

Package ‘bayesianVARs’

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Title MCMC Estimation of Bayesian Vectorautoregressions

Version 0.1.8

Description Efficient Markov Chain Monte Carlo (MCMC) algorithms for the fully Bayesian estimation of vectorautoregressions (VARs) featuring stochastic volatility (SV). Implements state-of-the-art shrinkage priors following Gruber & Kastner (2025) <[doi:10.1016/j.ijforecast.2025.02.001](https://doi.org/10.1016/j.ijforecast.2025.02.001)>. Efficient equation-per-equation estimation following Kastner & Huber (2020) <[doi:10.1002/for.2680](https://doi.org/10.1002/for.2680)> and Carrerio et al. (2021) <[doi:10.1016/j.jeconom.2021.11.010](https://doi.org/10.1016/j.jeconom.2021.11.010)>.

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<https://luisgruber.github.io/bayesianVARs/>

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bvar	<i>Markov Chain Monte Carlo Sampling for Bayesian Vectorautoregressions</i>
------	---

Description

bvar simulates from the joint posterior distribution of the parameters and latent variables and returns the posterior draws.

Usage

```
bvar(
  data,
  lags = 1L,
  draws = 1000L,
  burnin = 1000L,
  thin = 1L,
  prior_intercept = 10,
  prior_phi = specify_prior_phi(data = data, lags = lags, prior = "HS"),
  prior_sigma = specify_prior_sigma(data = data, type = "factor", quiet = TRUE),
  sv_keep = "last",
  expert_huge = FALSE,
  quiet = FALSE,
  startvals = list()
)
```

Arguments

data	Data matrix (can be a time series object). Each of M columns is assumed to contain a single time-series of length T .
lags	Integer indicating the order of the VAR, i.e. the number of lags of the dependent variables included as predictors.
draws	single integer indicating the number of draws after the burnin
burnin	single integer indicating the number of draws discarded as burnin
thin	single integer. Every <i>thin</i> th draw will be stored. Default is thin=1L.
prior_intercept	Either prior_intercept=FALSE and no constant term (intercept) will be included. Or a numeric vector of length M indicating the (fixed) prior standard deviations on the constant term. A single number will be recycled accordingly. Default is prior_intercept=10.
prior_phi	bayesianVARs_prior_phi object specifying prior for the reduced form VAR coefficients. Best use constructor specify_prior_phi .
prior_sigma	bayesianVARs_prior_sigma object specifying prior for the variance-covariance matrix of the VAR. Best use constructor specify_prior_sigma .

sv_keep	String equal to "all" or "last". In case of sv_keep = "last", the default, only draws for the very last log-variance h_T are stored.
expert_huge	logical value indicating whether to use a specific algorithm for huge VARs. Only possible if a factor structure for the errors is specified. See section MCMC algorithm below for more details.
quiet	logical value indicating whether information about the progress during sampling should be displayed during sampling (default is TRUE).
startvals	optional list with starting values.

Details

The VAR(p) model is of the following form: $\mathbf{y}'_t = \boldsymbol{\iota}' + \mathbf{x}'_t \boldsymbol{\Phi} + \boldsymbol{\epsilon}'_t$, where \mathbf{y}_t is a M -dimensional vector of dependent variables and $\boldsymbol{\epsilon}_t$ is the error term of the same dimension. \mathbf{x}_t is a $K = pM$ -dimensional vector containing lagged/past values of the dependent variables \mathbf{y}_{t-l} for $l = 1, \dots, p$ and $\boldsymbol{\iota}$ is a constant term (intercept) of dimension $M \times 1$. The reduced-form coefficient matrix $\boldsymbol{\Phi}$ is of dimension $K \times M$.

bvar offers two different specifications for the errors: The user can choose between a factor stochastic volatility structure or a cholesky stochastic volatility structure. In both cases the disturbances $\boldsymbol{\epsilon}_t$ are assumed to follow a M -dimensional multivariate normal distribution with zero mean and variance-covariance matrix $\boldsymbol{\Sigma}_t$. In case of the cholesky specification $\boldsymbol{\Sigma}_t = \mathbf{U}'^{-1} \mathbf{D}_t \mathbf{U}^{-1}$, where \mathbf{U}^{-1} is upper unitriangular (with ones on the diagonal). The diagonal matrix \mathbf{D}_t depends upon latent log-variances, i.e. $\mathbf{D}_t = \text{diag}(\exp(h_{1t}), \dots, \exp(h_{Mt}))$. The log-variances follow a priori independent autoregressive processes $h_{it} \sim N(\mu_i + \phi_i(h_{i,t-1} - \mu_i), \sigma_i^2)$ for $i = 1, \dots, M$. In case of the factor structure, $\boldsymbol{\Sigma}_t = \boldsymbol{\Lambda} \mathbf{V}_t \boldsymbol{\Lambda}' + \mathbf{G}_t$. The diagonal matrices \mathbf{V}_t and \mathbf{G}_t depend upon latent log-variances, i.e. $\mathbf{G}_t = \text{diag}(\exp(h_{1t}), \dots, \exp(h_{Mt}))$ and $\mathbf{V}_t = \text{diag}(\exp(h_{M+1,t}), \dots, \exp(h_{M+r,t}))$. The log-variances follow a priori independent autoregressive processes $h_{it} \sim N(\mu_i + \phi_i(h_{i,t-1} - \mu_i), \sigma_i^2)$ for $i = 1, \dots, M$ and $h_{M+j,t} \sim N(\phi_j h_{M+j,t-1}, \sigma_{M+j}^2)$ for $j = 1, \dots, r$.

Value

An object of type `bayesianVARs_bvar`, a list containing the following objects:

- `PHI`: A `bayesianVARs_coef` object, an array, containing the posterior draws of the VAR coefficients (including the intercept).
- `U`: A `bayesianVARs_draws` object, a matrix, containing the posterior draws of the contemporaneous coefficients (if cholesky decomposition for sigma is specified).
- `logvar`: A `bayesianVARs_draws` object containing the log-variance draws.
- `sv_para`: A `bayesianVARs_draws` object containing the posterior draws of the stochastic volatility related parameters.
- `phi_hyperparameter`: A matrix containing the posterior draws of the hyperparameters of the conditional normal prior on the VAR coefficients.
- `u_hyperparameter`: A matrix containing the posterior draws of the hyperparameters of the conditional normal prior on U (if cholesky decomposition for sigma is specified).
- `bench`: `proc_time` object containing the run time of the sampler.

- `V_prior`: An array containing the posterior draws of the variances of the conditional normal prior on the VAR coefficients.
- `facload`: A `bayesianVARs_draws` object, an array, containing draws from the posterior distribution of the factor loadings matrix (if factor decomposition for `sigma` is specified).
- `fac`: A `bayesianVARs_draws` object, an array, containing factor draws from the posterior distribution (if factor decomposition for `sigma` is specified).
- `Y`: Matrix containing the dependent variables used for estimation.
- `X` matrix containing the lagged values of the dependent variables, i.e. the covariates.
- `lags`: Integer indicating the lag order of the VAR.
- `intercept`: Logical indicating whether a constant term is included.
- `heteroscedastic` logical indicating whether heteroscedasticity is assumed.
- `Yraw`: Matrix containing the dependent variables, including the initial 'lags' observations.
- `Traw`: Integer indicating the total number of observations.
- `sigma_type`: Character specifying the decomposition of the variance-covariance matrix.
- `datamat`: Matrix containing both 'Y' and 'X'.
- `config`: List containing information on configuration parameters.

MCMC algorithm

To sample efficiently the reduced-form VAR coefficients assuming a **factor structure for the errors**, the equation per equation algorithm in Kastner & Huber (2020) is implemented. The factor structure has the advantage that an algorithm for sampling from high-dimensional Gaussian distributions can be exploited by setting `expert_huge = TRUE`. However, this speeds up computations only if $K > T$, i.e. the number of coefficients per equations exceeds the number of observations. All parameters and latent variables associated with the factor-structure are sampled using package `factorstochvol-package`'s function `update_fsv` callable on the C-level only.

To sample efficiently the reduced-form VAR coefficients, assuming a **cholesky-structure for the errors**, the corrected triangular algorithm in Carriero et al. (2021) is implemented. The SV parameters and latent variables are sampled using package `stochvol`'s `update_fast_sv` function. The precision parameters, i.e. the free off-diagonal elements in U , can be sampled using Bayesian linear regression methods, see Cogley and Sargent (2005).

References

- Gruber, L. and Kastner, G. (2025). Forecasting macroeconomic data with Bayesian VARs: Sparse or dense? It depends! *International Journal of Forecasting*. doi:10.1016/j.ijforecast.2025.02.001.
- Kastner, G. and Huber, F. Sparse (2020). Bayesian vector autoregressions in huge dimensions. *Journal of Forecasting*, **39**, 1142–1165, doi:10.1002/for.2680.
- Kastner, G. (2019). Sparse Bayesian Time-Varying Covariance Estimation in Many Dimensions *Journal of Econometrics*, **210**(1), 98–115, doi:10.1016/j.jeconom.2018.11.007.
- Carriero, A. and Chan, J. and Clark, T. E. and Marcellino, M. (2021). Corrigendum to “Large Bayesian vector autoregressions with stochastic volatility and non-conjugate priors” [J. Econometrics 212 (1) (2019) 137–154]. *Journal of Econometrics*, doi:10.1016/j.jeconom.2021.11.010.

Cogley, S. and Sargent, T. (2005). Drifts and volatilities: monetary policies and outcomes in the post WWII US. *Review of Economic Dynamics*, **8**, 262–302, doi:10.1016/j.red.2004.10.009.

Hosszejni, D. and Kastner, G. (2021). Modeling Univariate and Multivariate Stochastic Volatility in R with stochvol and factorstochvol. *Journal of Statistical Software*, **100**, 1–34. doi:10.18637/jss.v100.i12.

See Also

- Helpers for prior configuration: `specify_prior_phi()`, `specify_prior_sigma()`.
- Plotting: `plot.bayesianVARs_bvar()`.
- Extractors: `coef.bayesianVARs_bvar()`, `vcov.bayesianVARs_bvar()`.
- 'stable' bvar: `stable_bvar()`.
- summary method: `summary.bayesianVARs_bvar()`.
- predict method: `predict.bayesianVARs_bvar()`.
- fitted method: `fitted.bayesianVARs_bvar()`.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Plot
plot(mod)

# Summary
summary(mod)
```

coef	<i>Extract VAR coefficients</i>
------	---------------------------------

Description

Extracts posterior draws of the VAR coefficients from a VAR model estimated with `bvar()`.

Usage

```
## S3 method for class 'bayesianVARs_bvar'
coef(object, ...)
```

Arguments

object	A <code>bayesianVARs_bvar</code> object obtained from <code>bvar()</code> .
...	Currently ignored.

Value

Returns a numeric array of dimension $M \times K \times draws$, where M is the number of time-series, K is the number of covariates per equation (including the intercept) and $draws$ is the number of stored posterior draws.

See Also

[summary.bayesianVARs_draws\(\)](#), [vcov.bayesianVARs_bvar\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Extract posterior draws of VAR coefficients
bvar_coefs <- coef(mod)
```

extractB0

Retrieve the structural parameter B_0 samples from an IRF object.

Description

Retrieve the structural parameter B_0 samples from an IRF object.

Usage

```
extractB0(x)
```

Arguments

x a bayesianVARs_irf object

Author(s)

Stefan Haan <sthaan@edu.aau.at>

See Also

[specify_structural_restrictions](#)

Examples

```

train_data <- 100 * usmacro_growth[,c("GDPC1", "GDPCTPI", "GS1", "M2REAL", "CPIAUCSL")]
prior_sigma <- specify_prior_sigma(train_data, type="cholesky", cholesky_heteroscedastic=FALSE)
mod <- bvar(train_data, lags=5L, prior_sigma=prior_sigma, quiet=TRUE)

structural_restrictions <- specify_structural_restrictions(
  mod,
  restrictions_B0=rbind(
    c(1 ,NA,0 ,NA,NA),
    c(0 ,1 ,0 ,NA,NA),
    c(0 ,NA,1 ,NA,NA),
    c(0 ,0 ,NA,1 ,NA),
    c(0 ,0 ,0 ,0 ,1 )
  )
)
irf_structural <- irf(
  mod, ahead=8,
  structural_restrictions=structural_restrictions
)
B0 <- extractB0(irf_structural)

# Visually check that the restriction B0[1,1] >= 0 has been satisfied
hist(
  B0[1,1,],
  xlim=range(0, B0),
  main = paste0("Posterior B0[", 1, ", ", 1, "]")
)
abline(v=0, col=2, lwd=2)

```

fitted.bayesianVARs_bvar

Simulate fitted/predicted historical values for an estimated VAR model

Description

Simulates the fitted/predicted (in-sample) values for an estimated VAR model.

Usage

```

## S3 method for class 'bayesianVARs_bvar'
fitted(object, error_term = TRUE, ...)

```

Arguments

object	A bayesianVARs_bvar object estimated via <code>bvar()</code> .
error_term	logical indicating whether to include the error term or not.
...	Currently ignored.

Value

An object of class `bayesianVARs_fitted`.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Simulate predicted historical values including the error term.
pred <- fitted(mod, error_term = TRUE)

# Simulate fitted historical values not including the error term.
fit <- fitted(mod, error_term = FALSE)

# Visualize
plot(pred)
plot(fit)
```

 irf

Impulse response functions

Description

Effect of the structural (factor) shocks over time.

Usage

```
irf(
  x,
  ahead = 8,
  structural_restrictions = NULL,
  shocks = NULL,
  hairy = FALSE,
  ...
)
```

Arguments

`x` An object of type `bayesianVARs_bvar`.

`ahead` maximum number of time horizons.

`structural_restrictions` an object generated by [specify_structural_restrictions](#). If specified, the IRFs to the structural shocks identified by the restrictions given in this argument will be calculated. If not specified, the orthogonal IRFs based on the Cholesky decomposition will be calculated. Note that the orthogonal IRFs depend on the

	ordering of the variables and do not necessarily correspond to shocks with a meaningful interpretation.
shocks	an matrix with r rows, where r is the number of shocks (see 'Details'). Each column specifies a shock. Default: <code>diag(r)</code> , will calculate the responses to all structural (or factor) shocks of one standard deviation.
hairy	set to TRUE in order to plot each path seperately. To show valid quantiles, an Bayes optimal order of the posterior samples will be calculated which can take a long time even for moderately many samples. Default: FALSE.
...	Following expert arguments can be specified: solver: "randomized" or "lp". If some columns have more than one sign restriction, "lp" might find a solution, even if "randomized" is unable to. However "lp" can produce artificially narrow confidence bands which do not properly reflect the uncertainty in the identification scheme. Default: "randomized" randomized_max_rotations_per_sample: if using the "randomized" solver, how many rotations are drawn for each sample of the reduced form parameters in x . Default: 2.

Details

If a factor model was used, then the number of shocks is equal to the number of factors.

If the Cholesky model was used, then the number of shocks is equal to the number of variables.

Value

Returns a `bayesianVARs_irf` object.

Author(s)

Stefan Haan <sthaan@edu.aau.at>

References

Arias, J. and Rubio-Ramírez, J. and Waggoner, D. (2014). Inference Based on SVARs Identified with Sign and Zero Restrictions: Theory and Applications. *FRB Atlanta Working Paper Series*, doi:10.2139/ssrn.2580264.

See Also

[specify_structural_restrictions](#), [extractB0](#)

Examples

```
train_data <- 100 * usmacro_growth[,c("GDPC1", "GDPCTPI", "GS1", "M2REAL", "CPIAUCSL")]
prior_sigma <- specify_prior_sigma(train_data, type="cholesky", cholesky_heteroscedastic=FALSE)
mod <- bvar(train_data, lags=5L, prior_sigma=prior_sigma, quiet=TRUE)

structural_restrictions <- specify_structural_restrictions(
  mod,
```

```

restrictions_B0=rbind(
  c(1 ,NA,0 ,NA,NA),
  c(0 ,1 ,0 ,NA,NA),
  c(0 ,NA,1 ,NA,NA),
  c(0 ,0 ,NA,1 ,NA),
  c(0 ,0 ,0 ,0 ,1 )
)
)
irf_structural <- irf(
  mod, ahead=8,
  structural_restrictions=structural_restrictions
)
plot(irf_structural)

```

my_gig

Draw from generalized inverse Gaussian

Description

Vectorized version of [rgig](#)

Usage

```
my_gig(n, lambda, chi, psi)
```

Arguments

n	A single integer indicating the number of draws to generate.
lambda	vector of shape parameters.
chi	vector of shape/scale parameters. Must be nonnegative for positive lambdas and positive else.
psi	vector of shape/scale parameters. Must be nonnegative for negative lambdas and positive else.

Value

Matrix of dimension $c(n,m)$, where m is the maximum length of λ , ψ and χ .

Examples

```
gigsamples <- my_gig(2, c(1,1), c(1,1), c(1,1))
```

pairs_predict

Pairwise visualization of out-of-sample posterior predictive densities.

Description

Pairwise visualization of out-of-sample posterior predictive densities.

Usage

```
## S3 method for class 'bayesianVARs_predict'
pairs(x, vars, ahead, ...)
```

Arguments

x	An object of class <code>bayesianVARs_predict</code> obtained via <code>predict.bayesianVARs_bvar()</code> .
vars	Integer vector (or coercible to such) indicating which variables to plot.
ahead	Integer vector (or coercible to such) indicating which step ahead to plot. <code>max(ahead)</code> must be smaller equal to <code>dim(x\$predictions)[1]</code> .
...	Currently ignored!

Value

Returns x invisibly.

Note

Note that that `bayesianVARs_predict` can also be used withing `plot.bayesianVARs_bvar()`.

See Also

Other plotting `plot.bayesianVARs_bvar()`, `plot.bayesianVARs_fitted()`, `plot.bayesianVARs_predict()` `posterior_heatmap()`.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Simulate from posterior predictive
predictions <- predict(mod, ahead = 1:3)

# Visualize
pairs(predictions, vars = 1:3, ahead = 1:3)
```

plot.bayesianVARs_bvar

Plot method for bayesianVARs_bvar

Description

Visualization of in-sample fit. Can also be used to display prediction intervals of future values.

Usage

```
## S3 method for class 'bayesianVARs_bvar'
plot(
  x,
  predictions = NULL,
  quantiles = c(0.05, 0.5, 0.95),
  dates = NULL,
  n_col = 1,
  ...
)
```

Arguments

x	An object of class bayesianVARs_bvar obtained via bvar() .
predictions	Optional array of out of sample predictions, e.g. obtained via predict.bayesianVARs_bvar() .
quantiles	numeric vector indicating which quantiles to plot.
dates	optional vector of dates for labelling the x-axis. The default values is NULL; in this case, the axis will be labeled with numbers.
n_col	integer indicating the number of columns to use for plotting.
...	Currently ignored!

Value

Returns x invisibly.

See Also

Other plotting [plot.bayesianVARs_fitted\(\)](#), [plot.bayesianVARs_predict\(\)](#), [pairs.bayesianVARs_predict\(\)](#), [posterior_heatmap\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)
```

```
# Simulate from posterior predictive
predictions <- predict(mod, ahead = 1:3)

# Visualize
plot(mod, predictions = predictions)
```

```
plot.bayesianVARs_fitted
```

Visualization of in-sample fit of an estimated VAR.

Description

Visualization of in-sample fit of an estimated VAR.

Usage

```
## S3 method for class 'bayesianVARs_fitted'
plot(
  x,
  dates = NULL,
  vars = "all",
  quantiles = c(0.05, 0.5, 0.95),
  n_col = 1L,
  ...
)
```

Arguments

x	A bayesianVARs_fitted object.
dates	optional vector of dates for labelling the x-axis. The default values is NULL; in this case, the axis will be labeled with numbers.
vars	character vector containing the names of the variables to be visualized. The default is "all" indicating that the fit of all variables is visualized.
quantiles	numeric vector indicating which quantiles to plot.
n_col	integer indicating the number of columns to use for plotting.
...	Currently ignored.

Value

returns x invisibly

See Also

- fitted method for class 'bayesianVARs_bvar': [fitted.bayesianVARs_bvar\(\)](#).
- Other plotting [plot.bayesianVARs_bvar\(\)](#), [plot.bayesianVARs_fitted\(\)](#), [plot.bayesianVARs_predict\(\)](#), [pairs.bayesianVARs_predict\(\)](#), [posterior_heatmap\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Simulate predicted historical values including the error term.
pred <- fitted(mod, error_term = TRUE)

# Visualize
plot(pred)
```

plot.bayesianVARs_irf *Impulse Responses Plot*

Description

Visualization of the impulse responses. Responses are plotted on a grid, where rows correspond to variables and columns correspond to shocks.

Usage

```
## S3 method for class 'bayesianVARs_irf'
plot(
  x,
  vars = "all",
  quantiles = c(0.05, 0.25, 0.5, 0.75, 0.95),
  default_hair_color = "#FF000003",
  true_irf = NULL,
  ...
)
```

Arguments

x	An object of type bayesianVARs_irf obtained via irf .
vars	character vector containing the names of the variables to be visualized. The default is "all" indicating that all variables are visualized.
quantiles	numeric vector indicating which quantiles to plot. If hairy=TRUE was specified when calling irf , a proportion of max(quantiles) IRFs will be plotted. Specify 0 to plot a point-estimate only. If hairy=FALSE was specified (the default), point-wise quantiles will be plotted. Note that the curve of point-wise medians is not necessarily in the set of IRFs (see Inoue 2022).
default_hair_color	the color of the IRF samples, if hairy=TRUE was specified.

`true_irf` If the true IRFs are known (because the data was simulated) they can be plotted alongside the estimates, such that the quality of the estimates may be judged. `true_irf` should be a numeric array with dimensions variables, shocks and time, in that order.
`...` Currently ignored!

Author(s)

Stefan Haan <sthaan@edu.aau.at>

References

Inoue, A. and Kilian, L. (2022). Joint Bayesian inference about impulse responses in VAR models. *Journal of Econometrics*, doi:10.1016/j.jeconom.2021.05.010.

See Also

[irf](#)

plot.bayesianVARs_predict

Fan chart

Description

Visualization of (out-of-sample) predictive distribution.

Usage

```
## S3 method for class 'bayesianVARs_predict'
plot(
  x,
  dates = NULL,
  vars = "all",
  ahead = NULL,
  quantiles = c(0.05, 0.25, 0.5, 0.75, 0.95),
  n_col = 1L,
  first_obs = 1L,
  ...
)
```

Arguments

`x` An object of type `bayesianVARs_predict` obtained via `predict.bayesianVARs_bvar()`.
`dates` optional vector of dates for labeling the x-axis. The default values is `NULL`; in this case, the axis will be labeled with numbers.

vars	character vector containing the names of the variables to be visualized. The default is "all" indicating that all variables are visualized.
ahead	Integer vector (or coercible to such) indicating which step ahead to plot. max(ahead) must be smaller equal to dim(x\$predictions)[1].
quantiles	numeric vector indicating which quantiles to plot.
n_col	integer indicating the number of columns to use for plotting.
first_obs	integer indicating the first observation to be used for plotting.
...	Currently ignored!

Value

Returns x invisibly!

See Also

Other plotting [plot.bayesianVARs_bvar\(\)](#), [plot.bayesianVARs_fitted\(\)](#), [pairs.bayesianVARs_predict\(\)](#) [posterior_heatmap\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Simulate from posterior predictive
predictions <- predict(mod, ahead = 1:3)

# Visualize
plot(predictions, vars = 1:3, ahead = 1:3)
```

plot.bayesianVARs_residuals

Visualization of the residuals of an estimated VAR.

Description

Visualization of the residuals of an estimated VAR.

Usage

```
## S3 method for class 'bayesianVARs_residuals'
plot(
  x,
  dates = NULL,
  vars = "all",
```

```

    quantiles = c(0.05, 0.5, 0.95),
    n_col = 1L,
    ...
)

```

Arguments

x	A <code>bayesianVARs_residuals</code> object.
dates	optional vector of dates for labelling the x-axis. The default values is NULL; in this case, the axis will be labeled with numbers.
vars	character vector containing the names of the variables to be visualized. The default is "all" indicating that the fit of all variables is visualized.
quantiles	numeric vector indicating which quantiles to plot.
n_col	integer indicating the number of columns to use for plotting.
...	Currently ignored.

Value

returns x invisibly

See Also

- residuals method for class 'bayesianVARs_bvar': [residuals.bayesianVARs_bvar\(\)](#).
- Other plotting [plot.bayesianVARs_bvar\(\)](#), [plot.bayesianVARs_fitted\(\)](#), [plot.bayesianVARs_predict\(\)](#), [pairs.bayesianVARs_predict\(\)](#), [posterior_heatmap\(\)](#).

Examples

```

# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

mod.resids <- residuals(mod)

# Visualize
plot(mod.resids)

```

posterior_heatmap

Posterior heatmaps for matrix valued parameters

Description

`posterior_heatmap()` generates a heatmap for draws of matrix values parameters visualizing point wise summaries, such as mean, median, variance, standard deviation, interquartile range etc. etc.

Usage

```
posterior_heatmap(
  x,
  FUN,
  ...,
  transpose = FALSE,
  colorbar = TRUE,
  colorbar_width = 0.1,
  gap_width = 0.25,
  xlabels = NULL,
  ylabels = NULL,
  add_numbers = FALSE,
  zlim = NULL,
  colspace = NULL,
  border_color = NA,
  zero_color = NA,
  main = "",
  detect_lags = TRUE,
  cex.axis = 0.75,
  cex.colbar = 1,
  cex.numbers = 1,
  asp = NULL
)
```

Arguments

x	An array of dimension $a \times b \times draws$, where $a \times b$ is the dimension of the parameter to visualize and draws is the number of posterior draws.
FUN	The summary function to be applied to margins $c(1,2)$ of x. E.g. "median", "mean", "IQR", "sd" or "var". <code>apply(x, 1:2, FUN, ...)</code> must return a matrix!
...	optional arguments to FUN.
transpose	logical indicating whether to transpose the matrix or not, i.e. whether to plot an $a \times b$ or an $b \times a$ matrix. Default is FALSE.
colorbar	logical indicating whether to display a colorbar or not. Default is TRUE.
colorbar_width	numeric. A value between 0 and 1 indicating the proportion of the width of the plot for the colorbar.
gap_width	numeric. A value between 0 and 1 indicating the width of the gap between the heatmap and the colorbar. The width is computed as <code>gap_width*colorbar_width</code> .
xlabels	<code>ylabels=NULL</code> , the default, indicates that <code>colnames(x)</code> will be displayed. <code>ylabels=""</code> indicates that no ylabels will be displayed.
ylabels	<code>xlabels=NULL</code> , the default, indicates that <code>rownames(x)</code> will be displayed. <code>xlabels=""</code> indicates that no xlabels are displayed.
add_numbers	logical. <code>add_numbers=TRUE</code> , the default indicates that the actual values of summary will be displayed.

<code>zlim</code>	numeric vector of length two indicating the minimum and maximum values for which colors should be plotted. By default this range is determined by the maximum of the absolute values of the selected summary.
<code>colspace</code>	Optional argument indicating the color palette to be used. If not specified, <code>colorspace::diverge_hcl()</code> will be used, or, if FUN returns only positive values <code>colorspace::sequential_hcl()</code> . See below for a more detailed description of the default usage.
<code>border_color</code>	The color of the rectangles borders. If not specified no borders will be displayed.
<code>zero_color</code>	The color of exact zero elements. By default this is not specified and then will depend on <code>colspace</code> .
<code>main</code>	main title for the plot.
<code>detect_lags</code>	logical. If <code>class(x)</code> is "bayesianVARs_coef", then <code>detect_lags=TRUE</code> will separate the sub matrices corresponding to the lags with black lines.
<code>cex.axis</code>	The magnification to be used for y-axis annotation relative to the current setting of <code>cex</code> .
<code>cex.colbar</code>	The magnification to be used for colorbar annotation relative to the current setting of <code>cex</code> .
<code>cex.numbers</code>	The magnification to be used for the actual values (if <code>add_numbers=TRUE</code>) relative to the current setting of <code>cex</code> .
<code>asp</code>	aspect ratio.

Details

colspace:

If not specified either `colorspace::diverge_hcl(1001, alpha = alpha, palette = "Blue-Red")` or `colorspace::sequential_hcl(1001, alpha = alpha, rev = TRUE, palette = "Reds 2")` will be used.

Value

Returns `x` invisibly.

See Also

Other plotting [plot.bayesianVARs_bvar\(\)](#), [plot.bayesianVARs_fitted\(\)](#), [plot.bayesianVARs_predict\(\)](#), [pairs.bayesianVARs_predict\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(100*data, sv_keep = "all", quiet = TRUE)

# Extract posterior draws of VAR coefficients
phi_post <- coef(mod)
```

```
# Visualize posterior median of VAR coefficients
posterior_heatmap(phi_post, median, detect_lags = TRUE, border_color = rgb(0, 0, 0, alpha = 0.2))

# Extract posterior draws of variance-covariance matrices (for each point in time)
sigma_post <- vcov(mod)
# Visualize posterior interquartile-range of variance-covariance matrix of the first observation
posterior_heatmap(sigma_post[1,,], IQR)
```

predict.bayesianVARs_bvar

Predict method for Bayesian VARs

Description

Simulates from (out-of-sample) predictive density for Bayesian VARs estimated via `bvar()` and computes log predictive likelihoods if ex-post observed data is supplied.

Usage

```
## S3 method for class 'bayesianVARs_bvar'
predict(
  object,
  ahead = 1L,
  each = 1L,
  stable = TRUE,
  simulate_predictive = TRUE,
  LPL = FALSE,
  Y_obs = NA,
  LPL_VoI = NA,
  ...
)
```

Arguments

<code>object</code>	A <code>bayesianVARs_bvar</code> object, obtained from <code>bvar()</code> .
<code>ahead</code>	Integer vector (or coercible to such), indicating the number of steps ahead at which to predict.
<code>each</code>	Single integer (or coercible to such) indicating how often should be drawn from the posterior predictive distribution for each draw that has been stored during MCMC sampling.
<code>stable</code>	logical indicating whether to consider only those draws from the posterior that fulfill the 'stable' criterion. Default is TRUE.
<code>simulate_predictive</code>	logical, indicating whether the posterior predictive distribution should be simulated.

LPL	logical indicating whether ahead-step-ahead log predictive likelihoods should be computed. If LPL=TRUE, Y_obs has to be specified.
Y_obs	Data matrix of observed values for computation of log predictive likelihood. Each of ncol(object\$Yraw) columns is assumed to contain a single time-series of length length(ahead).
LPL_VoI	either integer vector or character vector of column-names indicating for which subgroup of time-series in object\$Yraw a joint log predictive likelihood shall be computed.
...	Currently ignored!

Value

Object of class `bayesianVARs_predict`, a list that may contain the following elements:

- `predictions` array of dimensions `c(length(ahead), ncol(object$Yraw), each * dim(object$PHI)[3])` containing the simulations from the predictive density (if `simulate_predictive=TRUE`).
- `LPL` vector of length `length(ahead)` containing the log-predictive-likelihoods (taking into account the joint distribution of all variables) (if `LPL=TRUE`).
- `LPL_univariate` matrix of dimension `c(length(ahead), ncol(object$Yraw))` containing the marginalized univariate log-predictive-likelihoods of each series (if `LPL=TRUE`).
- `LPL_VoI` vector of length `length(ahead)` containing the log-predictive-likelihoods for a subset of variables (if `LPL=TRUE` and `LPL_VoI != NA`).
- `Yraw` matrix containing the data used for the estimation of the VAR.
- `LPL_draws` matrix containing the simulations of the log-predictive-likelihood (if `LPL=TRUE`).
- `PL_univariate_draws` array containing the simulations of the univariate predictive-likelihoods (if `LPL=TRUE`).
- `LPL_sub_draws` matrix containing the simulations of the log-predictive-likelihood for a subset of variables (if `LPL=TRUE` and `LPL_VoI != NA`).

See Also

[stable_bvar\(\)](#), [plot.bayesianVARs_predict\(\)](#), [pairs.bayesianVARs_predict\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Split data in train and test
train <- data[1:(nrow(data)-4),]
test <- data[-c(1:(nrow(data)-4)),]

# Estimate model using train data only
mod <- bvar(train, quiet = TRUE)

# Simulate from 1-step to 4-steps ahead posterior predictive and compute
# log-predictive-likelihoods
```

```
predictions <- predict(mod, ahead = 1:4, LPL = TRUE, Y_obs = test)

# Summary
summary(predictions)

# Visualize via fan-charts
plot(predictions)

# In order to evaluate the joint predictive density of a subset of the
# variables (variables of interest), consider specifying 'LPL_VoI':
predictions <- predict(mod, ahead = 1:4, LPL = TRUE, Y_obs = test, LPL_VoI = c("GDPC1", "FEDFUNDS"))
predictions$LPL_VoI
```

```
print.bayesianVARs_bvar
```

Pretty printing of a bvar object

Description

Pretty printing of a bvar object

Usage

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

Arguments

x	Object of class bayesianVARs_bvar, usually resulting from a call of <code>bvar()</code> .
...	Ignored.

Value

Returns x invisibly.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Print model
mod
```

```
print.bayesianVARs_predict
      Print method for bayesianVARs_predict objects
```

Description

Print method for bayesianVARs_predict objects.

Usage

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

Arguments

x	A bayesianVARs_predict object obtained via predict.bayesianVARs_bvar() .
...	Currently ignored!

Value

Returns x invisibly.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Split data in train and test
train <- data[1:(nrow(data)-4),]
test <- data[-c(1:(nrow(data)-4)),]

# Estimate model using train data only
mod <- bvar(train, quiet = TRUE)

# Simulate from 1-step ahead posterior predictive
predictions <- predict(mod, ahead = 1L)
print(predictions)
```

```
print.summary.bayesianVARs_bvar
      Print method for summary.bayesianVARs_bvar objects
```

Description

Print method for summary.bayesianVARs_bvar objects.

Usage

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

Arguments

x A `summary.bayesianVARs_bvar` object obtained via `summary.bayesianVARs_bvar()`.
... Currently ignored!

Value

Returns x invisibly!

Examples

```
# Access a subset of the usmacro_growth dataset  
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]  
  
# Estimate model  
mod <- bvar(data, quiet = TRUE)  
  
# Print summary  
summary(mod)
```

```
print.summary.bayesianVARs_predict  
                                  Print method for summary.bayesianVARs_predict objects
```

Description

Print method for `summary.bayesianVARs_predict` objects.

Usage

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

Arguments

x A `summary.bayesianVARs_predict` object obtained via `summary.bayesianVARs_predict()`.
... Currently ignored.

Value

Returns x invisibly.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Split data in train and test
train <- data[1:(nrow(data)-4),]
test <- data[-c(1:(nrow(data)-4)),]

# Estimate model using train data only
mod <- bvar(train, quiet = TRUE)

# Simulate from 1-step ahead posterior predictive
predictions <- predict(mod, ahead = 1L)
sum <- summary(predictions)
print(sum)
```

residuals.bayesianVARs_bvar

Extract Model Residuals

Description

Extract model residuals, defined as the difference between the observed time-series and the in-sample predictions of the VAR model. Because in-sample prediction is subject to uncertainty of the VAR parameter estimates, this uncertainty carries over to the model residuals.

Usage

```
## S3 method for class 'bayesianVARs_bvar'
residuals(object, ...)
```

Arguments

object	A <code>bayesianVARs_bvar</code> object estimated via <code>bvar()</code> .
...	Passed to <code>fitted.bayesianVARs_bvar()</code> .

Value

An object of class `bayesianVARs_residuals`.

See Also

[fitted.bayesianVARs_bvar](#)

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

mod.resids <- residuals(mod)
plot(mod.resids)
```

specify_prior_phi *Specify prior on PHI*

Description

Configures prior on PHI, the matrix of reduced-form VAR coefficients.

Usage

```
specify_prior_phi(
  data = NULL,
  M = ncol(data),
  lags = 1L,
  prior = "HS",
  priormean = 0,
  PHI_tol = 1e-18,
  DL_a = "1/K",
  DL_tol = 0,
  R2D2_a = 0.1,
  R2D2_b = 0.5,
  R2D2_tol = 0,
  NG_a = 0.1,
  NG_b = 1,
  NG_c = 1,
  NG_tol = 0,
  SSVS_c0 = 0.01,
  SSVS_c1 = 100,
  SSVS_semiautomatic = TRUE,
  SSVS_p = 0.5,
  HMP_lambda1 = c(0.01, 0.01),
  HMP_lambda2 = c(0.01, 0.01),
  normal_sds = 10,
  global_grouping = "global",
  ...
)
```

Arguments

data	Optional. Data matrix (can be a time series object). Each of M columns is assumed to contain a single time-series of length T .
M	positive integer indicating the number of time-series of the VAR.
lags	positive integer indicating the order of the VAR, i.e. the number of lags of the dependent variables included as predictors.
prior	character, one of "HS", "R2D2", "NG", "DL", "SSVS", "HMP" or "normal".
priormean	real numbers indicating the prior means of the VAR coefficients. One single number means that the prior mean of all own-lag coefficients w.r.t. the first lag equals priormean and \emptyset else. A vector of length M means that the prior mean of the own-lag coefficients w.r.t. the first lag equals priormean and \emptyset else. If priormean is a matrix of dimension $c(\text{lags} * M, M)$, then each of the M columns is assumed to contain $\text{lags} * M$ prior means for the VAR coefficients of the respective VAR equations.
PHI_tol	Minimum number that the absolute value of a VAR coefficient draw can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero.
DL_a	(Single) positive real number. The value is interpreted as the concentration parameter for the local scales. Smaller values enforce heavier shrinkage. If the argument <code>global_grouping</code> specifies e.g. k groups, then <code>DL_a</code> can be a numeric vector of length k and the elements indicate the shrinkage in each group. A matrix of dimension $c(s, 2)$ specifies a discrete hyperprior, where the first column contains s support points and the second column contains the associated prior probabilities. <code>DL_a</code> has only to be specified if <code>prior="DL"</code> .
DL_tol	Minimum number that a parameter draw of one of the shrinking parameters of the Dirichlet Laplace prior can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero. <code>DL_tol</code> has only to be specified if <code>prior="DL"</code> .
R2D2_a	(Single) positive real number. The value is interpreted as the concentration parameter for the local scales. Smaller values enforce heavier shrinkage. If the argument <code>global_grouping</code> specifies e.g. k groups, then <code>R2D2_a</code> can be a numeric vector of length k and the elements indicate the shrinkage in each group. A matrix of dimension $c(s, 2)$ specifies a discrete hyperprior, where the first column contains s support points and the second column contains the associated prior probabilities. <code>R2D2_a</code> has only to be specified if <code>prior="R2D2"</code> .
R2D2_b	(Single) positive real number. The value indicates the shape parameter of the inverse gamma prior on the (semi-)global scales. If the argument <code>global_grouping</code> specifies e.g. k groups, then <code>NG_b</code> can be a numeric vector of length k and the elements determine the shape parameter in each group. <code>R2D2_b</code> has only to be specified if <code>prior="R2D2"</code> .
R2D2_tol	Minimum number that a parameter draw of one of the shrinking parameters of the R2D2 prior can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero. <code>R2D2_tol</code> has only to be specified if <code>prior="R2D2"</code> .

NG_a	(Single) positive real number. The value is interpreted as the concentration parameter for the local scales. Smaller values enforce heavier shrinkage. If the argument <code>global_grouping</code> specifies e.g. <code>k</code> groups, then <code>NG_a</code> can be a numeric vector of length <code>k</code> and the elements indicate the shrinkage in each group. A matrix of dimension <code>c(s, 2)</code> specifies a discrete hyperprior, where the first column contains <code>s</code> support points and the second column contains the associated prior probabilities. <code>NG_a</code> has only to be specified if <code>prior="NG"</code> .
NG_b	(Single) positive real number. The value indicates the shape parameter of the inverse gamma prior on the (semi-)global scales. If the argument <code>global_grouping</code> specifies e.g. <code>k</code> groups, then <code>NG_b</code> can be a numeric vector of length <code>k</code> and the elements determine the shape parameter in each group. <code>NG_b</code> has only to be specified if <code>prior="NG"</code> .
NG_c	(Single) positive real number. The value indicates the scale parameter of the inverse gamma prior on the (semi-)global scales. If the argument <code>global_grouping</code> specifies e.g. <code>k</code> groups, then <code>NG_c</code> can be a numeric vector of length <code>k</code> and the elements determine the scale parameter in each group. Expert option would be to set the scale parameter proportional to <code>NG_a</code> . E.g. in the case where a discrete hyperprior for <code>NG_a</code> is chosen, a desired proportion of let's say <code>0.2</code> is achieved by setting <code>NG_c="0.2*a"</code> (character input!). <code>NG_c</code> has only to be specified if <code>prior="NG"</code> .
NG_tol	Minimum number that a parameter draw of one of the shrinking parameters of the normal-gamma prior can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero. <code>NG_tol</code> has only to be specified if <code>prior="NG"</code> .
SSVS_c0	single positive number indicating the (unscaled) standard deviation of the spike component. <code>SSVS_c0</code> has only to be specified if <code>prior="SSVS"</code> . It should be that $SSVS_{c0} \ll SSVS_{c1}$! <code>SSVS_c0</code> has only to be specified if <code>prior="SSVS"</code> .
SSVS_c1	single positive number indicating the (unscaled) standard deviation of the slab component. <code>SSVS_c0</code> has only to be specified if <code>prior="SSVS"</code> . It should be that $SSVS_{c0} \ll SSVS_{c1}$!
SSVS_semiautomatic	logical. If <code>SSVS_semiautomatic=TRUE</code> both <code>SSVS_c0</code> and <code>SSVS_c1</code> will be scaled by the variances of the posterior of PHI under a FLAT conjugate (dependent Normal-Wishart prior). <code>SSVS_semiautomatic</code> has only to be specified if <code>prior="SSVS"</code> .
SSVS_p	Either a single positive number in the range $(0, 1)$ indicating the (fixed) prior inclusion probability of each coefficient. Or numeric vector of length 2 with positive entries indicating the shape parameters of the Beta distribution. In that case a Beta hyperprior is placed on the prior inclusion probability. <code>SSVS_p</code> has only to be specified if <code>prior="SSVS"</code> .
HMP_lambda1	numeric vector of length 2. Both entries must be positive. The first indicates the shape and the second the rate of the Gamma hyperprior on own-lag coefficients. <code>HMP_lambda1</code> has only to be specified if <code>prior="HMP"</code> .
HMP_lambda2	numeric vector of length 2. Both entries must be positive. The first indicates the shape and the second the rate of the Gamma hyperprior on cross-lag coefficients. <code>HMP_lambda2</code> has only to be specified if <code>prior="HMP"</code> .

normal_sds	numeric vector of length n , where $n = lagsM^2$ is the number of all VAR coefficients (excluding the intercept), indicating the prior variances. A single number will be recycled accordingly! Must be positive. normal_sds has only to be specified if prior="normal".
global_grouping	One of "global", "equation-wise", "covariate-wise", "olcl-lagwise" "fol" indicating the sub-groups of the semi-global(-local) modifications to HS, R2D2, NG, DL and SSVS prior. Works also with user-specified indicator matrix of dimension $c(lags*M,M)$. Only relevant if prior="HS", prior="DL", prior="R2D2", prior="NG" or prior="SSVS".
...	Do not use!

Details

For details concerning prior-elicitation for VARs please see Gruber & Kastner (2025).

Currently one can choose between six hierarchical shrinkage priors and a normal prior: prior="HS" stands for the Horseshoe-prior, prior="R2D2" for the R^2 -induced-Dirichlet-decomposition-prior, prior="NG" for the normal-gamma-prior, prior="DL" for the Dirichlet-Laplace-prior, prior="SSVS" for the stochastic-search-variable-selection-prior, prior="HMP" for the semi-hierarchical Minnesota prior and prior=normal for the normal-prior.

Semi-global shrinkage, i.e. group-specific shrinkage for pre-specified subgroups of the coefficients, can be achieved through the argument global_grouping.

Value

A bayesianVARs_prior_phi-object.

References

Gruber, L. and Kastner, G. (2025). Forecasting macroeconomic data with Bayesian VARs: Sparse or dense? It depends! *International Journal of Forecasting*. doi:10.1016/j.ijforecast.2025.02.001.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Horseshoe prior for a VAR(2)
phi_hs <- specify_prior_phi(data = data, lags = 2L ,prior = "HS")

# Semi-global-local Horseshoe prior for a VAR(2) with semi-global shrinkage parameters for
# cross-lag and own-lag coefficients in each lag
phi_hs_sg <- specify_prior_phi(data = data, lags = 2L, prior = "HS",
global_grouping = "olcl-lagwise")

# Semi-global-local Horseshoe prior for a VAR(2) with equation-wise shrinkage
# construct indicator matrix for equation-wise shrinkage
semi_global_mat <- matrix(1:ncol(data), 2*ncol(data), ncol(data),
byrow = TRUE)
phi_hs_ew <- specify_prior_phi(data = data, lags = 2L, prior = "HS",
```

```

global_grouping = semi_global_mat)
# (for equation-wise shrinkage one can also use 'global_grouping = "equation-wise"')

# Estimate model with your prior configuration of choice
mod <- bvar(data, lags = 2L, prior_phi = phi_hs_sg, quiet = TRUE)

```

specify_prior_sigma *Specify prior on Sigma*

Description

Configures prior on the variance-covariance of the VAR.

Usage

```

specify_prior_sigma(
  data = NULL,
  M = ncol(data),
  type = c("factor", "cholesky"),
  factor_factors = 1L,
  factor_restrict = c("none", "upper"),
  factor_priorfacloadtype = c("rowwiseng", "colwiseng", "normal"),
  factor_priorfacload = 0.1,
  factor_facloadtol = 1e-14,
  factor_priorng = c(1, 1),
  factor_priormu = c(0, 10),
  factor_priorphiidi = c(10, 3),
  factor_priorphifac = c(10, 3),
  factor_priorsigmaidi = 1,
  factor_priorsigmafac = 1,
  factor_priorh0idi = "stationary",
  factor_priorh0fac = "stationary",
  factor_heteroskedastic = TRUE,
  factor_priorhomoskedastic = NA,
  factor_interweaving = 4,
  cholesky_U_prior = c("HS", "DL", "R2D2", "NG", "SSVS", "normal", "HMP"),
  cholesky_U_tol = 1e-18,
  cholesky_heteroscedastic = TRUE,
  cholesky_priormu = c(0, 100),
  cholesky_priorphi = c(20, 1.5),
  cholesky_priorsigma2 = c(0.5, 0.5),
  cholesky_priorh0 = "stationary",
  cholesky_priorhomoscedastic = as.numeric(NA),
  cholesky_DL_a = "1/n",
  cholesky_DL_tol = 0,

```

```

cholesky_R2D2_a = 0.4,
cholesky_R2D2_b = 0.5,
cholesky_R2D2_tol = 0,
cholesky_NG_a = 0.5,
cholesky_NG_b = 0.5,
cholesky_NG_c = 0.5,
cholesky_NG_tol = 0,
cholesky_SSVS_c0 = 0.001,
cholesky_SSVS_c1 = 1,
cholesky_SSVS_p = 0.5,
cholesky_HMP_lambda3 = c(0.01, 0.01),
cholesky_normal_sds = 10,
expert_sv_offset = 0,
quiet = FALSE,
...
)

```

Arguments

data	Optional. Data matrix (can be a time series object). Each of M columns is assumed to contain a single time-series of length T .
M	positive integer indicating the number of time-series of the VAR.
type	character, one of "factor" (the default) or "cholesky", indicating which decomposition to be applied to the covariance-matrix.
factor_factors	Number of latent factors to be estimated. Only required if type="factor".
factor_restrict	Either "upper" or "none", indicating whether the factor loadings matrix should be restricted to have zeros above the diagonal ("upper") or whether all elements should be estimated from the data ("none"). Setting restrict to "upper" often stabilizes MCMC estimation and can be important for identifying the factor loadings matrix, however, it generally is a strong prior assumption. Setting restrict to "none" is usually the preferred option if identification of the factor loadings matrix is of less concern but covariance estimation or prediction is the goal. Only required if type="factor".
factor_priorfacloadtype	Can be "normal", "rowwiseng", "colwiseng". Only required if type="factor". "normal": Normal prior. The value of priorfacload is interpreted as the standard deviations of the Gaussian prior distributions for the factor loadings. "rowwiseng": Row-wise Normal-Gamma prior. The value of priorfacload is interpreted as the shrinkage parameter a . "colwiseng": Column-wise Normal-Gamma prior. The value of priorfacload is interpreted as the shrinkage parameter a . For details please see Kastner (2019).
factor_priorfacload	Either a matrix of dimensions M times factor_factors with positive elements or a single number (which will be recycled accordingly). Only required if

type="factor". The meaning of factor_priorfacload depends on the setting of factor_priorfacloadtype and is explained there.

- factor_facloadtol
Minimum number that the absolute value of a factor loadings draw can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero. Only required if type="factor".
- factor_priorng
Two-element vector with positive entries indicating the Normal-Gamma prior's hyperparameters c and d (cf. Kastner (2019)). Only required if type="factor".
- factor_priormu
Vector of length 2 denoting prior mean and standard deviation for unconditional levels of the idiosyncratic log variance processes. Only required if type="factor".
- factor_priorphiidi
Vector of length 2, indicating the shape parameters for the Beta prior distributions of the transformed parameters $(\phi+1)/2$, where ϕ denotes the persistence of the idiosyncratic log variances. Only required if type="factor".
- factor_priorphifac
Vector of length 2, indicating the shape parameters for the Beta prior distributions of the transformed parameters $(\phi+1)/2$, where ϕ denotes the persistence of the factor log variances. Only required if type="factor".
- factor_priorsigmaidi
Vector of length M containing the prior volatilities of log variances. If factor_priorsigmaidi has exactly one element, it will be recycled for all idiosyncratic log variances. Only required if type="factor".
- factor_priorsigmafac
Vector of length factor_factors containing the prior volatilities of log variances. If factor_priorsigmafac has exactly one element, it will be recycled for all factor log variances. Only required if type="factor".
- factor_priorh0idi
Vector of length 1 or M, containing information about the Gaussian prior for the initial idiosyncratic log variances. Only required if type="factor". If an element of factor_priorh0idi is a nonnegative number, the conditional prior of the corresponding initial log variance h_0 is assumed to be Gaussian with mean 0 and standard deviation factor_priorh0idi times *sigma*. If an element of factor_priorh0idi is the string 'stationary', the prior of the corresponding initial log volatility is taken to be from the stationary distribution, i.e. h_0 is assumed to be Gaussian with mean 0 and variance $\sigma^2/(1 - \phi^2)$.
- factor_priorh0fac
Vector of length 1 or factor_factors, containing information about the Gaussian prior for the initial factor log variances. Only required if type="factor". If an element of factor_priorh0fac is a nonnegative number, the conditional prior of the corresponding initial log variance h_0 is assumed to be Gaussian with mean 0 and standard deviation factor_priorh0fac times *sigma*. If an element of factor_priorh0fac is the string 'stationary', the prior of the corresponding initial log volatility is taken to be from the stationary distribution, i.e. h_0 is assumed to be Gaussian with mean 0 and variance $\sigma^2/(1 - \phi^2)$.
- factor_heteroskedastic
Vector of length 1, 2, or M + factor_factors, containing logical values indicating whether time-varying (factor_heteroskedastic = TRUE) or constant

(`factor_heteroskedastic = FALSE`) variance should be estimated. If `factor_heteroskedastic` is of length 2 it will be recycled accordingly, whereby the first element is used for all idiosyncratic variances and the second element is used for all factor variances. Only required if `type="factor"`.

`factor_priorhomoskedastic`

Only used if at least one element of `factor_heteroskedastic` is set to `FALSE`. In that case, `factor_priorhomoskedastic` must be a matrix with positive entries and dimension $c(M, 2)$. Values in column 1 will be interpreted as the shape and values in column 2 will be interpreted as the rate parameter of the corresponding inverse gamma prior distribution of the idiosyncratic variances. Only required if `type="factor"`.

`factor_interweaving`

The following values for interweaving the factor loadings are accepted (Only required if `type="factor"`):

- 0:** No interweaving.
- 1:** Shallow interweaving through the diagonal entries.
- 2:** Deep interweaving through the diagonal entries.
- 3:** Shallow interweaving through the largest absolute entries in each column.
- 4:** Deep interweaving through the largest absolute entries in each column.

For details please see Kastner et al. (2017). A value of 4 is the highly recommended default.

`cholesky_U_prior`

character, one of "HS", "R2D2", "NG", "DL", "SSVS", "HMP" or "normal". Only required if `type="cholesky"`.

`cholesky_U_tol`

Minimum number that the absolute value of a free off-diagonal element of an U -draw can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero. Only required if `type="cholesky"`.

`cholesky_heteroscedastic`

single logical indicating whether time-varying (`cholesky_heteroscedastic = TRUE`) or constant (`cholesky_heteroscedastic = FALSE`) variance should be estimated. Only required if `type="cholesky"`.

`cholesky_priormu`

Vector of length 2 denoting prior mean and standard deviation for unconditional levels of the log variance processes. Only required if `type="cholesky"`.

`cholesky_priorphi`

Vector of length 2, indicating the shape parameters for the Beta prior distributions of the transformed parameters $(\phi+1)/2$, where ϕ denotes the persistence of the log variances. Only required if `type="cholesky"`.

`cholesky_priorsigma2`

Vector of length 2, indicating the shape and the rate for the Gamma prior distributions on the variance of the log variance processes. (Currently only one global setting for all M processes is supported). Only required if `type="cholesky"`.

`cholesky_priorh0`

Vector of length 1 or M , containing information about the Gaussian prior for the initial idiosyncratic log variances. Only required if `type="cholesky"`. If an element of `cholesky_priorh0` is a nonnegative number, the conditional prior

of the corresponding initial log variance h_0 is assumed to be Gaussian with mean 0 and standard deviation $\text{cholesky_prior}h_0$ times sigma . If an element of $\text{cholesky_prior}h_0$ is the string 'stationary', the prior of the corresponding initial log volatility is taken to be from the stationary distribution, i.e. h_0 is assumed to be Gaussian with mean 0 and variance $\text{sigma}^2/(1 - \text{phi}^2)$.

`cholesky_priorhomoscedastic`

Only used if `cholesky_heteroscedastic=FALSE`. In that case, `cholesky_priorhomoscedastic` must be a matrix with positive entries and dimension $c(M, 2)$. Values in column 1 will be interpreted as the shape and values in column 2 will be interpreted as the scale parameter of the corresponding inverse gamma prior distribution of the variances. Only required if `type="cholesky"`.

`cholesky_DL_a`

(Single) positive real number. The value is interpreted as the concentration parameter for the local scales. Smaller values enforce heavier shrinkage. A matrix of dimension $c(s, 2)$ specifies a discrete hyperprior, where the first column contains s support points and the second column contains the associated prior probabilities. `cholesky_DL_a` has only to be specified if `cholesky_U_prior="DL"`.

`cholesky_DL_tol`

Minimum number that a parameter draw of one of the shrinking parameters of the Dirichlet Laplace prior can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero. `DL_tol` has only to be specified if `cholesky_U_prior="DL"`.

`cholesky_R2D2_a`

(Single) positive real number. The value is interpreted as the concentration parameter for the local scales. Smaller values enforce heavier shrinkage. A matrix of dimension $c(s, 2)$ specifies a discrete hyperprior, where the first column contains s support points and the second column contains the associated prior probabilities. `cholesky_R2D2_a` has only to be specified if `cholesky_U_prior="R2D2"`.

`cholesky_R2D2_b`

single positive number, where greater values indicate heavier regularization. `cholesky_R2D2_b` has only to be specified if `cholesky_U_prior="R2D2"`.

`cholesky_R2D2_tol`

Minimum number that a parameter draw of one of the shrinking parameters of the R2D2 prior can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero. `cholesky_R2D2_tol` has only to be specified if `cholesky_U_prior="R2D2"`.

`cholesky_NG_a`

(Single) positive real number. The value is interpreted as the concentration parameter for the local scales. Smaller values enforce heavier shrinkage. A matrix of dimension $c(s, 2)$ specifies a discrete hyperprior, where the first column contains s support points and the second column contains the associated prior probabilities. `cholesky_NG_a` has only to be specified if `cholesky_U_prior="NG"`.

`cholesky_NG_b`

(Single) positive real number. The value indicates the shape parameter of the inverse gamma prior on the global scales. `cholesky_NG_b` has only to be specified if `cholesky_U_prior="NG"`.

`cholesky_NG_c`

(Single) positive real number. The value indicates the scale parameter of the inverse gamma prior on the global scales. Expert option would be to set the scale parameter proportional to `NG_a`. E.g. in the case where a discrete hyperprior

for NG_a is chosen, a desired proportion of let's say 0.2 is achieved by setting NG_c="0.2a" (character input!). cholesky_NG_c has only to be specified if cholesky_U_prior="NG".

cholesky_NG_tol

Minimum number that a parameter draw of one of the shrinking parameters of the normal-gamma prior can take. Prevents numerical issues that can appear when strong shrinkage is enforced if chosen to be greater than zero. cholesky_NG_tol has only to be specified if cholesky_U_prior="NG".

cholesky_SSVS_c0

single positive number indicating the (unscaled) standard deviation of the spike component. cholesky_SSVS_c0 has only to be specified if cholesky_U_prior="SSVS". It should be that $SSVS_{c0} \ll SSVS_{c1}$!

cholesky_SSVS_c1

single positive number indicating the (unscaled) standard deviation of the slab component. cholesky_SSVS_c1 has only to be specified if cholesky_U_prior="SSVS". It should be that $SSVS_{c0} \ll SSVS_{c1}$!

cholesky_SSVS_p

Either a single positive number in the range $(0, 1)$ indicating the (fixed) prior inclusion probability of each coefficient. Or numeric vector of length 2 with positive entries indicating the shape parameters of the Beta distribution. In that case a Beta hyperprior is placed on the prior inclusion probability. cholesky_SSVS_p has only to be specified if cholesky_U_prior="SSVS".

cholesky_HMP_lambda3

numeric vector of length 2. Both entries must be positive. The first indicates the shape and the second the rate of the Gamma hyperprior on the contemporaneous coefficients. cholesky_HMP_lambda3 has only to be specified if cholesky_U_prior="HMP".

cholesky_normal_sds

numeric vector of length $\frac{M^2-M}{2}$, indicating the prior variances for the free off-diagonal elements in U . A single number will be recycled accordingly! Must be positive. cholesky_normal_sds has only to be specified if cholesky_U_prior="normal".

expert_sv_offset

... Do not use!

quiet

logical indicating whether informative output should be omitted.

...

Do not use!

Details

bvar offers two different specifications for the errors: The user can choose between a factor stochastic volatility structure or a cholesky stochastic volatility structure. In both cases the disturbances ϵ_t are assumed to follow a M -dimensional multivariate normal distribution with zero mean and variance-covariance matrix Σ_t . In case of the cholesky specification $\Sigma_t = U'^{-1} D_t U^{-1}$, where U^{-1} is upper unitriangular (with ones on the diagonal). The diagonal matrix D_t depends upon latent log-variances, i.e. $D_t = \text{diag}(\exp(h_{1t}), \dots, \exp(h_{Mt}))$. The log-variances follow a priori independent autoregressive processes $h_{it} \sim N(\mu_i + \phi_i(h_{i,t-1} - \mu_i), \sigma_i^2)$ for $i = 1, \dots, M$. In case of the factor structure, $\Sigma_t = \Lambda V_t \Lambda' + G_t$. The diagonal matrices V_t and G_t depend upon latent log-variances, i.e. $G_t = \text{diag}(\exp(h_{1t}), \dots, \exp(h_{Mt}))$ and $V_t =$

$\text{diag}(\exp(h_{M+1,t}), \dots, \exp(h_{M+r,t}))$. The log-variances follow a priori independent autoregressive processes $h_{it} \sim N(\mu_i + \phi_i(h_{i,t-1} - \mu_i), \sigma_i^2)$ for $i = 1, \dots, M$ and $h_{M+j,t} \sim N(\phi_j h_{M+j,t-1}, \sigma_{M+j}^2)$ for $j = 1, \dots, r$.

Value

Object of class `bayesianVARs_prior_sigma`.

References

Kastner, G. (2019). Sparse Bayesian Time-Varying Covariance Estimation in Many Dimensions *Journal of Econometrics*, **210**(1), 98–115, doi:10.1016/j.jeconom.2018.11.007

Kastner, G., Frühwirth-Schnatter, S., and Lopes, H.F. (2017). Efficient Bayesian Inference for Multivariate Factor Stochastic Volatility Models. *Journal of Computational and Graphical Statistics*, **26**(4), 905–917, doi:10.1080/10618600.2017.1322091.

See Also

[specify_prior_phi\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# examples with stochastic volatility (heteroscedasticity) -----
# factor-decomposition with 2 factors and colwise normal-gamma prior on the loadings
sigma_factor_cng_sv <- specify_prior_sigma(data = data, type = "factor",
factor_factors = 2L, factor_priorfacloadtype = "colwiseng", factor_heteroskedastic = TRUE)

# cholesky-decomposition with Dirichlet-Laplace prior on U
sigma_cholesky_dl_sv <- specify_prior_sigma(data = data, type = "cholesky",
cholesky_U_prior = "DL", cholesky_DL_a = 0.5, cholesky_heteroscedastic = TRUE)

# examples without stochastic volatility (homoscedasticity) -----
# factor-decomposition with 2 factors and colwise normal-gamma prior on the loadings
sigma_factor_cng <- specify_prior_sigma(data = data, type = "factor",
factor_factors = 2L, factor_priorfacloadtype = "colwiseng",
factor_heteroskedastic = FALSE, factor_priorhomoskedastic = matrix(c(0.5,0.5),
ncol(data), 2))

# cholesky-decomposition with Horseshoe prior on U
sigma_cholesky_dl <- specify_prior_sigma(data = data, type = "cholesky",
cholesky_U_prior = "HS", cholesky_heteroscedastic = FALSE)

# Estimate model with your prior configuration of choice
mod <- bvar(data, prior_sigma = sigma_factor_cng_sv, quiet = TRUE)
```

```
specify_structural_restrictions
```

Set identifying restrictions for the structural VAR parameters.

Description

Set identifying restrictions for the structural VAR parameters.

Usage

```
specify_structural_restrictions(  
  x,  
  restrictions_facload = NULL,  
  restrictions_B0_inv_t = NULL,  
  restrictions_B0 = NULL,  
  restrictions_structural_coeff = NULL,  
  restrictions_long_run_ir = NULL  
)
```

Arguments

x An object of type `bayesianVARs_bvar`.

restrictions_facload an M times r matrix of restrictions on the factor loadings. This is equivalent to the instantaneous effects of the factor shocks. Can only be used, if the factor decomposition for σ was specified.

restrictions_B0_inv_t an M times M matrix of restrictions on the instantaneous effects of the structural shocks. Columns correspond to shocks, rows to variables. Can only be used, if the cholesky decomposition for σ was specified.

restrictions_B0 an M times M matrix of restrictions on the structural matrix. Can only be used, if the cholesky decomposition for σ was specified.

restrictions_structural_coeff a matrix of restrictions on the structural VAR coefficients. Its size should match the dimensions of `x$PHI`. Can only be used, if the cholesky decomposition for σ was specified.

restrictions_long_run_ir a matrix of restrictions on the long run impulse responses. The long run impulse responses are the sum of the impulse responses summed over all time horizons. Restrictions on the long run impulse responses can be specified for both the factor and the cholesky decomposition of σ . In the case of a factor decomposition its size is expected to be M times r . In the case of a cholesky decomposition the size must be M times M .

Details

All `restrictions_*` entries have following meaning

NA: an unrestricted entry.

0: The entry at this position is restricted to be zero (i.e. an exclusion restriction).

A positive number: The sign of this entry should be positive.

A negative number: The sign of this entry should be negative.

The structural VAR(p) model is of the following form:

$$y'_t B_0 = x'_t \Phi B_0 + \omega'_t$$

Author(s)

Stefan Haan <sthaan@edu.aau.at>

See Also

[irf](#), [extractB0](#), [specify_prior_sigma](#)

Examples

```
train_data <- 100 * usmacro_growth[,c("GDPC1", "GDPCTPI", "GS1", "M2REAL", "CPIAUCSL")]
prior_sigma <- specify_prior_sigma(train_data, type="cholesky", cholesky_heteroscedastic=FALSE)
mod <- bvar(train_data, lags=5L, prior_sigma=prior_sigma, quiet=TRUE)

structural_restrictions <- specify_structural_restrictions(
  mod,
  restrictions_B0=rbind(
    c(1 ,NA,0 ,NA,NA),
    c(0 ,1 ,0 ,NA,NA),
    c(0 ,NA,1 ,NA,NA),
    c(0 ,0 ,NA,1 ,NA),
    c(0 ,0 ,0 ,0 ,1 )
  )
)
irf_structural <- irf(
  mod, ahead=8,
  structural_restrictions=structural_restrictions
)
plot(irf_structural)
```

stable_bvar	<i>Stable posterior draws</i>
-------------	-------------------------------

Description

stable_bvar() detects and discards all posterior draws of an bayesianVARs_bvar object that do not fulfill the stability condition: A VAR(p) model is considered as stable only if the eigenvalues of the companion form matrix lie inside the unit circle.

Usage

```
stable_bvar(object, quiet = FALSE)
```

Arguments

object	A bayesianVARs_bvar object obtained via <code>bvar()</code> .
quiet	logical indicating whether informative output should be omitted.

Value

An object of type bayesianVARs_bvar.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Discard "unstable" draws
stable_mod <- stable_bvar(mod)
```

summary.bayesianVARs_bvar	<i>Summary method for bayesianVARs_bvar objects</i>
---------------------------	---

Description

Summary method for bayesianVARs_bvar objects.

Usage

```
## S3 method for class 'bayesianVARs_bvar'
summary(object, quantiles = c(0.025, 0.25, 0.5, 0.75, 0.975), digits = 3L, ...)
```

Arguments

object	A bayesianVARs_bvar object obtained via <code>bvar()</code> .
quantiles	numeric vector which quantiles to compute.
digits	Single integer indicating the number of decimal places to be used for rounding the summary statistics. Negative values are not allowed.
...	Currently ignored!

Value

An object of type `summary.bayesianVARs_bvar`.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate model
mod <- bvar(data, quiet = TRUE)

# Summary
sum <- summary(mod)
```

```
summary.bayesianVARs_draws
```

Summary statistics for bayesianVARs posterior draws.

Description

Summary statistics for bayesianVARs posterior draws.

Usage

```
## S3 method for class 'bayesianVARs_draws'
summary(object, quantiles = c(0.25, 0.5, 0.75), ...)
```

Arguments

object	An object of class <code>bayesianVARs_draws</code> usually obtained through extractors like <code>coef.bayesianVARs_bvar()</code> and <code>vcov.bayesianVARs_bvar()</code> .
quantiles	A vector of quantiles to evaluate.
...	Currently ignored.

Value

A list object of class bayesianVARs_draws_summary holding

- mean: Vector or matrix containing the posterior mean.
- sd: Vector or matrix containing the posterior standard deviation .
- quantiles: Array containing the posterior quantiles.

See Also

Available extractors: [coef.bayesianVARs_bvar\(\)](#), [vcov.bayesianVARs_bvar\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]
# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Extract posterior draws of VAR coefficients
bvar_coefs <- coef(mod)

# Compute summary statistics
summary_stats <- summary(bvar_coefs)

# Compute summary statistics of VAR coefficients without using coef()
summary_stats <- summary(mod$PHI)

# Test which list elements of 'mod' are of class 'bayesianVARs_draws'.
names(mod)[sapply(names(mod), function(x) inherits(mod[[x]], "bayesianVARs_draws"))]
```

```
summary.bayesianVARs_predict
```

Summary method for bayesianVARs_predict objects

Description

Summary method for bayesianVARs_predict objects.

Usage

```
## S3 method for class 'bayesianVARs_predict'
summary(object, ...)
```

Arguments

object	A bayesianVARs_predict object obtained via predict.bayesianVARs_bvar() .
...	Currently ignored!

Value

A summary.bayesianVARs_predict object.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Split data in train and test
train <- data[1:(nrow(data)-4),]
test <- data[-c(1:(nrow(data)-4)),]

# Estimate model using train data only
mod <- bvar(train, quiet = TRUE)

# Simulate from 1-step ahead posterior predictive
predictions <- predict(mod, ahead = 1L)
summary(predictions)
```

usmacro_growth *Data from the US-economy*

Description

21 selected quarterly time-series from 1953:Q1 to 2021:Q2. From FRED-QD data base (McCracken and Ng, 2021). Release date 2021-07. Data is transformed to be interpreted as growth-rates (first log-differences with the exception of interest rates, which are already growth rates).

Usage

```
usmacro_growth
```

Format

A matrix with 247 rows and 21 columns.

Source

Raw (untransformed) data available at <https://www.stlouisfed.org/research/economists/mccracken/fred-databases>, <https://www.stlouisfed.org/-/media/project/frbstl/stlouisfed/research/fred-md/historical-vintages-of-fred-qd-2018-05-to-2024-12.zip>.

References

McCracken, M. W. and Ng, S. (2021). FRED-QD: A Quarterly Database for Macroeconomic Research, *Review, Federal Reserve Bank of St. Louis*, **103**(1), 1–44, doi:10.20955/r.103.144.

 vcov.bayesianVARs_bvar

Extract posterior draws of the (time-varying) variance-covariance matrix for a VAR model

Description

Returns the posterior draws of the possibly time-varying variance-covariance matrix of a VAR estimated via `bvar()`. Returns the full paths if `sv_keep="all"` when calling `bvar()`. Otherwise, the draws of the variance-covariance matrix for the last observation are returned, only.

Usage

```
## S3 method for class 'bayesianVARs_bvar'
vcov(object, t = seq_len(nrow(object$logvar)), ...)
```

Arguments

<code>object</code>	An object of class <code>bayesianVARs_bvar</code> obtained via <code>bvar()</code> .
<code>t</code>	Vector indicating which points in time should be extracted, defaults to all.
<code>...</code>	Currently ignored.

Value

An array of class `bayesianVARs_draws` of dimension $T \times M \times M \times draws$, where T is the number of observations, M the number of time-series and $draws$ the number of stored posterior draws.

See Also

[summary.bayesianVARs_draws](#), [coef.bayesianVARs_bvar\(\)](#).

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]
# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Extract posterior draws of the variance-covariance matrix
bvar_vcov <- vcov(mod)
```

[.bayesianVARs_coef *Extract or Replace Parts of a bayesianVARs_coef object*

Description

Extract or replace parts of a bayesianVARs_coef object.

Usage

```
## S3 method for class 'bayesianVARs_coef'  
x[i, j, ...]
```

Arguments

x	An object of type bayesianVARs_coef.
i	indices
j	indices
...	further indices

Value

An object of type bayesianVARs_coef.

Examples

```
# Access a subset of the usmacro_growth dataset  
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]  
  
# Estimate a model  
mod <- bvar(data, sv_keep = "all", quiet = TRUE)  
  
# Extract coefficients, which are of class bayesianVARs_coef  
phi <- coef(mod)  
phi[1,1,1]
```

[.bayesianVARs_draws *Extract or Replace Parts of a bayesianVARs_draws object*

Description

Extract or replace parts of a bayesianVARs_draws object.

Usage

```
## S3 method for class 'bayesianVARs_draws'  
x[i, j, ...]
```

Arguments

x	An object of type bayesianVARs_draws.
i	indices
j	indices
...	further indices

Value

An object of type bayesianVARs_draws.

Examples

```
# Access a subset of the usmacro_growth dataset
data <- usmacro_growth[,c("GDPC1", "CPIAUCSL", "FEDFUNDS")]

# Estimate a model
mod <- bvar(data, sv_keep = "all", quiet = TRUE)

# Extract coefficients, which are of class bayesianVARs_draws
phi <- coef(mod)
phi[1,1,1]
```

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