic shrinkage processes, which extend global-local shrinkage priors to the time series setting by allowing shrinkage to depend on its own past. These priors yield locally adaptive estimates, useful for time series and regression functions with irregular features. The package includes full MCMC implementations for trend filtering using dynamic shrinkage on signal differences, producing locally constant or linear fits with adaptive credible bands. Also included are models with static shrinkage and normal-inverse-Gamma priors for comparison. Additional tools cover dynamic regression with time-varying coefficients and B-spline models with shrinkage on basis differences, allowing for flexible curve-fitting with unequally spaced data. Some support for heteroscedastic errors, outlier detection, and change point estimation.

Methods in this package are described in Kowal et al. (2019) <[doi:10.1111/rssb.12325](https://doi.org/10.1111/rssb.12325)>, Wu et al. (2024) <[doi:10.1080/07350015.2024.2362269](https://doi.org/10.1080/07350015.2024.2362269)>, Schafer and Matteson (2024) <[doi:10.1080/00401706.2024.2407316](https://doi.org/10.1080/00401706.2024.2407316)>, and Cho and Matteson (2024) <[doi:10.48550/arXiv.2408.11315](https://doi.org/10.48550/arXiv.2408.11315)>.

License GPL (>= 3)

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Description

Run the MCMC sampler for ABCO with a penalty on first ($D = 1$), or second ($D = 2$) differences of the conditional expectation. The penalty utilizes the dynamic horseshoe prior on the evolution errors. Sampling is accomplished with a (parameter-expanded) Gibbs sampler, mostly relying on a dynamic linear model representation.

Usage

```
abco(
  y,
  D = 1,
  useAnom = TRUE,
  obsSV = "const",
  nsave = 1000,
  nburn = 1000,
  nskip = 4,
  mcmc_params = list("mu", "omega", "ypred", "evol_sigma_t2", "gamma", "zeta",
    "obs_sigma_t2", "zeta_sigma_t2", "dhs_phi", "dhs_mean", "h_smooth"),
  verbose = TRUE,
  D_asv = 1,
  evol_error_asv = "HS",
  computeDIC = TRUE,
  nugget_asv = TRUE
)
```

Arguments

<code>y</code>	the T vector of time series observations
<code>D</code>	degree of differencing ($D = 1$, or $D = 2$)
<code>useAnom</code>	logical; if TRUE, include an anomaly component in the observation equation
<code>obsSV</code>	Options for modeling the error variance. It must be one of the following: <ul style="list-style-type: none"> • <code>const</code>: Constant error variance for all time points. • <code>SV</code>: Stochastic Volatility model. • <code>ASV</code>: Adaptive Stochastic Volatility model.
<code>nsave</code>	number of MCMC iterations to record
<code>nburn</code>	number of MCMC iterations to discard (burnin)
<code>nskip</code>	number of MCMC iterations to skip between saving iterations, i.e., save every (<code>nskip + 1</code>)th draw
<code>mcmc_params</code>	named list of parameters for which we store the MCMC output; must be one or more of:

	<ul style="list-style-type: none"> • "mu" (conditional mean) • "omega" (Dth difference of mu) • "ypred" (posterior predictive distribution) • "evol_sigma_t2" (evolution error variance) • "gamma": (Threshold parameter in the thresholded DHS log-volatility dynamics). • "zeta": (Outlier component) • "obs_sigma_t2" (observation error variance) • "zeta_sigma_t2" (outlier error variance) • "dhs_phi" (DHS AR(1) coefficient) • "dhs_mean" (DHS AR(1) unconditional mean) • "h" (log variances or log of "obs_sigma_t2". Only used when obsSV = "ASV") • "h_smooth" (smooth estimate of log variances. Only used when obsSV = "ASV" and nugget_asv = TRUE)
verbose	logical; should R report extra information on progress?
D_asv	integer; degree of differencing (0, 1, or 2) for the ASV model. Only used when obsSV = "ASV".
evol_error_asv	character; evolution error distribution for the ASV model. Must be one of the five options used in evol_error. Only used when obsSV = "ASV".
computeDIC	logical; if TRUE (default), compute the deviance information criterion DIC and the effective number of parameters p_d
nugget_asv	logical; if TRUE, fits the nugget variant of the ASV model. Only used when obsSV = "ASV".

Value

A named list of the nsave MCMC samples for the parameters named in mcmc_params

btf

MCMC Sampler for Bayesian Trend Filtering

Description

Run the MCMC for Bayesian trend filtering with a penalty on zeroth ($D = 0$), first ($D = 1$), or second ($D = 2$) differences of the conditional expectation. The penalty is determined by the prior on the evolution errors, which include:

- the dynamic horseshoe prior ('DHS');
- the horseshoe prior ('HS');
- the Bayesian lasso ('BL');
- the normal stochastic volatility model ('SV');
- the normal-inverse-gamma prior ('NIG').

In each case, the evolution error is a scale mixture of Gaussians. Sampling is accomplished with a (parameter-expanded) Gibbs sampler, mostly relying on a dynamic linear model representation.

Usage

```

btf(
  y,
  evol_error = "DHS",
  D = 2,
  obsSV = "const",
  nsave = 1000,
  nburn = 1000,
  nskip = 4,
  mcmc_params = list("mu", "ypred", "evol_sigma_t2", "obs_sigma_t2", "dhs_phi",
    "dhs_mean", "h_smooth"),
  computeDIC = TRUE,
  verbose = TRUE,
  D_asv = 1,
  evol_error_asv = "HS",
  nugget_asv = TRUE
)

```

Arguments

y	the T x 1 vector of time series observations
evol_error	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior; the default), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
D	degree of differencing (D = 0, D = 1, or D = 2)
obsSV	Options for modeling the error variance. It must be one of the following: <ul style="list-style-type: none"> • const: Constant error variance for all time points. • SV: Stochastic Volatility model. • ASV: Adaptive Stochastic Volatility model. for the observation error variance
nsave	number of MCMC iterations to record
nburn	number of MCMC iterations to discard (burnin)
nskip	number of MCMC iterations to skip between saving iterations, i.e., save every (nskip + 1)th draw
mcmc_params	named list of parameters for which we store the MCMC output; must be one or more of: <ul style="list-style-type: none"> • "mu" (conditional mean) • "ypred" (posterior predictive distribution) • "evol_sigma_t2" (evolution error variance) • "obs_sigma_t2" (observation error variance) • "dhs_phi" (DHS AR(1) coefficient) • "dhs_mean" (DHS AR(1) unconditional mean) • "h_smooth" (smooth estimate of log variances. Only used when obsSV = "ASV" and nugget_asv = TRUE)

computeDIC	logical; if TRUE, compute the deviance information criterion DIC and the effective number of parameters p_d
verbose	logical; should R report extra information on progress?
D_asv	integer; degree of differencing (0, 1, or 2) for the ASV model. Only used when <code>obsSV = "ASV"</code> .
evol_error_asv	character; evolution error distribution for the ASV model. Must be one of the five options used in <code>evol_error</code> . Only used when <code>obsSV = "ASV"</code> .
nugget_asv	logical; if TRUE, fits the nugget variant of the ASV model. Only used when <code>obsSV = "ASV"</code> .

Value

A named list of the `nsave` MCMC samples for the parameters named in `mcmc_params`

Note

The data y may contain NAs, which will be treated with a simple imputation scheme via an additional Gibbs sampling step. In general, rescaling y to have unit standard deviation is recommended to avoid numerical issues.

btf0

MCMC Sampler for Bayesian Trend Filtering: $D = 0$

Description

Run the MCMC for Bayesian trend filtering with a penalty on the conditional expectation. The penalty is determined by the prior on the evolution errors, which include:

- the dynamic horseshoe prior ('DHS');
- the horseshoe prior ('HS');
- the Bayesian lasso ('BL');
- the normal stochastic volatility model ('SV');
- the normal-inverse-gamma prior ('NIG').

In each case, the evolution error is a scale mixture of Gaussians. Sampling is accomplished with a (parameter-expanded) Gibbs sampler, mostly relying on a dynamic linear model representation.

Usage

```
btf0(
  y,
  evol_error = "DHS",
  obsSV = "const",
  nsave = 1000,
  nburn = 1000,
  nskip = 4,
```

```

mcmc_params = list("mu", "ypred", "evol_sigma_t2", "obs_sigma_t2", "dhs_phi",
  "dhs_mean", "h_smooth"),
computeDIC = TRUE,
verbose = TRUE,
D_asv = 1,
evol_error_asv = "HS",
nugget_asv = TRUE
)

```

Arguments

<code>y</code>	the $T \times 1$ vector of time series observations
<code>evol_error</code>	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
<code>obsSV</code>	Options for modeling the error variance. It must be one of the following: <ul style="list-style-type: none"> • <code>const</code>: Constant error variance for all time points. • <code>SV</code>: Stochastic Volatility model. • <code>ASV</code>: Adaptive Stochastic Volatility model.
<code>nsave</code>	number of MCMC iterations to record
<code>nburn</code>	number of MCMC iterations to discard (burnin)
<code>nskip</code>	number of MCMC iterations to skip between saving iterations, i.e., save every $(n\text{skip} + 1)$ th draw
<code>mcmc_params</code>	named list of parameters for which we store the MCMC output; must be one or more of: <ul style="list-style-type: none"> • "mu" (conditional mean) • "ypred" (posterior predictive distribution) • "evol_sigma_t2" (evolution error variance) • "obs_sigma_t2" (observation error variance) • "dhs_phi" (DHS AR(1) coefficient) • "dhs_mean" (DHS AR(1) unconditional mean) • "h_smooth" (smooth estimate of log variances. Only used when <code>obsSV = "ASV"</code> and <code>nugget_asv = TRUE</code>)
<code>computeDIC</code>	logical; if TRUE, compute the deviance information criterion DIC and the effective number of parameters p_d
<code>verbose</code>	logical; should R report extra information on progress?
<code>D_asv</code>	integer; degree of differencing (0, 1, or 2) for the ASV model. Only used when <code>obsSV = "ASV"</code> .
<code>evol_error_asv</code>	character; evolution error distribution for the ASV model. Must be one of the five options used in <code>evol_error</code> . Only used when <code>obsSV = "ASV"</code> .
<code>nugget_asv</code>	logical; if TRUE, fits the nugget variant of the ASV model. Only used when <code>obsSV = "ASV"</code> .

Value

A named list of the `nsave` MCMC samples for the parameters named in `mcmc_params`

Note

The data `y` may contain NAs, which will be treated with a simple imputation scheme via an additional Gibbs sampling step. In general, rescaling `y` to have unit standard deviation is recommended to avoid numerical issues.

btf_bspline

MCMC Sampler for B-spline Bayesian Trend Filtering

Description

Run the MCMC for B-spline fitting with a Bayesian trend filtering model on the coefficients, i.e., a penalty on zeroth ($D=0$), first ($D=1$), or second ($D=2$) differences of the B-spline basis coefficients. The penalty is determined by the prior on the evolution errors, which include:

- the dynamic horseshoe prior ('DHS');
- the horseshoe prior ('HS');
- the Bayesian lasso ('BL');
- the normal stochastic volatility model ('SV');
- the normal-inverse-gamma prior ('NIG').

In each case, the evolution error is a scale mixture of Gaussians. Sampling is accomplished with a (parameter-expanded) Gibbs sampler, mostly relying on a dynamic linear model representation.

Usage

```
btf_bspline(
  y,
  times = NULL,
  num_knots = NULL,
  evol_error = "DHS",
  D = 2,
  nsave = 1000,
  nburn = 1000,
  nskip = 4,
  mcmc_params = list("mu", "ypred", "beta", "evol_sigma_t2", "obs_sigma_t2", "dhs_phi",
    "dhs_mean"),
  computeDIC = TRUE,
  verbose = TRUE
)
```

Arguments

<code>y</code>	the $T \times 1$ vector of time series observations
<code>times</code>	the $T \times 1$ vector of observation points; if NULL, assume equally spaced
<code>num_knots</code>	the number of knots; if NULL, use the default of $\max(20, \min(\text{ceiling}(T/4), 150))$
<code>evol_error</code>	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
<code>D</code>	degree of differencing ($D = 0$, $D = 1$, or $D = 2$)
<code>nsave</code>	number of MCMC iterations to record
<code>nburn</code>	number of MCMC iterations to discard (burn-in)
<code>nskip</code>	number of MCMC iterations to skip between saving iterations, i.e., save every $(\text{nskip} + 1)$ th draw
<code>mcmc_params</code>	named list of parameters for which we store the MCMC output; must be one or more of: <ul style="list-style-type: none"> • "mu" (conditional mean) • "beta" (B-spline basis coefficients) • "ypred" (posterior predictive distribution) • "evol_sigma_t2" (evolution error variance) • "obs_sigma_t2" (observation error variance) • "dhs_phi" (DHS AR(1) coefficient) • "dhs_mean" (DHS AR(1) unconditional mean)
<code>computeDIC</code>	logical; if TRUE, compute the deviance information criterion DIC and the effective number of parameters p_d
<code>verbose</code>	logical; should R report extra information on progress?

Value

A named list of the `nsave` MCMC samples for the parameters named in `mcmc_params`

Note

The data `y` may contain NAs, which will be treated with a simple imputation scheme via an additional Gibbs sampling step. In general, rescaling `y` to have unit standard deviation is recommended to avoid numerical issues.

The primary advantages of `btf_bspline` over `btf` are

1. Unequally-spaced points are handled automatically and
2. Computations are linear in the number of basis coefficients, which may be substantially fewer than the number of time points.

btf_bspline0

MCMC Sampler for B-spline Bayesian Trend Filtering: $D = 0$ **Description**

Run the MCMC for B-spline fitting with a penalty the B-spline basis coefficients. The penalty is determined by the prior on the evolution errors, which include:

- the dynamic horseshoe prior ('DHS');
- the horseshoe prior ('HS');
- the Bayesian lasso ('BL');
- the normal stochastic volatility model ('SV');
- the normal-inverse-gamma prior ('NIG').

In each case, the evolution error is a scale mixture of Gaussians. Sampling is accomplished with a (parameter-expanded) Gibbs sampler, mostly relying on a dynamic linear model representation.

Usage

```
btf_bspline0(
  y,
  times = NULL,
  num_knots = NULL,
  evol_error = "DHS",
  nsave = 1000,
  nburn = 1000,
  nskip = 4,
  mcmc_params = list("mu", "ypred", "beta", "evol_sigma_t2", "obs_sigma_t2", "dhs_phi",
    "dhs_mean"),
  computeDIC = TRUE,
  verbose = TRUE
)
```

Arguments

<code>y</code>	the $T \times 1$ vector of time series observations
<code>times</code>	the $T \times 1$ vector of observation points; if NULL, assume equally spaced
<code>num_knots</code>	the number of knots; if NULL, use the default of $\max(20, \min(\text{ceiling}(T/4), 150))$
<code>evol_error</code>	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
<code>nsave</code>	number of MCMC iterations to record
<code>nburn</code>	number of MCMC iterations to discard (burn-in)

nskip	number of MCMC iterations to skip between saving iterations, i.e., save every (nskip + 1)th draw
mcmc_params	named list of parameters for which we store the MCMC output; must be one or more of: <ul style="list-style-type: none"> • "mu" (conditional mean) • "beta" (B-spline basis coefficients) • "ypred" (posterior predictive distribution) • "evol_sigma_t2" (evolution error variance) • "obs_sigma_t2" (observation error variance) • "dhs_phi" (DHS AR(1) coefficient) • "dhs_mean" (DHS AR(1) unconditional mean)
computeDIC	logical; if TRUE, compute the deviance information criterion DIC and the effective number of parameters p_d
verbose	logical; should R report extra information on progress?

Value

A named list of the nsave MCMC samples for the parameters named in mcmc_params

Note

The data y may contain NAs, which will be treated with a simple imputation scheme via an additional Gibbs sampling step. In general, rescaling y to have unit standard deviation is recommended to avoid numerical issues.

The primary advantages of btf_bspline over btf are

1. Unequally-spaced points are handled automatically and
2. Computations are linear in the number of basis coefficients, which may be substantially fewer than the number of time points.

btf_reg

MCMC Sampler for Bayesian Trend Filtering: Regression

Description

Run the MCMC for Bayesian trend filtering regression with a penalty on first ($D=1$) or second ($D=2$) differences of each dynamic regression coefficient. The penalty is determined by the prior on the evolution errors, which include:

- the dynamic horseshoe prior ('DHS');
- the horseshoe prior ('HS');
- the Bayesian lasso ('BL');
- the normal stochastic volatility model ('SV');
- the normal-inverse-gamma prior ('NIG').

In each case, the evolution error is a scale mixture of Gaussians. Sampling is accomplished with a (parameter-expanded) Gibbs sampler, mostly relying on a dynamic linear model representation.

Usage

```

btf_reg(
  y,
  X = NULL,
  evol_error = "DHS",
  D = 1,
  obsSV = "const",
  nsave = 1000,
  nburn = 1000,
  nskip = 4,
  mcmc_params = list("mu", "ypred", "beta", "evol_sigma_t2", "obs_sigma_t2", "dhs_phi",
    "dhs_mean", "h_smooth"),
  use_backfitting = FALSE,
  computeDIC = TRUE,
  verbose = TRUE,
  D_asv = 1,
  evol_error_asv = "HS",
  nugget_asv = TRUE
)

```

Arguments

y	the T x 1 vector of time series observations
X	the T x p matrix of time series predictors
evol_error	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
D	degree of differencing (D = 1 or D = 2)
obsSV	Options for modeling the error variance. It must be one of the following: <ul style="list-style-type: none"> • const: Constant error variance for all time points. • SV: Stochastic Volatility model. • ASV: Adaptive Stochastic Volatility model.
nsave	number of MCMC iterations to record
nburn	number of MCMC iterations to discard (burn-in)
nskip	number of MCMC iterations to skip between saving iterations, i.e., save every (nskip + 1)th draw
mcmc_params	named list of parameters for which we store the MCMC output; must be one or more of: <ul style="list-style-type: none"> • "mu" (conditional mean) • "ypred" (posterior predictive distribution) • "beta" (dynamic regression coefficients) • "evol_sigma_t2" (evolution error variance) • "obs_sigma_t2" (observation error variance) • "dhs_phi" (DHS AR(1) coefficient)

- "dhs_mean" (DHS AR(1) unconditional mean)
- "h_smooth" (smooth estimate of log variances. Only used when obsSV = "ASV" and nugget_asv = TRUE)

use_backfitting	logical; if TRUE, use backfitting to sample the predictors $j=1,\dots,p$ (faster, but usually less MCMC efficient)
computeDIC	logical; if TRUE, compute the deviance information criterion DIC and the effective number of parameters p_d
verbose	logical; should R report extra information on progress?
D_asv	integer; degree of differencing (0, 1, or 2) for the ASV model. Only used when obsSV = "ASV".
evol_error_asv	character; evolution error distribution for the ASV model. Must be one of the five options used in evol_error. Only used when obsSV = "ASV".
nugget_asv	logical; if TRUE, fits the nugget variant of the ASV model. Only used when obsSV = "ASV".

Value

A named list of the nsave MCMC samples for the parameters named in mcmc_params

Note

The data y may contain NAs, which will be treated with a simple imputation scheme via an additional Gibbs sampling step. In general, rescaling y to have unit standard deviation is recommended to avoid numerical issues.

btf_sparse

Run the MCMC for sparse Bayesian trend filtering

Description

Sparse Bayesian trend filtering has two penalties: (1) a penalty on the first ($D = 1$) or second ($D = 2$) differences of the conditional expectation and (2) a penalty on the conditional expectation, i.e., shrinkage to zero.

Usage

```
btf_sparse(
  y,
  evol_error = "DHS",
  zero_error = "DHS",
  D = 2,
  obsSV = "const",
  nsave = 1000,
  nburn = 1000,
```

```

nskip = 4,
mcmc_params = list("mu", "ypred", "evol_sigma_t2", "obs_sigma_t2", "zero_sigma_t2",
  "dhs_phi", "dhs_mean", "dhs_phi_zero", "dhs_mean_zero", "h_smooth"),
computeDIC = TRUE,
verbose = TRUE,
D_asv = 1,
evol_error_asv = "HS",
nugget_asv = TRUE
)

```

Arguments

y	the $T \times 1$ vector of time series observations
evol_error	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
zero_error	the shrinkage-to-zero distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
D	degree of differencing ($D = 1$, or $D = 2$)
obsSV	Options for modeling the error variance. It must be one of the following: <ul style="list-style-type: none"> • const: Constant error variance for all time points. • SV: Stochastic Volatility model. • ASV: Adaptive Stochastic Volatility model.
nsave	number of MCMC iterations to record
nburn	number of MCMC iterations to discard (burnin-in)
nskip	number of MCMC iterations to skip between saving iterations, i.e., save every (nskip + 1)th draw
mcmc_params	named list of parameters for which we store the MCMC output; must be one or more of: <ul style="list-style-type: none"> • "mu" (conditional mean) • "ypred" (posterior predictive distribution) • "evol_sigma_t2" (evolution error variance) • "zero_sigma_t2" (shrink-to-zero error variance) • "obs_sigma_t2" (observation error variance) • "dhs_phi" (DHS AR(1) coefficient for evolution error) • "dhs_mean" (DHS AR(1) unconditional mean for evolution error) • "dhs_phi_zero" (DHS AR(1) coefficient for shrink-to-zero error) • "dhs_mean_zero" (DHS AR(1) unconditional mean for shrink-to-zero error) • "h_smooth" (smooth estimate of log variances. Only used when obsSV = "ASV" and nugget_asv = TRUE)
computeDIC	logical; if TRUE, compute the deviance information criterion DIC and the effective number of parameters p_d

verbose	logical; should R report extra information on progress?
D_asv	integer; degree of differencing (0, 1, or 2) for the ASV model. Only used when obsSV = "ASV".
evol_error_asv	character; evolution error distribution for the ASV model. Must be one of the five options used in evol_error. Only used when obsSV = "ASV".
nugget_asv	logical; if TRUE, fits the nugget variant of the ASV model. Only used when obsSV = "ASV".

Details

Each penalty is determined by a prior, which include:

- the dynamic horseshoe prior ('DHS');
- the horseshoe prior ('HS');
- the Bayesian lasso ('BL');
- the normal stochastic volatility model ('SV');
- the normal-inverse-gamma prior ('NIG').

In each case, the prior is a scale mixture of Gaussians. Sampling is accomplished with a (parameter-expanded) Gibbs sampler, mostly relying on a dynamic linear model representation.

Value

A named list of the nsave MCMC samples for the parameters named in mcmc_params

Note

The data y may contain NAs, which will be treated with a simple imputation scheme via an additional Gibbs sampling step. In general, rescaling y to have unit standard deviation is recommended to avoid numerical issues.

build_Q

Compute the quadratic term in Bayesian trend filtering

Description

Compute the quadratic term arising in the full conditional distribution of a Bayesian trend filtering model with $D = 1$ or $D = 2$. This function exploits the known D -banded structure of Q to compute the matrix directly, using objects in the Matrix package.

Usage

```
build_Q(obs_sigma_t2, evol_sigma_t2, D = 1)
```

Arguments

obs_sigma_t2 the T x 1 vector of observation error variances
 evol_sigma_t2 the T x 1 vector of evolution error variances
 D the degree of differencing (one or two)

Value

Banded T x T Matrix (object) Q

build_XtX	<i>Compute X'X</i>
-----------	--------------------

Description

Build the $T_p \times T_p$ matrix XtX using the `Matrix()` package

Usage

`build_XtX(X)`

Arguments

X T x p matrix of predictors

Value

Block diagonal $T_p \times T_p$ Matrix (object) where each $p \times p$ block is `tcrossprod(matrix(X[t,]))`

Note

$X'X$ is a one-time computing cost. Special cases may have more efficient computing options, but the Matrix representation is important for efficient computations within the sampler.

<code>computeDIC_ASV</code>	<i>Function for calculating DIC and Pb (Bayesian measures of model complexity and fit by Spiegelhalter et al. 2002)</i>
-----------------------------	---

Description

Function for calculating DIC and Pb (Bayesian measures of model complexity and fit by Spiegelhalter et al. 2002)

Usage

`computeDIC_ASV(y, beta, post_sigma2, post_loglike)`

Arguments

y	the $T \times 1$ vector of time series observations.
beta	the known mean of the process. 0 by default.
post_sigma2	posterior samples of the variance, i.e. $\exp(h)$
post_loglike	log likelihood based on the posterior sample.

Value

a list containing DIC and p_d. Two options for estimating both DIC and p_d, which are both included.

credBands	<i>Compute Simultaneous Credible Bands</i>
-----------	--

Description

Compute $(1-\alpha)\%$

Usage

```
credBands(sampFuns, alpha = 0.05)
```

Arguments

sampFuns	Nsims \times m matrix of Nsims MCMC samples and m points along the curve
alpha	confidence level

Value

m \times 2 matrix of credible bands; the first column is the lower band, the second is the upper band

Note

The input needs not be curves: the simultaneous credible "bands" may be computed for vectors. The resulting credible intervals will provide joint coverage at the $(1-\alpha)\%$ level across all components of the vector.

Description

Wrapper function for fitting models with Dynamic Shrinkage Processes (DSP), including:

- Adaptive Bayesian Changepoint analysis and local Outlier (ABCO),
- Bayesian Trend Filter for Gaussian Data
- Time-varying Regression
- Bayesian Trend Filter with B-spline for irregularly spaced or functional time-series.
- Bayesian Smoothing for Count Data

Method for printing basic information about the MCMC sampling settings for the fitted model

Usage

```
dsp_fit(
  y,
  model_spec,
  nsave = 1000,
  nburn = 1000,
  nskip = 4,
  computeDIC = TRUE,
  verbose = TRUE,
  ...
)

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

Arguments

<code>y</code>	a numeric vector of the $T \times 1$ vector of time series observations
<code>model_spec</code>	a list containing model specification generated from <code>dsp_spec()</code> .
<code>nsave</code>	integer scalar (default = 1000); number of MCMC iterations to record
<code>nburn</code>	integer scalar (default = 1000); number of MCMC iterations to discard (burn-in)
<code>nskip</code>	integer scalar (default = 4); number of MCMC iterations to skip between saving iterations, i.e., save every $(nskip + 1)$ th draw
<code>computeDIC</code>	logical; if TRUE (default), compute the deviance information criterion DIC and the effective number of parameters p_d
<code>verbose</code>	logical; should extra information on progress be printed to the console? Defaults to FALSE
<code>...</code>	currently not used
<code>x</code>	object of class <code>dsp</code> from <code>dsp_fit()</code>

Details

A brief summary of the settings used to fit the model including number of iterations, burn in, and thinning rates.

Value

dsp_fit returns an object of class "dsp".

An object of class "dsp" is defined as a list containing at least the following components:

mcmc_output	a list of the nsave MCMC samples for the parameters named in mcmc_params
DIC	Deviance Information Criterion
mcpair	named vector of supplied nsave, nburn, and nskip
model_spec	the object supplied for model_spec argument

MCMC Output Parameters

Shared parameters across wrappers include:

- mu: Conditional mean.
- ypred: Posterior predictive \tilde{y}_t .
- evol_sigma_t2: Variance of the state variable.
- obs_sigma_t2: Observation variance σ_ϵ^2 (family = "gaussian").
- dhs_phi, dhs_mean: DHS hyperparameter draws (evol_error = "DHS").

Model-specific parameters include:

- changepoint (Gaussian):
 - omega: D -th differenced posterior mean draws, μ_t .
 - zeta: anomaly component (when useAnom = TRUE).
 - zeta_sigma_t2: anomaly variance (when useAnom = TRUE).
 - gamma: threshold parameter in the thresholded DHS log-volatility dynamics.
- regression (Gaussian):
 - beta: time-varying regression coefficient draws.
- bspline (Gaussian):
 - beta: B-spline basis coefficient draws.
- smoothing (Negative Binomial):
 - r: overdispersion parameter draws.

Note

The data y may contain NAs, which will be treated with a simple imputation scheme via an additional Gibbs sampling step. In general, rescaling y to have unit standard deviation is recommended to avoid numerical issues when family is "gaussian".

Examples

```

set.seed(200)
signal = c(rep(0, 50), rep(10, 50))
noise = rep(1, 100)
noise_var = rep(1, 100)
for (k in 2:100){
  noise_var[k] = exp(0.9*log(noise_var[k-1]) + rnorm(1, 0, 0.5))
  noise[k] = rnorm(1, 0, sqrt(noise_var[k])) }

y = signal + noise
model_spec = dsp_spec(family = "gaussian", model = "changepoint",
                      D = 1, useAnom = TRUE, obsSV = "SV")
mcmc_output = dsp_fit(y, model_spec = model_spec, nsave = 500, nburn = 500)

print(mcmc_output)

```

 dsp_spec

Model Specification

Description

Method for creating dsp specification object prior to fitting.

Method for printing basic information about the model specification

Usage

```

dsp_spec(family, model, ...)

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

```

Arguments

- | | |
|--------|--|
| family | A character string specifying the model family. Must be one of: <ul style="list-style-type: none"> • "gaussian": Gaussian family. • "negbinomial": Negative binomial family. |
| model | A character string specifying the model type: <ul style="list-style-type: none"> • family = "gaussian": <ul style="list-style-type: none"> – "changepoint": Change point detection with Adaptive Bayesian Change-point analysis and local Outlier (ABCO), – "smoothing": Bayesian smoothing, – "regression": Time-varying regression, – "bspline": Bayesian smoothing with B-spline for irregularly spaced or functional time-series. • family = "negbinomial": |

- "smoothing": Bayesian smoothing.
- ... currently not used
- x object of class `dsp_spec` from `dsp_spec()`

Value

A list containing the model specification.

Examples

```
model_spec <- dsp_spec(family = "gaussian",
                      model = "changepoint")

print(model_spec)
```

ergMean	<i>Compute the ergodic (running) mean.</i>
---------	--

Description

Compute the ergodic (running) mean.

Usage

```
ergMean(x)
```

Arguments

x vector for which to compute the running mean

Value

A vector `y` with each element defined by $y[i] = \text{mean}(x[1:i])$

Description

The penalty is determined by the prior on the evolution errors, which include:

- the dynamic horseshoe prior ('DHS');
- the horseshoe prior ('HS');
- the Bayesian lasso ('BL');
- the normal-inverse-gamma prior ('NIG').

In each case, the evolution error is a scale mixture of Gaussians. Sampling is accomplished with a (parameter-expanded) Gibbs sampler, mostly relying on a dynamic linear model representation.

Usage

```
fit_ASV(
  y,
  beta = 0,
  evol_error = "DHS",
  D = 1,
  nsave = 1000,
  nburn = 1000,
  nskip = 4,
  mcmc_params = list("h", "logy2hat", "sigma2", "evol_sigma_t2", "dhs_phi", "dhs_mean"),
  nugget = FALSE,
  computeDIC = TRUE,
  verbose = TRUE
)
```

Arguments

<code>y</code>	the $T \times 1$ vector of time series observations.
<code>beta</code>	the mean of the observed process y . If not provided, they are assumed to be 0.
<code>evol_error</code>	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
<code>D</code>	degree of differencing ($D = 1$, or $D = 2$)
<code>nsave</code>	number of MCMC iterations to record
<code>nburn</code>	number of MCMC iterations to discard (burn-in)
<code>nskip</code>	number of MCMC iterations to skip between saving iterations, i.e., save every (<code>nskip + 1</code>)th draw
<code>mcmc_params</code>	named list of parameters for which we store the MCMC output; must be one or more of:

- "h" (Log variance)
- "h_smooth" (smooth estimate of log variances. Only used when nugget_asv = TRUE)
- "logy2hat" (posterior predictive distribution of $\log(y^2)$)
- "sigma2" (Variance, i.e. $\exp(h)$)
- "evol_sigma_t2" (evolution error variance)
- "dhs_phi" (DHS AR(1) coefficient)
- "dhs_mean" (DHS AR(1) unconditional mean)

nugget	logical; if TRUE, fits the nugget variant of the ASV model
computeDIC	logical; if TRUE, compute the deviance information criterion DIC and the effective number of parameters p_d
verbose	logical; should R report extra information on progress?

Value

A named list of the `nsave` MCMC samples for the parameters named in `mcmc_params`

Note

The data `y` may contain NAs, which will be treated with a simple imputation scheme via an additional Gibbs sampling step. In general, rescaling `y` to have unit standard deviation is recommended to avoid numerical issues.

`fit_paramsASV`
Helper function for Sampling parameters for ASV model

Description

Helper function for Sampling parameters for ASV model

Usage

```
fit_paramsASV(data, sParams, evol_error, D)
```

Arguments

<code>data</code>	the $T \times 1$ vector of time series observations.
<code>sParams</code>	list from the previous run of <code>fit_paramsASV</code> function or <code>init_paramsASV</code> function
<code>evol_error</code>	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
<code>D</code>	degree of differencing ($D = 1$, or $D = 2$)

Value

a list containing 4 sets of parameters

- `s_p_error_term`: matrix containing mean and the variance from 10-component gaussian mixture (Omori et al. 2007)
- `s_mu`: a vector containing the posterior sample of log variance h ,
- `s_evolParams0`: a list containing posterior samples of parameters associated with the variance of first D observation of the log variance term, h .
- `s_evolParams`: a list containing posterior samples parameters associated with the variance of D to the last observations of the log variance term, h .

<code>fit_paramsASV_n</code>	<i>Helper function for Sampling parameters for ASV model with a nugget Effect</i>
------------------------------	---

Description

Helper function for Sampling parameters for ASV model with a nugget Effect

Usage

```
fit_paramsASV_n(data, sParams, evol_error, D)
```

Arguments

<code>data</code>	the $T \times 1$ vector of time series observations.
<code>sParams</code>	list from the previous run of <code>fit_paramsASV</code> function or <code>init_paramsASV</code> function
<code>evol_error</code>	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
<code>D</code>	degree of differencing ($D = 1$, or $D = 2$)

Value

a list containing 4 sets of parameters

- `s_p_error_term`: matrix containing mean and the variance from 10-component gaussian mixture (Omori et al. 2007)
- `s_mu`: a vector containing the posterior sample of log variance h ,
- `s_evolParams0`: a list containing posterior samples of parameters associated with the variance of first D observation of the log variance term, h .
- `s_evolParams`: a list containing posterior samples parameters associated with the variance of D to the last observations of the log variance term, h .

generate_ly2hat	<i>Posterior predictive sampler on the transformed y (log(y^2))</i>
-----------------	---

Description

Posterior predictive sampler on the transformed y (log(y^2))

Usage

```
generate_ly2hat(h, p_error_term)
```

Arguments

h	the log varaince term h
p_error_term	2 dimensional data frame containing mean and the variance from the 10 componet Gaussian mixture in Omori et al 2007 paper.

Value

a vector containing posterior predictive on log(y^2)

getARpXmat	<i>Compute the design matrix X for AR(p) model</i>
------------	--

Description

Compute the design matrix X for AR(p) model

Usage

```
getARpXmat(y, p = 1, include_intercept = FALSE)
```

Arguments

y	(T x 1) vector of responses
p	order of AR(p) model
include_intercept	logical; if TRUE, first column of X is ones

getEffSize	<i>Summarize of effective sample size</i>
------------	---

Description

Compute the summary statistics for the effective sample size (ESS) across posterior samples for possibly many variables

Usage

```
getEffSize(postX)
```

Arguments

postX	An array of arbitrary dimension (nsims × ... × ...), where nsims is the number of posterior samples
-------	---

Value

Table of summary statistics using the function `summary()`.

getNonZeros	<i>Compute Non-Zeros (Signals)</i>
-------------	------------------------------------

Description

Estimate the location of non-zeros (signals) implied by horseshoe-type thresholding.

Usage

```
getNonZeros(post_evol_sigma_t2, post_obs_sigma_t2 = NULL)
```

Arguments

post_evol_sigma_t2	the Nsims × T or Nsims × T × p matrix/array of posterior draws of the evolution error variances.
post_obs_sigma_t2	the Nsims × 1 or Nsims × T matrix of posterior draws of the observation error variances.

Details

Thresholding is based on $\kappa[t] > 1/2$, where $\kappa = 1/(1 + \text{evol_sigma_t2}/\text{obs_sigma_t2})$, `evol_sigma_t2` is the evolution error variance, and `obs_sigma_t2` is the observation error variance. In particular, the decision rule is based on the posterior mean of κ .

Value

A vector (or matrix) of indices identifying the signals according to the horseshoe-type thresholding rule.

Note

The thresholding rule depends on whether the prior variance for the state variable μ (i.e., `evol_sigma_t2`) is scaled by the observation standard deviation, `obs_sigma_t2`. Explicitly, if $\mu[t] \sim N(0, \text{evol_sigma_t2}[t])$ then the correct thresholding rule is based on $\kappa = 1/(1 + \text{evol_sigma_t2}/\text{obs_sigma_t2})$. However, if $\mu[t] \sim N(0, \text{evol_sigma_t2}[t]*\text{obs_sigma_t2}[t])$ then the correct thresholding rule is based on $\kappa = 1/(1 + \text{evol_sigma_t2})$. The latter case may be implemented by omitting the input for `post_obs_sigma_t2` (or setting it to NULL).

<code>initCholReg_spam</code>	<i>Compute initial Cholesky decomposition for TVP Regression</i>
-------------------------------	--

Description

Computes the Cholesky decomposition for the quadratic term in the (Gaussian) posterior of the TVP regression coefficients. The sparsity pattern will not change during the MCMC, so we can save computation time by computing this up front.

Usage

```
initCholReg_spam(obs_sigma_t2, evol_sigma_t2, XtX, D = 1)
```

Arguments

<code>obs_sigma_t2</code>	the $T \times 1$ vector of observation error variances
<code>evol_sigma_t2</code>	the $T \times p$ matrix of evolution error variances
<code>XtX</code>	the $Tp \times Tp$ matrix of $X'X$ (one-time cost; see <code>?build_XtX</code>)
<code>D</code>	the degree of differencing (one or two)

<code>initChol_spam</code>	<i>Compute initial Cholesky decomposition for Bayesian Trend Filtering</i>
----------------------------	--

Description

Computes the Cholesky decomposition for the quadratic term in the (Gaussian) posterior of the Bayesian Trend Filtering coefficients. The sparsity pattern will not change during the MCMC, so we can save computation time by computing this up front.

Usage

```
initChol_spam(nT, D = 1)
```

Arguments

nT	number of time points
D	degree of differencing (D = 1 or D = 2)

initDHS	<i>Initialize the evolution error variance parameters</i>
---------	---

Description

Compute initial values for evolution error variance parameters under the dynamic horseshoe prior

Usage

```
initDHS(omega)
```

Arguments

omega	T x p matrix of evolution errors
-------	----------------------------------

Value

List of relevant components: the T x p evolution error SD `sigma_wt`, the T x p log-volatility `ht`, the p x 1 log-vol unconditional mean(s) `dhs_mean`, the p x 1 log-vol AR(1) coefficient(s) `dhs_phi`, the T x p log-vol innovation SD `sigma_eta_t` from the PG priors, the p x 1 initial log-vol SD `sigma_eta_0`, and the mean of log-vol means `dhs_mean0` (relevant when $p > 1$)

initEvol0	<i>Initialize the parameters for the initial state variance</i>
-----------	---

Description

The initial state SDs are assumed to follow half-Cauchy priors, $C+(0,A)$, where the SDs may be common or distinct among the states.

Usage

```
initEvol0(mu0, commonSD = TRUE)
```

Arguments

mu0	p x 1 vector of initial values (undifferenced)
commonSD	logical; if TRUE, use common SDs (otherwise distinct)

Details

This function initializes the parameters for a PX-Gibbs sampler.

Value

List of relevant components: the $p \times 1$ evolution error SD σ_{w0} , the $p \times 1$ parameter-expanded RV's px_sigma_w0 , and the corresponding global scale parameters σ_{00} and px_sigma_00 (ignore if commonSD)

initEvolParams	<i>Initialize the evolution error variance parameters</i>
----------------	---

Description

Compute initial values for evolution error variance parameters under the various options: dynamic horseshoe prior ('DHS'), horseshoe prior ('HS'), Bayesian lasso ('BL'), normal stochastic volatility ('SV'), or normal-inverse-gamma prior ('NIG').

Usage

```
initEvolParams(omega, evol_error = "DHS")
```

Arguments

omega	T x p matrix of evolution errors
evol_error	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), or 'NIG' (normal-inverse-gamma prior)

Value

List of relevant components: σ_{wt} , the $T \times p$ matrix of evolution standard deviations, and additional parameters associated with the DHS and HS priors.

initSV	<i>Initialize the stochastic volatility parameters</i>
--------	--

Description

Compute initial values for normal stochastic volatility parameters. The model assumes an AR(1) for the log-volatility.

Usage

```
initSV(omega)
```

Arguments

omega	T x p matrix of errors
-------	------------------------

Value

List of relevant components: `sigma_wt`, the $T \times p$ matrix of standard deviations, and additional parameters (unconditional mean, AR(1) coefficient, and standard deviation).

<code>init_paramsASV</code>	<i>Helper function for initializing parameters for ASV model</i>
-----------------------------	--

Description

Helper function for initializing parameters for ASV model

Usage

```
init_paramsASV(data, evol_error, D)
```

Arguments

<code>data</code>	the $T \times 1$ vector of time series observations.
<code>evol_error</code>	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
<code>D</code>	degree of differencing ($D = 1$, or $D = 2$)

Value

a list containing 4 sets of parameters

- `s_p_error_term`: matrix containing mean and the variance from 10-component gaussian mixture (Omori et al. 2007)
- `s_mu`: a vector containing the posterior sample of log variance h ,
- `s_evolParams0`: a list containing posterior samples of parameters associated with the variance of first D observation of the log variance term, h .
- `s_evolParams`: a list containing posterior samples parameters associated with the variance of D to the last observations of the log variance term, h .

init_paramsASV_n	<i>Helper function for initializing parameters for ASV model with a nugget effect</i>
------------------	---

Description

Helper function for initializing parameters for ASV model with a nugget effect

Usage

```
init_paramsASV_n(data, evol_error, D)
```

Arguments

data	the $T \times 1$ vector of time series observations.
evol_error	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), 'BL' (Bayesian lasso), or 'NIG' (normal-inverse-gamma prior)
D	degree of differencing ($D = 1$, or $D = 2$)

Value

a list containing 4 sets of parameters

- s_p_error_term: matrix containing mean and the variance from 10-component gaussian mixture (Omori et al. 2007)
- s_mu: a vector containing the posterior sample of log variance h ,
- s_evolParams0: a list containing posterior samples of parameters associated with the variance of first D observation of the log variance term, h .
- s_evolParams: a list containing posterior samples parameters associated with the variance of D to the last observations of the log variance term, h .

invlogit	<i>Compute the inverse log-odds</i>
----------	-------------------------------------

Description

Compute the inverse log-odds

Usage

```
invlogit(x)
```

Arguments

x scalar or vector for which to compute the (componentwise) inverse log-odds

Value

A scalar or vector of values in (0,1)

logit	<i>Compute the log-odds</i>
-------	-----------------------------

Description

Compute the log-odds

Usage

logit(x)

Arguments

x scalar or vector in (0,1) for which to compute the (componentwise) log-odds

Value

A scalar or vector of log-odds

ncind	<i>Sample components from a discrete mixture of normals</i>
-------	---

Description

Sample Z from 1,2,...,k, with $P(Z=i)$ proportional to $q_i N(\mu_i, \text{sig}_i^2)$.

Usage

ncind(y, mu, sig, q)

Arguments

y vector of data
 mu vector of component means
 sig vector of component standard deviations
 q vector of component weights

Value

Sample from {1,...,k}

plot.dsp

Plot the Bayesian trend filtering fitted values

Description

Plot the BTF posterior mean of the conditional expectation with posterior credible intervals (point-wise and joint), the observed data, and true curves (if known)

Usage

```
## S3 method for class 'dsp'
plot(
  x,
  type,
  true_values = NULL,
  times = NULL,
  y_obs = NULL,
  include_joint_bands = FALSE,
  alpha = 0.05,
  xlab = NULL,
  ylab = NULL,
  main = NULL,
  xlim = NULL,
  ylim = NULL,
  mar = NULL,
  par_args = list(),
  legend = TRUE,
  legend_cex = 1,
  legend_pt_cex = 2,
  nr = NULL,
  nc = NULL,
  cp_thres = 0.5,
  ...
)
```

Arguments

<code>x</code>	an object of class <code>dsp</code> from <code>dsp_fit()</code> .
<code>type</code>	character string giving the parameter name to visualize; must be one of the entries in <code>x\$mcmc_output</code> .
<code>true_values</code>	optional ground-truth values to overlay on the plot. For scalar parameters, this should be a length-1 numeric value; for time-varying parameters, a $T \times 1$ vector; and for multi-parameter time-varying quantities, a $T \times p$ matrix matching the plotted parameter dimensions.
<code>times</code>	optional vector of observation points. If <code>NULL</code> , the function assumes T equally spaced points on $[0, 1]$.

y_obs	optional vector of observed data point of length T. Only for 2-dimensional parameters.
include_joint_bands	logical; if TRUE, include simultaneous credible bands in addition to pointwise credible intervals when available. Joint bands are currently supported only for time-varying parameters: zeta, omega, ypred, and mu for 2D outputs, and zeta, omega, ypred, mu, and beta for 3D outputs.
alpha	numeric credibility level used to construct posterior intervals and bands. Defaults to 0.05, corresponding to 95% intervals/bands.
xlab	optional x-axis label. If NULL, defaults to "t".
ylab	optional y-axis label. If NULL, defaults to type.
main	optional plot title. For multi-panel plots, this may be either a single title or a vector of titles of length equal to the number of panels.
xlim	optional x-axis limits passed to the plotting routine.
ylim	optional y-axis limits passed to the plotting routine.
mar	optional numeric vector of length 4 giving plot margins, passed to <code>graphics::par()</code> .
par_args	optional named list of additional graphical parameters passed to <code>graphics::par()</code> , such as <code>cex.axis</code> , <code>cex.lab</code> , <code>cex.main</code> , <code>las</code> , or <code>mgp</code> .
legend	logical; if TRUE, add a legend to the plot.
legend_cex	numeric scaling factor for legend text size.
legend_pt_cex	numeric scaling factor for legend symbol size.
nr	optional number of rows in the plotting layout for multi-panel (3-dimensional) parameters.
nc	optional number of columns in the plotting layout for multi-panel (3-dimensional) parameters.
cp_thres	(default 0.5) cutoff proportion for percentage of posterior samples exceeding the threshold needed to label a changepoint
...	additional graphical arguments passed to the main plotting call, such as <code>graphics::plot()</code> or <code>graphics::hist()</code> depending on the parameter dimension.

Details

The plotting behavior depends on the dimension of the posterior samples stored in `x$mcmc_output[[type]]`:

- **1D (scalar parameter):** A density-style summary is produced using a histogram with an overlaid kernel density estimate. The posterior mean and equal-tailed $(1 - \alpha)100\%$ credible interval are marked, and the true value is added if supplied through `true_values`.
- **2D (time-varying parameter):** A time-series plot is produced showing the posterior mean together with equal-tailed pointwise credible intervals. If `include_joint_bands = TRUE` and the parameter is one of `zeta`, `omega`, `ypred`, or `mu`, simultaneous credible bands are also displayed. If `true_values` is provided, the ground truth is overlaid.

- **3D (multi-parameter time-varying quantity):** A multi-panel collection of time-series plots is produced, one panel for each slice of the third dimension. Each panel shows the posterior mean, pointwise credible intervals, optional joint bands when supported, and optional ground-truth values. The panel layout is controlled by `nr` and `nc`; if omitted, a near-square layout is chosen automatically.

Axis labels, titles, plotting limits, margins, legend display, and additional graphical settings can be customized through `xlab`, `ylab`, `main`, `xlim`, `ylim`, `mar`, `legend`, `legend_cex`, `legend_pt_cex`, and `par_args`.

The x-axis values are taken from `times`. If `times` is not supplied, evenly spaced points on $[0, 1]$ are used. For differenced variance parameters such as `"evol_sigma_t2"` and `"zeta_sigma_t2"`, the initial time points associated with prior initialization are automatically removed before plotting.

For fitted changepoint models, changepoint annotations may be added when supported by the plotted parameter and the corresponding latent components are present in the MCMC output.

Value

No return value, called for side effects

References

Krivobokova, T., Kneib, T., and Claeskens, G. (2010). Simultaneous confidence bands for penalized spline estimators. *Journal of the American Statistical Association*, **105**(490), 852–863. doi:10.1198/jasa.2010.tm09165

Examples

```
set.seed(200)
signal = c(rep(0, 50), rep(10, 50))
noise = rep(1, 100)
noise_var = rep(1, 100)
for (k in 2:100){
  noise_var[k] = exp(0.9*log(noise_var[k-1]) + rnorm(1, 0, 0.5))
  noise[k] = rnorm(1, 0, sqrt(noise_var[k])) }

y = signal + noise
model_spec = dsp_spec(family = "gaussian", model = "changepoint",
                      D = 1, useAnom = TRUE, obsSV = "SV")
mcmc_output = dsp_fit(y, model_spec = model_spec, nsave = 500, nburn = 500)
# Estimated posterior mean vs ground truth
plot(mcmc_output, type = "mu", true_values = signal)
# Estimated innovation variance vs ground truth for illustration only
plot(mcmc_output, type = "obs_sigma_t2", true_values = noise^2)
```

predict.dsp

Predict changepoints from the output of ABCO

Description

Predict changepoints from the output of ABCO

Usage

```
## S3 method for class 'dsp'
predict(object, cp_thres = 0.5, cp_prop = FALSE, ...)
```

Arguments

object	object of class dsp from <code>dsp_fit()</code>
cp_thres	(default 0.5) cutoff proportion for percentage of posterior samples exceeding the threshold needed to label a changepoint
cp_prop	(default FALSE) logical flag determining if the posterior proportions of threshold exceedance is to be returned.
...	currently unused

Details

The changepoint model uses a thresholding mechanism with a latent indicator variable. This function calculates the proportion of samples where the increment exceeds the threshold.

Value

If `cp_prop = FALSE`, a numeric vector of indices that correspond to indices of the observed data. If `cp_prop = TRUE`, a list containing:

- 'cp_t': a numeric vector of indices that correspond to indices of the observed data.
- 'cp_prop': a numeric vector of length (T - D) with the pointwise proportion of samples where the increment exceeds the threshold.

If no proportions exceed `cp_thres`, then the vector will be a length 0 integer vector.

Examples

```
set.seed(200)
signal = c(rep(0, 50), rep(10, 50))
noise = rep(1, 100)
noise_var = rep(1, 100)
for (k in 2:100){
  noise_var[k] = exp(0.9*log(noise_var[k-1]) + rnorm(1, 0, 0.5))
  noise[k] = rnorm(1, 0, sqrt(noise_var[k])) }

y = signal + noise
```

```

model_spec = dsp_spec(family = "gaussian", model = "changeoint",
                      D = 1, useAnom = TRUE)
mcmc_output = dsp_fit(y, model_spec = model_spec, nsave = 500, nburn = 500)
predict(mcmc_output)

```

sampleAR1

Sample the AR(1) coefficient(s)

Description

Compute one draw of the AR(1) coefficient in a model with Gaussian innovations and time-dependent innovation variances. In particular, we use the sampler for the log-volatility AR(1) process with the parameter-expanded Polya-Gamma sampler. The sampler also applies to a multivariate case with independent components.

Usage

```
sampleAR1(h_yc, h_phi, h_sigma_eta_t, prior_dhs_phi = NULL)
```

Arguments

h_yc	the $T \times p$ matrix of centered log-volatilities (i.e., the log-vols minus the unconditional means <code>dhs_mean</code>)
h_phi	the $p \times 1$ vector of previous AR(1) coefficient(s)
h_sigma_eta_t	the $T \times p$ matrix of log-vol innovation standard deviations
prior_dhs_phi	the parameters of the prior for the log-volatility AR(1) coefficient <code>dhs_phi</code> ; either NULL for uniform on $[-1, 1]$ or a 2-dimensional vector of (shape1, shape2) for a Beta prior on $[(dhs_phi + 1)/2]$

Value

$p \times 1$ vector of sampled AR(1) coefficient(s)

Note

For the standard AR(1) case, $p = 1$. However, the function applies more generally for sampling $p > 1$ independent AR(1) processes (jointly).

sampleBTF	<i>Sampler for first or second order random walk (RW) Gaussian dynamic linear model (DLM)</i>
-----------	---

Description

Compute one draw of the $T \times 1$ state variable μ in a DLM using back-band substitution methods. This model is equivalent to the Bayesian trend filtering (BTF) model, assuming appropriate (shrinkage/sparsity) priors for the evolution errors.

Usage

```
sampleBTF(
  y,
  obs_sigma_t2,
  evol_sigma_t2,
  D = 1,
  loc_obs = NULL,
  chol0 = NULL,
  prior_mean = NULL
)
```

Arguments

<code>y</code>	the $T \times 1$ vector of time series observations
<code>obs_sigma_t2</code>	the $T \times 1$ vector of observation error variances
<code>evol_sigma_t2</code>	the $T \times 1$ vector of evolution error variances
<code>D</code>	the degree of differencing (one or two)
<code>loc_obs</code>	list of the row and column indices to fill in a band-sparse matrix
<code>chol0</code>	(optional) the $m \times m$ matrix of initial Cholesky factorization; if NULL, use the Matrix package for sampling, otherwise use the spam package
<code>prior_mean</code>	optional (default is NULL); numeric $T \times 1$ vector specifying the prior mean of μ

Value

$T \times 1$ vector of simulated states

Note

Missing entries (NAs) are not permitted in y . Imputation schemes are available.

sampleBTF_bspline	<i>Sampler for first or second order random walk (RW) Gaussian dynamic linear model (DLM)</i>
-------------------	---

Description

Compute one draw of the $p \times 1$ B-spline basis coefficients β in a DLM using back-band substitution methods. The coefficients are penalized with a prior on the $D = 0$, $D = 1$, or $D = 2$ differences. This model is equivalent to the Bayesian trend filtering (BTF) model applied to $p \times 1$ vector of equally-spaced B-spline coefficients, with the basis matrix serving as a design matrix in the observation equation.

Usage

```
sampleBTF_bspline(
  y,
  X,
  obs_sigma2,
  evol_sigma_t2,
  XtX_bands,
  Xty = NULL,
  D = 1
)
```

Arguments

<code>y</code>	the $T \times 1$ vector of time series observations
<code>X</code>	the $T \times p$ basis matrix
<code>obs_sigma2</code>	the scalar observation error variance
<code>evol_sigma_t2</code>	the $p \times 1$ vector of evolution error variances
<code>XtX_bands</code>	list with 4 vectors consisting of the 4-bands of $XtX = \text{crossprod}(X)$ (one-time cost)
<code>Xty</code>	the $p \times 1$ matrix $\text{crossprod}(X,y)$, which is a one-time cost (assuming no missing entries in y)
<code>D</code>	the degree of differencing (zero, one, or two)

Value

$p \times 1$ vector of simulated basis coefficients β

Note

Missing entries (NAs) are not permitted in y . Imputation schemes are available.

sampleBTF_reg	<i>Sampler for first or second order random walk (RW) Gaussian dynamic linear model (DLM)</i>
---------------	---

Description

Compute one draw of the $T \times p$ state variable β in a DLM using back-band substitution methods. This model is equivalent to the Bayesian trend filtering (BTF) model applied to p dynamic regression coefficients corresponding to the design matrix X , assuming appropriate (shrinkage/sparsity) priors for the evolution errors.

Usage

```
sampleBTF_reg(y, X, obs_sigma_t2, evol_sigma_t2, XtX, D = 1, chol0 = NULL)
```

Arguments

<code>y</code>	the $T \times 1$ vector of time series observations
<code>X</code>	the $T \times p$ matrix of time series predictors
<code>obs_sigma_t2</code>	the $T \times 1$ vector of observation error variances
<code>evol_sigma_t2</code>	the $T \times p$ matrix of evolution error variances
<code>XtX</code>	the $Tp \times Tp$ matrix of $X'X$ (one-time cost; see <code>?build_XtX</code>)
<code>D</code>	the degree of differencing (one or two)
<code>chol0</code>	(optional) the $m \times m$ matrix of initial Cholesky factorization; if <code>NULL</code> , use the <code>Matrix</code> package for sampling, otherwise use the <code>spam</code> package

Value

$T \times p$ matrix of simulated dynamic regression coefficients β

Note

Missing entries (NAs) are not permitted in `y`. Imputation schemes are available.

sampleBTF_reg_backfit	<i>(Backfitting) Sampler for first or second order random walk (RW) Gaussian dynamic linear model (DLM)</i>
-----------------------	---

Description

Compute one draw of the $T \times p$ state variable β in a DLM using back-band substitution methods. This model is equivalent to the Bayesian trend filtering (BTF) model applied to p dynamic regression coefficients corresponding to the design matrix X , assuming appropriate (shrinkage/sparsity) priors for the evolution errors. The sampler here uses a backfitting method that draws each predictor $j=1, \dots, p$ conditional on the other predictors (and coefficients), which leads to a faster $O(Tp)$ algorithm. However, the MCMC may be less efficient.

Usage

```
sampleBTF_reg_backfit(y, X, beta, obs_sigma_t2, evol_sigma_t2, D = 1)
```

Arguments

`y` the $T \times 1$ vector of time series observations
`X` the $T \times p$ matrix of time series predictors
`beta` the $T \times p$ matrix of previous dynamic regression coefficients
`obs_sigma_t2` the $T \times 1$ vector of observation error variances
`evol_sigma_t2` the $T \times p$ matrix of evolution error variances
`D` the degree of differencing (one or two)

Value

$T \times p$ matrix of simulated dynamic regression coefficients `beta`

Note

Missing entries (NAs) are not permitted in `y`. Imputation schemes are available.

sampleBTF_sparse	<i>Sampler for first or second order random walk (RW) Gaussian dynamic linear model (DLM) with additional shrinkage to zero</i>
------------------	---

Description

Compute one draw of the $T \times 1$ state variable `mu` in a DLM using back-band substitution methods. This model is equivalent to the Bayesian trend filtering (BTF) model, assuming appropriate (shrinkage/sparsity) priors for the evolution errors, with an additional shrinkage-to-zero prior.

Usage

```
sampleBTF_sparse(  
  y,  
  obs_sigma_t2,  
  evol_sigma_t2,  
  zero_sigma_t2,  
  D = 1,  
  chol0 = NULL  
)
```

Arguments

y	the T x 1 vector of time series observations
obs_sigma_t2	the T x 1 vector of observation error variances
evol_sigma_t2	the T x 1 vector of evolution error variances
zero_sigma_t2	the T x 1 vector of shrink-to-zero variances
D	the degree of differencing (one or two)
chol0	(optional) the m x m matrix of initial Cholesky factorization; if NULL, use the Matrix package for sampling, otherwise use the spam package

Value

T x 1 vector of simulated states

Note

Missing entries (NAs) are not permitted in y. Imputation schemes are available.

sampleDSP

Sample the dynamic shrinkage process parameters

Description

Compute one draw for each of the parameters in the dynamic shrinkage process for the special case in which the shrinkage parameter $\kappa \sim \text{Beta}(\alpha, \beta)$ with $\alpha = \beta$. The primary example is the dynamic horseshoe process with $\alpha = \beta = 1/2$.

Usage

```
sampleDSP(
  omega,
  evolParams,
  sigma_e = 1,
  loc = NULL,
  prior_dhs_phi = c(10, 2),
  alphaPlusBeta = 1
)
```

Arguments

omega	T x p matrix of evolution errors
evolParams	list of parameters to be updated (see Value below)
sigma_e	the observation error standard deviation; for (optional) scaling purposes
loc	list of the row and column indices to fill in a band-sparse matrix

prior_dhs_phi	the parameters of the prior for the log-volatility AR(1) coefficient dhs_phi; either NULL for uniform on [-1,1] or a 2-dimensional vector of (shape1, shape2) for a Beta prior on [(dhs_phi + 1)/2]
alphaPlusBeta	For the symmetric prior kappa ~ Beta(alpha, beta) with alpha=beta, specify the sum [alpha + beta]

Value

List of relevant components:

- the $T \times p$ evolution error standard deviations `sigma_wt`,
- the $T \times p$ log-volatility `ht`, the $p \times 1$ log-vol unconditional mean(s) `dhs_mean`,
- the $p \times 1$ log-vol AR(1) coefficient(s) `dhs_phi`,
- the $T \times p$ log-vol innovation standard deviations `sigma_eta_t` from the Polya-Gamma priors,
- the $p \times 1$ initial log-vol SD `sigma_eta_0`,
- and the mean of log-vol means `dhs_mean0` (relevant when $p > 1$)

Note

The priors induced by `prior_dhs_phi` all imply a stationary (log-) volatility process.

sampleEvol0

Sample the parameters for the initial state variance

Description

The initial state SDs are assumed to follow half-Cauchy priors, $C+(0,A)$, where the SDs may be common or distinct among the states.

Usage

```
sampleEvol0(mu0, evolParams0, commonSD = FALSE, A = 1)
```

Arguments

<code>mu0</code>	$p \times 1$ vector of initial values (undifferenced)
<code>evolParams0</code>	list of relevant components (see below)
<code>commonSD</code>	logical; if TRUE, use common SDs (otherwise distinct)
<code>A</code>	prior scale parameter from the half-Cauchy prior, $C+(0,A)$

Details

This function samples the parameters for a PX-Gibbs sampler.

Value

List of relevant components: the $p \times 1$ evolution error SD `sigma_w0` and the $p \times 1$ parameter-expanded RV's `px_sigma_w0`

sampleEvolParams	<i>Sampler evolution error variance parameters</i>
------------------	--

Description

Compute one draw of evolution error variance parameters under the various options:

- dynamic horseshoe prior ('DHS');
- horseshoe prior ('HS');
- normal-inverse-gamma prior ('NIG').

Usage

```
sampleEvolParams(
  omega,
  evolParams,
  sigma_e = 1,
  evol_error = "DHS",
  loc = NULL
)
```

Arguments

omega	T x p matrix of evolution errors
evolParams	list of parameters pertaining to each evol_error type to be updated
sigma_e	the observation error standard deviation; for (optional) scaling purposes
evol_error	the evolution error distribution; must be one of 'DHS' (dynamic horseshoe prior), 'HS' (horseshoe prior), or 'NIG' (normal-inverse-gamma prior)
loc	list of the row and column indices to fill in a band-sparse matrix

Value

List of relevant components in evolParams: sigma_wt, the T x p matrix of evolution standard deviations, and additional parameters associated with the DHS and HS priors.

Note

The list evolParams is specific to each evol_error type, but in each case contains the evolution error standard deviations sigma_wt.

To avoid scaling by the observation standard deviation sigma_e, simply use sigma_e = 1 in the functional call.

sampleFastGaussian	<i>Sample a Gaussian vector using the fast sampler of BHATTACHARYA et al.</i>
--------------------	---

Description

Sample from $N(\mu, \Sigma)$ where $\Sigma = \text{solve}(\text{crossprod}(\Phi) + \text{solve}(D))$ and $\mu = \Sigma * \text{crossprod}(\Phi, \alpha)$:

Usage

```
sampleFastGaussian(Phi, Ddiag, alpha)
```

Arguments

Phi	$n \times p$ matrix (of predictors)
Ddiag	$p \times 1$ vector of diagonal components (of prior variance)
alpha	$n \times 1$ vector (of data, scaled by variance)

Value

Draw from $N(\mu, \Sigma)$, which is $p \times 1$, and is computed in $O(n^2 * p)$

Note

Assumes D is diagonal, but extensions are available

sampleLogVolMu	<i>Sample the AR(1) unconditional means</i>
----------------	---

Description

Compute one draw of the unconditional means in an AR(1) model with Gaussian innovations and time-dependent innovation variances. In particular, we use the sampler for the log-volatility AR(1) process with the parameter-expanded Polya-Gamma sampler. The sampler also applies to a multivariate case with independent components.

Usage

```
sampleLogVolMu(h, h_mu, h_phi, h_sigma_eta_t, h_sigma_eta_0, h_log_scale = 0)
```

Arguments

h	the $T \times p$ matrix of log-volatilities
h_mu	the $p \times 1$ vector of previous means
h_phi	the $p \times 1$ vector of AR(1) coefficient(s)
h_sigma_eta_t	the $T \times p$ matrix of log-vol innovation standard deviations
h_sigma_eta_0	the standard deviations of initial log-vols
h_log_scale	prior mean from scale mixture of Gaussian (Polya-Gamma) prior, e.g. $\log(\sigma_e^2)$ or dhs_mean0

Value

a list containing

- the sampled mean(s) dhs_mean and
- the sampled precision(s) dhs_mean_prec_j from the Polya-Gamma parameter expansion

sampleLogVolMu0	<i>Sample the mean of AR(1) unconditional means</i>
-----------------	---

Description

Compute one draw of the mean of unconditional means in an AR(1) model with Gaussian innovations and time-dependent innovation variances (for $p > 1$). More generally, the sampler applies to the "mean" parameter (on the log-scale) for a Polya-Gamma parameter expanded hierarchical model.

Usage

```
sampleLogVolMu0(h_mu, h_mu0, dhs_mean_prec_j, h_log_scale = 0)
```

Arguments

h_mu	the $p \times 1$ vector of means
h_mu0	the previous mean of unconditional means
dhs_mean_prec_j	the $p \times 1$ vector of precisions (from the Polya-Gamma parameter expansion)
h_log_scale	prior mean from scale mixture of Gaussian (Polya-Gamma) prior, e.g. $\log(\sigma_e^2)$

Value

The sampled mean parameter dhs_mean0

Note

This sampler is particularly for $p > 1$ and the setting in which we want hierarchical shrinkage effects, e.g. predictor- and time-dependent shrinkage, predictor-dependent shrinkage, and global shrinkage, with a natural hierarchical ordering.

sampleLogVols

Sample the latent log-volatilities

Description

Compute one draw of the log-volatilities using a discrete mixture of Gaussians approximation to the likelihood (see Omori, Chib, Shephard, and Nakajima, 2007) where the log-vols are assumed to follow an AR(1) model with time-dependent innovation variances. More generally, the code operates for p independent AR(1) log-vol processes to produce an efficient joint sampler in $O(Tp)$ time.

Usage

```
sampleLogVols(
  h_y,
  h_prev,
  h_mu,
  h_phi,
  h_sigma_eta_t,
  h_sigma_eta_0,
  loc = NULL
)
```

Arguments

<code>h_y</code>	the $T \times p$ matrix of data, which follow independent SV models
<code>h_prev</code>	the $T \times p$ matrix of the previous log-vols
<code>h_mu</code>	the $p \times 1$ vector of log-vol unconditional means
<code>h_phi</code>	the $p \times 1$ vector of log-vol AR(1) coefficients
<code>h_sigma_eta_t</code>	the $T \times p$ matrix of log-vol innovation standard deviations
<code>h_sigma_eta_0</code>	the $p \times 1$ vector of initial log-vol innovation standard deviations
<code>loc</code>	list of the row and column indices to fill in a band-sparse matrix

Value

$T \times p$ matrix of simulated log-vols

Note

For Bayesian trend filtering, $p = 1$. More generally, the sampler allows for $p > 1$ but assumes (contemporaneous) independence across the log-vols for $j = 1, \dots, p$.

sampleSVparams	<i>Sampler for the stochastic volatility parameters</i>
----------------	---

Description

Compute one draw of the normal stochastic volatility parameters. The model assumes an AR(1) for the log-volatility.

Usage

```
sampleSVparams(omega, svParams)
```

Arguments

omega	T x p matrix of errors
svParams	list of parameters to be updated

Value

List of relevant components in svParams: sigma_wt, the T x p matrix of standard deviations, and additional parameters associated with SV model.

sampleSVparams0	<i>Sampler for the stochastic volatility parameters using same functions as DHS prior</i>
-----------------	---

Description

Compute one draw of the normal stochastic volatility parameters. The model assumes an AR(1) for the log-volatility.

Usage

```
sampleSVparams0(omega, svParams)
```

Arguments

omega	T x p matrix of errors
svParams	list of parameters to be updated

Value

List of relevant components in svParams: sigma_wt, the T x p matrix of standard deviations, and additional parameters associated with SV model.

sample_j_wrap	<i>Sampling from 10-component Gaussian Mixture component described in Omori et al. 2007</i>
---------------	---

Description

Samples from the conditional posterior distribution of the $\log(\chi^2)$ distribution by approximating it with the mixture Gaussian distribution described in Omori et al. 2007.

Usage

```
sample_j_wrap(Td, obs = NULL)
```

Arguments

Td	length of the vector
obs	Td x 1 vector for the data.

Value

Dataframe containing the posterior samples: mean and variance for the mixture component.

Note

When the obs is not not specified, the components are samples from the prior distribution.

sample_mat_c	<i>Wrapper function for C++ call for sample mat, check pre-conditions to prevent crash</i>
--------------	--

Description

Wrapper function for C++ call for sample mat, check pre-conditions to prevent crash

Usage

```
sample_mat_c(row_ind, col_ind, mat_val, mat_l, num_inp, linht, rd, D)
```

Arguments

row_ind	list of the row indices to fill in the bandsparse matrix
col_ind	list of the columns indices to fill in the bandsparse matrix
mat_val	list of the values to fill in the bandsparse matrix
mat_l	dimension of the band-sparse matrix
num_inp	number of non-zero elements in the bandsparse matrix
linht	T-D vector of linear term in the sampler
rd	T-D vector of standard normal noise samples
D	the degree of differencing for changepoint

 simBaS

Compute Simultaneous Band Scores (SimBaS)

Description

Compute simultaneous band scores (SimBaS) from Meyer et al. (2015, Biometrics). SimBaS uses MC(MC) simulations of a function of interest to compute the minimum alpha such that the joint credible bands at the alpha level do not include zero. This quantity is computed for each grid point (or observation point) in the domain of the function.

Usage

```
simBaS(sampFuns)
```

Arguments

sampFuns	Nsims x m matrix of Nsims MCMC samples and m points along the curve
----------	---

Value

m x 1 vector of simBaS

Note

The input needs not be curves: the simBaS may be computed for vectors to achieve a multiplicity adjustment.

The minimum of the returned value, P_{simBaS_t} , over the domain t is the Global Bayesian P-Value (GBP) for testing whether the function is zero everywhere.

<code>simRegression</code>	<i>Simulate noisy observations from a dynamic regression model</i>
----------------------------	--

Description

Simulates data from a time series regression with dynamic regression coefficients. The dynamic regression coefficients are simulated as a Gaussian random walk, where jumps occur with a pre-specified probability sparsity. The coefficients are initialized by a $N(0,1)$ simulation.

Usage

```
simRegression(
  nT = 200,
  p = 20,
  p_0 = 15,
  sparsity = 0.05,
  RSNR = 5,
  ar1 = 0,
  include_plot = FALSE
)
```

Arguments

<code>nT</code>	number of time points
<code>p</code>	number of predictors (total)
<code>p_0</code>	number of true zero regression terms
<code>sparsity</code>	the probability of a jump (i.e., a change in the dynamic regression coefficient)
<code>RSNR</code>	root-signal-to-noise ratio
<code>ar1</code>	the AR(1) coefficient for the predictors X; default is zero for iid $N(0,1)$ predictors
<code>include_plot</code>	logical; if TRUE, include a plot of the simulated data and the true curve

Value

a list containing

- the simulated function `y`
- the simulated predictors `X`
- the simulated dynamic regression coefficients `beta_true`
- the true function `mu_true`
- the true observation standard deviation `sigma_true`

Note

The root-signal-to-noise ratio is defined as $RSNR = (\text{sd of true function})/(\text{sd of noise})$.

<code>simRegression0</code>	<i>Simulate noisy observations from a dynamic regression model</i>
-----------------------------	--

Description

Simulates data from a time series regression with dynamic regression coefficients. The dynamic regression coefficients are selected using the options from the `simUnivariate()` function in the `wmtsa` package.

Usage

```
simRegression0(
  signalNames = c("bumps", "blocks"),
  nT = 200,
  RSNR = 10,
  p_0 = 5,
  include_intercept = TRUE,
  scale_all = TRUE,
  include_plot = TRUE,
  ar1 = 0
)
```

Arguments

<code>signalNames</code>	vector of strings matching the "name" argument in the <code>simUnivariate()</code> function, e.g. "bumps" or "doppler"
<code>nT</code>	number of points
<code>RSNR</code>	root-signal-to-noise ratio
<code>p_0</code>	number of true zero regression terms to include
<code>include_intercept</code>	logical; if TRUE, the first column of X is 1's
<code>scale_all</code>	logical; if TRUE, scale all regression coefficients to [0,1]
<code>include_plot</code>	logical; if TRUE, include a plot of the simulated data and the true curve
<code>ar1</code>	the AR(1) coefficient for the predictors X; default is zero for iid N(0,1) predictors

Value

a list containing

- the simulated function `y`
- the simulated predictors `X`
- the simulated dynamic regression coefficients `beta_true`
- the true function `mu_true`
- the true observation standard deviation `sigma_true`

Note

The number of predictors is $p = \text{length}(\text{signalNames}) + p_0$.

The root-signal-to-noise ratio is defined as $\text{RSNR} = (\text{sd of true function})/(\text{sd of noise})$.

simUnivariate *Generate univariate signals of different type*

Description

Using code from the archived wmtsa package

Usage

```
simUnivariate(name, n = 1024, snr = Inf)
```

Arguments

name	character string of name of the test wavelet signal to be generated; one of "dirac", "kronecker", "heavisine", "bumps", "blocks", "doppler", "ramp", "cusp", "crease", "sing", "hisine", "losine", "linchirp", "twochirp", "quadchirp", "mishmash1", "mishmash2", "mishmash3", "levelshift", "jumpsine", "gauss", "patches", "linear", "quadratic", "cubic";
n	length of the series; defaults to 1024 points; increasing n infills the time series
snr	desired signal-to-noise ratio; default Inf corresponds to 0 noise

Value

A numeric vector the same length as n.

Examples

```
nms <- c("blocks", "linchirp", "mishmash1", "bumps")
z <- lapply(nms, simUnivariate)
```

spec_dsp	<i>Compute the spectrum of an AR(p) model</i>
----------	---

Description

Compute the spectrum of an AR(p) model

Usage

```
spec_dsp(ar_coefs, sigma_e, n.freq = 500)
```

Arguments

ar_coefs	(p x 1) vector of AR(p) coefficients
sigma_e	observation standard deviation
n.freq	number of frequencies at which to evaluate the spectrum

Value

A (n.freq x 2) matrix where the first column is the frequencies and the second column is the spectrum evaluated at that frequency

summary.dsp	<i>Summarize DSP MCMC chains</i>
-------------	----------------------------------

Description

Summarize DSP MCMC chains

Usage

```
## S3 method for class 'dsp'
summary(object, pars, probs = c(0.025, 0.25, 0.5, 0.75, 0.975), ...)
```

Arguments

object	object of class dsp from <code>dsp_fit()</code>
pars	parameter names specified for summaries; currently defaults to all parameters named in <code>object\$mcmc_output</code>
probs	numeric vector of <code>quantile()</code> s requested for posterior summary of pars. Defaults to <code>c(0.025, 0.25, 0.50, 0.75, 0.975)</code>
...	currently not being used

Value

Returns a named list of the same length as pars where within each element of the list is a numeric matrix (vector parameters) or vector (scalar parameters). For matrices, each row is a time point (or dimension) of the parameter and each column is a named summary. The names are accessible with colnames. For vectors (scalar parameters), each element is a named summary.

Examples

```
set.seed(200)
signal = c(rep(0, 50), rep(10, 50))
noise = rep(1, 100)
noise_var = rep(1, 100)
for (k in 2:100){
  noise_var[k] = exp(0.9*log(noise_var[k-1]) + rnorm(1, 0, 0.5))
  noise[k] = rnorm(1, 0, sqrt(noise_var[k])) }

y = signal + noise
model_spec = dsp_spec(family = "gaussian", model = "changepoint",
                      D = 1, useAnom = TRUE, obsSV = "SV")
mcmc_output = dsp_fit(y, model_spec = model_spec, nsave = 500, nburn = 500)

summary_fit <- summary(mcmc_output)
summary_fit$mu["mean"]
summary_fit$evol_sigma_t2["mean"]
```

t_create_loc

Initializer for location indices for filling in band-sparse matrix

Description

Create row and column indices for locations of symmetric band-sparse matrix. Starts with the locations of the diagonal, proceed with upper-diagonals, followed by lower-diagonals.

Usage

```
t_create_loc(len, D)
```

Arguments

len	length of the diagonal of the band-sparse matrix
D	number of super-diagonals to include for the band-sparse

Value

a list containing

- the row indices r and
- the column indices c

t_initEvolParams_no *Initialize the evolution error variance parameters*

Description

Compute initial values for evolution error variance parameters under the dynamic horseshoe prior

Usage

t_initEvolParams_no(y, D, omega)

Arguments

y	the T vector of time series observations
D	degree of differencing (D = 1, or D = 2)
omega	T vector of evolution errors

Value

List of relevant components: sigma_wt, the T vector of evolution standard deviations, and additional parameters associated with the DHS priors.

t_initEvolZeta_ps *Initialize the anomaly component parameters*

Description

Compute initial values for either a horseshoe prior or horseshoe+ prior for the anomaly component.

Usage

t_initEvolZeta_ps(zeta)

Arguments

zeta	T vector of initial estimates.
------	--------------------------------

Value

List of relevant components: sigma_wt, the T vector of standard deviations, and additional parameters for inverse gamma priors (shape and scale).

t_initSV	<i>Initialize the stochastic volatility parameters</i>
----------	--

Description

Compute initial values for normal stochastic volatility parameters. The model assumes an AR(1) for the log-volatility.

Usage

```
t_initSV(omega)
```

Arguments

omega	T vector of errors
-------	--------------------

Value

List of relevant components: sigma_wt, the T vector of standard deviations, and additional parameters (unconditional mean, AR(1) coefficient, and standard deviation).

t_sampleAR1	<i>Sample the TAR(1) coefficients</i>
-------------	---------------------------------------

Description

Compute one draw of the TAR(1) coefficients in a model with Gaussian innovations and time-dependent innovation variances. In particular, we use the sampler for the log-volatility TAR(1) process with the parameter-expanded Polya-Gamma sampler. The sampler also applies to a multivariate case with independent components.

Usage

```
t_sampleAR1(h_yc, h_phi, h_phi2, h_sigma_eta_t, h_st, prior_dhs_phi = NULL)
```

Arguments

h_yc	the T vector of centered log-volatilities (i.e., the log-vols minus the unconditional means dhs_mean)
h_phi	the 1 vector of previous AR(1) coefficient(s)
h_phi2	the 1 vector of previous penalty coefficient(s)
h_sigma_eta_t	the T vector of log-vol innovation standard deviations
h_st	the T vector of indicators on whether each time-step exceed the estimated threshold
prior_dhs_phi	the parameters of the prior for the log-volatility AR(1) coefficient dhs_phi; either NULL for uniform on [-1,1] or a 2-dimensional vector of (shape1, shape2) for a Beta prior on [(dhs_phi + 1)/2]

Value

2 vector of sampled TAR(1) coefficient(s)

t_sampleBTF	<i>Sampler for first or second order random walk (RW) Gaussian dynamic linear model (DLM)</i>
-------------	---

Description

Compute one draw of the T state variable μ in a DLM using back-band substitution methods. This model is equivalent to the Bayesian trend filtering (BTF) model, assuming appropriate (shrinkage/sparsity) priors for the evolution errors.

Usage

```
t_sampleBTF(y, obs_sigma_t2, evol_sigma_t2, D = 1, loc_obs)
```

Arguments

y	the T x 1 vector of time series observations
obs_sigma_t2	the T x 1 vector of observation error variances
evol_sigma_t2	the T x 1 vector of evolution error variances
D	the degree of differencing (one or two)
loc_obs	list of the row and column indices to fill in a band-sparse matrix

Value

T x 1 vector of simulated states

Note

Missing entries (NAs) are not permitted in y. Imputation schemes are available.

t_sampleEvolParams *Sample the thresholded dynamic shrinkage process parameters*

Description

Compute one draw for each of the parameters in the thresholded dynamic shrinkage process for the special case in which the shrinkage parameter $\kappa \sim \text{Beta}(\alpha, \beta)$ with $\alpha = \beta = 1/2$.

Usage

```
t_sampleEvolParams(
  omega,
  evolParams,
  D = 1,
  sigma_e = 1,
  lower_b,
  upper_b,
  loc,
  prior_dhs_phi = c(20, 1),
  alphaPlusBeta = 1
)
```

Arguments

omega	T vector of evolution errors
evolParams	list of parameters to be updated (see Value below)
D	the degree of differencing (one or two)
sigma_e	the observation error standard deviation; for (optional) scaling purposes
lower_b	the lower bound in the uniform prior of the threshold variable
upper_b	the upper bound in the uniform prior of the threshold variable
loc	list of the row and column indices to fill in a band-sparse matrix
prior_dhs_phi	the parameters of the prior for the log-volatility AR(1) coefficient dhs_phi; either NULL for uniform on [-1,1] or a 2-dimensional vector of (shape1, shape2) for a Beta prior on [(dhs_phi + 1)/2]
alphaPlusBeta	For the symmetric prior $\kappa \sim \text{Beta}(\alpha, \beta)$ with $\alpha = \beta$, specify the sum [alpha + beta]

Value

List of relevant components:

- the T evolution error standard deviations sigma_wt,
- the T log-volatility ht,

- the 1 log-vol unconditional mean(s) dhs_mean,
- the 1 log-vol AR(1) coefficient(s) dhs_phi,
- the 1 log-vol correction coefficient(s) dhs_phi2,
- the T log-vol innovation standard deviations sigma_eta_t from the Polya-Gamma priors,
- the 1 initial log-vol SD sigma_eta_0,
- the 1 threshold parameter r

Note

The priors induced by prior_dhs_phi all imply a stationary (log-) volatility process.

t_sampleEvolZeta_ps *Sampler for the anomaly component parameters*

Description

Compute one draw of the anomaly component parameters.

Usage

```
t_sampleEvolZeta_ps(omega, evolParams)
```

Arguments

omega	T vector of errors
evolParams	list of parameters to be updated

Value

List of relevant components in evolParams: sigma_wt, the T vector of standard deviations, and additional parameters for inverse gamma priors (shape and scale).

t_sampleLogVolMu *Sample the TAR(1) unconditional means*

Description

Compute one draw of the unconditional means in an TAR(1) model with Gaussian innovations and time-dependent innovation variances. In particular, we use the sampler for the log-volatility TAR(1) process with the parameter-expanded Polya-Gamma sampler. The sampler also applies to a multivariate case with independent components.

Usage

```
t_sampleLogVolMu(
  h,
  h_mu,
  h_phi,
  h_phi2,
  h_sigma_eta_t,
  h_sigma_eta_0,
  h_st,
  h_log_scale = 0
)
```

Arguments

h	the T vector of log-volatilities
h_mu	the 1 vector of previous means
h_phi	the 1 vector of AR(1) coefficient(s)
h_phi2	the 1 vector of previous penalty coefficient(s)
h_sigma_eta_t	the T vector of log-vol innovation standard deviations
h_sigma_eta_0	the standard deviations of initial log-vols
h_st	the T vector of indicators on whether each time-step exceed the estimated threshold
h_log_scale	prior mean from scale mixture of Gaussian (Polya-Gamma) prior, e.g. $\log(\sigma_e^2)$ or dhs_mean0

Value

the sampled mean(s) dhs_mean

t_sampleLogVols	<i>Sample the latent log-volatilities</i>
-----------------	---

Description

Compute one draw of the log-volatilities using a discrete mixture of Gaussians approximation to the likelihood (see Omori, Chib, Shephard, and Nakajima, 2007) where the log-vols are assumed to follow an TAR(1) model with time-dependent innovation variances. More generally, the code operates for p independent TAR(1) log-vol processes to produce an efficient joint sampler in $O(Tp)$ time.

Usage

```
t_sampleLogVols(
  h_y,
  h_prev,
  h_mu,
  h_phi,
  h_phi2,
  h_sigma_eta_t,
  h_sigma_eta_0,
  h_st,
  loc
)
```

Arguments

h_y	the T vector of data, which follow independent SV models
h_prev	the T vector of the previous log-vols
h_mu	the 1 vector of log-vol unconditional means
h_phi	the 1 vector of log-vol AR(1) coefficients
h_phi2	the 1 vector of previous penalty coefficient(s)
h_sigma_eta_t	the T vector of log-vol innovation standard deviations
h_sigma_eta_0	the 1 vector of initial log-vol innovation standard deviations
h_st	the T vector of indicators on whether each time-step exceed the estimated threshold
loc	list of the row and column indices to fill in the band-sparse matrix in the sampler

Value

T x p vector of simulated log-vols

Note

For Bayesian trend filtering, $p = 1$. More generally, the sampler allows for $p > 1$ but assumes (contemporaneous) independence across the log-vols for $j = 1, \dots, p$.

t_sampleR_mh

Sample the threshold parameter

Description

Compute one draw of the threshold parameter in the TAR(1) model with Gaussian innovations and time-dependent innovation variances. The sampler utilizes metropolis hasting to draw from uniform prior.

Usage

```
t_sampleR_mh(
  h_yc,
  h_phi,
  h_phi2,
  h_sigma_eta_t,
  h_sigma_eta_0,
  h_st,
  h_r,
  lower_b,
  upper_b,
  omega,
  D
)
```

Arguments

h_yc	the T vector of centered log-volatilities (i.e., the log-vols minus the unconditional means dhs_mean)
h_phi	the 1 vector of previous AR(1) coefficient(s)
h_phi2	the 1 vector of previous penalty coefficient(s)
h_sigma_eta_t	the T vector of log-vol innovation standard deviations
h_sigma_eta_0	the 1 vector of initial log-vol innovation standard deviations
h_st	the T vector of indicators on whether each time-step exceed the estimated threshold
h_r	1 the previous draw of the threshold parameter
lower_b	the lower bound in the uniform prior of the threshold variable
upper_b	the upper bound in the uniform prior of the threshold variable
omega	T vector of evolution errors
D	the degree of differencing (one or two)

Value

the sampled threshold value r

t_sampleSVparams	<i>Sampler for the stochastic volatility parameters</i>
------------------	---

Description

Compute one draw of the normal stochastic volatility parameters. The model assumes an AR(1) for the log-volatility.

Usage

```
t_sampleSVparams(omega, svParams)
```

Arguments

omega	T vector of errors
svParams	list of parameters to be updated

Value

List of relevant components in svParams: sigma_wt, the T vector of standard deviations, and additional parameters associated with SV model.

uni.slice

Univariate Slice Sampler from Neal (2008)

Description

Compute a draw from a univariate distribution using the code provided by Radford M. Neal. The documentation below is also reproduced from Neal (2008).

Usage

```
uni.slice(x0, g, w = 1, m = Inf, lower = -Inf, upper = +Inf, gx0 = NULL)
```

Arguments

x0	Initial point
g	Function returning the log of the probability density (plus constant)
w	Size of the steps for creating interval (default 1)
m	Limit on steps (default infinite)
lower	Lower bound on support of the distribution (default -Inf)
upper	Upper bound on support of the distribution (default +Inf)
gx0	Value of g(x0), if known (default is not known)

Value

The point sampled, with its log density attached as an attribute.

Note

The log density function may return -Inf for points outside the support of the distribution. If a lower and/or upper bound is specified for the support, the log density function will not be called outside such limits.

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