Package ‘MARSS’
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Description The MARSS package provides maximum-likelihood parameter estimation for constrained and unconstrained linear multivariate autoregressive state-space (MARSS) models fit to multivariate time-series data. Fitting is primarily via an Expectation-Maximization (EM) algorithm, although fitting via the BFGS algorithm (using the optim function) is also provided. MARSS models are a class of dynamic linear model (DLM) and vector autoregressive model (VAR) model. Functions are provided for parametric and innovations bootstrapping, Kalman filtering and smoothing, bootstrap model selection criteria (AICb), confidences intervals via the Hessian approximation and via bootstrapping and calculation of auxiliary residuals for detecting outliers and shocks. The user guide shows examples of using MARSS for parameter estimation for a variety of applications, model selection, dynamic factor analysis, outlier and shock detection, and addition of covariates. Type RShowDoc("UserGuide", package="MARSS") at the R command line to open the MARSS user guide. Online workshops (lectures and computer labs) at <https://nwfsc-timeseries.github.io/> See the NEWS file for update information.
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Description

The MARSS package fits constrained and unconstrained multivariate autoregressive time-series models to multivariate time series data. To open the user guide from the command line, type `rShowDoc("UserGuide", package="MARSS")`. To open an overview page with package information and links to the R scripts in the User Guide, type `rShowDoc("index", package="MARSS")`.

The main function is `marss` which is used to fit a specified model to data and estimate the model parameters. MARSS model specification is based on "form" (an argument to a MARSS() call). The form tells MARSS() what to expect in the model list (model is a MARSS argument) and how to translate that into the base model form used in the fitting algorithms.

The default MARSS model form is "marxss", which is a model of the following form:

\[
\begin{align*}
  x(t+1) &= B x(t) + U + C c(t) + G w(t), \text{ where } w(t) \sim \text{MVN}(0, Q) \\
  y(t) &= Z x(t) + A + D d(t) + H v(t), \text{ where } v(t) \sim \text{MVN}(0, R) \\
  x(1) &\sim \text{MVN}(x_0, V_0) \text{ or } x(0) \sim \text{MVN}(x_0, V_0)
\end{align*}
\]

The parameters, hidden state processes (x), and observations (y) are matrices:

- x(t) is m x 1
- y(t) is n x 1 (m≤=n)
- Z is n x m
- B is m x m
- U is m x 1
- Q is g x g (typically m x m)
- G is m x g
- A is n x 1
- R is h x h (typically n x n)
- H is n x h
MARSS-package

- C is m x q
- D is n x p
- c(t) is q x 1
- d(t) is q x 1
- x0 is m x 1
- V0 is m x m

All parameters can be time-varying.

The package functions estimate the model parameters using an EM algorithm (primarily but see `MARSSoptim`). Parameters may be constrained to have shared elements (elements which are constrained to have the same value) or fixed elements (with the other elements estimated). The states and smoothed state estimates are provided via a Kalman filter and smoother. Bootstrapping, confidence interval estimation, bias estimation, model selection and simulation functions are provided. The main user interface to the package is the top-level function `MARSS`.

Details

Important MARSS functions:

- `MARSS` Top-level function for specifying and fitting MARSS models.
- `MARSSsimulate` Produces simulated data from a MARSS model.
- `MARSSkem` Estimates MARSS parameters using an EM algorithm.
- `MARSSkf` Kalman filter and smoother.
- `MARSSoptim` Estimates MARSS parameters using a quasi-Newton algorithm via `optim`.
- `MARSSaic` Calculates AICc, AICc, and various bootstrap AICs.
- `MARSSboot` Creates bootstrap MARSS parameter estimates.
- `MARSSparamCIs` Computes confidence intervals for maximum-likelihood estimates of MARSS parameters.

Author(s)

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References

Type `RshowDoc("index",package="MARSS")` to see all the package documentation, tutorials, and R scripts from the User Guide.
**MARSS Function Defaults and Allowed Methods**

**Description**

Defaults and allowed fitting methods for the MARSS function are specified in the file MARSSsettings.R. These are hidden thus to see them preface with MARSS:::

**Details**

allowed.methods is a vector with the allowed method arguments for the MARSS function. kem.methods and optim.methods are vectors of method arguments that fall in each of these two categories; used by MARSS. alldefaults is a list that specifies the defaults for MARSS arguments if they are not passed in.

**augment.marssMLE**

*Return the model predicted values, residuals, and optionally confidence intervals*

**Description**

Return a data.frame with the observations or states fitted values, residuals, and upper and lower confidence intervals if requested.

**Usage**

```r
augment.marssMLE(x, type.predict = c("observations", "states"),
                 interval = c("none", "confidence"),
                 conf.level = 0.95, form=attr(x["model"], "form"))
```

**Arguments**

- `x` a marssMLE object
- `type.predict` Type of prediction: for the observations (y) or for the states (x). Read the details below for states. tidy would be the more common function for returning state estimates.
- `interval` Type of interval: none or confidence interval. If the latter, approximate intervals from the standard errors of the fitted values is given.
- `conf.level` Confidence level.
- `form` If you want the augment function to use a different augment function than augment_form. This might be useful if you manually specified a DFA model and want to use augment.dfa for rotating.
Details

See `residuals.marssMLE` for a discussion of the residuals calculations for MARSS models. The reported CIs are the approximate CIs computed using the standard deviations: \( \text{qnorm}(\alpha/2) \times \text{se.fit} + \text{fitted} \).

**type.predict observations**

This returns a familiar model predicted value of the response \( (y) \) and the difference between the model prediction and the actual data \( y(t) \) is the residual. If there is no data point, then the residual is NA. The standard errors help visualize how well the model fits to the data. See `fitted.marssMLE` for a discussion of the calculation of the fitted values for the observations (the modeled values). The standardized residuals can be used for outlier detection. See `residuals.marssMLE` and the chapter on shock detection in the user guide.

In the literature on state-space models, it is very common to use the one-step ahead predicted values of the data. The fitted values returned by augment are NOT the one-step ahead values and the residuals are not the one-step ahead residuals (called Innovations in the state-space literature). If you want the one-step ahead fitted values, you can use `fitted(x, one.step.ahead=TRUE)`. The innovations are returned by `MARSSkf`.

**type.predict states**

The states are estimated. If you want the expected value of the states and an estimate of their standard errors (for confidence intervals), then augment is not what you want to use. You want to use tidy to return the estimate of the state.

`augment(MLEobj, type.predict="states")` returns a model prediction of \( \hat{x}(t) \) given \( \hat{x}(t - 1) \).

The residuals returned are for \( w(t) \), the difference between \( \hat{x}(t) \) and the prediction of \( \hat{x}(t) \). These types of residuals are used for outlier detection or shock detection in the state process. They are also used for model diagnostics. See `residuals.marssMLE` and read the references cited.

Examples

```r
# Make a plot of the observations and model fits
dat <- t(harborSeal)
dat <- dat[c(2,11,12),]
MLEobj <- MARSS(dat, model=list(Z=factor(c("WA","OR","OR"))))
library(broom)
library(ggplot2)
theme_set(theme_bw())

# Make a plot of the observations and model fits
# Don't use augment. States are not data.
d <- tidy(MLEobj, type="states")
```
checkMARSSInputs

checkMARSSInputs( MARSS.inputs, silent=FALSE )

Arguments

MARSS.inputs A list comprised of the needed inputs to a MARSS call: data, inits, model, control, method, form)
silent Suppresses printing of progress bars, error messages, warnings and convergence information.

Details

This is a helper function to check that all the inputs to a MARSS call are properly specified.
If arguments inits or control are not provided by the user, they will be set by the alldefaults[[method]] object specified in MARSSsettings. Argument model specifies the model structure using a list of matrices; see MARSS or the User Guide for instructions on how to specify model structure. If model is left off, then the function MARSS.form() is used to determine the default model structure.

Value

If the function does not stop due to errors, it returns an updated list with elements
data Data supplied by user.
model Not changed. Will be updated by the MARSS.form function (e.g. MARSS.marxss).
inits A list specifying initial values for parameters to be used at iteration 1 in iterative maximum-likelihood algorithms.
method The method used for estimation.
form The equation form used to convert wrapper object to a marssMODEL object.
control See Arguments.

Author(s)

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checkModelList

**See Also**

MARSS marssMODEL checkModelList

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checkModelList  
*Check model List Passed into MARSS Call*

**Description**

This is a helper function to check the model list passed in to a MARSS() call for any errors. Not exported.

**Usage**

checkModelList( model, defaults, this.form.allows)

**Arguments**

- **model**: A list from which a marssMODEL model will be constructed.
- **defaults**: A list with the defaults for the elements in the model list in case the user leaves out any.
- **this.form.allows**: A list of what inputs are allowed for each element in the model list.

**Details**

This is a helper function to check that all the model list that will be passed to a MARSS form function to make the marssMODEL object. If elements in the list are left off, they will be filled in by defaults.

**Value**

If the function does not stop due to errors, it returns an updated model list with missing elements filled in by the defaults.

**Author(s)**

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**See Also**

MARSS marssMODEL checkModelList
coeff.marssMLE

## Description

The MARSS fitting function, MARSS, outputs marssMLE objects. `coeff(marssMLE)`, where marssMLE is one’s output from a MARSS call, will print out the estimated parameters. The default output is a list with the estimated parameters for each MARSS parameter; however, `coeff` can be altered using the `type` argument to output a vector of all the estimated values (`type="vector"`) or a list with the full parameter matrix with the estimated and fixed elements (`type="matrix"`).

## Usage

```r
## S3 method for class 'marssMLE'
coef(object, ..., type="list", form=NULL, what="par")
```

## Arguments

- **object**: A marssMLE object.
- **...**: Other arguments for `coef`.
- **type**: What to print. Default is "list". If you input `type` as a vector, `coef` returns a list of output. See examples.
  - "list": A list of only the estimated values in each matrix. Each model matrix has its own list element.
  - "vector": A vector of all the estimated values in each matrix.
  - "matrix": A list of the parameter matrices each parameter with fixed values at their fixed values and the estimated values at their estimated values. Time-varying parameters, including d and c in a marxss form model, are returned as an array with time in the 3rd dimension.
  - parameter name: Returns the parameter matrix for that parameter with fixed values at their fixed values and the estimated values at their estimated values. Note, time-varying parameters, including d and c in a marxss form model, are returned as an array with time in the 3rd dimension.
- **form**: By default, `coeff` uses the model form specified in the call to MARSS to determine how to display the coefficients. This information is in `attr(marssMLE$model, "form")`, however you can specify a different form. The default form is "marxss" which is a MARSS model with inputs, see MARSS.marxss. However, the MARSS EM algorithm transforms the model into form "marss"; see MARSS.marss. The marss form of the model is stored in `marssMLE$marss`. You can look at the coefficients in marss form by passing in `form="marss"` (this will return `marssMLE$par`). This is mainly useful for debugging numerical problems since the error reports will be for the model in marss form.
- **what**: By default, `coeff` shows the parameter estimates. Other options are "par.se", "par.lowCI", "par.upCI", "par.bias", and "start".
Value

A list of the estimated parameters for each model matrix.

Author(s)

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See Also

augment.marssMLE tidy.marssMLE print.marssMLE

Examples

dat <- t(harborSeal)
dat <- dat[,c(2,11),]
MLEobj <- MARSS(dat)

coef(MLEobj)
coef(MLEobj,type="vector")
coef(MLEobj,type="matrix")
#to retrieve just the Q matrix
coef(MLEobj,type="matrix")$Q

CSEGriskfigure  Plot Extinction Risk Metrics

Description

Generates a six-panel plot of extinction risk metrics used in Population Viability Analysis (PVA). This is a function used by one of the vignettes in the MARSS-package.

Usage

CSEGriskfigure(data, te = 100, absolutethresh = FALSE, threshold = 0.1,
datalogged = FALSE, silent = FALSE, return.model = FALSE,
CI.method = "hessian", CI.sim = 1000)

Arguments

data       A data matrix with 2 columns; time in first column and counts in second column. Note time is down rows, which is different than the base MARSS-package functions.
te         Length of forecast period (positive integer)
absolutethresh Is extinction threshold an absolute number? (T/F)
### CSEGriskfigure

<table>
<thead>
<tr>
<th><strong>threshold</strong></th>
<th>Extinction threshold either as an absolute number, if <code>absolutethresh=TRUE</code>, or as a fraction of current population count, if <code>absolutethresh=FALSE</code>.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>datalogged</strong></td>
<td>Are the data already logged? (T/F)</td>
</tr>
<tr>
<td><strong>silent</strong></td>
<td>Suppress printed output? (T/F)</td>
</tr>
<tr>
<td><strong>return.model</strong></td>
<td>Return state-space model as <code>marssMLE</code> object? (T/F)</td>
</tr>
<tr>
<td><strong>CI.method</strong></td>
<td>Confidence interval method: &quot;hessian&quot;, &quot;parametric&quot;, &quot;innovations&quot;, or &quot;none&quot;. See <code>MARSSparamCIs</code>.</td>
</tr>
<tr>
<td><strong>CI.sim</strong></td>
<td>Number of simulations for bootstrap confidence intervals (positive integer).</td>
</tr>
</tbody>
</table>

#### Details


#### Value

If `return.model=TRUE`, an object of class `marssMLE`.

#### Author(s)

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#### References


#### See Also

`MARSSboot marssMLE CSEGtmufigure`

#### Examples

```r
d <- harborSeal[,1:2]
kem <- CSEGriskfigure(d, datalogged = TRUE)
```
Description

Plot the uncertainty in the probability of hitting a percent threshold (quasi-extinction) for a single random walk trajectory. This is the quasi-extinction probability used in Population Viability Analysis. The uncertainty is shown as a function of the forecast, where the forecast is defined in terms of the forecast length (number of time steps) and forecasted decline (percentage). This is a function used by one of the vignettes in the MARSS-package.

Usage

CSEGtmufiure(N = 20, u = -0.1, s2p = 0.01, make.legend = TRUE)

Arguments

N  Time steps between the first and last population data point (positive integer)
u  Per time-step decline (-0.1 means a 10% decline per time step; 1 means a doubling per time step.)
s2p  Process variance (Q). (a positive number)
make.legend  Add a legend to the plot? (T/F)

Details

This figure shows the region of high uncertainty in dark grey. In this region, the minimum 95 percent confidence intervals on the probability of quasi-extinction span 80 percent of the 0 to 1 probability. Green hashing indicates where the 95 percent upper bound does not exceed 5% probability of quasi-extinction. The red hashing indicates, where the 95 percent lower bound is above 95% probability of quasi-extinction. The light grey lies between these two certain/uncertain extremes. The extinction calculation is based on Dennis et al. (1991). The minimum theoretical confidence interval is based on Fieberg and Ellner (2000). This figure was developed in Ellner and Holmes (2008).

Examples using this figure are shown in the user guide (RShowDoc("UserGuide",package="MARSS")) in the PVA chapter.

Author(s)

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References


See Also
CSEGriskfigure

Examples
CSEGtmufigure(N = 20, u = -0.1, s2p = 0.01)

fitted.marssMLE fitted function for MARSS MLE objects

Description
The MARSS fitting function, MARSS, outputs marssMLE objects. fitted(marssMLE), where marssMLE is the output from a MARSS call, will return the modeled value of y(t) or x(t). See details.

Usage
## S3 method for class 'marssMLE'
fitted(object, ..., one.step.ahead=FALSE, type=c("observations", "states"))

Arguments
object
... one.step.ahead
type

Details
observation fitted values
The model predicted (fitted) y at time t is \( Z(t)\dot{x}(t) + a(t) \), where the model is written in marss form. See MARSS.marxss for a discussion of the conversion of MARSS models with covariates (c and d) into marss form (which is how models are written in the internal MARSS algorithms). \( \dot{x}(t) \) is the expected value of the states at time t. If one.step.ahead=FALSE, \( \dot{x}(t) \) is the expected value conditioned on all the data, i.e. xtt returned by MARSSkf. If one.step.ahead=TRUE, \( \dot{x}(t) \) is the expected value conditioned on the data up to time t-1, i.e. xtt1 returned by MARSSkf. These are commonly known as the one step ahead predictions for a state-space model.

state fitted values
The model predicted \( \dot{x}(t) \) given \( \dot{x}(t-1) \) is \( B(t)\dot{x}(t-1) + u(t) \), where the model is written in marss form.
\( \hat{x}(t) \) is the expected value of the states at time \( t \). If one \_step\_ahead=FALSE, \( \hat{x}(t) \) is the expected value conditioned on all the data, i.e. \( \text{xtT} \) returned by \texttt{MARSSkf}. If one \_step\_ahead=TRUE, \( \hat{x}(t) \) is the expected value conditioned on the data up to time \( t-1 \), i.e. \( \text{xtt1} \) returned by \texttt{MARSSkf}. This type of state fitted value is used for process outlier detection and shock detection. See \texttt{residuals.marssMLE} and read the references cited.

If you want estimates of the states, which is want one typically wants not the fitted values, then you’ll want either the states estimate conditioned on all the data or conditioned on the data up to time \( t-1 \). These are returned by \texttt{MARSSkf} in \( \text{xtT} \) and \( \text{xtt1} \) respectively. Which one you want depends on your objective and application. You can also use the \texttt{tidy.marssMLE} function to return a \texttt{data.frame} with the \( \text{xtT} \), standard errors and confidence intervals.

**Value**

A \( T \) column matrix of the fitted values with one row for each observation (or state) time series.

**Author(s)**

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**Description**

This returns a \texttt{data.frame} with brief summary information.

- **coef.det** The coefficient of determination. This is the squared correlation between the fitted values and the original data points. This is simply a metric for the difference between the data points and the fitted values and should not be used for formal model comparison.
- **sigma** The sample variance (unbiased) of the data residuals (fitted minus data). This is another simple metric of the difference between the data and fitted values. This is different than the sigma returned by an \texttt{arima()} call for example. That sigma would be akin to \( \sqrt{\text{t}(Q)} \) in the MARSS parameters; 'akin' because MARSS models are multivariate and the sigma returned by \texttt{arima} is for a univariate model.
- **df** The number of estimated parameters. Denoted \texttt{num.params} in a \texttt{marssMLE} object.
- **logLik** The log-likelihood.
- **AIC** Akaike information criterion.
- **AICc** Akaike information criterion corrected for small sample size.
- **AICbb** Non-parametric bootstrap Akaike information criterion if .
- **AICbp** Parametric bootstrap Akaike information criterion if .
- **convergence** 0 if converged according to the convergence criteria set. Note the default convergence criteria are high in order to speed up fitting. A number other than 0 means the model did not meet the convergence criteria.
- **errors** 0 if no errors. 1 if some type of error or warning returned.
graywhales

Usage

\texttt{glance.marssMLE(x, \ldots)}

Arguments

\begin{itemize}
  \item \texttt{x} A marssMLE object
  \item \texttt{\ldots} Not used.
\end{itemize}

Examples

\begin{verbatim}
dat <- t(harborSeal)
dat <- dat[,c(2,11,12),]
MLEobj <- MARSS(dat, model=list(Z=factor(c("WA","OR","OR"))))

library(broom)
glance(MLEobj)
\end{verbatim}

\begin{longtable}{ll}
graywhales & \textit{Population Data Sets} \\
\end{longtable}

Description

Example data sets for use in the \texttt{MARSS-package} User Guide. Some are logged and some unlogged population counts. See the details below on each dataset.

The data sets are matrices with year in the first column and counts in other columns. Since MARSS functions require time to be across columns, these data matrices must be transposed before passing into MARSS functions.

Usage

\begin{verbatim}
data(graywhales)
data(grouse)
data(prairiechicken)
data(wilddogs)
data(kestrel)
data(-okanaganRedds)
data(rockfish)
data(redstart)
\end{verbatim}

Format

The data are supplied as a matrix with years in the first column and counts in the second (and higher) columns.
Source

- okanaganRedds A dataset of Chinook salmon redd (egg nest) surveys. This data comes from the Okanagan River in Washington state, a major tributary of the Columbia River (headwaters in British Columbia). Unlogged.
- rockfish LOGGED catch per unit effort data for Puget Sound total total rockfish (mix of species) from a series of different types of surveys.
- kestrel Three time series of American kestrel logged abundance from adjacent Canadian provinces along a longitudinal gradient (British Columbia, Alberta, Saskatchewan). Data have been collected annually, and corrected for changes in observer coverage and detectability. LOGGED.

Examples

str(graywhales)
str(grouse)
str(prairiechicken)
str(wilddogs)
str(kestrel)
str(OKanaganRedds)
str(rockfish)

Harbor Seal Population Count Data (Log counts)

Description

Data sets used in MARSS vignettes in the MARSS-package. These are data sets based on LOGGED count data from Oregon, Washington and California sites where harbor seals were censused while hauled out on land. "harborSealnomiss" is an extrapolated data set where missing values in the original dataset have been extrapolated so that the data set can be used to demonstrate fitting population models with different underlying structures.
Usage

data(harborSeal)
data(harborSealWA)

Format


Details

Matrix "harborSealWA" contains the original 1978-1999 LOGGED count data for five inland WA sites. Matrix "harborSealnomiss" contains 1975-2003 data for the same sites as well as four coastal sites, where missing values have been replaced with extrapolated values. Matrix "harborSeal" contains the original 1975-2003 LOGGED data (with missing values) for the WA and OR sites as well as a CA Mainland and CA ChannelIslands time series.

Source


Examples

str(harborSealWA)
str(harborSeal)
Description
These are model objects and utility functions for model objects in the package MARSS-package. Users would not normally work directly with these functions. is.marssMODEL() ensures model consistency. MARSS_formname() translates a model list as passed in call to MARSS() into a marssMODEL model object.

Usage
is.marssMODEL(MODELobj, method="kem")

Arguments

  MODELobj | An object of class marssMODEL.
  method   | Method used for fitting in case there are special constraints for that method.

Details
A marssMODEL object is an R representation of a MARSS model along with the data. Data in a marssMODEL object consists of multivariate time series data in which time is across columns and the n observed time series are in the n different rows.

The base MARSS model (form=marss) is

\[ x(t+1) = B(t) \cdot x(t) + U(t) + w(t), \text{ where } w(t) \sim MVN(0,Q(t)) \]
\[ y(t) = Z(t) \cdot x(t) + A(t) + v(t), \text{ where } v(t) \sim MVN(0,R(t)) \]
\[ x(1) \sim MVN(x0, V0) \]

The marssMODEL(form=marss) object describes this MARSS model but written in vec form:

\[ x(t+1) = \text{kron}(x(t),I)(f_b(t)+D_b(t)b) + (f_u(t)+D_u(t)u) + w(t), \text{ where } w(t) \sim MVN(0,Q) \]

\[ \text{vec}(Q) = f_q(t)+D_q(t)q \]
\[ y(t) = \text{kron}(x(t),I)(f_z(t)+D_z(t)z) + (f_a(t)+D_a(t)a) + v(t), \text{ where } v(t) \sim MVN(0,R) \]

\[ \text{vec}(R) = f_r(t)+D_r(t)r \]
\[ x(1) \sim MVN(f_p+D_p p, V0) \]

\[ \text{vec}(V0) = f_l+D_l l \]

In the marssMODEL(form=marss) object, f(t) + D(t)m, is the vec of a matrix M(t), so f_b(t)+D_b(t)b would be vec(B(t)). The estimated parameters are in the column vectors: b, u, q, z, a, r, p, and l. Each matrix M(t) is f(t) + D(t)m so is the sum of a fixed part f(t) and the linear combination, D(t), of the free (or estimated) part m.

The vec form of the MARSS model is specified by 3D matrices for each f and D for each parameter: B, U, Q, Z, A, R, x0, V0. The number of columns in the D matrix for a parameter determines the number of estimated values for that parameter. The first dimension for f (fixed) and D (free) must be:
The third dimension of \( f \) (fixed) and \( D \) (free) is either 1 (if not time-varying) or TT (if time-varying). The second dimension of \( f \) (fixed) is always 1, while the second dimension of \( D \) (free) depends on how many values are being estimated for a matrix. It can be 0 (if the matrix is fixed) or up to the size of the matrix (if all elements are being estimated).

**Value**

A vector of error messages or NULL is no errors.

**Author(s)**

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**See Also**

MARSS, MARSS.marxss, marssMODEL

---

**isleRoyal**

*Isle Royale Wolf and Moose Data*

**Description**

Example data set for estimation of species interaction strengths. These are data on the number of wolves and moose on Isle Royal, Michigan. The data are unlogged. The covariate data are the average Jan-Feb, average Apr-May and average July-Sept temperature (F) and precipitation (inches). Also included are 3-year running means of these covariates. The choice of covariates is based on those presented in the Isle Royale 2012 annual report.

**Usage**

data(isleRoyal)

**Format**

The data are supplied as a matrix with years in the first column.
Source


The source for the covariate data is the Western Regional Climate Center (http://www.wrcc.dri.edu) using their data for the NE Minnesota division. The website used was http://www.wrcc.dri.edu/cgi-bin/divplot1_form.pl?2103 and www.wrcc.dri.edu/spi/divplot1map.html.

Examples

str(isleRoyal)

Description

Data used in MARSS vignettes in the MARSS-package. Tracking data from ARGOS tags on eight individual loggerhead turtles, 1997-2006.

Usage

data(loggerhead)
data(loggerheadNoisy)

Format

Data frames "loggerhead" and "loggerheadNoisy" contain the following columns:

- **turtle**: Turtle name.
- **day**: Day of the month (character).
- **month**: Month number (character).
- **year**: Year (character).
- **lon**: Longitude of observation.
- **lat**: Latitude of observation.

Details

Data frame "loggerhead" contains the original latitude and longitude data. Data frame "loggerheadNoisy" has noise added to the lat and lon data to represent data corrupted by errors.
MARSS

**Source**

Gray’s Reef National Marine Sanctuary (Georgia) and WhaleNet: http://whale.wheelock.edu/whalenet-stuff/stop_cover_archive.html

**Examples**

```r
str(loggerhead)
str(loggerheadNoisy)
```

---

**MARSS Model Specification and Estimation**

**Description**

This is the main MARSS function for fitting multivariate autoregressive state-space (MARSS) models. Scroll down to the bottom to see some short examples. To open the user guide from the command line, type `RshowDoc("UserGuide",package="MARSS")`. To open a guide to show you how to get started quickly, type `RshowDoc("Quick_Start",package="MARSS")`. To open an overview page with package information and how to find all the R code for the user guide chapters, type `RshowDoc("index",package="MARSS")`. To get info on the auxiliary functions (like for bootstrapping and confidence intervals) go to `MARSS-package`. To see a discussion of how to get output from your model fits, go to `print.MARSS`. If MARSS() is throwing errors or warnings that you don’t understand, try the Troubleshooting section of the user guide or type `MARSSinfo()` at the command line.

The background section on this page is focused on fitting MARSS models in vectorized form. This form will almost certainly look unfamiliar. MARSS works by converting the users’ (more familiar model form) into the vectorized form which allows general linear constraints. You should go to the help page for the form of the model you are fitting to get background on that model form. Currently the MARSS package has two model forms: marxss and dfa.

**marxss** This is the default form. \( x(t) = B_t x(t-1) + U_t + C_t c_t + w_t; \) \( y(t) = Z_t x(t) + A_t + D_t d_t + v_t \). Any parameters can be set to zero. Most users will want this help page.

**dfa** This is a model form to allow easier specification of models for Dynamic Factor Analysis. \( x(t) = X_{t-1} + U_t + w_t; \) \( y(t) = Z_t x(t) + A_t + D_t d_t + v_t \). The \( Z \) parameters has a specific form and the \( Q \) is set at i.i.d with variance of 1.

The rest of this help page discusses the vectorized form of a MARSS model.

The MARSS package fits time-varying state-space models that can be transformed into the form (termed form=marxss):

\[
x(t) = (f(t(x(t-1)) (x) I_m(f_b(t)+D_b(t))beta) + (f_u(t)+D_u(t))upsilon) + w(t), \text{ where } w(t) \sim MVN(0,Q(t))
\]

\[
y(t) = (f(t(x(t)) (x) I_n(f_z(t)+D_z(t))zeta) + (f_a(t)+D_a(t))alpha) + v(t), \text{ where } v(t) \sim MVN(0,R(t))
\]

\( x(1) \sim MVN(x0, V0) \) or \( x(0) \sim MVN(x0, V0) \)
where beta, upsilon, zeta, and alpha are column vectors of estimated values, the \( f \) are column vectors of inputs, and the \( D \) are matrices of inputs. The \( f \) and \( D \) are potentially time-varying. \( (x) \) means kronecker product and \( I_p \) is a \( p \times p \) identity matrix. The function \texttt{MARSS()} \ is used to fit MARSS models using the argument \texttt{form} to specify the type of state-space model being fit.

Most commonly used multivariate autoregressive state-space models can be reformulated into the form above. The user is not required to specify their model in the marss form (which is unfamiliar and unwieldy). Instead \texttt{MARSS()} \ uses the \texttt{form} argument to specify a more familiar state-space form. The user specifies their model in that (more familiar) form. \texttt{MARSS()} \ calls a helper function \texttt{MARSS_form} to translate the user’s model into \texttt{form=}marss.

The default MARSS form is "marxss" which is the state-space model:

\[
x(t) = B(t) x(t-1) + u(t) + C(t)c(t) + G(t)w(t), \text{ where } w(t) \sim \text{MVN}(0,Q(t))
\]

\[
y(t) = Z(t) x(t) + a(t) + D(t)d(t) + H(t)v(t), \text{ where } v(t) \sim \text{MVN}(0,R(t))
\]

\[
x(1) \sim \text{MVN}(x0, V0) \text{ or } x(0) \sim \text{MVN}(x0, V0)
\]

See \texttt{MARSS.marxss} for arguments and defaults information.

If you are working with models with time-varying parameters, it is important to notice the time-index for the parameters in the process equation (the \( x \) equation). In some formulations (e.g. in the \texttt{KFAS}), the process equation is \( x(t)=B(t-1)x(t-1)+w(t-1) \) so \( B(t-1) \) goes with \( x(t) \) not \( B(t) \). Thus one needs to be careful to line up the time indices when passing in time-varying parameters to \texttt{MARSS()}.

**Usage**

```r
MARSS(y,
    model=NULL,
    inits=NULL,
    miss.value=as.numeric(NA),
    method = "kem",
    form = "marxss",
    fit=TRUE,
    silent = FALSE,
    control = NULL,
    fun.kf = "MARSSkfas",
    ...)
```

**Arguments**

The default settings for the optional arguments are set in \texttt{MARSSsettings.R} and are given below in the details section. For form specific defaults see the form help file (e.g. \texttt{MARSS.marxss} or \texttt{MARSS.dfa}).

\( A n \times T \) matrix of \( n \) time series over \( T \) time steps.

\texttt{inits} \quad A \text{ list with the same form as the list outputted by } \texttt{coef(fit)} \text{ that specifies initial values for the parameters. See also } \texttt{MARSS.marxss}.

\texttt{model} \quad \text{Model specification using parameter model text shortcuts or matrices. See Details and } \texttt{MARSS.marxss} \text{ for the default form. Or better yet open the Quick Start Guide } \texttt{RShowDoc("Quick_Start",package="MARSS").}

\texttt{miss.value} \quad \text{Deprecated. Denote missing values by NAs in your data.}
method: Estimation method. MARSS provides an EM algorithm (method="kem") (see MARSSkem) and the BFGS algorithm (method="BFGS") (see MARSSoptim).

form: The equation form used in the MARSS() call. The default is "marxss". See MARSS.marxss or MARSS.dfa.

fit: TRUE/FALSE Whether to fit the model to the data. If FALSE, a marssMLE object with only the model is returned.

silent: TRUE/FALSE Suppresses printing of full error messages, warnings, progress bars and convergence information. Setting silent=2 will produce more verbose error messages and progress information.

control: Estimation options for the maximization algorithm. The typically used control options for method="kem" are below but see marssMLE for the full list of control options. Note many of these are not allowed if method="BFGS"; see MARSSoptim for the allowed control options for this method.

- minit: The minimum number of iterations to do in the maximization routine (if needed by method). If method="kem", this is an easy way to up the iterations and see how your estimates are converging. (positive integer)
- maxit: Maximum number of iterations to be used in the maximization routine (if needed by method) (positive integer).
- min.iter.conv.test: Minimum iterations to run before testing convergence via the slope of the log parameter versus log iterations.
- conv.test.deltaT=9: Number of iterations to use for the testing convergence via the slope of the log parameter versus log iterations.
- conv.test.slope.tol: The slope of the log parameter versus log iteration to use as the cut-off for convergence. The default is 0.5 which is a bit high. For final analyses, this should be set lower. If you want to only use abstol as your convergence test, then to something very large, for example conv.test.slope.tol=1000. Type MARSSinfo(11) to see some comments of when you might want to do this.
- abstol: The logLik.(iter-1)-logLik.(iter) convergence tolerance for the maximization routine. To meet convergence both the abstol and slope tests must be passed.
- allow.degen: Whether to try setting Q or R elements to zero if they appear to be going to zero.
- trace: An integer specifying the level of information recorded and error-checking run during the algorithms. trace=0, specifies basic error-checking and brief error-messages; trace>0 will print full error messages. In addition if trace>0, the Kalman filter output will be added to the outputted marssMLE object. Additional information recorded depends on the method of maximization. For the EM algorithm, a record of each parameter estimate for each EM iteration will be added. See optim for trace output details for the BFGS method. trace=-1 will turn off most internal error-checking and most error messages. The internal error checks are time expensive so this can speed up MARSS. This is particularly useful for bootstrapping and simulation studies.
- silent: TRUE/FALSE (default), Suppresses all printing including progress bars, error messages and convergence information. 0, Turns on all printing
of progress bars, fitting information and error messages. 2. Prints a brief success/failure message.

- **safe TRUE/FALSE**(default), Setting safe=TRUE runs the Kalman smoother after each parameter update rather than running the smoother only once after updated all parameters. The latter is faster but is not a strictly correct EM algorithm. In most cases, safe=FALSE (default) will not change the fits. If this setting does cause problems, you will know because you will see an error regarding the log-likelihood dropping and it will direct you to set safe=TRUE.

**fun.kf**

What Kalman filter function to use. MARSS has two: MARSSkfas which is based on the Kalman filter in the KFAS package based on Koopman and Durbin and MARSSkfs which is a native R implementation of the Kalman filter and smoother in Shumway and Stoffer. The KFAS filter is much faster. MARSSkfas modifies the input and output in order to output the lag-one covariance smoother needed for the EM algorithm (per page 321 in Shumway and Stoffer (2000)).

Optional arguments passed to function specified by form.

**Details**

MARSS provides an interface to the base MARSS-package functions and allows specification and fitting of MARSS models. The available estimation methods are maximum-likelihood via an EM algorithm (**method**="kem") or via a quasi-Newton algorithm provided by function **optim**(**method**="BFGS"). The function **MARSS()** allows the user to specify models using the model argument. See **MARSS.marxss** for the format of the model argument for the default marxss form. See also the User Guide (reference and link below) or Quick Start Guide.

A call to **MARSS()** returns an object of class marssMLE. The MARSS package has print, coef, and residuals functions that will handle marssMLE objects. See **print.marssMLE**, **coef.marssMLE**, and **residuals.marssMLE**. The help page for **print.marssMLE** summarizes all the different output available for marssMLE objects and describes what the output is mathematically. Thus, this page is the first place to start and will direct you to the appropriate other method functions (like coef).

Many different types of multivariate time-series models can be converted to the MARSS form (see the User Guide). **MARSS()** allows the user to specify the form of their model using the argument **form**. The default form is a multivariate lag-1 (time-varying) state-space model which is denoted using **form**="marxss". Look at **MARSS.marxss**) to see how the model argument for the marxss form is specified. The model argument is a list, but the elements of that and how it is converted to a marssMODEL(form=marxss) object (needed for the internal algorithms) depends intimately on the equation form. Thus you will need to refer to that appropriate help page (**MARSS.formname**) for your equation form.

The likelihood surface for MARSS models can be multimodal or with strong ridges. It is recommended that for final analyses the ML estimates are checked by using a Monte Carlo initial conditions search; see the chapter on initial conditions searches in the User Guide. This requires more computation time, but reduces the chance of the algorithm terminating at a local maximum and not reaching the true MLEs. Also it is wise to check the EM results against the BFGS results (if possible) since if there are strong ridges in the likelihood. Such ridges seems to slow down the EM algorithm considerably and can cause the algorithm to report convergence far from the ML values. EM steps up the likelihood and the convergence test is based on the rate of change of the LL in each step; once on a strong ridge, the steps can slow dramatically. You can force the algorithm to keep
working by setting minit. BFGS seems less hindered by the ridges but can be prodigiously slow for some multivariate problems.

Value

An object of class marssMLE. The structure of this object is discussed below, but if you want to know how to get specific output (like residuals, coefficients, smoothed states, confidence intervals, etc), go here print.MARSS.

The outputted marssMLE object has the following components:

- model: MARSS model specification. It is a marssMODEL object in the form specified by the user in the MARSS() call. This is used by print functions so that the user sees the expected form.
- marss: The marssMODEL object in marss form. This form is needed for all the internal algorithms, thus is a required part of a marssMLE object.
- call: All the information passed in in the MARSS() call.
- start: List with specifying initial values that were used for each parameter matrix.
- control: A list of estimation options, as specified by arguments control.
- method: Estimation method.

If fit=TRUE, the following are also added to the marssMLE object. If fit=FALSE, an marssMLE object ready for fitting via the specified method is returned.

- par: A list of estimated parameter values Z, A, R, B, U, Q, x0, V0. See print.marssMLE or coef.marssMLE for information on outputting the model estimates. This will be in form "marss". Use print or coef to output the estimated parameters in the form in the MARSS() call (e.g. the default "marxss" form).
- states: The expected value of x conditioned on the data.
- states.se: The standard errors of the expected value of x.
- yt: The expected value of y conditioned on the data. Note this is just y for those y that are not missing.
- yt.se: The standard errors of the expected value of y. Note this is 0 for any non-missing y.
- numIter: Number of iterations required for convergence.
- convergence: Convergence status. 0 means converged successfully. Anything else is a warning or error. 2 means the MLEobj has an error; the MLEobj is returned so you can debug it. The other numbers are errors during fitting. The error code depends on the fitting method. See MARSSkem and MARSSoptim.
- logLik: Log-likelihood.
- AIC: Akaike’s Information Criterion.
- AICC: Sample size corrected AIC.

If control$trace is set to 1 or greater, the following are also added to the marssMLE object.

- kf: A list containing Kalman filter/smoother output from MARSSkf. This isn’t normally added to a marssMLE object since it is verbose, but can be computed using MARSSkf(marssMLE).
Ey

A list containing output from MARSSHatyt. This isn’t normally added to a marssMLE object since it is verbose, but can be computed using MARSSHatyt(marssMLE).

Author(s)

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References


See Also

marssMLE MARSSkemMARSSoptimMARSS-packageprint.MARSS MARSS.marxssMARSS.dfa augment.marssMLE tidy.marssMLE coef.marssMLE

Examples

dat <- t(harborSealWA)
dat <- dat[2:4,] # remove the year row
# fit a model with 1 hidden state and 3 observation time series
kemfit <- MARSS(dat, model=list(Z=matrix(1,3,1),
R="diagonal and equal"))
kemfit$model # This gives a description of the model
print(kemfit$model) # same as kemfit$model
summary(kemfit$model) # This shows the model structure

# add CIs to a marssMLE object
# default uses an estimated Hessian matrix
kem.with.hess.CIs <- MARSSparamCIs(kemfit)
kem.with.hess.CIs

# fit a model with 3 hidden states (default)
kemfit <- MARSS(dat, silent=TRUE) # suppress printing
kemfit

# fit a model with 3 correlated hidden states
# with one variance and one covariance
# maxit set low to speed up example, but more iters are needed for convergence
kemfit <- MARSS(dat, model=list(Q="equalvarcov"), control=list(maxit=50))
# use Q="unconstrained" to allow different variances and covariances
#fit a model with 3 independent hidden states
#where each observation time series is independent
#the hidden trajectories 2-3 share their U parameter
kemfit <- MARSS(dat, model=list(U=matrix(c("N","S","S"),3,1)))

#same model, but with fixed independent observation errors
#and the 3rd x processes are forced to have a U=0
#Notice how a list matrix is used to combine fixed and estimated elements
#all parameters can be specified in this way using list matrices
kemfit <- MARSS(dat, model=list(U=matrix(list("N","N",0,3,1), R=diag(0.01,3)))

#fit a model with 2 hidden states (north and south)
#where observation time series 1-2 are north and 3 is south
#Make the hidden state process independent with same process var
#Make the observation errors different but independent
#Make the growth parameters (U) the same
#Create a Z matrix as a design matrix that assigns the "N" state to the first 2 rows of dat
#and the "S" state to the 3rd row of data
Z <- matrix(c(1,1,0,0,0,1),3,2)
#You can use factor is a shortcut making the above design matrix for Z
#Z <- factor(c("N","N","S"))
#name the state vectors
colnames(Z) <- c("N","S")
kemfit <- MARSS(dat, model=list(Z=Z,
Q="diagonal and equal",R="diagonal and unequal",U="equal"))

#Print the model followed by the marssMLE object
kemfit$model
kemfit

## Not run:
## simulate some new data from our fitted model
sim.data=MARSSsimulate(kemfit, nsim=10, tSteps=10)

#Compute bootstrap AIC for the model; this takes a long, long time
kemfit.with.AIC <- MARSSaic(kemfit, output = "AICbp")
kemfit.with.AIC

## End(Not run)

## Not run:
## Many more short examples can be found in the
## Quick Examples chapter in the User Guide
RShowDoc("UserGuide",package="MARSS")

## You can find the R scripts from the chapters by
## going to the index page
RShowDoc("index",package="MARSS")

## End(Not run)
Convert Model Objects between Forms

Description

These are utility functions for model objects in the package MARSS-package. Users would not normally work directly with these functions.

Usage

marss_to_marxss(x, C.and.D.are.zero=FALSE)
m Marxss_to_marss(x, only.par = FALSE)

Arguments

x An object of class marxMLE.
C.and.D.are.zero If the C and D matrices are all 0, then a marss model can be converted to marxss without further information besides the marss model.
only.par If only.par=TRUE then only the par element is changed and marss is used for the marss object.

Details

As the name of the functions imply, these convert marxMLE objects of different forms into other forms. form=marss is the base form needed for the internal algorithms, thus other (more user friendly forms) must have a form_to_marss function to convert to the base form. The printing functions are customized to show output in the user-friendly form, thus a marss_to_form function is needed for print and coef methods for marxMLE objects.

Value

A marxMLE object of the appropriate form.

Author(s)

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See Also

marxMLE
Multivariate Dynamic Factor Analysis

Description

The Dynamic Factor Analysis model in MARSS is

\[ x(t+1) = x(t) + w(t), \text{ where } w(t) \sim \text{MVN}(0,I) \]
\[ y(t) = Z(t) x(t) + D(t)d(t) + v(t), \text{ where } v(t) \sim \text{MVN}(0,R(t)) \]
\[ x(1) \sim \text{MVN}(0, 5*I) \]

Passing in form="dfa" to MARSS() invokes a helper function to create that model and creates the Z matrix for the user. Q is by definition identity, x0 is zero and V0 is diagonal with large variance (5). U is zero, A is zero, and covariates only enter the Y equation. Because U and A are 0, the data should have mean 0 (de-meaned) otherwise one is likely to be creating a structurally inadequate model (i.e. the model implies that the data have mean = 0, yet data do not have mean = 0).

Arguments

Some arguments are common to all forms: "data", "inits", "control", "method", "form", "fit", "silent", "fun.kf". See MARSS for information on these arguments. In addition to these, form="dfa" has some special arguments that can be passed in:

- `demean` Logical. Default is TRUE, which means the data will be demeaned.
- `zScore` Logical. Default is TRUE, which means the data will be z-scored (demeaned and variance standardized to 1).

The `model` argument of the MARSS call is constrained in terms of what parameters can be changed and how they can be changed. See details below. An additional element, m, can be passed into the `model` argument that specifies the number of hidden state variables. It is not necessarily for the user to specify Z as the helper function will create a Z appropriate for a DFA model.

Details

The `model` argument is a list. The following details what list elements can be passed in:

- `B"Identity"`. Can be "identity", "diagonal and equal", or "diagonal and unequal".
- `U"Zero"`. Cannot be changed or passed in via model argument.
- `Q"Identity"`. Can be "identity", "diagonal and equal", or "diagonal and unequal".
- `Z` Can be passed in as a (list) matrix if the user does not want a default DFA Z matrix. There are many equivalent ways to construct a DFA Z matrix. The default is Zuur et al.'s form (see User Guide).
- `A"Default="zero"`. Can be "unequal", "zero" or a matrix.
- `R"Default="diagonal and equal"`. Can be set to "identity", "zero", "unconstrained", "diagonal and unequal", "diagonal and equal", "equalvarcov", or a (list) matrix to specify general forms.
• $x_0$ Default="zero". Can be "unconstrained", "unequal", "zero", or a (list) matrix.
• $\phi_0$ Default=diagonal matrix with 5 on the diagonal. Can be "identity", "zero", or a matrix.
• $t_{initx}$ Default=0. Can be 0 or 1. Tells MARSS whether $x_0$ is at $t=0$ or $t=1$.
• $m$ Default=1. Can be 1 to $n$ (the number of y time-series). Must be integer.

See the User Guide chapter on Dynamic Factor Analysis for examples of using form="dfa".

Value

A object of class marssMLE. See print.marssMLE for a discussion of the various output available for marssMLE objects (coefficients, residuals, Kalman filter and smoother output, imputed values for missing data, etc.). See MARSSsimulate for simulating from marssMLE objects, MARSSboot for bootstrapping, MARSSaic for calculation of various AIC related model selection metrics, and MARSSparamCIs for calculation of confidence intervals and bias.

Usage

```
MARSS(y, inits=NULL, model=NULL, miss.value=as.numeric(NA), method = "kem", form = "dfa",
```

Author(s)

Eli Holmes, NOAA, Seattle, USA.

References


See Also

MARSS MARSS.marxss

Examples

```R
## Not run:
#See the Dynamic Factor Analysis chapter in the User Guide
RShowDoc("UserGuide",package="MARSS")

## End(Not run)
```
Description

The form of MARSS models for users is marxss, the MARSS models with inputs. See `MARSS.marxss`. In the internal algorithms, the marss form is used and the D and d are incorporated into the A matrix and C and c are incorporated into the U matrix.

This is a MARSS(1) model of the marss form:

\[
\begin{align*}
    x(t) &= B(t) x(t-1) + u(t) + G(t)w(t), \text{ where } w(t) \sim \text{MVN}(0,Q(t)) \\
    y(t) &= Z(t) x(t) + a(t) + H(t)v(t), \text{ where } v(t) \sim \text{MVN}(0,R(t)) \\
    x(0) &\sim \text{MVN}(x0, V0)
\end{align*}
\]

Note, marss is a model form. A model form is defined by a collection of form functions discussed in `marssMODEL`. These functions are not exported to the user, but are called by MARSS() using the argument form. These internal functions convert the users model list into the vec form of a MARSS model and do extensive error-checking.

Details

See the help page for the `MARSS.marxss` form for details.

Value

A object of class `marssMLE`.

Usage

`MARSS(y, inits=NULL, model=NULL, miss.value=as.numeric(NA), method="kem", form="marxss")`

Author(s)

Eli Holmes, NOAA, Seattle, USA.

See Also

`marssMODEL` `MARSS.marxss`

Examples

```r
## Not run:
#See the MARSS man page for examples
?MARSS

#and the Quick Examples chapter in the User Guide
RShowDoc("UserGuide",package="MARSS")

## End(Not run)
```
Multivariate AR-1 State-space Model with Inputs

Description

The argument form="marxss" in a MARSS() function call specifies a MAR-1 model with exogenous variables model. This is a MARSS(1) model of the form:

\[
x(t) = B(t) x(t-1) + u(t) + C(t)c(t) + G(t)w(t), \text{ where } w(t) \sim \text{MVN}(0,Q(t)) \\
y(t) = Z(t) x(t) + a(t) + D(t)d(t) + H(t)v(t), \text{ where } v(t) \sim \text{MVN}(0,R(t)) \\
x(0) \sim \text{MVN}(x0, V0)
\]

Note, marxss is a model form. A model form is defined by a collection of form functions discussed in marssMODEL. These functions are not exported to the user, but are called by MARSS() using the argument form.

Details

The allowed arguments when form="marxss" are 1) the arguments common to all forms: "data", "inits", "control", "method", "form", "fit", "silent", "fun.kf" (see marss for information on these arguments) and 2) the argument "model" which is a list describing the MARXSS model (the model list is described below). See the Quick Start guide (RShowDoc("Quick_Start",package="MARSS")) or the User Guide (RShowDoc("UserGuide",package="MARSS")) for examples.

The argument model must be a list. The elements in the list specify the structure for the B, u, C, c, Q, Z, a, D, d, R, x0, and V0 in the MARXSS model (above). The list elements can have the following values:

- **Z** Default="identity". A text string, "identity","unconstrained", "diagonal and unequal", "diagonal and equal", "equalvarcov", or "onestate", or a length n vector of factors specifying which of the m hidden state time series correspond to which of the n observation time series. May be specified as a n x m list matrix for general specification of both fixed and shared elements within the matrix. May also be specified as a numeric n x m matrix to use a custom fixed Z. "onestate" gives a n x 1 matrix of 1s. "identity","unconstrained", "diagonal and unequal", "diagonal and equal", and "equalvarcov" all specify n x n matrices.

- **B** Default="identity". A text string, "identity", "unconstrained", "diagonal and unequal", "diagonal and equal", "equalvarcov", "zero". Can also be specified as a list matrix for general specification of both fixed and shared elements within the matrix. May also be specified as a numeric m x m matrix to use a custom fixed B, but in this case all the eigenvalues of B must fall in the unit circle.

- **U, x0** Default="unconstrained". A text string, "unconstrained", "equal", "unequal" or "zero". May be specified as a m x 1 list matrix for general specification of both fixed and shared elements within the matrix. May also be specified as a numeric m x 1 matrix to use a custom fixed U or x0. Notice that U is capitalized.

- **A** Default="scaling". A text string, "scaling","unconstrained", "equal", "unequal" or "zero". May be specified as a n x 1 list matrix for general specification of both fixed and shared elements within the matrix. May also be specified as a numeric n x 1 matrix to use a custom
fixed A. Care must be taken when specifying A so that the model is not under-constrained and unsolvable model. The default "scaling" only applies to Z matrices that are design matrices (only 1s and 0s and all rows sum to 1). When a column in Z has multiple 1s, the first row with a 1 is assigned A=0 and the rows with 1s for that column have an estimated A. This is used to treat A as an intercept where one A for each X (hidden state) is fixed at 0 and any other Ys associated with that X have an estimated A value. This ensures a solvable model (when Z is a design matrix). A is capitalized.

- **Q** Default="diagonal and unequal". A text string, "identity", "unconstrained", "diagonal and unequal", "diagonal and equal", "equalvarcov", "zero". May be specified as a list matrix for general specification of both fixed and shared elements within the matrix. May also be specified as a numeric g x g matrix to use a custom fixed matrix. Default value of g is m, so Q is a m x m matrix. g is the num of columns in G (below).

- **R** Default="diagonal and equal". A text string, "identity", "unconstrained", "diagonal and unequal", "diagonal and equal", "equalvarcov", "zero". May be specified as a list matrix for general specification of both fixed and shared elements within the matrix. May also be specified as a numeric h x h matrix to use a custom fixed matrix. Default value of h is n, so R is a n x n matrix. h is the num of columns in H (below).

- **v0** Default="zero". A text string, "identity", "unconstrained", "diagonal and unequal", "diagonal and equal", "equalvarcov", "zero". May be specified as a list matrix for general specification of both fixed and shared elements within the matrix. May also be specified as a numeric h x h matrix to use a custom fixed matrix.

- **d and c** Default="zero". A text string, "identity", "unconstrained", "diagonal and unequal", "diagonal and equal", "equalvarcov", "zero". Can be specified as a list matrix for general specification of both fixed and shared elements within the matrix. May also be specified as a numeric m x m matrix to use a custom fixed matrix.

- **d and c** Default="zero". Numeric matrix. No missing values allowed. Must have 1 column or the same number of columns as the data, y. The numbers of rows in d must be the same number of columns in D; similarly for c and C. c and d are lower case.

- **g and h** Default="identity". A text string, "identity", Can be specified as a numeric matrix or array for time-varying cases. Must have m rows and g columns (G) or n rows and h columns (H). g is the dim of Q and h is the dim of R.

- **tinitx** Default=0. Whether the initial state is specified at t=0 (default) or t=1.

All parameters except x0 and V0 may be time-varying. If time-varying, then text shortcuts cannot be used. Enter as an array with the 3rd dimension being time. Time dimension must be 1 or equal to the number of time-steps in the data. See Quick Start guide (RShowDoc("Quick_Start",package="MARSS")) or the User Guide (RShowDoc("UserGuide",package="MARSS")) for examples. Valid model structures for method="BFGS" are the same as for method="kem". See MARSSoptim for the allowed options for this method.

The default estimation method, method="kem", is the EM algorithm described in the user guide. The default settings for the control and inits arguments are set via MARSS:::alldefaults$kem in MARSSsettings.R. The defaults for the model argument are set in MARSS_marxss.R. For this method, they are:

- **inits** = list(B=1, U=0, Q=0.05, Z=1, A=0, R=0.05, x0=-99, V0=0.05, G=0, H=0, L=0, C=0, D=0, c=0, d=0)
• model = list(Z="identity", A="scaling", R="diagonal and equal", B="identity", U="unconstrained", Q="diagonal and unequal", x0="unconstrained", V0="zero", C="zero", D="zero", c=matrix(0,0,1), d=matrix(0,0,1), tinitx=0, diffuse=FALSE)
• control=list(minit=15, maxit=500, abstol=0.001, trace=0, sparse=FALSE, safe=FALSE, allow.degen=TRUE, min.degen.iter=50, degen_lim=1.0e-04, min.iter.conv.test=15, conv.test.deltaT=9, conv.test.slope.tol= 0.5, demean.states=FALSE) You can read about these in MARSS. If you want to speed up your fits, you can turn off most of the model checking using trace=-1.
• fun.kf = "MARSSkfas"; This sets the Kalman filter function to use. MARSSkfas() is generally more stable as it uses Durban & Koopman’s algorithm. But it may dramatically slow down when the dataset is large (more than 10 rows of data). Try the classic Kalman filter algorithm to see if it runs faster by setting fun.kf="MARSSkffs". You can read about the two algorithms in MARSSk.

For method="BFGS", type MARSS::alldefaults$BFGS to see the defaults.

Value
A object of class marssMLE. See print.marssMLE for a discussion of the various output available for marssMLE objects (coefficients, residuals, Kalman filter and smoother output, imputed values for missing data, etc.). See MARSSsimulate for simulating from marssMLE objects. MARSSboot for bootstrapping, MARSSaic for calculation of various AIC related model selection metrics, and MARSSparamCIs for calculation of confidence intervals and bias. See plot.marssMLE for some default plots of a model fit.

Usage
MARSS(y, inits=NULL, model=NULL, miss.value=as.numeric(NA), method = "kem", form = "marxss"

Author(s)
Eli Holmes, NOAA, Seattle, USA.

See Also
marssMODEL MARSS.dfa

Examples
## Not run:
#See the MARSS man page for examples
?MARSS

#and the Quick Examples chapter in the User Guide
RShowDoc("UserGuide",package="MARSS")

## End(Not run)
Description

Calculates AIC, AICc, a parametric bootstrap AIC (AICbp) and a non-parametric bootstrap AIC (AICbb).

Usage

MARSSaic(MLEobj, output = c("AIC", "AICc"),
          Options = list(nboot = 1000, return.logL.star = FALSE,
                         silent = FALSE))

Arguments

MLEobj An object of class marssMLE. This object must have a $par element containing MLE parameter estimates from e.g., MARSSkem().
output A vector containing one or more of the following: "AIC", "AICc", "AICbp", "AICbb", "AICi", "boot.params". See Details.
Options A list containing:
          • nboot Number of bootstraps (positive integer)
          • return.logL.star Return the log-likelihoods for each bootstrap? (T/F)
          • silent Suppress printing of the progress bar during AIC bootstraps? (T/F)

Details

When sample size is small, Akaike's Information Criterion (AIC) under-penalizes more complex models. The most commonly used small sample size corrector is AICc, which uses a penalty term of Kn/(n-K-1), where K is the number of estimated parameters. However, for time series models, AICc still under-penalizes complex models; this is especially true for MARSS models.

Two small-sample estimators specific for MARSS models have been developed. Cavanaugh and Shumway (1997) developed a variant of bootstrapped AIC using Stoffer and Wall’s (1991) bootstrap algorithm ("AICbb"). Holmes and Ward (2010) developed a variant on AICb ("AICbp") using a parametric bootstrap. The parametric bootstrap permits AICb calculation when there are missing values in the data, which Cavanaugh and Shumway’s algorithm does not allow. More recently, Bengtsson and Cavanaugh (2006) developed another small-sample AIC estimator, AICi, based on fitting candidate models to multivariate white noise.

When the output argument passed in includes both "AICbp" and "boot.params", the bootstrapped parameters from "AICbp" will be added to MLEobj.

Value

Returns the marssMLE object that was passed in with additional AIC components added on top as specified in the 'output' argument.
Author(s)

Eli Holmes and Eric Ward, NOAA, Seattle, USA.
eli(dot)holmes(at)noaa(dot)gov, eric(dot)ward(at)noaa(dot)gov

References


See Also

MARSSboot

Examples

dat <- t(harborSealWA)
dat <- dat[2:3,]
kem <- MARSS(dat, model=list(Z=matrix(1,2,1),
                    R="diagonal and equal"))
kemAIC = MARSSaic(kem, output=c("AIC","AICc"))

MARSSapplynames

Names for marssMLE Object Components

Description

Puts names on the par, start, par.se, init components of marssMLE objects. This is a utility function in the MARSS-package and is not exported.

Usage

MARSSapplynames(MLEobj)

Arguments

MLEobj An object of class marssMLE.

Details

The X.names and Y.names are attributes of marssMODEL objects (which would be in $marss and $model in the marssMLE object). These names are applied to the par elements in the marssMLE object.
**Value**

The object passed in, with row and column names on matrices as specified.

**Author(s)**

Eli Holmes, NOAA, Seattle, USA.
eli(dot)holmes(at)noaa(dot)gov

**See Also**

marssMLE marssMODEL

---

**MARSSboot**

**Bootstrap MARSS Parameter Estimates**

**Description**

Creates bootstrap parameter estimates and simulated (or bootstrapped) data (if appropriate). This is a base function in the MARSS-package.

**Usage**

MARSSboot(MLEobj, nboot = 1000,
  output = "parameters", sim = "parametric",
  param.gen = "MLE", control = NULL, silent = FALSE)

**Arguments**

- **MLEobj**
  An object of class marssMLE. Must have a $par$ element containing MLE parameter estimates.

- **nboot**
  Number of bootstraps to perform.

- **output**
  Output to be returned: "data", "parameters" or "all".

- **sim**
  Type of bootstrap: "parametric" or "innovations". See Details.

- **param.gen**
  Parameter generation method: "hessian" or "MLE".

- **control**
  The options in MLEobj$control are used by default. If supplied here, must contain all of the following:
  - **max.iter** Maximum number of EM iterations.
  - **tol** Optional tolerance for log-likelihood change. If log-likelihood decreases less than this amount relative to the previous iteration, the EM algorithm exits.
  - **allow.degen** Whether to try setting Q or R elements to zero if they appear to be going to zero.

- **silent**
  Suppresses printing of progress bar.
Details

Approximate confidence intervals (CIs) on the model parameters can be calculated by numerically estimating the Hessian matrix (the matrix of partial 2nd derivatives of the parameter estimates). The Hessian CIs (param.gen="hessian") are based on the asymptotic normality of ML estimates under a large-sample approximation. CIs that are not based on asymptotic theory can be calculated using parametric and non-parametric bootstrapping (param.gen="MLE"). In this case, parameter estimates are generated by the ML estimates from each bootstrapped data set. The MLE method (kem or BFGS) is determined by MLEobj$method.

Stoffer and Wall (1991) present an algorithm for generating CIs via a non-parametric bootstrap for state-space models (sim = "innovations"). The basic idea is that the Kalman filter can be used to generate estimates of the residuals of the model fit. These residuals are then standardized and resampled and used to generate bootstrapped data using the MARSS model and its maximum-likelihood parameter estimates. One of the limitations of the Stoffer and Wall algorithm is that it cannot be used when there are missing data, unless all data at time t are missing. An alternative approach is a parametric bootstrap (sim = "parametric"), in which the ML parameter estimates are used to produce bootstrapped data directly from the state-space model.

Value

A list with the following components:

- boot.params: Matrix (number of params x nboot) of parameter estimates from the bootstrap.
- boot.data: Array (n x t x nboot) of simulated (or bootstrapped) data (if requested and appropriate).
- marss: The marssMODEL object (form="marss") that was passed in via MLEobj$marss.
- nboot: Number of bootstraps performed.
- output: Type of output returned.
- sim: Type of bootstrap.
- param.gen: Parameter generation method: "hessian" or "KalmanEM".

Author(s)

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References


See Also

marssMLE marssMODEL MARSSaic

Examples

# nboot is set low in these examples in order to run quickly
# normally nboot would be >1000 at least
dat <- t(kestrel)
dat <- dat[2:3,]
# maxit set low to speed up the example
kem <- MARSS(dat, model=list(U="equal",Q=diag(.01,2)),
                   control=list(maxit=50))

# bootstrap parameters from a Hessian matrix
hess.list <- MARSSboot(kem, param.gen="hessian", nboot=4)

# from resampling the innovations (no missing values allowed)
boot.innov.list <- MARSSboot(kem, output="all", sim="innovations", nboot=4)

# bootstrapped parameter estimates
hess.list$boot.params

MARSSharveyobsFI MARSS Hessian Matrix via the Harvey (1989) Recursion

Description

Calculates the observed Fisher Information analytically via the recursion by Harvey (1989). This is the same as the Hessian of the log-likelihood function at the MLEs. This is a utility function in the MARSS-package and is not exported. Use MARShessian to access.

Usage

MARSSharveyobsFI( MLEobj )

Arguments

MLEobj An object of class marssMLE. This object must have a $par element containing MLE parameter estimates from e.g. MARSSkem.

Value

The observed Fisher Information matrix computed via equation 3.4.69 in Harvey (1989). The differentials in the equation are computed in the recursion in equations 3.4.73a to 3.4.74b. Harvey (1989) discusses missing observations in section 3.4.7. However, the MARSSharveyobsFI() function implements the approach of Shumway and Stoffer (2006) in section 6.4 for the missing values. See Holmes (2012) for a full discussion of the missing values modifications.
Author(s)

Eli Holmes, NOAA, Seattle, USA.
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References


See Also

MARSShessian MARSSparamCIs

Examples

dat <- t(harborSeal)
dat <- dat[c(2,11),]
MLEobj <- MARSS(dat)
MARSS::MARSSharveyobsFI(MLEobj)

Description

Computes the expected value of random variables involving Y for the EM algorithm. Users can also use print(MLEobj, what="Ey") to access this output. See print.marssMLE.

Usage

MARSShatyt( MLEobj )

Arguments

MLEobj A marssMLE object with the par element of estimated parameters, model element with the model description and data.
Details

For state space models, `MARSShessian()` computes the expectations involving Y. If Y is completely observed, this entails simply replacing Y with the observed y. When Y is only partially observed, the expectation involves the conditional expectation of a multivariate normal.

Value

A list with the following components (n is the number of state processes). Following the notation in Holmes (2012), y(1) is the observed data (for t=1:TT) while y(2) is the unobserved data. y(1,1:t) is the observed data from time 1 to t.

- `ytt` Estimates E[Y(t) | Y(1,1:TT)=y(1,1:TT)] (n x T matrix).
- `yttQ` Estimates E[Y(t) | Y(1,1:t-1)=y(1,1:t-1)] (n x T matrix).
- `yttP` Estimates E[Y(t) | Y(t) | Y(1)=y(1)] (n x n x T array).
- `yxttt` Estimates E[Y(t) | X(t) | Y(1)=y(1)] (n x m x T array).
- `yxtttQ` Estimates E[Y(t) | X(t-1) | Y(1)=y(1)] (n x m x T array).
- `yxtttP` Estimates E[Y(t) | X(t+1) | Y(1)=y(1)] (n x m x T array).
- `errors` Any error messages due to ill-conditioned matrices.
- `ok` (T/F) Whether errors were generated.

Author(s)

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References


See Also

`MARSS marssMODEL MARSSkem`

**MARSShessian**

*MARSS Parameter Variance-Covariance Matrix from the Hessian Matrix*

Description

Calculates an approximate parameter variance-covariance matrix for the parameters using an inverse of the Hessian of the log-likelihood function at the MLEs. It appends `$Hessian`, `$parMean`, `$parSigma` to the marssMLE object.
MARSShessian

Usage

MARSShessian( MLEobj, method="Harvey1989" )
MARSSFisherI( MLEobj, method="Harvey1989" )

Arguments

MLEobj An object of class marssMLE. This object must have a $par element containing MLE parameter estimates from e.g. MARSSkem.

method The method to use for computing the Hessian. Options are 'Harvey1989' to use the Harvey (1989) recursion, which is an analytical solution, 'fdHess' or 'optim' which are two numerical methods. Although 'optim' can be passed to the function, 'fdHess' is used for all numerical estimates used in the package.

Details

Method 'fdHess' uses fdHess from package nlme to numerically estimate the Hessian matrix (the matrix of partial 2nd derivatives of the parameter estimates). Method 'optim' uses optim with hessian=TRUE and list(maxit=0) to ensure that the Hessian is computed at the values in the par element of the MLE object. Method 'Harvey1989' (the default) uses the recursion in Harvey (1989) to compute the observed Fisher Information of a MARSS model analytically.

Hessian CIs are based on the asymptotic normality of ML estimates under a large-sample approximation.

MARSSFisherI() is a (non-exported) utility function called by MARSShessian(). It calls either MARSShessian.numerical or MARSSharveyobsFI to return the Hessian based on the value of method.

Value

MARSShessian() attaches Hessian, parMean and parSigma to the MLE object.

Author(s)

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See Also

MARSSharveyobsFI MARSShessian.numerical MARSSparamCIs marssMLE

Examples

dat = t(harborSeal)
dat = dat[c(2,11),]
MLEobj = MARSS(dat)
MLEobj.hessian = MARSShessian(MLEobj)

#show the approx Hessian
MLEobj.hessian$Hessian
#generate a parameter sample using the Hessian
#this uses the rmvnorm function in the mvtnorm package
hess.params = rmvnorm(1, mean=MLEobj.hessian$parMean,
                      sigma=MLEobj.hessian$parSigma)

MARSShessian.numerical

**MARSS Hessian Matrix via Numerical Approximation**

**Description**

Calculates the Hessian of the log-likelihood function at the MLEs using either the `fdHess` function in the `nlme` package or the `optim` function. This is a utility function in the MARSS-package and is not exported. Use `MARSShessian` to access.

**Usage**

```r
MARSShessian.numerical(MLEobj, fun="fdHess")
```

**Arguments**

- **MLEobj**: An object of class `marssMLE`. This object must have a `$par` element containing MLE parameter estimates from e.g. `MARSSkem`.
- **fun**: The function to use for computing the Hessian. Options are `"fdHess"` or `"optim"`.

**Details**

Method `"fdHess"` uses `fdHess` from package `nlme` to numerically estimate the Hessian matrix (the matrix of partial 2nd derivatives of the parameter estimates). Method `"optim"` uses `optim` with `hessian=TRUE` and `list(maxit=10)` to ensure that the Hessian is computed at the values in the `par` element of the MLE object.

**Value**

The numerically estimated Hessian of the log-likelihood function at the MLEs.

**Author(s)**

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**See Also**

`MARSSharveyobsFI` `MARSShessian` `MARSSparamCIs`
**MARSSinfo**  
*Information for MARSS Error Messages and Warnings*

**Description**

Prints out more information for MARSS error messages and warnings.

**Usage**

```
MARSSinfo(number)
```

**Arguments**

- `number`: An error or warning message number.

**Value**

A print out of information.

**Author(s)**

Eli Holmes, NOAA, Seattle, USA.  
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---

**MARSSinits**  
*Initial Values for MLE*

**Description**

Sets up generic starting values for parameters for maximum-likelihood estimation algorithms that use an iterative maximization routine needing starting values. Examples of such algorithms are the EM algorithm in MARSSkem and Newton methods in MARSSoptim. This is a utility function in the MARSS-package. It is not exported to the user. Users looking for information on specifying initial conditions should look at MARSS and look at the help file for their model form.

The function assumes that the user passed in the inits list using the parameter names in whatever form was specified in the MARSS() call. The default is form="marxss". The MARSSinits() function calls MARSSinits_foo, where foo is the form specified in the MARSS() call. MARSSinits_foo translates the inits list in form foo into form marss.

**Examples**

```r
dat <- t(harborSeal)
dat <- dat[2:11,]
MLEobj <- MARSS(dat)
MARSS::MARSShessian.numerical(MLEobj)
```
**Usage**

`MARSSinits(MLEobj, inits=list(B=1, U=0, Q=0.05, Z=1, A=0, R=0.05, x0=-99, V0=5, G=0, H=0, L=0))`

**Arguments**

- **MLEobj**
  An object of class `marssMLE`.

- **inits**
  A list of column vectors (matrices with one column) of the estimated values in each parameter matrix.

**Details**

Creates an `inits` parameter list for use by iterative maximization algorithms.

Default values for `inits` are supplied in `MARSSsettings.R`. The user can alter these and supply any of the following (m is the dim of X and n is the dim of Y in the MARSS model):

- **elem=A,U**
  A numeric vector or matrix which will be constructed into `inits$elem` by the command `array(inits$elem, dim=c(n or m, 1))`. If elem is fixed in the model, any `inits$elem` values will be overridden and replaced with the fixed value. Default is `array(0, dim=c(n or m, 1))`.

- **elem=Q,R,B**
  A numeric vector or matrix. If length equals the length `MODELobj$fixed$elem` then `inits$elem` will be constructed by `array(inits$elem, dim=dim(MODELobj$fixed$elem))`. If length is 1 or equals dim of Q or dim of R then `inits$elem` will be constructed into a diagonal matrix by the command `diag(inits$elem)`. If elem is fixed in the model, any `inits$elem` values will be overridden and replaced with the fixed value. Default is `diag(0.05, dim of Q or R)` for Q and R. Default is `diag(1, m)` for B.

- **x0**
  If `inits$x0=-99`, then starting values for x0 are estimated by a linear regression through the count data assuming A=0. This will be a poor start if `inits$A` is not 0. If `inits$x0` is a numeric vector or matrix, `inits$x0` will be constructed by the command `array(inits$x0, dim=c(m, 1))`. If x0 is fixed in the model, any `inits$x0` values will be overridden and replaced with the fixed value. Default is `inits$x0=-99`.

- **Z**
  If Z is fixed in the model, `inits$Z` set to the fixed value. If Z is not fixed, then the user must supply `inits$Z`. There is no default.

- **elem=V0**
  V0 is never estimated, so this is never used.

**Value**

A list with specifying initial values for the estimated values for each parameter matrix in a MARSS model in marss form. So this will be a list with elements B, U, Q, Z, A, R, x0, V0, G, H, L.

**Author(s)**

Eli Holmes, NOAA, Seattle, USA.

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**See Also**

`marssMODEL` `MARSSkem` `MARSSoptim`
MARSSinnovationsboot Bootstrapped Data using Stoffer and Wall’s Algorithm

Description

Creates bootstrap data via sampling from the standardized innovations matrix. This is a base function in the MARSS-package. Users should access this with MARSSboot.

Usage

MARSSinnovationsboot(MLEobj, nboot = 1000, minIndx = 3)

Arguments

MLEobj An object of class marssMLE. This object must have a $par element containing MLE parameter estimates from e.g. MARSSkem or MARSS. This algorithm may not be used if there are missing datapoints in the data.
nboot Number of bootstraps to perform.
minIndx Number of innovations to skip. Stoffer & Wall suggest not sampling from innovations 1-3.

Details

Stoffer and Wall (1991) present an algorithm for generating CIs via a non-parametric bootstrap for state-space models. The basic idea is that the Kalman filter can be used to generate estimates of the residuals of the model fit. These residuals are then standardized and resampled and used to generate bootstrapped data using the MARSS model and its maximum-likelihood parameter estimates. One of the limitations of the Stoffer and Wall algorithm is that it cannot be used when there are missing data, unless all data at time t are missing.

Value

A list containing the following components:

- boot.states Array (dim is m x tSteps x nboot) of simulated state processes.
- boot.data Array (dim is n x tSteps x nboot) of simulated data.
- marss marssMODEL object element of the marssMLE object (marssMLE$marss).
- nboot Number of bootstraps performed.

m is the number state processes (x in the MARSS model) and n is the number of observation time series (y in the MARSS model).

Author(s)

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MARSSkem

References

See Also
stdInnov MARSSparamCIs MARSSboot

Examples

dat <- t(kestrel)
dat <- dat[2:3,]
MLEobj <- MARSS(dat, model=list(U="equal",Q=diag(.01,2))
boot.obj <- MARSSinnovationsboot(MLEobj)

MARSSkem  Maximum Likelihood Estimation for Multivariate Autoregressive State-Space Models

Description
MARSSkem() performs maximum-likelihood estimation, using an EM algorithm for constrained and unconstrained MARSS models. Users would not call this function directly normally. The function MARSS calls MARSSkem. However users might want to used MARSSkem directly if they need to avoid some of the error-checking overhead associated with the MARSS function.

Usage
MARSSkem(MLEobj)

Arguments
MLEobj An object of class marssMLE.

Details
Objects of class marssMLE may be built from scatch but are easier to construct using MARSS with MARSS(..., fit=FALSE).
Options for MARSSkem() may be set using MLEobj$control. The commonly used elements of control are follows (see marssMLE:

minit Minimum number of EM iterations. You can use this to force the algorithm to do a certain number of iterations. This is helpful if your soln is not converging.
maxit Maximum number of EM iterations.
min.iter.conv.test The minimum number of iterations before the log-log convergence test will be computed. If maxit is set less than this, then convergence will not be computed (and the algorithm will just run for maxit iterations).
Whether to set the prior at t=0 ("x00") or at t=1 ("x10"). The default is "x00".

The number of iterations to use in the log-log convergence test. This defaults to 9.

Tolerance for log-likelihood change for the delta logLik convergence test. If log-likelihood changes less than this amount relative to the previous iteration, the EM algorithm exits. This is normally (default) set to NULL and the log-log convergence test is used instead.

Whether to try setting Q or R elements to zero if they appear to be going to zero.

A positive integer. If not 0, a record will be created of each variable over all EM iterations and detailed warning messages (if appropriate) will be printed.

If TRUE, `MARSSkem` will rerun `MARSSkf` after each individual parameter update rather than only after all parameters are updated. The latter is slower and unnecessary for many models, but in some cases, the safer and slower algorithm is needed because the ML parameter matrices have high condition numbers.

Suppresses printing of progress bars, error messages, warnings and convergence information.

The `marssMLE` object which was passed in, with additional components:

- **method**: String "kem".
- **kf**: Kalman filter output.
- **iter.record**: If `mleobj$control$trace = TRUE`, a list with `par` = a record of each estimated parameter over all EM iterations and `logLik` = a record of the log likelihood at each iteration.
- **numIter**: Number of iterations needed for convergence.
- **convergence**: Did estimation converge successfully?
  - **convergence=0**: Converged in both the abstol test and the log-log plot test.
  - **convergence=1**: Some of the parameter estimates did not converge (based on the log-log plot test AND abstol tests) before `mleobj$control$maxit` was reached. This is not an error per se.
  - **convergence=2**: No convergence diagnostics were computed because the MLE object had problems and was not fit. This isn’t a convergence error just information.
  - **convergence=3**: No convergence diagnostics were computed because the MLE object was not fit. This isn’t a convergence error just information.
  - **convergence=10**: Abstol convergence only. Some of the parameter estimates did not converge (based on the log-log plot test) before `mleobj$control$maxit` was reached. However `mleobj$control$abstol` was reached.
  - **convergence=11**: Log-log convergence only. Some of the parameter estimates did not converge (based on the abstol test) before `mleobj$control$maxit` was reached. However the log-log convergence test was passed.
  - **convergence=12**: Abstol convergence only. Log-log convergence test was not computed because `mleobj$control$maxit` was set to less than `control=min.iter.conv.test`.
convergence=13 Lack of convergence info. Parameter estimates did not converge based on the abstol test before MLEobj$control$maxit was reached. No log-log information since control$min.iter.conv.test is less than MLEobj$control$maxit so no log-log plot test could be done.

convergence=42 MLEobj$control$abstol was reached but the log-log plot test returned NAs. This is an odd error and you should set control$trace=TRUE and look at the outputted $iter.record to see what is wrong.

convergence=52 The EM algorithm was abandoned due to numerical errors. Usually this means one of the variances either went to zero or to all elements being equal. This is not an error per se. Most likely it means that your model is not very good for your data (too inflexible or too many parameters). Try setting control$trace=1 to view a detailed error report.

convergence=62 The algorithm was abandoned due to errors in the log-log convergence test. You should not get this error (it is included for debugging purposes to catch improper arguments passed into the log-log convergence test).

convergence=63 The algorithm was run for control$maxit iterations, control$abstol not reached, and the log-log convergence test returned errors. You should not get this error (it is included for debugging purposes to catch improper arguments passed into the log-log convergence test).

convergence=72 Other convergence errors. This is included for debugging purposes to catch misc. errors.

logLik Log-likelihood.

states State estimates from the Kalman filter.

states.se Confidence intervals based on state standard errors, see caption of Fig 6.3 (p. 337) Shumway & Stoffer.

errors Any error messages.

Discussion

To ensure that the global maximum-likelihood values are found, it is recommended that you test the fit under different initial parameter values, particularly if the model is not a good fit to the data. This requires more computation time, but reduces the chance of the algorithm terminating at a local maximum and not reaching the true MLEs. For many models and for draft analyses, this is unnecessary, but answers should be checked using an initial conditions search before reporting final values. See the chapter on initial conditions in the User Guide for a discussion on how to do this.

MARSSkem() calls a Kalman filter/smooother (MARSSkf) for hidden state estimation. The algorithm allows two options for the initial state conditions: fixed but unknown or a prior. In the first case, x0 (whether at t=0 or t=1) is treated as fixed but unknown (estimated); in this case, fixed$V0=0 and x0 is estimated. This is the default behavior. In the second case, the initial conditions are specified with a prior and V0!=0. In the later case, x0 or V0 may be estimated. MARSS will allow you to try to estimate both, but many researchers have noted that this is not robust so you should fix one or the other.

If you get errors, it generally means that the solution involves an ill-conditioned matrix. For example, your Q or R matrix is going to a value in which all elements have the same value, for example zero. If for example, you tried to fit a model with fixed and high R matrix and the variance in
that R matrix was much higher than what is actually in the data, then you might drive Q to zero. Also if you try to fit a structurally inadequate model, then it is not unusual that Q will be driven to zero. For example, if you fit a model with 1 hidden state trajectory to data that clearly have 2 quite different hidden state trajectories, you might have this problem. Comparing the likelihood of this model to a model with more structural flexibility should reveal that the structurally inflexible model is inadequate (much lower likelihood).

Convergence testing is done via a combination of two tests. The first test (abstol test) is the test that the change in the absolute value of the log-likelihood from one iteration to another is less than some tolerance value (abstol). The second test (log-log test) is that the slope of a plot of the log of the parameter value or log-likelihood versus the log of the iteration number is less than some tolerance. Both of these must be met to generate the Success! parameters converged output. If you want to circumvent one of these tests, then set the tolerance for the unwanted test to be high. That will guarantee that that test is met before the convergence test you want to use is met. The tolerance for the abstol test is set by `controlDabstol` and the tolerance for the log-log test is set by `controlDconvNtestNslopeNtol`. Anything over 1 is huge for both of these.

### Author(s)
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### References


### See Also
MARSSkf, marssMLE, MARSSoptim

### Examples
```r
dat <- t(harborSeal)
dat <- dat[2:4,]
#you can use MARSS to construct a proper marssMLE object.
MLEobj <- MARSS(dat, model=list(Q="diagonal and equal", U="equal"), fit=FALSE)
#Pass this MLEobj to MARSSkem to do the fit.
```
MARSSkemcheck

Model Checking for MLE objects Passed to MARSSkem

Description

This is a helper function in the MARSS-package that checks that the model can be handled by the MARSSkem algorithm. It also returns the structure of the model as a list of text strings.

Usage

MARSSkemcheck( MLEobj )

Arguments

MLEobj An object of class marssMLE.

Value

A list with of the model elements A, B, Q, R, U, x0, Z, V0 specifying the structure of the model using text strings).

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See Also

marssMODEL MARSSkem

MARSSkf

Kalman Filtering and Smoothing for Time-varying MARSS models

Description

Provides Kalman filter and smoother output for MARSS models with (or without) time-varying parameters. This is a base function in the MARSS-package. MARSSkf is a small helper function to select which Kalman filter/smooher function to use based on which function was requested (in MLEobj$fun.kf). The default function is MARSSkfas.

Usage

MARSSkf( MLEobj, only.logLik=FALSE, return.lag.one=TRUE, return.kfas.model=FALSE )
MARSSkfss( MLEobj )
MARSSkfas( MLEobj, only.logLik=FALSE, return.lag.one=TRUE, return.kfas.model=FALSE )
Arguments

MLEobj  A marssMLE object with the `par` element of estimated parameters, `marss` element with the model description (in marss form) and data, and `control` element for the fitting algorithm specifications. `control$debugkf` specifies that detailed error reporting will be returned (only used by MARSSkf). `model$diffuse=TRUE` specifies that a diffuse prior be used (only used by MARSSkfas). See KFAS documentation. When the diffuse prior is set, V0 should be non-zero since the diffuse prior variance is V0*kappa, where kappa goes to infinity.

only.logLik Used by MARSSkfas. If set, only the log-likelihood is returned using the KFAS function `logLik.SSModel`. This is much faster if only the log-likelihood is needed.

return.lag.one Used by MARSSkfas. If set to FALSE, the smoothed lag-one covariance values are not returned (Vtt1T is set to NULL). This speeds up MARSSkfas because to return the smoothed lag-one covariance a stacked MARSS model is used with twice the number of state vectors—thus the state matrices are larger and take more time to work with.

return.kfas.model Used by MARSSkfas. If set to TRUE, it returns the MARSS model in KFAS model form (class `SSModel`). This is useful if you want to use other KFAS functions or write your own functions to work with `optim` to do optimization. This can speed things up since there is a bit of code overhead in `MARSSoptim` associated with the `marssMODELMODEL` model specification needed for the constrained EM algorithm (but not strictly needed for `optim`; useful but not required.).

Details

For state-space models, the Kalman filter and smoother provide optimal (minimum mean square error) estimates of the hidden states. The Kalman filter is a forward recursive algorithm which computes estimates of the states x(t) conditioned on the data up to time t (xtt). The Kalman smoother is a backward recursive algorithm which starts at time T and works backwards to t = 1 to provide estimates of the states conditioned on all data (xtT). The data may contain missing values (NAs). All parameters may be time varying.

The expected value of the initial state, x0, is an estimated parameter (or treated as a prior). This E(initial state) can be treated in two different ways. One can treat it as x00, meaning E(x at t=0 | y at t=0), and then compute x10, meaning E(x at t=1 | y at t=0), from x00. Or one can simply treat the initial state as x10, meaning E(x at t=1 | y at t=0). The approaches lead to the same parameter estimates, but the likelihood is written slightly differently in each case and you need your likelihood calculation to correspond to how the initial state is treated in your model (either x00 or x10). The EM algorithm in the MARSS package (`MARSSkem`) provides both Shumway and Stoffer’s derivation that uses tinitx=0 and Ghahramani et al algorithm which uses tinitx=1. The `MLEobj$model$tinitx` argument specifies whether the initial states (specified with x0 and V0) is at t=0 (tinitx=0) or t=1 (tinitx=1).

MARSSkfss() is a native R implementation based on the traditional Kalman filter and smoother equation as shown in Shumway and Stoffer (sec 6.2, 2006). The equations have been altered slightly to the initial state distribution to be to be specified at t=0 or t=1 (data starts at t=1) per per Ghahramani and Hinton (1996). In addition, the filter and smoother equations have been altered to allow
partially deterministic models (some or all elements of the Q diagonal equal to 0), partially perfect
observation models (some or all elements of the R diagonal equal to 0) and fixed (albeit unknown)
initial states (some or all elements of the V0 diagonal equal to 0) (per Holmes 2012). The code in-
cludes numerous checks to alert the user if matrices are becoming ill-conditioned and the algorithm
unstable.

MARSSkfas() uses the (Fortran-based) Kalman filter and smoother function (kfs) in the KFAS
Koopman and Durbin algorithm is faster and more stable since it avoids matrix inverses. Exact dif-
fuse priors are also allowed in the KFAS Kalman filter function. The standard output from the KFAS
functions do not include the lag-one covariance smoother needed for the EM algorithm. MARSSkfas
computes the smoothed lag-one covariance using the Kalman filter applied to a stacked MARSS
model as described on page 321 in Shumway and Stoffer (2000). Also the KFAS model specifi-
cation only has the initial state at t=1 (as x(1) conditioned on y(0), which is missing). When the
initial state is specified at t=0 (as x(0) conditioned on y(0), which is missing), MARSSkfas computes
the required E(x(1)|y(0)) and var(x(1)|y(0)) using the Kalman filter equations per Ghahramani and

The likelihood returned for both functions is the exact likelihood when there are missing values
rather than the approximate likelihood sometimes presented in texts for the missing values case.
The functions return the same filter, smoother and log-likelihood values. The differences are that
MARSSkfas is faster (and more stable) but MARSSkf has many internal checks and error messages
which can help debug numerical problems (but slow things down). Also MARSSkf returns some
output specific to the traditional filter algorithm (J and Kt).

Value

A list with the following components (m is the number of state processes). "V" elements are called
"P" in Shumway and Stoffer (S&S eqn 6.17 with s=T). The output is referenced against equations
in Shumway and Stoffer (2006); the Kalman filter and smoother implemented in MARSS is for a
more general MARSS model than that shown in S&S but the output has the same meaning. In the
expectations below, the parameters are left off, so E[x | y] is really E[x | theta, y] where theta is the
parameter list.

xtT State first moment conditioned on y(1:T): E[x(t) | y(1:T)] (m x T matrix). Kalman
smoother output.

VttT State variance conditioned on y(1:T): E[(x(t)-xtT(t))(x(t)-xtT(t))'| | y(1:T)] (m x
m x T array). Kalman smoother output. P_tT in S&S (S&S eqn 6.18 with s=T,
t1=t2=t). State second moment E[x(t)x(t)' | y(1:T)] = VttT(t)+xtT(t)xtT(t)'.

Vtt1T State lag-one covariance E[(x(t)-xtT(t))(x(t-1)-xtT(t-1))'| | y(1:T)] (m x m x T).
Kalman smoother output. P_t,t-1T in S&S (S&S eqn 6.18 with s=T, t1=t, t2=t-1).
State lag-one second moments E[x(t)x(t-1)' | y(1:T)] = Vtt1T(t)+xtT(t)xtT(t-1)'.

x0T Initial state estimate E[x(i) | y(1:T)] (m x 1). If control$kf.x0="x00", i=0; if
="x10", i=1. Kalman smoother output.

V0T Estimate of initial state covariance matrix E[x(i)x(i)' | y(1:T)] (m x m). If
model$initx=0, i=0; if =1, i=1. Kalman smoother output. P_0T in S&S.

J (m x m x T) Kalman smoother output. Only for MARSSkfss. (ref S&S eqn 6.49)
J0  J at init time (t=0 or t=1) (m x m x T). Kalman smoother output. Only for MARSSkfss.

xtt State first moment conditioned on y(1:t): E[x(t) | y(1:t)] (m x T). Kalman filter output. (S&S eqn 6.17 with s=t)

xtt1 State first moment conditioned on y(1:t-1): E[x(t) | y(1:t-1)] (m x T). Kalman filter output. (S&S eqn 6.17 with s=t-1)

Vtt State variance conditioned on y(1:t): E[(x(t)-xtt(t))(x(t)-xtt(t))' | y(1:t)] (m x m x T array). Kalman filter output. P_t^t in S&S (S&S eqn 6.18 with s=t, t1=t2=t). State second moment E[x(t)x(t)' | y(1:t)] = Vtt(t)+xtt(t)xtt(t)'

Vtt1 State variance conditioned on y(1:t-1): E[(x(t)-xtt1(t))(x(t)-xtt1(t))' | y(1:t-1)] (m x m x T array). Kalman filter output. P_{t1}^t in S&S (S&S eqn 6.18 with s=t-1, t1=t2=t). State second moment E[x(t)x(t)' | y(1:t-1)] = Vtt1(t)+xtt1(t)xtt1(t)'

Kt Kalman gain (m x m x T). Kalman filter output (ref S&S eqn 6.23). Only for MARSSkfss.

Innov Innovations y(t) - E[y(t) | y(1:t-1)] (n x T). Kalman filter output. Only returned with MARSSkfss. (ref page S&S 339).

Sigma Innovations covariance matrix. Kalman filter output. Only returned with MARSSkfss. (ref S&S eqn 6.61)

logLik Log-likelihood logL(y(1:T) | theta) (ref S&S eqn 6.62)

kfas.model The model in KFAS model form (class SSMModel). Only for MARSSkfss.

errors Any error messages.

Author(s)

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References


See Also

MARSS marssMODEL MARSSkem

Examples

dat <- t(harborSeal)
dat <- dat[2:nrow(dat),]
# you can use MARSS to construct a MLEobj
#MARSS calls MARSSinits to construct default initial values
MLEobj <- MARSS(dat, fit=FALSE)
#MARSSkf needs a marss MLE object with the par element set
MLEobj$par <- MLEobj$start
#Compute the kf output at the params used for the inits
kflist <- MARSSkf(as(MLEobj )

---

**marssMLE**

*Maximum Likelihood MARSS Estimation Object*

**Description**

An object in the **MARSS-package** that has all the elements needed for maximum-likelihood estimation of multivariate autoregressive state-space model: the data, model, estimation methods, and any control options needed for the method. If the model has been fit and parameters estimated, the object will also have the MLE parameters. Other functions add other elements to the marssMLE object, such as CIs, s.e.'s, AICs, and the hessian. There are print, summary, coef, residuals, predict and simulate methods for marssMLE objects and a bootstrap function. Rather than working directly with the elements of a marssMLE object, use **print.marssMLE** to extract output.

**Usage**

`is.marssMLE(MLEobj)`

**Arguments**

- **MLEobj**: An object of class **marssMLE**. See Details.
Details

The `is.marssMLE()` function checks components `marss`, `start` and `control`, which must be present for estimation by functions e.g. `MARSSkem`. Components returned from estimation must include at least `method`, `par`, `kf`, `numIter`, `convergence` and `logLik`. Additional components (e.g. AIC) may be returned, as described in function help files.

**model**  An object of class `marssModel` in whatever form the user specified in the call to `MARSS()`. Default is form `marxss`.

**marss**  An object of class `marssModel` in `marss` forms, needed for all the base `MARSS` functions.

**start**  List with 8 matrices `Z`, `A`, `R`, `B`, `U`, `Q`, `x0`, `V0`, specifying initial values for parameters to be used (if needed) by the maximization algorithm.

- `B` Initial value(s) for `B` matrix (`m x m`).
- `U` Initial value(s) for `U` matrix (`m x 1`).
- `Q` Initial value(s) for `Q` variance-covariance matrix (`m x m`).
- `Z` Initial value(s) for `Z` matrix (`n x m`).
- `A` Initial value(s) for `A` matrix (`n x 1`).
- `R` Initial value(s) for `R` variance-covariance matrix (`n x n`).
- `x0` Initial value(s) for initial state vector (`m x 1`).
- `V0` Initial variance(s) for initial state variance (`m x m`).

**control**  A list specifying estimation options. The following options are needed by `MARSSkem`. Other control options can be set if needed for other estimation methods, e.g. the control options listed for `optim` for use with `MARSSoptim`. The default values for control options are set in `alldefaults[[method]]` which is specified in `MARSSsettings.R`.

- `minit`  The minimum number of iterations to do in the maximization routine (if needed by method).
- `maxit`  Maximum number of iterations to be used in the maximization routine (if needed by method).
- `min.iter.conv.test`  Minimum iterations to run before testing convergence via the slope of the log parameter versus log iterations.
- `conv.test.deltaT=9`  Number of iterations to use for the testing convergence via the slope of the log parameter versus log iterations.
- `conv.test.slope.tol`  The slope of the log parameter versus log iteration to use as the cutoff for convergence. The default is 0.5 which is a bit high. For final analyses, this should be set lower.
- `abstol`  The `logLik.(iter-1)-logLik.(iter)` convergence tolerance for the maximization routine. Both the abstol and the slope of the log of the parameters versus the log iteration tests must be met for convergence.
- `trace`  A positive integer. If not 0, a record will be created during maximization iterations (what’s recorded depends on method of maximization). -1 turns off most internal error checking.
- `safe`  Logical. If TRUE, then the Kalman filter is run after each update equation in the EM algorithm. This slows down the algorithm. The default is FALSE.
- `allow.degen`  If TRUE, replace `Q` or `R` diagonal elements by 0 when they become very small.
- `min.degen.iter`  Number of iterations before trying to set a diagonal element of `Q` or `R` to zero.
**degen.lim** How small the Q or R diagonal element should be before attempting to replace it with zero.

**silent** Suppresses printing of progress bar, error messages and convergence information.

**method** A string specifying the estimation method. MARSS allows "kem" for EM and "BFGS" for quasi-Newton. Once the model has been fitted, additional elements are added.

**par** A list with 8 matrices of estimated parameter values Z, A, R, B, U, Q, x0, V0.

**states** Expected values of the x (hidden states).

**states.se** Standard errors on the estimates states.

**ytT** Expected values of the y. This is just y for non-missing y.

**ytT.se** Standard errors on the ytT. This will be 0 for non-missing y.

**kf** A list containing Kalman filter/smoother output if control$trace is > 0.

**Ey** A list containing expectations involving y. Output if control$trace is > 0.

**numIter** Number of iterations which were required for convergence.

**convergence** Convergence status and errors. 0 means converged successfully. Anything else means an error or warning.

**logLik** Log-likelihood.

**AIC** AIC

**AICc** Corrected AIC.

**call** A list of all the arguments passed into the MARSS call. Not required for most functions, but is a record of what was used to call MARSS for checking and can be used to customize the printing of MARSS output.

---

**Value**

TRUE if no problems; otherwise a message describing the problems.

**Author(s)**

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**See Also**

marssMODEL MARSSkem
marssMLE-class

Class "marssMLE"

Description

marssMLE objects specify fitted multivariate autoregressive state-space models (maximum-likelihood) in the package MARSS-package.

Methods

- **print** signature(x = "marssMLE"): ...
- **summary** signature(object = "marssMLE"): ...
- **coef** signature(object = "marssMLE"): ...
- **predict** signature(object = "marssMLE"): ...
- **simulate** signature(object = "marssMLE"): ...

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marssMODEL-class

Class "marssMODEL"

Description

marssMODEL objects describe a vectorized form for the multivariate autoregressive state-space models used in the package MARSS-package.

The object has the following attributes:

- **form** The form that the model object is in.
- **par.names** The names of each parameter matrix in the model.
- **model.dims** A list with the dimensions of all the matrices in non-vectorized form.
- **X.names** Names for the X rows.
- **Y.names** Names for the Y rows.
- **equation** The model equation. Used to write the model in LaTeX.
These attributes are set in the MARSS_form.R file, in the MARSS.form() function and must be internally consistent with the elements of the model. These attributes are used for internal error checking.

Each parameter matrix in a MARSS equation can be written in vectorized form: vec(P) = f + Dp, where f is the fixed part, p are the estimated parameters, and D is the matrix that transforms the p into a vector to be added to f.

An object of class "marssMODEL" is a list with elements:

- data Data supplied by user.
- fixed A list with the f row vectors for each parameter matrix.
- free A list with the D matrices for each parameter matrix.
- tinitx At what t, 0 or 1, is the initial x defined at?
- diffuse Whether a diffuse initial prior is used. TRUE or FALSE. Not used unless method="BFGS" was used.

For the marss form, the matrices are called: Z, A, R, B, U, Q, x0, V0. This is the form used by all internal algorithms, thus alternate forms must be transformed to marss form before fitting. For the marxss form (the default form in a MARSS() call), the matrices are called: Z, A, R, B, U, Q, D, C, d, c, x0, V0.

Each form, should have a file called MARSS_form.R, with the following functions. Let foo be some form.

- MARSS.foo(MARSS.call) This is called in MARSS() and takes the input from the MARSS() call (a list called MARSS.call) and returns that list with two model objects added. First is a model object in marss form in the $marss element and a model object in the form foo.
- marss_to_foo(marssMLE or marssMODEL) If called with marssMODEL (in form marss), marss_to_foo returns a model in form foo. If marss_to_foo is called with a marssMLE object (which must have a $marss element by definition), it returns a $model element in form foo and all if the marssMLE object has par, par.se, par.CI, par.bias, start elements, these are also converted to foo form. The function is mainly used by print.foo which needs the par (and related) elements of a marssMLE object to be in foo form for printing.
- foo_to_marss(marssMODEL or marssMLE) This converts marssMODEL(form=foo) to marssMODEL(form=marss). This transformation is always possible since MARSS only works for models for which this is possible. If called with marssMODEL, it returns only a marssMODEL. If called with marssMLE, it adds the $marss element with a marssMODEL in marss form and if the par (or related) elements exists, these are converted to marss form.
- print_foo(marssMLE or marssMODEL) print.marssMLE prints the par (and par.se and start) element of a marssMLE object but does not make assumptions about its form. Normally this element is in form=marss. print.marssMLE checks for a print_foo function and runs that on the marssMLE object first. This allows one to call foo_to_marss() to covert the par (and related) element to foo form so they look familiar to the user (the marss form will look strange). If called with marssMLE, print_foo returns a marssMLE object with the par (and related) elements in foo form. If called with a marssMODEL, print_foo returns a marssMODEL in foo form.
- coef_foo(marssMLE) See print_foo. Coef.marssMLE also uses the par (and related) elements.
• predict_foo(marssMLE) Called by predict.marssMLE to do any needed conversions. Typically a form will want the newdata element in a particular format and this will need to be converted to marss form. This returns an updated marssMLE object and newdata.

• describe_foo(marssMODEL) Called by describe.marssMODEL to do allow custom description output.

• is.marssMODEL_foo(marssMODEL) Check that the model object in foo form has all the parts it needs and that these have the proper size and form.

• MARSSinits_foo(marssMLE, inits.list) Allows customization of the inits used by the form. Returns an inits list in marss form.

Methods

print signature(x = "marssMODEL"): ...

summary signature(object = "marssMODEL"): ...

Author(s)

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Description

Parameter estimation for MARSS models using R’s optim function. This allows access to R’s quasi-Newton algorithms available via the optim function. The MARSSoptim function is called when MARSS is called with method="BFGS". This is a base function in the MARSS-package.

Usage

MARSSoptim(MLEobj)

Arguments

MLEobj An object of class marssMLE.

Details

Objects of class marssMLE may be built from scratch but are easier to construct using MARSS with MARSS(..., fit=FALSE, method="BFGS").

Options for optim are passed in using MLEobj$control. See optim for a list of that function’s control options. If lower and upper for optim need to be passed in, they should be passed in as part of control as control$lower and control$upper. Additional control arguments affect printing and initial conditions.
The initial condition is at $t=0$ if kf.x0="x00". The initial condition is at $t=1$ if kf.x0="x10".

If diffuse=TRUE, a diffuse initial condition is used. MLEobj$par$V0 is then the scaling function for the diffuse part of the prior. Thus the prior is V0*kappa where kappa→Inf. Note that setting a diffuse prior does not change the correlation structure within the prior. If diffuse=FALSE, a non-diffuse prior is used and MLEobj$par$V0 is the non-diffuse prior variance on the initial states. The the prior is V0.

Suppresses printing of progress bars, error messages, warnings and convergence information.

The marssMLE object which was passed in, with additional components:

- **method**: String "BFGS".
- **kf**: Kalman filter output.
- **iter.record**: If MLEobj$control$trace = TRUE, then this is the $message value from optim.
- **numIter**: Number of iterations needed for convergence.
- **convergence**: Did estimation converge successfully?
  - **convergence=0**: Converged in less than MLEobj$control$maxit iterations and no evidence of degenerate solution.
  - **convergence=1**: Maximum number of iterations MLEobj$control$maxit was reached before MLEobj$control$abstol condition was satisfied.
  - **convergence=10**: Some of the variance elements appear to be degenerate. T
  - **convergence=52**: The algorithm was abandoned due to errors from the "L-BFGS-B" method.
  - **convergence=53**: The algorithm was abandoned due to numerical errors in the likelihood calculation from MARSSkf. If this happens with "BFGS", it can sometimes be helped with a better initial condition. Try using the EM algorithm first (method="kem"), and then using the parameter estimates from that to as initial conditions for method="BFGS".
- **logLik**: Log-likelihood.
- **states**: State estimates from the Kalman filter.
- **states.se**: Confidence intervals based on state standard errors, see caption of Fig 6.3 (p. 337) Shumway & Stoffer.
- **errors**: Any error messages.

The function only returns parameter estimates. To compute CIs, use MARSSparamCIs but if you use parametric or non-parametric bootstrapping with this function, it will use the EM algorithm to compute the bootstrap parameter estimates! The quasi-Newton estimates are too fragile for the bootstrap routine since one often needs to search to find a set of initial conditions that work (i.e. don’t lead to numerical errors).
Estimates from MARSSoptim (which come from optim) should be checked against estimates from the EM algorithm. If the quasi-Newton algorithm works, it will tend to find parameters with higher likelihood faster than the EM algorithm. However, the MARSS likelihood surface can be multimodal with sharp peaks at degenerate solutions where a Q or R diagonal element equals 0. The quasi-Newton algorithm sometimes gets stuck on these peaks even when they are not the maximum. Neither an initial conditions search nor starting near the known maximum (or from the parameters estimates after the EM algorithm) will necessarily solve this problem. Thus it is wise to check against EM estimates to ensure that the BFGS estimates are close to the MLE estimates (and visa-versa, it’s wise to rerun with method="BFGS" after using method="kem"). Conversely, there is a strong flat ridge in your likelihood, the EM algorithm can report early convergence while the BFGS may continue much further along the ridge and find very different parameter values. Of course a likelihood surface with strong flat ridges makes the MLEs less informative...

Note this is mainly a problem if the time series are short or very gappy. If the time series are long, then the likelihood surface should be nice with a single interior peak. In this case, the quasi-Newton algorithm works well but it can still be sensitive (and slow) if not started with a good initial condition. Thus starting it with the estimates from the EM algorithm is often desirable.

One should be aware that the prior set on the variance of the initial states at t=0 or t=1 can have catastrophic effects on one’s estimates if the presumed prior covariance structure conflicts with the structure implied by the MARSS model. For example, if you use a diagonal variance-covariance matrix for the prior but the model implies a matrix with non-zero covariances, your MLE estimates can be strongly influenced by the prior variance-covariance matrix. Setting a diffuse prior does not help because the diffuse prior still has the correlation structure specified by V0. One way to detect priors effects is to compare the BFGS estimates to the EM estimates. Persistent differences typically signify a problem with the correlation structure in the prior conflicting with the implied correlation structure in the MARSS model. If this is the case, using V0=0 and estimating the x0 elements (with control$kf.x0="x10") can often help.

**Author(s)**

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**See Also**

MARSS MARSSkem marssMLE optim

**Examples**

```r
dat <- t(harborSealWA)
dat <- dat[2:4,] #remove the year row

#fit a model with EM and then use that fit as the start for BFGS
#fit a model with 1 hidden state where obs errors are iid
#R="diagonal and equal" is the default so not specified
#Q is fixed
kemfit <- MARSS(dat, model=list(Z=matrix(1,3,1),Q=matrix(.01)))
bfgsfit <- MARSS(dat, model=list(Z=matrix(1,3,1),Q=matrix(.01)),
    inits=coef(kemfit,form="marss"), method="BFGS")
```
MARSSparamCIs

Description
Computes standard errors, confidence intervals and bias for the maximum-likelihood estimates of MARSS model parameters. If you want confidence intervals on the estimated hidden states, see `print.marssMLE` and look for "states.cis".

Usage
```r
MARSSparamCIs(MLEobj, method = "hessian", alpha = 0.05, nboot = 1000, silent = TRUE, hessian.fun = "Harvey1989")
```

Arguments
- `MLEobj`: An object of class `marssMLE`. Must have a `$par` element containing the MLE parameter estimates.
- `method`: Method for calculating the standard errors: "hessian", "parametric", and "innovations" implemented currently.
- `alpha`: Alpha level for the 1-alpha confidence intervals.
- `nboot`: Number of bootstraps to use for "parametric" and "innovations" methods.
- `hessian.fun`: The function to use for computing the Hessian. See `MARSShessian`.
- `silent`: If false, a progress bar is shown for "parametric" and "innovations" methods.

Details
Approximate confidence intervals (CIs) on the model parameters may be calculated from the Hessian matrix (the matrix of partial 2nd derivatives of the parameter estimates) or parametric or non-parametric (innovations) bootstrapping using `nboot` bootstraps. The Hessian CIs are based on the asymptotic normality of MLE parameters under a large-sample approximation. The Hessian computation for variance-covariance matrices is a symmetric approximation and the lower CIs for variances might be negative. Using a Hessian approximation for variances is approximate. Bootstrap estimates of parameter bias are reported if method "parametric" or "innovations" is specified.

Note, these are added to the `par` (etc) elements of a marssMLE object but are in marss form not marxss form. Thus the `MLEobj$par.upCI` and related elements that are added to the marssMLE object may not look familiar to the user. Instead the user should extract these elements using `print(MLEobj)` and passing in the argument `what` set to "par.se", "par.bias", "par.lowCIs", or "par.upCIs". See `print.marssMLE`.

Value
`MARSSparamCIs` returns the `marssMLE` object passed in, with additional components `par.se`, `par.upCI`, `par.lowCI`, `par.CI.alpha`, `par.CI.method`, `par.CI.nboot` and `par.bias` (if method is "parametric" or "innovations").
**MARSSsimulate**

Simulate Data from a MARSS Model

Description

Generates simulated data from a MARSS model with specified parameter estimates. This is a base function in the `MARSS-package`.

Usage

```r
MARSSsimulate(MLEobj, tSteps = NULL, nsim = 1, silent = TRUE, miss.loc = NULL)
```

Arguments

- **MLEobj**: A fitted `marssMLE` object, as output by `MARSS`.
- **tSteps**: Number of time steps in each simulation. If left off, it is taken to be consistent with `MLEobj`.
- **nsim**: Number of simulated data sets to generate.
- **silent**: Suppresses progress bar.
- **miss.loc**: Optional matrix specifying where to put missing values. See Details.
Details

Optional argument miss.loc is an array of dimensions n x tSteps x nsim, specifying where to put missing values in the simulated data. If missing, this would be constructed using MLEobj.marss$data. If the locations of the missing values are the same for all simulations, miss.loc can be a matrix of dim=c(n, tSteps) (the original data for example). The default, if miss.loc is left off, is that there are no missing values even if MLEobj.marss$data has missing values.

Value

| sim.states | Array (dim m x tSteps x nsim) of state processes simulated from parameter estimates. m is the number of states (rows in X). |
| sim.data   | Array (dim n x tSteps x nsim) of data simulated from parameter estimates. n is the number of rows of data (Y). |
| MLEobj     | The marssMLE object from which the data were simulated. |
| miss.loc   | Matrix identifying where missing values were placed. |
| tSteps     | Number of time steps in each simulation. |
| nsim       | Number of simulated data sets generated. |

Author(s)

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See Also

marssMODEL marssMLE MARSSboot

Examples

d = harborSeal[,]c(2,11)
dat = t(d)
MLEobj = MARSS(dat)

#simulate data that are the
#same length as original data and no missing data
sim.obj = MARSSsimulate(MLEobj, tSteps=dim(d)[1], nsim=5)

#simulate data that are the
#same length as original data and have missing data in the same location
sim.obj = MARSSsimulate(MLEobj, tSteps=dim(d)[1], nsim=5, miss.loc=dat)
**MARSSvectorizeparam**  
*Vectorize or Replace the par List*

**Description**

Converts `MLEobj[["what"]]]` to a vector or assigns a vector to `MLEobj[["what"]]]`. This is a utility function in the **MARSS-package** for marssMODEL objects of form="marss" and is not exported. Users achieve this functionality with `coef.marssMLE`.

**Usage**

```r
MARSSvectorizeparam(MLEobj, parvec = NA, what = "par")
```

**Arguments**

- **MLEobj**  
  An object of class **marssMLE**.

- **parvec**  
  NA or a vector. See Value.

- **what**  
  What part of the MLEobj is being replaced or vectorized. Need to be a par list.

**Details**

Utility function to generate parameter vectors for optimization functions, and to set `MLEobj[[what]]` using a vector of values. The function bases the unlisting and naming order on `names(MLEobj$marss$fixed)`. Appends matrix name to the row names in the par list.

**Value**

If parvec=NA, a vector of the elements of the what element. Otherwise, a **marssMLE** object with `MLEobj[["what"]]]` set by parvec.

**Author(s)**

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**See Also**

`marssMLE`

**Examples**

```r
dat <- t(harborSealWA)
dat <- dat[2:4,]
kem <- MARSS(dat)
paramvec = MARSS::MARSSvectorizeparam(kem)
paramvec
```
model.frame.marssMODEL

model.frame method for marssMLE and marssMODEL objects

Description

model.frame(marssMLE) or model.frame(marssMODEL), where marssMLE is the output from a
MARSS call marssMODEL is the model element of a marssMLE object, will return a data.frame with
the data (y) and inputs/covariates (c and d elements) for a MARXSS model. See MARSS.marxss.
This is mainly a utility function to help with the broom functions (tidy, augment and glance).

Usage

## S3 method for class 'marssMODEL'
model.frame(formula, ...)

Arguments

formula A marssMODEL object.

... Other arguments not used.

Value

A data.frame with the data and inputs (c and d) in a MARXSS model. See MARSS.marxss.

Author(s)

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plankton Plankton Data Sets

Description

Example data sets for use in MARSS vignettes for the MARSS-package.
The lakeWAplankton dataset consists for two datasets: lakeWAplanktonRaw and a dataset derived
from the raw dataset, lakeWAplanktonTrans. lakeWAplanktonRaw is a 32-year time series (1962-
1994) of monthly plankton counts from Lake Washington, Washington, USA. Columns 1 and 2 are
year and month. Column 3 is temperature (C), column 4 is total phosphorous, and column 5 is
pH. The next columns are the plankton counts in units of cells per mL for the phytoplankton and
organisms per L for the zooplankton. Since MARSS functions require time to be across columns,
these data matrices must be transposed before passing into MARSS functions.
lakeWAplanktonTrans is a transformed version of lakeWAplanktonRaw. Zeros have been replaced with NAs (missing). The logged (natural log) raw plankton counts have been standardized to a mean of zero and variance of 1 (so logged and then z-scored). Temperature, TP & pH were also z-scored but not logged (so z-score of the untransformed values for these covariates). The single missing temperature value was replaced with -1 and the single missing TP value was replaced with -0.3.

The Ives data are from Ives et al. (2003) for West Long Lake (the low planktivory case). The Ives data are unlogged. ivesDataLP and ivesDataByWeek are the same data with LP having the missing weeks in winter removed while in ByWeek, the missing values are left in. The phosphorous column is the experimental input rate + the natural input rate for phosphorous, and Ives et al. used 0.1 for the natural input rate when no extra phosphorous was added. The phosphorous input rates for weeks with no sampling (and no experimental phosphorous input) have been filled with 0.1 in the "by week" data.

**Usage**

```r
data(ivesDataLP)
data(ivesDataByWeek)
data(lakeWAplankton)
```

**Format**

The data are provided as a matrix with time running down the rows.

**Source**

- lakeWAplanktonRaw Adapted from the Lake Washington database of Dr. W. T. Edmondson, as funded by the Andrew Mellon Foundation; data courtesy of Dr. Daniel Schindler, University of Washington, Seattle, WA.

**Examples**

```r
str(ivesDataLP)
str(ivesDataByWeek)
```
plot.marssMLE

plotting functions for MARSS MLE objects

Description

Plots fitted observations and estimated states with confidence intervals using base R graphics (plot) and ggplot2 (autoplot). Diagnostic plots also shown. By default all plots are plotted. Individual plots can be plotted by passing in type.plot. If an individual plot is made using autoplot, the ggplot object is returned which can be further manipulated.

Usage

## S3 method for class 'marssMLE'
plot(x, plot.type=c("observations", "states", "model.residuals", "state.residuals", "model.residuals.qqplot", "state.residuals.qqplot"), form=c("marxss", "marss", "dfa"), conf.int=TRUE, conf.level=0.95, decorate=TRUE, ...)

## S3 method for class 'marssMLE'
autoplot(x, plot.type=c("observations", "states", "model.residuals", "state.residuals", "model.residuals.qqplot", "state.residuals.qqplot"), form=c("marxss", "marss", "dfa"), conf.int=TRUE, conf.level=0.95, decorate=TRUE, ...)

Arguments

x A marssMLE object.

plot.type Type of plot. If not passed in, all plots are drawn. Options for arguments include 'observations' (fits to the raw data), 'states' (estimates of the hidden or latent trends), 'model.residuals' (residuals for the observation error), 'state.residuals' (residuals associated with the process model), 'model.residuals.qqplot' (qq plot for the observation residuals), 'state.residuals.qqplot' (qq plot for the state residuals).

form Optional. Form of the model. This is normally taken from the form attribute of the MLE object (x), but the user can specify a different form.

conf.int TRUE/FALSE. Whether to include a confidence interval.

conf.level Confidence level for CIs.

decorate TRUE/FALSE. Add smoothing lines to residuals plots or qqline to qqplots.

... Other arguments, not used.
Value

If an individual plot is selected using plot.type and autoplot is called, then the ggplot object is returned invisibly.

Author(s)

Eric Ward and Eli Holmes

Examples

data(harborSealWA)
fit <- MARSS(t(harborSealWA[-1]), model=list(Z=as.factor(c(1,1,1,1,2)), R="diagonal and equal"))
plot(fit, plot.type="observations")

require(ggplot2)
autoplot(fit, plot.type="observations")

## Not run:
# DFA example
dfa <- MARSS(t(harborSealWA[-1]), model=list(m=2), form="dfa")
plot(dfa, plot.type="states")

## End(Not run)

print.marssMLE

Printing functions for MARSS MLE objects

Description

The MARSS fitting function, MARSS, outputs marssMLE objects. print(marssMLE), where marssMLE is a marssMLE object, will print out information on the fit. However, print can be used to print a variety of information (residuals, smoothed states, imputed missing values, etc) from a marssMLE object using the what argument in the print call.

Usage

## S3 method for class 'marssMLE'
print(x, digits = max(3,getOption("digits")-4), ..., what="fit", form=NULL, silent=FALSE)

Arguments

x A marssMLE object.
digits Number of digits for printing.
... Other arguments for print.
what What to print. Default is "fit". If you input what as a vector, print returns a list. See examples.
• "model" The model parameters with names for the estimated parameters. The output is customized by the form of the model that was fit. This info is in \texttt{attr(x$\text{model}$, "form")}.
• "par" A list of only the estimated values in each matrix. Each model matrix has its own list element. Standard function: \texttt{coef(x)}
• "start" or "inits" The values that the optimization algorithm was started at. \texttt{x$start$} shows this in form="marss" while \texttt{print} shows it in whatever form is in \texttt{attr(x$\text{model}$, "form")}.
• "paramvector" A vector of all the estimated values in each matrix. Standard function: \texttt{coef(x, type="vector")}. See \texttt{coef.marssMLE}.
• "par.se","par.bias","par.lowCIs","par.upCIs" A vector the estimated parameter standard errors, parameter bias, lower and upper confidence intervals. Standard function: \texttt{MARSSparamCIs(x)} See \texttt{MARSSparamCIs}.
• "xtT" or "states" The estimated states conditioned on all the data. \texttt{x$states$}
• "data" The data. This is in \texttt{x$\text{model}$}$data$
• "logLik" The log-likelihood. Standard function: \texttt{x$logLik$}. See \texttt{MARSSkf} for a discussion of the computation of the log-likelihood for MARSS models.
• "ytT" The expected value of the data conditioned on all the data. Returns the data if present and the expected value if missing. This is in \texttt{x$ytT$} (ytT is analogous to xtT).
• "states.se" The state standard errors. \texttt{x$states.se$}
• "states.cis" Approximate confidence intervals for the states. See \texttt{MARSSparamCIs}.
• "model.residuals" The smoothed model residuals. \texttt{x(t)-E(y(t)|xtT(t))}, aka actual data at time $t$ minus the expected value of the data conditioned on the smoothed states estimate at time $t$. Standard function: \texttt{residuals(x)$\text{model.residuals}$} See \texttt{residuals.marssMLE} for a discussion of residuals in the context of MARSS models.
• "state.residuals" The smoothed state residuals. \texttt{E(xtT(t))-E(x(t)|xtT(t-1))}, aka the expected value of x at t conditioned on all the data minus the expected value of x at t conditioned on (x(t-1) conditioned on all the data). Standard function: \texttt{residuals(x)$\text{state.residuals}$} See \texttt{residuals.marssMLE}.
• parameter name Returns the parameter matrix for that parameter with fixed values at their fixed values and the estimated values at their estimated values. Standard function: \texttt{coef(x, type="matrix")$elem$}
• "kfs" The Kalman filter and smoother output. See \texttt{MARSSkf} for a description of the output. The full kf output is not normally attached to the output from a MARSS() call. This will run the filter/smoother if needed and return the list INVISIBLY. So assign the output as \texttt{foo=print(x,what="kfs")}
• "Ey" The expectations involving y conditioned on all the data. See \texttt{MARSShatyt} for a discussion of these expectations. This output is not normally attached to the output from a MARSS() call except ytT which is the predicted value of any missing y. The list is returned INVISIBLY. So assign the output as \texttt{foo=print(x,what="Ey")}.

\texttt{form} By default, print uses the model form specified in the call to \texttt{MARSS}. This information is in \texttt{attr(marssMLE$\text{model}$, "form")}, however you can specify a
different form. `form="marss"` should always work since this is the model form in which the model objects are stored (in `marssMLE$marss`).

`silent` If TRUE, do not print just return the object. If print call is assigned, nothing will be printed. See examples. If `what="fit"`, there is always output printed.

**Value**

A print out of information. If you assign the print call to a value, then you can reference the output. See the examples.

**Author(s)**

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**Examples**

```r
dat = t(harborSeal)
dat = dat[,c(2,11),]
MLEobj = MARSS(dat)

print(MLEobj)

print(MLEobj,what="model")

print(MLEobj,what="par")

# silent doesn't mean silent unless the print output is assigned
print(MLEobj,what="paramvector", silent=TRUE)
tmp=print(MLEobj,what="paramvector", silent=TRUE)
# silent means some info on what you are printing is shown whether
# or not the print output is assigned
print(MLEobj,what="paramvector", silent=FALSE)
tmp=print(MLEobj,what="paramvector", silent=FALSE)

cis=print(MLEobj,what="states.cis")
cis$up95CI

vars=print(MLEobj, what=c("R","Q"))
```

---

**Description**

`print(marssMODEL)`, where `marssMODEL` is a marssMODEL object, will print out information on the model in short form (e.g. ‘diagonal and equal’).

`summary(marssMODEL)`, where `marssMODEL` is a marssMODEL object, will print out detailed information on each parameter matrix showing where the estimated values (and their names) occur.
residuals.marssMLE

Usage

## S3 method for class 'marssMODEL'
print(x, ...)
## S3 method for class 'marssMODEL'
summary(object, ..., silent = FALSE)

Arguments

x
A marssMODEL object.

object
A marssMODEL object.

...
Other arguments.

silent
TRUE/FALSE Whether to print output.

Value

print(marssMODEL) prints out of the structure of each parameter matrix in 'English' (e.g. 'diagonal and unequal') and returns invisibly the list. If you assign the print call to a value, then you can reference the output.

summary(marssMODEL) prints out of the structure of each parameter matrix in as list matrices showing where each estimated value occurs in each matrix and returns invisibly the list. The output can be verbose, especially if parameter matrices are time-varying. Pass in silent=TRUE and assign output (a list with each parameter matrix) to a variable. Then specific parameters can be looked at.

Author(s)

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Examples

dat <- t(harborSeal)
dat <- dat[c(2,11),]
MLEobj <- MARSS(dat)

print(MLEobj$model)
#this is identical to
print(MLEobj, what="model")

---

residuals.marssMLE MARSS Standardized Residuals

Description

Calculates the standardized (or auxiliary) residuals sensu Harvey, Koopman and Penzer (1998).
Usage

```r
## S3 method for class 'marssMLE'
residuals(object, ..., Harvey=FALSE, normalize=FALSE)
```

Arguments

- `object`: An object of class `marssMLE`.
- `...`: Not used.
- `Harvey`: TRUE/FALSE
- `normalize`: TRUE/FALSE

Details

This function returns the conditional expected value (mean) and variance of the model and state residuals. 'conditional' means in this context, conditioned on the observed data and a set of parameters. If there are no missing values in the data, \( E(Y|y)=y \), and the model residual is the difference between \( y \) and the model prediction. If the data are missing, then the model residual is 0 because \( E(Y)=\)model prediction and the difference between these is 0. The variance however is non-zero.

If Harvey=TRUE, the function uses the algorithm on page 112 of Harvey, Koopman and Penzer (1998) to compute the conditional residuals and variance of the residuals. If Harvey=FALSE, the function uses the equations in the technical report Holmes (2014). The difference in the algorithm only concerns model residuals and their variance for missing values. The residual variance for missing data is not normally used but is needed for leave-one-out analyses when a non-missing \( y(t) \) is removed and then its residual is compared the model residual variance at time \( t \).

The residuals matrix (and se and variance) has a value for each time step. The residuals in column \( t \) is the residual associated with the data at time \( t \) and the process (\( x \)) transition from \( x(t) \) to \( x(t+1) \). In the MARSS package, the process equation is written \( x(t) = f(x(t-1)) + w(t) \), and the residual in column \( t \) is then \( v(t) w(t+1) \). In many other texts, the process equation is written \( x(t) = f(x(t-1)) + w(t-1) \) and the residual in column \( t \) is denoted \( v(t) w(t) \). Regardless, the meaning is the same; the state residual in column \( t \) is associated with the transition from \( x(t) \) to \( x(t+1) \) not the transition from \( x(t-1) \) to \( x(t) \).

**model.residuals**

The model residuals \( v_t \) are the difference between the data and the predicted data at time \( t \):

\[
v_t = y_t - Zx_t - a
\]

In a state-space model, \( x_t \) is stochastic and the model residuals are a random variable. \( y_t \) is also stochastic, though often observed unlike \( x_t \). The model residual random variable is:

\[
V_t = Y_t - ZX_t - a
\]

The unconditional mean and variance of \( V_t \) is 0 and \( R \). This function (residuals(MLEobj)) returns the conditional mean and variance of \( V_t \).

**model.residuals** is the expected value of \( V_t \) conditioned on the data and parameter set \( \Theta \) (all the parameters including \( Z, a, B, u, R, Q \)).

\[
E(V_t|y_T^T, \Theta) = E(Y_t|y_T^T, \Theta) - ZE(X_t|y_T^T, \Theta) - a
\]
If there are no missing data, this becomes

\[ E(V_t | y_t^T, \Theta) = y_t - Z E(X_t | y_1^T, \Theta) - \alpha \]

\( y_t \) are data at time t and \( E(X_t | y_1^T, \Theta) \) is the Kalman smoother estimate of the states \( x_t \) at time t, i.e. the expected value of the states conditioned on all the data and the parameter set \( \Theta \). Thus \( res1 \) and \( res2 \) in the code below will be the same.

```r
dat = t(harborSeal)[2:3,]
MLEobj = MARSS(dat)
Z = coef(MLEobj, type="matrix")%*%Z
A = coef(MLEobj, type="matrix")%*%A
res1 = dat - Z %*% MLEobj$states - A %*% matrix(1,1,ncol(dat))
res2 = residuals(MLEobj)$model.residuals
```

The model.residuals for the missing data is 0 (in \( res2 \)) because that is the expected value of \( V_t \) when data are missing.

\( \text{var.residuals} \) returned by the function is the conditional variance of the residuals. Rows 1 to n are the conditional variance of the model residuals. This is the variance of \( V_t \) conditioned on the data and the parameter set \( \Theta \). The unconditional variance (no data) would just be \( R \). See Holmes 2014.

If \( \text{Harvey}=\text{TRUE} \), there will be no variance calculation for the missing \( y \). If \( \text{Harvey}=\text{FALSE} \), the variance of the model residuals at the t with missing \( y \) are computed (via Holmes 2014). The interpretation is that although the data are missing, you can still imagine that these data exist and you need the variance of residuals. For example, if you are doing a leave-one-out cross-validation, the data exist and you need their variance because you are going to compute some diagnostics using the left-out data. For outlier diagnostics and shock detection, the variances for the missing values are not needed.

**state.residuals**

The state residuals \( w_t \) are the difference between the state at time t and the expected value of the state at time t given the state at time t-1:

\[ w_t = x_t - Bx_{t-1} - u \]

Like the model residual, the state residual \( w_t \) is a random variable since \( x_t \) is a random variable:

\[ W_t = X_t - BX_{t-1} - u \]

The unconditional mean and variance of \( W_t \) is 0 and \( Q \). \( \text{residuals(MLEobj)} \) returns the conditional mean and variance of \( W_t \).

\( \text{state.residuals} \) is the expected value of \( W_t \) conditioned on the data (all the data 1 to T) and parameter set \( \Theta \).

\[ E(W_t | y_1^T, \Theta) = E(X_t | y_1^T, \Theta) - BE(X_{t-1} | y_1^T, \Theta) - u \]

Thus \( res1 \) and \( res2 \) in the code below will be the same.

```r
dat = t(harborSeal)[2:3,]
TT = ncol(dat)
```
residuals.marssMLE

ML Obj = MARSS(dat)
B = coef(MLEobj, type="matrix")$B
U = coef(MLEobj, type="matrix")$U
state$t = MLEobj$states[,2:TT]
state$t1 = MLEobj$states[,1:(TT-1)]
res1 = state$t - B %*% state$t1 - U %*% matrix(1,1,TT-1)
res2 = residuals(MLEobj)$state.residuals

The state residuals always exist since the expected value of the states exist without data and will be identical with Harvey=TRUE or Harvey=FALSE. Generally speaking, \( E(W_t | y_T^T) \) is not 0 even if there are missing data. Note that the state residual at the last time step (T) will be NA because it is the residual associated with \( x(T) \) to \( x(T+1) \) and \( T+1 \) is beyond the data. Similarly, the variance matrix at the last time step will have NAs for the same reason.

**standardized residuals**

residuals.marssMLE will return the standardized residuals sensu Harvey et al. (1998) for you in std.residuals for outlier and shock detection. These are the model and state residuals scaled by the inverse square root of the missing values corrected variance of the residuals. Note the standardized model residuals are set to NA when there are missing data (if there is no data point, there is no model residual). The standardized state residuals however always exist since the expected value of the states exist without data.

The interpretation of the standardized residuals is not straight-forward when the Q and R variance-covariance matrices are non-diagonal. The residuals which were generated by a non-diagonal variance-covariance matrices are transformed into orthogonal residuals in MVN(0,I) space. For example, if \( v \) is 2x2 correlated errors with variance-covariance matrix \( R \). The transformed residuals (from this function) for the i-th row of \( v \) is a combination of the row 1 effect and the row 1 effect plus the row 2 effect. So in this case, row 2 of the transformed residuals would not be regarded as solely the row 2 residual but rather how different row 2 is from row 1, relative to expected. If the errors are highly correlated, then the transformed residuals can look rather non-intuitive.

**normalized residuals**

If normalize=FALSE, the unconditional variance of \( V_t \) and \( W_t \) are R and Q and the model is assumed to be written as

\[
\begin{align*}
  y_t &= Z x_t + a + v_t \\
  x_t &= B x_{t-1} + u + w_t
\end{align*}
\]

Harvey et al (1998) writes the model as

\[
\begin{align*}
  y_t &= Z x_t + a + Hv_t \\
  x_t &= B x_{t-1} + u + Gw_t
\end{align*}
\]

with the variance of \( V_t \) and \( W_t \) equal to I (identity).

residuals.marssMLE returns the residuals defined as in the first equations. To get the residuals defined as Harvey et al. (1998) define them (second equations), then use normalize=TRUE. In that case the unconditional variance of residuals will be I instead of R and Q. Note, that the ‘normalized’ residuals are not the same as the ‘standardized’ residuals. In former, the unconditional residuals have a variance of I while in the latter it is the conditional residuals that have a variance of I.
Value

A list with the following components

model.residuals
  The smoothed model residuals $E(V(t)|y(1:T),\Theta)$, where $\Theta$ is the set of model parameters. Sometimes called the smoothations. This is different than the Kalman filter innovations which are $E(V(t)|y(1:t-1),\Theta)$.

state.residuals
  The smoothed state residuals $E(X(t)|y(1:T))-E(X(t)|E(x(t-1)|y(1:T)))$.

residuals
  The model residuals as a $(n+m) \times TT$ matrix with `model.residuals` on top and `state.residuals` below. `model.residuals` is $\hat{\eta}_t$ on page 112 of Harvey, Koopman and Penzer (1998).

var.residuals
  The variance of the model residuals and state residuals as a $(n+m) \times (n+m) \times TT$ matrix with the model residuals in rows 1 to n.

std.residuals
  The standardized model residuals as a $(n+m) \times TT$ matrix. This is `residuals` divided by the square root of `var.residuals`—although the matrix equivalent of that equation.

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References

Harvey, A., S. J. Koopman, and J. Penzer. 1998. Messy time series: a unified approach. Advances in Econometrics 13: 103-144 (see page 112-113). Eqn 21 is the Kalman eqns. Eqn 23 and 24 is the backward recursion to compute the smoothations. This function uses the MARSSkf output for eqn 21 and then implements the backwards recursion in eqn 23 and eqn 24. Pages 120-134 discuss the use of standardized residuals for outlier and structural break detection.

de Jong, P. and J. Penzer. 1998. Diagnosing shocks in time series. Journal of the American Statistical Association 93: 796-806. This one shows the same equations; see eqn 6. This paper mentions the scaling based on the inverse of the sqrt (chol) of the variance-covariance matrix for the residuals (model and state together). This is in the right column, half-way down on page 800.


See Also

MARSSkem marssMLE
Examples

```r
dat <- t(harborSeal)
dat <- dat[c(2,11),]
MLEobj <- MARSS(dat)

# state residuals
state.resids1 <- residuals(MLEobj)$state.residuals
# this is the same as
states <- MLEobj$states
Q <- coef(MLEobj,type="matrix")$Q
state.resids2 <- states[2:30]-states[1:29]-matrix(coef(MLEobj,type="matrix")$U,2,29)
# standardize to variance of 1
state.resids2 <- (solve(t(chol(Q))) %*% state.resids2)
# compare the two
cbind(t(state.resids1[-30]),t(state.resids2))

# standardized (by variance) model & state residuals
residuals(MLEobj)$std.residuals
```

Description

Example data set for use in MARSS vignettes for the DLM chapter in the MARSS-package User Guide. This is a 42-year time-series of the logit of juvenile salmon survival along with an index of April coastal upwelling. See the source for details.

Usage

```r
data(SalmonSurvCUI)
```

Format

The data are provided as a matrix with time running down the rows. Column 1 is year, column 2 is the logit of the proportion of juveniles that survive to adulthood, column 3 is an index of the April coastal upwelling index.

Source

Scheuerell, Mark D., and John G. Williams. "Forecasting climate-induced changes in the survival of Snake River spring/summer Chinook salmon (Oncorhynchus tshawytscha)." Fisheries Oceanography 14.6 (2005): 448-457.

Examples

```r
str(SalmonSurvCUI)
```
**Description**

Standardizes Kalman filter innovations. This is a helper function called by `MARSSinnovationsboot` in the `MARSS-package`. Not exported.

**Usage**

```
stdInnov(SIGMA, INNOV)
```

**Arguments**

- `SIGMA`: n x n x T array of Kalman filter innovations variances. This is output from `MARSSkf`.
- `INNOV`: n x T matrix of Kalman filter innovations. This is output from `MARSSkf`.

**Details**

n = number of observation (y) time series. T = number of time steps in the time series.

**Value**

n x T matrix of standardized innovations.

**Author(s)**

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**References**


**See Also**

`MARSSboot` `MARSSkf` `MARSSinnovationsboot`

**Examples**

```r
## Not run:
std.innovations <- stdInnov(kflist$Sigma, kflist$Innov)

## End(Not run)
```
tidy.marssMLE | Return estimated parameters and states with summary information

**Description**

This returns a data.frame with the estimated parameters (or states) of a MARSS model with optionally standard errors and confidence intervals. This assembles information available via the print and coef functions into a data.frame that summarizes the estimates.

If conf.int=TRUE. For parameters, `MARSSparamCIs` will be run to add confidence intervals to the fitted model object if these are not already added. The default CIs are calculated using an analytically computed Hessian matrix. This can be changed by passing in optional arguments for `MARSSparamCIs`. For states, the approximate CIs using the standard deviation of the states is used to compute the confidence intervals ($\text{qnorm}(\alpha/2) \times \text{se.fit} + \text{fitted}$).

If you have a DFA model (form='dfa'), you can pass in rotate=TRUE to return the rotated trends. If you want the rotated loadings, you will need to compute those yourself:

```r
dfa <- MARSS(t(harborSealWA[,1]), model=list(m=2), form='dfa')
Z.est <- coef(dfa, type="matrix")$Z
H.inv <- varimax(coef(dfa, type="matrix")$Z)$rotmat
Z.rot <- Z.est \%\% H.inv
```

The tidy function is compatible with the broom package.

**Usage**

```r
tidy.marssMLE(x, type = c("parameters", "states"),
conf.int = TRUE, conf.level = 0.95,
form=attr(x["model"], "form")[[1], ...)
```

**Arguments**

- `x` | a marssMLE object
- `type` | Estimates for the parameters or for the states.
- `conf.int` | Whether to include a confidence interval.
- `conf.level` | Confidence level if interval is returned.
- `form` | If you want the augment function to use a different form than that specified in `attr(x$model, "form")`. Useful if you have a DFA model that you manually set up, which does not have the form attribute set.
- `...` | Optional arguments. If `conf.int=TRUE`, then arguments to specify how CIs are computed can be passed in. See details and `MARSSparamCIs`. If `form="dfa"`, `rotate=TRUE` can be passed in to rotate the trends (only trends not Z matrix).
utility.functions

Examples

dat <- t(harborSeal)
dat <- dat[c(2,11,12),]
MLEobj <- MARSS(dat, model=list(Z=factor(c("WA","OR","OR"))))

library(broom)
library(ggplot2)

# A data frame of the estimated parameters
tidy(MLEobj)

# Make a plot of the estimated states
# Don't use augment. States are not data.
d <- tidy(MLEobj, type="states")
ggplot(data = d) +
  geom_line(aes(t, estimate)) +
  geom_ribbon(aes(x=t, ymin=conf.low, ymax=conf.high), linetype=2, alpha=0.1) +
  facet_grid(~term) +
  xlab("Time Step") + ylab("Count")

utility.functions  Matrix Utilities

Description

Matrix utilities for MARSS functions in the MARSS-package. These are not exported but can be accessed using the MARSS:: prefix.

Usage

is.blockdiag(x)
is.validvarcov(x, method="kem")
is.identity(x, dim=NULL)
is.diagonal(x, na.rm=FALSE)
is.equaltri(x)
makediag(x, nrow=NA)
takediag(x)
is.design(x, strict=TRUE, dim=NULL, zero.rows.ok=FALSE, zero.cols.ok=FALSE)
is.fixed(x, by.row=FALSE)
is.identity(x, dim=NULL)
is.zero(x)
vec(x)
unvec(x, dim=NULL)
is.wholenumber(x, tol = .Machine$double.eps*0.5)
imat(x)
rwishart(nu, V)
myststrsplit(x)
convert.model.mat(param.matrix)
fixed.free.to.formula(fixed,free,dim)
matrix.power(x, n)
sub3D(x,t=1)
pinv(x)
pcholinv(x)
pchol(x)
is.solvable(A,y=NULL)
all.equal.vector(x)
parmat(MLEobj, elem = c("B", "U", "Q", "Z", "A", "R", "x0", "V0", "G", "H", "L"),
    t = 1, dims = NULL, model.loc = "marss")

Arguments

x, A, y  A matrix (or vector for 'makediag' or string for 'mysrsplit').
na.rm  How to treat NAs in the block diag test.
dim, dims  Matrix dimensions. Some functions will take the vec of a matrix. In this case, the optional dim arg specifies the matrix dimensions.
fixed  A fixed matrix per the MARSS specification for fixed matrix syntax.
free  A free matrix per the MARSS specification for free matrix syntax.
nrow  Number of rows.
tol  Tolerance.
method  kem or BFGS. Used to add extra test for MARSSoptim().
t  The time index or third dimension of a 3D matrix
nu, V  Parameters of a Wishart distribution.
param.matrix  The list matrix version of a time-invariant MARSS model.
n  An integer for the power function.
zero.rows.ok, zero.cols.ok  Means the design matrix can have all zero rows or columns.
strict  Specifies whether the design matrix must be only 0s and 1s.
by.row  For is.fixed, reports whether is.fixed by row rather than for the whole matrix.
MLEobj  A marssMLE object.
elem  The parameter matrix of a marss model to return.
model.loc  Whether to use the marss or model marssMODEL in the marssMLE object.

Details

- is... tests for various matrix properties. isDiagonal() from the Matrix package is used to test numeric matrices for diagonality. is.diagonal() is only used to determine if list matrices (that combine numeric and character values) are diagonal. is.zero tests for near zeroness and give TRUE for is.zero((.5-.3)-(3.-1)) unlike ==0.
- vec(x) creates a column vector from a matrix per the standard vec math function.
- unvec(c,dim) takes the vector c and creates a matrix with the specified dimensions.
- Imat(nrow) returns the identity matrix of dimension nrow.
• fixed.free.to.formula takes a fixed and free pair and constructs a list matrix (or array if time-varying) with formulas in each matrix element.
• convert.model.mat takes a list matrix with formulas in each element and converts to a fixed/free pair.
• sub3D returns a 2D matrix after subsetting a 3D matrix on the third (time) dimension. Ensures that R always returns a matrix.
• mystrsplit is a customized string splitter used by convert.model.mat.
• rwishart generates random draws from a wishart distribution.
• matrix.power is a faster way to get the n-th power of a matrix.
• pinv is the pseudoinverse based on singular value decomposition PInv=UD^+V' where a diagonal matrix with non-zero diagonal values of D (from svd) replaced with 1/D.
• pcholinv is the inverse based on the Cholsky decomposition but modified to allow 0s on the diagonal of x (with corresponding 0 row/column). These appear as 0 row/columns in the returned inverse.
• pchol returns the Cholsky decomposition but modified to allow 0s on the diagonal of x (with corresponding 0 row/column).
• is.solvable returns information on the solvability of the linear system y=Ax using the SVD decomposition.
• all.equal.vector tests if the all the elements in a vector, matrix, or array are all equal. Works on list matrices too.
• parmat constructs the parameter matrix with both the fixed and free values from the vectorized form in a marssMLE object. Users should use coef(MLEobj) (See coef.marssMLE).

Value
See above.

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zscore z-score a vector or matrix

Description
Removes the mean and standardizes the variance to 1.

Usage
zscore(x)
Arguments

\( x \)  
\( n \times T \) matrix of numbers

Details

\( n = \) number of observation (\( y \)) time series. \( T = \) number of time steps in the time series.

The z-scored values (\( z \)) of a matrix of \( y \) values are

\[
z_i = \Sigma^{-1}(y_i - \bar{y})\]

where \( \Sigma \) is a diagonal matrix with the standard deviations of each time series (row) along the diagonal, and \( \bar{y} \) is a vector of the means.

Value

\( n \times T \) matrix of z-scored values.

Author(s)

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Examples

```r
zscore(1:10)  
x <- zscore(matrix(c(rnorm(6),NA),3,10))  
# mean is 0 and variance is 1  
apply(x, 1, mean, na.rm=TRUE)  
apply(x, 1, var, na.rm=TRUE)
```
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