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Analysis of multivariate time-series using the MARSS package

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Disclaimer: E. E. Holmes, E. J. Ward, and M. D. Scheuerell are NOAA scientists employed by the U.S. National Marine Fisheries Service. The views and opinions presented here are solely those of the authors and do not necessarily represent those of our employer.
Preface

The initial motivation for our work with MARSS models was a collaboration with Rich Hinrichsen. Rich developed a framework for analysis of multi-site population count data using MARSS models and bootstrap AICb (Hinrichsen and Holmes, 2009). Our work (EEH and EJW) extended Rich’s framework, made it more general, and led to the development of a parametric bootstrap AICb for MARSS models, which allows one to do model-selection using datasets with missing values (Ward et al., 2010; Holmes and Ward, 2010). Later, we developed additional algorithms for simulation and confidence intervals. Discussions with Mark Scheuerell led to an extensive revision of the EM algorithm and to the development of a general EM algorithm for constrained MARSS models (Holmes, 2012). Discussions with Mark also led to a complete rewrite of the model specification so that the package could be used for MARSS models in general—rather than simply the form of MARSS model used in our applications. Many collaborators have helped test the package; we thank especially Yasmin Lucero, Kevin See, and Brice Semmens. Development of the code into a R package would not have been possible without Kellie Wills, who wrote much of the original package code outside of the algorithm functions. Finally, we thank the participants of our MARSS workshops and courses and the MARSS users who have contacted us regarding issues that were unclear in the manual, errors, or suggestions regarding new applications. Discussions with these users have helped us improve the manual and go in new directions.

The application chapters were developed originally as part of workshops on analysis of multivariate time-series data given at the Ecological Society of America meetings since 2005 and taught by us along with Yasmin Lucero, Stephanie Hampton, and Brice Semmens. The chapter on extinction estimation and trend estimation was initially developed by Brice Semmens and later extended by us for this user guide. The algorithm behind the TMU figure in Chapter 7 was developed during a collaboration with Steve Ellner (Ellner and Holmes, 2008). Later we further developed the chapters as part of a course we taught on analysis of fisheries and environmental time-series data at the University of Washington, where we are affiliate faculty. You can find online versions of the workshops and the time-series analysis course on EEH’s website http://faculty.washington.edu/eeholmes.

The authors are research scientists at the Northwest Fisheries Science Center (NWFSC). This work was conducted as part of our jobs at the NWFSC, a research center for NOAA Fisheries which is a United States federal government agency. A CAMEO grant from the National Science Foundation and NOAA Fisheries provided the initial impetus for the development of the package as part of a research project with Stephanie Hampton, Lindsay Scheef, and Steven Katz on analysis of marine plankton time series. During the initial stages of this work, EJW was supported on a post-doctoral fellowship from the
National Research Council and MDS was partially supported by a PECASE award from the White House Office of Science and Technology Policy.

You are welcome to use the code and adapt it with full attribution. You should use citation \cite{Holmes et al. (2012)} for the MARSS package. It may not be used in any commercial applications nor may it be copyrighted. Use of the EM algorithm should cite \cite{Holmes (2012)}. Links to more code and publications on MARSS applications can be found by following the links at our academic websites:

- \url{http://faculty.washington.edu/eeholmes}
- \url{http://faculty.washington.edu/scheuerl}
- \url{http://faculty.washington.edu/warde}
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Part I

The MARSS package
Overview

MARSS stands for Multivariate Auto-Regressive(1) State-Space. The MARSS package is an R package for estimating the parameters of linear MARSS models with Gaussian errors. This class of model is extremely important in the study of linear stochastic dynamical systems, and these models are important in many different fields, including economics, engineering, genetics, physics and ecology (Appendix D). The model class has different names in different fields, for example in some fields they are termed dynamic linear models (DLMs) or vector autoregressive (VAR) state-space models. The MARSS package allows you to easily fit time-varying constrained and unconstrained MARSS models with or without covariates to multivariate time-series data via maximum-likelihood using primarily an EM algorithm\(^1\).

A full MARSS model, with Gaussian errors, takes the form:

\[
\begin{align*}
    x_t &= B_t x_{t-1} + u_t + C_t c_t + G_t w_t, \text{ where } w_t \sim \text{MVN}(0, Q_t) \quad (1.1a) \\
    y_t &= Z_t x_t + a_t + D_t d_t + H_t v_t, \text{ where } v_t \sim \text{MVN}(0, R_t) \quad (1.1b) \\
    x_1 &\sim \text{MVN}(\pi, \Lambda) \text{ or } x_0 \sim \text{MVN}(\pi, \Lambda) \quad (1.1c)
\end{align*}
\]

The \( x \) equation is termed the state process and the \( y \) equation is termed the observation process. Data enter the model as the \( y \); that is the \( y \) is treated as the data although there may be missing data. The \( c \) and \( d \) are inputs (aka, exogenous variables, covariates or indicator variables). The \( G \) and \( H \) are also typically inputs (fixed values with no missing values).

The bolded terms are matrices with the following definitions:

- \( x \) is a \( m \times T \) matrix of states. Each \( x_t \) is a realization of the random variable \( X_t \) at time \( t \).
- \( w \) is a \( m \times T \) matrix of the process errors. The process errors at time \( t \) are multivariate normal with mean 0 and covariance matrix \( Q_t \).
- \( y \) is a \( n \times T \) matrix of the observations. Some observations may be missing.

\(^1\) Fitting via the BFGS algorithm is also provided using R’s optim function, but this is not the focus of the package.
v is a $n \times T$ column vector of the non-process errors. The observation errors at
time $t$ are multivariate normal with mean 0 and covariance matrix $R_t$.
$B_t$ and $Z_t$ are parameters and are $m \times m$ and $n \times m$ matrices.
$u_t$ and $a_t$ are parameters and are $m \times 1$ and $n \times 1$ column vectors.
$Q_t$ and $R_t$ are parameters and are $g \times g$ (typically $m \times m$) and $h \times h$ (typically $n \times n$) variance-covariance matrices.
$\pi$ is either a parameter or a fixed prior. It is a $m \times 1$ matrix.
$\Lambda$ is either a parameter or a fixed prior. It is a $m \times m$ variance-covariance matrix.
$C_t$ and $D_t$ are parameters and are $m \times p$ and $n \times q$ matrices.
c and d are inputs (no missing values) and are $p \times T$ and $q \times T$ matrices.
$G_t$ and $H_t$ are inputs (no missing values) and are $m \times g$ and $n \times h$ matrices.

1.1 What does the MARSS package do?

Written in an unconstrained form$^2$, a MARSS model can be written out as follows. Two state processes ($x$) and three observation processes ($y$) are used here as an example.

---

$^2$ meaning all the elements in a parameter matrices are allowed to be different and none constrained to be equal or related.
### 1.1 What does the MARSS package do?

The MARSS package fits models via maximum likelihood. The MARSS package allows you to specify constraints by fixing elements in a parameter matrix or specifying that some elements are estimated—and have a linear relationship to other elements. Here is an example of a MARSS model with fixed and estimated parameter elements:

\[
\begin{bmatrix}
    x_1 \\
    x_2
\end{bmatrix}_t = \begin{bmatrix} b_{11} & b_{12} \\
    b_{21} & b_{22}
\end{bmatrix} \begin{bmatrix}
    x_1 \\
    x_2
\end{bmatrix}_{t-1} + \begin{bmatrix} w_1 \\
    w_2
\end{bmatrix}_t, \quad \begin{bmatrix} y_1 \\
    y_2 \\
    y_3
\end{bmatrix}_t = \begin{bmatrix} z_{11} & z_{12} \\
    z_{21} & z_{22} \\
    z_{31} & z_{32}
\end{bmatrix} \begin{bmatrix}
    x_1 \\
    x_2
\end{bmatrix}_t + \begin{bmatrix} v_1 \\
    v_2 \\
    v_3
\end{bmatrix}_t, \quad \begin{bmatrix} x_1 \\
    x_2
\end{bmatrix}_0 \sim \text{MVN}\left( \begin{bmatrix} \pi_1 \\
    \pi_2
\end{bmatrix}, \begin{bmatrix} \pi_{11} & \pi_{12} \\
    \pi_{21} & \pi_{22}
\end{bmatrix} \right)
\]

However, not all parameter elements can be estimated simultaneously. Constraints are required in order to specify a model with a unique solution. The MARSS package allows you to specify constraints by fixing elements in a parameter matrix or specifying that some elements are estimated—and have a linear relationship to other elements. Here is an example of a MARSS model with fixed and estimated parameter elements:

\[
\begin{bmatrix}
    x_1 \\
    x_2
\end{bmatrix}_t = \begin{bmatrix} a & 0 \\
    0 & a
\end{bmatrix} \begin{bmatrix}
    x_1 \\
    x_2
\end{bmatrix}_{t-1} + \begin{bmatrix} w_1 \\
    w_2
\end{bmatrix}_t, \quad \begin{bmatrix} y_1 \\
    y_2 \\
    y_3
\end{bmatrix}_t = \begin{bmatrix} d \\
    c \\
    1 + 2d + 3c + 2 + 3d
\end{bmatrix} \begin{bmatrix}
    x_1 \\
    x_2
\end{bmatrix}_t + \begin{bmatrix} v_1 \\
    v_2 \\
    v_3
\end{bmatrix}_t, \quad \begin{bmatrix} x_1 \\
    x_2
\end{bmatrix}_0 \sim \text{MVN}\left( \begin{bmatrix} \pi_1 \\
    \pi_2
\end{bmatrix}, \begin{bmatrix} 1 & 0 \\
    0 & 1
\end{bmatrix} \right)
\]

Notice that some elements are fixed (in this case to 0, but could be any fixed number), some elements are shared (have the same value), and some elements are linear combinations of other estimated values (\(c, 1 + 2d + 3c\) and \(2 + 3d\) are linear combinations of \(c\) and \(d\)).

The MARSS package is unusual among packages for fitting MARSS models in that fitting is performed via a constrained EM algorithm (Holmes 2012), based on a vectorized form of Equation 1.1 (See Chapter 3 for the vectorized form used in the algorithm). Although fitting via the BFGS algorithm is also provided using `method="BFGS"` and the `optim` function in \(\text{R}\), the examples in this guide use the EM algorithm primarily because it gives robust estimation for datasets replete with missing values and for high-dimensional models with various constraints. However, there are many models/datasets where BFGS is faster and we typically try both for problems. The EM algorithm is also often used to provide initial conditions for the BFGS algorithm (or an MCMC routine) in order to improve the performance of those algorithms. In addition to the main model
fitting function, the MARSS package supplies functions for bootstrap and
approximate confidence intervals, parametric and non-parametric bootstrapping, model selection (AIC and bootstrap AIC), simulation, and bootstrap
bias correction.

1.2 What does MARSS output and how do I get the output?

MARSS models are used in many different ways and different users will want
different types of output. Some users will want the parameter estimates while
others want the smoothed states and others want to use MARSS to interpolate
missing values and want the expected values of missing data.

The best way to find out how to get output is to type \texttt{?print.MARSS}
at the command line after installing MARSS. The print help page discusses
how to get parameter estimates in different forms, the smoothed and filtered
states, all the Kalman filter and smoother output, all the expectations of \( y \)
(missing data), confidence intervals and bias estimates for the parameters,
and standard errors of the states. If you are looking only for Kalman filter
and smoother output, see the relevant section in Chapter \ref{ch:filter} and see the help
page for the \texttt{MARSSkf} function (type \texttt{?MARSSkf} at the R command line).

You might also want to look at the \texttt{augment}, \texttt{tidy} and \texttt{glance} functions
which will summarize commonly needed output from a MARSS model fit.
Type \texttt{?augment.marssMLE} at the command line to see examples. These func-
tions work as they do in the \texttt{broom} R package.

1.3 How to get started (quickly)

If you already work with models in the form of Equation \ref{eq:SSE}
you can immediately fit your model with the MARSS package. Install the MARSS package
and then type \texttt{library(MARSS)} at the command line to load the package.
Look at the Quick Start Guide and then skim through Chapter \ref{ch:example}
Appendix \ref{app:example} also has many examples of how to specify different forms for your parameter
matrices.

1.4 Getting your data in right format

Your data need to be a matrix (not dataframe nor a ts object) with time across
the columns \((n \times T)\) matrix. The MARSS functions assume discrete time steps
and you will need a column for each time step. Replace any missing time steps
with NA. Write your model down on paper and identify which parameters
correspond to which parameter matrices in Equation \ref{eq:SSE} Call the \texttt{MARSS()}
function (Chapter \ref{ch:example}) using your data and using the \texttt{model} argument to specify
the structure of each parameter.
1.4.1 Getting a ts object into the right form

A R ts object (time series object) stores information about the time steps of the data and often seasonal information (the quarter or month). MARSS needs this information in matrix form. If you have your data in ts form, then you are probably using year and season (quarter, month) as covariates to estimate trend and seasonality. Here is how to get your ts into the form that MARSS wants.

*Univariate example.* This converts a univariate ts object with year and quarter into a matrix with a row for the response (here called Temp), year, and quarter.

\[ z = \text{ts}(\text{rnorm}(10), \text{frequency} = 4, \text{start} = \text{c}(1959, 2)) \]
\[ \text{dat} = \text{data.frame(Yr = floor(time(z) + .Machine$double.eps),} \]
\[ \quad \text{Qtr = cycle(z), Temp = z)} \]
\[ \text{dat} = t(\text{dat}) \]

When you call MARSS, \text{dat}["Temp",] is the data. \text{dat}[c("Yr","Qtr"),] are your covariates.

*Multivariate example.* In this example, we have two temperature readings and a salinity reading. The data are monthly.

\[ z <- \text{ts}(\text{matrix}(\text{rnorm}(300), 100, 3), \text{start} = \text{c}(1961, 1), \]
\[ \quad \text{frequency} = 12, \text{names= c("Temp1","Temp2","Sal")}) \]
\[ \text{dat} = \text{data.frame(Yr = floor(time(z) + .Machine$double.eps),} \]
\[ \quad \text{Month = cycle(z), z)} \]

When you call MARSS, \text{dat}[c("Temp1","Temp2"),] are the data and \text{dat}[c("Yr","Month","Sal"),] are your covariates.

See the chapters that discuss seasonality for examples of how to model seasonality. The brute force method of treating month or quarter as a factor requires estimation of more parameters than necessary in many cases.

1.5 Important notes about the algorithms

*Specification of a properly constrained model with a unique solution is the responsibility of the user because MARSS has no way to tell if you have specified an insufficiently constrained model—with correspondingly an infinite number of solutions.*

Specifying a properly constrained model with a unique solution is imperative. How do you know if the model is properly constrained? If you are using a MARSS model form that is widely used, then you can probably assume that it is properly constrained. If you go to papers where someone developed the model or method, the issue of constraints necessary to ensure “identifiability” will likely be addressed if it is an issue. Are you fitting novel MARSS models? Then you will need to do some study on identifiability in this class.
of models using textbooks (Appendix D). Often textbooks do not address identifiability explicitly. Rather it is addressed implicitly by only showing a model constructed in such a way that it is identifiable. In our work, if we suspect identification problems, we will often first do a Bayesian analysis with flat priors and look for oddities in the posteriors, such as ridges, plateaus or bimodality.

All the EM code in the MARSS package is currently in native R. Thus the model fitting is slow. The classic Kalman filter/smooth algorithm, as shown in Shumway and Stoffer (2006, p. 331-335), is based on the original smoother presented in Rauch (1963). This Kalman filter is provided in function MARSSkfas, but the default Kalman filter and smoother used in the MARSS package is based on the algorithm in Kohn and Ansley (1989) and papers by Koopman et al. This Kalman filter and smoother is provided in the KFAS package (Helske 2012). Table 2 in Kohn (1993) indicates that the classic algorithm is 40-100 times slower than the algorithm given in Kohn and Ansley (1989), Kohn (1993), and Koopman et al. (1999). The MARSS package function MARSSkfas provides a translator between the model objects in MARSS and those in KFAS so that the KFAS functions can be used. MARSSkfas also includes a lag-one covariance smoother algorithm as this is not output by the KFAS functions, and it provides proper formulation of the priors so that one can use the KFAS functions when the prior on the states is set at \( t = 0 \) instead of \( t = 1 \). Simply off-setting your data to start at \( t=2 \) and sending that value to \( t_{\text{init}} = 1 \) in the KFAS Kalman filter would not be mathematically correct!

EM algorithms will quickly get in the vicinity of the maximum likelihood, but the final approach to the maximum is generally slow relative to quasi-Newton methods. On the flip side, EM algorithms are quite robust to initial conditions choices and can be extremely fast at getting close to the MLE values for high-dimensional models. The MARSS package also allows one to use the BFGS method to fit MARSS models, thus one can use an EM algorithm to “get close” and then the BFGS algorithm to polish off the estimate. Restricted maximum-likelihood algorithms are also available for AR(1) state-space models, both univariate (Staples et al. 2004) and multivariate (Hinchsen and Holmes 2009). REML can give parameter estimates with lower variance than plain maximum-likelihood algorithms. However, the algorithms for REML when there are missing values are not currently available (although that will probably change in the near future). Another maximum-likelihood method is data-cloning which adapts MCMC algorithms used in Bayesian analysis for maximum-likelihood estimation (Lele et al. 2007).

Missing values are seamlessly accommodated with the MARSS package. Simply specify missing data with NAs. The likelihood computations are exact and will deal appropriately with missing values. However, no innovations\(^3\) referring to the non-parametric bootstrap developed by Stoffer and Wall (1991).
1.5 Important notes about the algorithms

Bootstrapping can be done if there are missing values. Instead parametric bootstrapping must be used.

You should be aware that maximum-likelihood estimates of variance in MARSS models are fundamentally biased, regardless of the algorithm used. This bias is more severe when one or the other of $R$ or $Q$ is very small, and the bias does not go to zero as sample size goes to infinity. The bias arises because variance is constrained to be positive. Thus if $R$ or $Q$ is essentially zero, the mean estimate will not be zero and thus the estimate will be biased high while the corresponding bias of the other variance will be biased low. You can generate unbiased variance estimates using a bootstrap estimate of the bias. The function `MARSSparamCIs()` will do this. However be aware that adding an estimated bias to a parameter estimate will lead to an increase in the variance of your parameter estimate. The amount of variance added will depend on sample size.

You should also be aware that mis-specification of the prior on the initial states ($\pi$ and $\Lambda$) can have catastrophic effects on your parameter estimates if your prior conflicts with the distribution of the initial states implied by the MARSS model. These effects can be very difficult to detect because the model will appear to be well-fitted. Unless you have a good idea of what the parameters should be, you might not realize that your prior conflicts.

The most common problems we have found with priors on $x_0$ are the following. Problem 1) The correlation structure in $\Lambda$ (whether the prior is diffuse or not) does not match the correlation structure in $x_0$ implied by your model. For example, you specify a diagonal $\Lambda$ (independent states), but the implied distribution has correlations. Problem 2) The correlation structure in $\Lambda$ does not match the structure in $x_0$ implied by constraints you placed on $\pi$. For example, you specify that all values in $\pi$ are shared, yet you specify that $\Lambda$ is diagonal (independent).

Unfortunately, using a diffuse prior does not help with these two problems because the diffuse prior still has a correlation structure and can still conflict with the implied correlation in $x_0$. One way to get around these problems is to set $\Lambda=0$ (a $m \times m$ matrix of zeros) and estimate $\pi \equiv x_0$ only. Now $\pi$ is a fixed but unknown (estimated) parameter, not the mean of a distribution. In this case, $\Lambda$ does not exist in your model and there is no conflict with the model. Be aware however that estimating $\pi$ as a parameter is not always robust. If you specify that $\Lambda=0$ and specify that $\pi$ corresponds to $x_0$, but your model “explodes” when run backwards in time, you cannot estimate $\pi$ because you cannot get a good estimate of $x_0$. Sometimes this can be avoided by specifying that $\pi$ corresponds to $x_1$ so that it can be constrained by the data $y_1$.

In summary, if the implied correlation structure of your initial states is independent (diagonal variance-covariance matrix), you should generally be ok with a diagonal and high variance prior or with treating the initial states as parameters (with $\Lambda = 0$). But if your initial states have an implied correlation structure that is not independent, then proceed with caution. ‘With caution’
means that you should assume you have problems and test how your model fits with simulated data.

There is a large class of models in the statistical finance literature that have the form

\[ x_{t+1} = Bx_t + \Gamma \eta_t \]
\[ y_t = Zx_t + \eta_t \]

For example, ARMA(p,q) models can be written in this form. The MARSS model framework in this package will not allow you to write models in that form. You can put the \( \eta_t \) into the \( x_t \) vector and set \( R = 0 \) to make models of this form using the MARSS form, but the EM algorithm in the MARSS package won’t let you estimate parameters because the parameters will drop out of the full likelihood being maximized in the algorithm. You can try using BFGS by passing in the method argument to the MARSS() call.

1.6 Troubleshooting

Numerical errors due to ill-conditioned matrices are not uncommon when fitting MARSS models. The Kalman and EM algorithms need inverses of matrices. If those matrices become ill-conditioned, for example all elements are close to the same value, then the algorithm becomes unstable. Warning messages will be printed if the algorithms are becoming unstable and you can set `control$trace=1`, to see details of where the algorithm is becoming unstable. Whenever possible, you should avoid using shared \( \pi \) values in your model. The way our algorithm deals with \( \Lambda \) tends to make this case unstable, especially if \( R \) is not diagonal. In general, estimation of a non-diagonal \( R \) is more difficult, more prone to ill-conditioning, and more data-hungry.

You may also see non-convergence warnings, especially if your MLE model turns out to be degenerate. This means that one of the elements on the diagonal of your Q or R matrix are going to zero (are degenerate). It will take the EM algorithm forever to get to zero. BFGS will have the same problem, although it will often get a bit closer to the degenerate solution. If you are using `method="kem"`, MARSS will warn you if it looks like the solution is degenerate. If you use `control=list(allow.degen=TRUE)`, the EM algorithm will attempt to set the degenerate variances to zero (instead of trying to get to zero using an infinite number of iterations). However, if one of the variances is going to zero, first think about why this is happening. This is typically caused by one of three problems: 1) you made a mistake in inputting your data, e.g. used -99 as the missing value in your data but did not replace these with NAs before passing to MARSS, 2) your data are not sufficient to estimate multiple variances or 3) your data are inconsistent with the model you are trying fit.

The algorithms in the MARSS package are designed for cases where the Q and R diagonals are all non-minuscule. For example, the EM update equation

\[ \pi = \begin{bmatrix} a_1 \\ \vdots \\ a_n \end{bmatrix} \]
for u will grind to a halt (not update u) if Q is tiny (like 1E-7). Conversely, the BFGS equations are likely to miss the maximum-likelihood when R is tiny because then the likelihood surface becomes hyper-sensitive to π. The solution is to use the degenerate likelihood function for the likelihood calculation and the EM update equations. MARSS will implement this automatically when Q or R diagonal elements are set to zero and will try setting Q and R terms to zero automatically if control$allow.degen=TRUE.

One odd case can occur when R goes to zero (a matrix of zeros), but you are estimating π. If model$tinitx=1, then π=x₀ and y₁−Zx₀ can go to 0 as well as var(y₁−Zx₀) by driving R to zero. But as this happens, the log-likelihood associated with y₁ will go (correctly) to infinity and thus the log-likelihood goes to infinity. But if you set R = 0, the log-likelihood will be finite. The reason is that R ≈ 0 and R = 0 specify different likelihoods associated with y₁−Zx₀. With R = 0, y₁−Zx₀ does not have a distribution; it is just a fixed value. So there is no likelihood to go to infinity. If some elements of the diagonal of R are going to zero, you should be suspect of the parameter estimates. Sometimes the structure of your data, e.g. one data value followed by a long string of missing values, is causing an odd spike in the likelihood at R ≈ 0. Try manually setting R equal to zero to get the correct log-likelihood\(^5\).

1.7 Other related packages

Packages that will do Kalman filtering and smoothing are many, but packages that estimate the parameters in a MARSS model, especially constrained MARSS models, are much less common. The following are those with which we are familiar, however there are certainly more packages for estimating MARSS models in engineering and economics of which we are unfamiliar. The MARSS package is unusual in that it uses an EM algorithm for maximizing the likelihood as opposed to a Newton-esque method (e.g. BFGS). The package is also unusual in that it allows you to specify the initial conditions at t = 0 or t = 1, allows degenerate models (with some of the diagonal elements of R or Q equal to zero). Lastly, model specification in the MARSS package has a one-to-one relationship between the model list in MARSS and the model as you would write it on paper as a matrix equation. This makes the learning curve a bit less steep. However, the MARSS package has not been optimized for speed and probably will be really slow if you have time-series data with a lot of time points.

\(^5\) The likelihood returned when R ≈ 0 is not incorrect. It is just not the likelihood that you probably want. You want the likelihood where the R term is dropped because it is zero.
Our book from our time-series class has example applications from Applied Time-Series Analysis for Fisheries and Environmental Sciences.

The stats package (part of base R) has functions for fitting univariate structural time series models (MARSS models with a univariate $y$). Read the help file at ?StructTS. The Kalman filter and smoother functions are described here: ?KalmanLike.

DLM is an R package for fitting MARSS models. Our impression is that it is mainly Bayesian focused but does allow MLE estimation via the optim() function. It has a book, Dynamic Linear Models with R by Petris et al., which has many examples of how to write MARSS models for different applications.

sspir is an R package for fitting ARSS (univariate) models with Gaussian, Poisson and binomial error distributions.

dse is an R package for multivariate Gaussian state-space models with a focus on ARMA models.

SsfPack is a package for Ox/Splus that fits constrained multivariate Gaussian state-space models using mainly (it seems) the BFGS algorithm but the newer versions support other types of maximization. SsfPack is very flexible and written in C to be fast. It has been used extensively on statistical finance problems and is optimized for dealing with large (financial) data sets. It is used and documented in Time Series Analysis by State Space Methods by Durbin and Koopman, An Introduction to State Space Time Series Analysis by Commandeur and Koopman, and Statistical Algorithms for Models in State Space Form: SsfPack 3.0, by Koopman, Shephard, and Doornik.

Brodgar The Brodgar software was developed by Alain Zuur to do (among many other things) dynamic factor analysis, which involves a special type of MARSS model. The methods and many example analyses are given in Analyzing Ecological Data by Zuur, Ieno and Smith. This is the one package that we are aware of that also uses an EM algorithm for parameter estimation.

eViews is a commercial economics software that will estimate at least some types of MARSS models.

KFAS is an R package that provides a fast Kalman filter and smoother. Examples in the package show how to estimate MARSS models using the KFAS functions and R's optim() function. The MARSS package uses the filter and smoother functions from the KFAS package.

S+FinMetrics is a S-plus module for fitting MAR models, which are called vector autoregressive (VAR) models in the economics and finance literature. It has some support for state-space VAR models, though we haven't used it so are not sure which parameters it allows you to estimate. It was developed by Andrew Bruce, Doug Martin, Jiahui Wang, and Eric Zivot, and it has a book associated with it: Modeling Financial Time Series with S-plus by Eric Zivot and Jiahui Wang.
1.7 Other related packages

kftrack The R package provides a suite of functions specialized for fitting MARSS models to animal tracking data.
The main package functions

The MARSS package is object-based. It has two main types of objects: a model object (class marssMODEL) and a maximum-likelihood fitted model object (class marssMLE). A marssMODEL object specifies the structure of the model to be fitted. It is an R code version of the MARSS equation (Equation 1.1). A marssMLE object specifies both the model and the information necessary for fitting (initial conditions, controls, method). If the model has been fitted, the marssMLE object will also have the parameter estimates and (optionally) confidence intervals and bias.

2.1 The MARSS() function: inputs

The function MARSS() is an interface to the core fitting functions in the MARSS package. It allows a user to fit a MARSS model using a list to describe the model structure. It returns marssMODEL and marssMLE objects which the user can later use in other functions, e.g. simulating or computing bootstrap confidence intervals.

MLEobj=MARSS(data, model=list(), ..., fit=TRUE) This function will fit a MARSS model to the data using a model list which is a list describing the structure of the model parameter matrices. In the default model, i.e. if you use MARSS(dat) with no model argument, Z and B are the identity matrix, R is a diagonal matrix with one variance, Q is a diagonal matrix with unique variances, u is unique, a is scaling, and C, c, D, and d are all zero. The output is a marssMLE object where the estimated parameter matrices are in MLEobj$par. If fit=FALSE, it returns a minimal marssMLE object that is ready for passing to a fitting function (below) but with no par element.
2.2 The \texttt{MARSS()} function: outputs

The marssMLE object returned by a \texttt{MARSS()} call includes the estimated parameters, states, and expected values of any missing data. Derived statistics, such as confidence intervals and standard errors, can be computed using the functions described below. The \texttt{print} method for marssMLE objects will print or compute all the frequently needed output using the \texttt{what=} argument in the print call. Type \texttt{?print.MARSS} at the R command line to see the print help file.

- **estimated parameters** \texttt{coef(MLEobj)} The \texttt{coef} function can output parameters in a variety of formats. See \texttt{?coef.marssMLE}.
- **residuals** \texttt{residuals(MLEobj)} See \texttt{?residuals.marssMLE} for a discussion of standardized residuals in the context of MARSS models.
- **Kalman filter and smoother output** The smoothed states are in \texttt{MLEobj$states} but the full Kalman filter and smoother output is available from \texttt{MARSSkf(MLEobj)}. See \texttt{?MARSSkf} for a discussion of the Kalman filter and smoother outputs. If you just want the estimated states conditioned on all the data, use \texttt{print(MLEobj, what="xtT")}. If you want all the Kalman filter and smoother output, use \texttt{print(MLEobj, what="kfs")}.
- **expected value of missing y** \texttt{MARSShatyt(MLEobj)} See \texttt{?MARSShatyt} for a discussion of the expectations involving y. If you just want the estimated missing y conditioned on all the data, use \texttt{print(MLEobj, what="ytT")}. If you want all the expectations involving y conditioned on the data, use \texttt{print(MLEobj, what="Ey")}.
- **log-likelihood** \texttt{logLik(MLEobj)} The \texttt{logLik} function returns the log-likelihood.
- **AIC** \texttt{AIC(MLEobj)} The \texttt{AIC} function (\texttt{stats} package) returns the uncorrected AIC. Use \texttt{MLEobj$AICc} to return the small sample size corrected AIC.

2.3 Core functions for fitting a MARSS model

The following core functions are designed to work with ‘unfitted’ marssMLE objects, that is a marssMLE object without the \texttt{par} element. Users do not normally need to call the \texttt{MARSSkem} or \texttt{MARSSoptim} functions since \texttt{MARSS()} will call those. Note, these functions can be called with a marssMLE object with a \texttt{par} element, but these functions will overwrite that element.

\texttt{MLEobj=MARSSkem(MLEobj)} This will fit a MARSS model via the EM algorithm to the data using a properly specified marssMLE object, which has data, the marssMODEL object and the necessary initial condition and control elements. See the appendix on the object structures in the MARSS package. \texttt{MARSSkem} does no error-checking. See \texttt{is.marssMLE()}. \texttt{MARSSkem} uses \texttt{MARSSkf} described below.
MLEobj=MARSSoptim(MLEobj) This will fit a MARSS model via the BFGS algorithm provided in optim(). This requires a properly specified marssMLE object, such as would be passed to MARSSkem.

is.marssMLE(MLEobj) This will check that a marssMLE object is properly specified and ready for fitting. This should be called before MARSSkem or MARSSoptim is called. This function is not typically needed if using MARSS() since MARSS() builds the model object for the user and does error-checking on model structure.

2.4 Functions for a fitted marssMLE object

The following functions use a marssMLE object that has a populated par element, i.e. a marssMLE object returned from one of the fitting functions (MARSS, MARSSkem, MARSSoptim). Below MODELobj means the argument is a marssMODEL object and MLEobj means the argument is a marssMLE object. Type ?function.name to see information on function usage and examples.

standard functions The standard R functions for fitted objects are provided: residuals(), fitted(), logLik(), AIC(), and coef().

summary functions Standard functions for printing output are also available: summary() and print() along with ‘tidyverse’ output using the broom R package and augment(), tidy(), and glance().

In addition, the following are special functions for MARSS fitted models:

kf=MARSSkf(MLEobj) This will compute the expected values of the hidden states given data via the Kalman filter (to produce estimates conditioned on $1:t−1$) and the Kalman smoother (to produce estimates conditioned on $1:T$). The function also returns the exact likelihood of the data conditioned on MLEobj$par. A variety of other Kalman filter/smoother information is also output (kf is a list of output); see ?MARSSkf for details.

MLEobj=MARSSaic(MLEobj) This adds model selection criteria, AIC, AICc, and AICb, to a marssMLE object. Note, AIC and AICc are added to marssMLE objects by the MARSS() function but AICb is not.

boot=MARSSboot(MLEobj) This returns a list containing bootstrapped parameters and data via parametric or innovations bootstrapping.

MLEobj=MARSShessian(MLEobj) This adds a numerically estimated Hessian matrix to a marssMLE object.

MLEobj=MARSSparamCIs(MLEobj) This adds standard errors, confidence intervals, and bootstrap estimated bias for the maximum-likelihood parameters using bootstrapping or the Hessian to the passed-in marssMLE object.

sim.data=MARSSsimulate(MLEobj) This returns simulated data from a MARSS model specified via a list of parameter matrices in MLEobj$parList (this is a list with elements Q, R, U, etc).
paramVec=MARSSvectorizeparam(MLEobj) This returns the estimated (and only the estimated) parameters as a vector. This is useful for storing the results of simulations and for writing functions that fit MARSS models using R’s optim function. print(MLEobj, what="paramvector") will return the same vector.

new.MLEobj=MARSSvectorizeparam(MLEobj, paramVec) This will return a marssMLE object in which the estimated parameters (which are in MLEobj$par along with the fixed values) are replaced with the values in paramVec.

2.5 Functions for marssMODEL objects

is.marssMODEL(MODELobj) This will check that the free and fixed matrices in a marssMODEL object are properly specified. This function is not typically needed if using MARSS() since MARSS() builds the marssMODEL object for the user and does error-checking on model structure.

summary(MODELobj) This will print the model parameter matrices showing the fixed values (in parentheses) and the location of the estimated elements. The estimated elements are shown as g1, g2, g3, ... which indicates which elements are shared (i.e., forced to have the same value). For example, an i.i.d. \( \mathbf{R} \) matrix would appear as a diagonal matrix with just g1 on the diagonal.
Algorithms used in the MARSS package

3.1 The full time-varying model used in the MARSS EM algorithm

In mathematical form, the model that is being fit with the package is

\[
\begin{align*}
\mathbf{x}_t &= (\mathbf{x}_{t-1} \otimes \mathbf{I}_m) \text{vec}(\mathbf{B}_t) + (\mathbf{u}_t \otimes \mathbf{I}_m) \text{vec}(\mathbf{U}_t) + \mathbf{w}_t, \quad \mathbf{W}_t \sim \text{MVN}(0, \mathbf{Q}_t) \\
\mathbf{y}_t &= (\mathbf{x}_t \otimes \mathbf{I}_n) \text{vec}(\mathbf{Z}_t) + (\mathbf{a}_t \otimes \mathbf{I}_n) \text{vec}(\mathbf{A}_t) + \mathbf{v}_t, \quad \mathbf{V}_t \sim \text{MVN}(0, \mathbf{R}_t) \\
\mathbf{x}_{00} &= \pi + \mathbf{F} \triangleq \mathbf{L}, \quad \mathbf{L} \sim \text{MVN}(0, \mathbf{\Lambda})
\end{align*}
\] (3.1)

Each model parameter matrix, \(\mathbf{B}_t, \mathbf{U}_t, \mathbf{Q}_t, \mathbf{Z}_t, \mathbf{A}_t,\) and \(\mathbf{R}_t,\) is written as a time-varying linear model, \(\mathbf{f}_t + \mathbf{D}_t \mathbf{m},\) where \(\mathbf{f}\) and \(\mathbf{D}\) are fully-known (not estimated and no missing values) and \(\mathbf{m}\) is a column vector of the estimates elements of the parameter matrix:

\[
\begin{align*}
\text{vec}(\mathbf{B}_t) &= \mathbf{f}_{t,\mathbf{B}} + \mathbf{D}_{t,\mathbf{B}} \mathbf{\beta} \\
\text{vec}(\mathbf{U}_t) &= \mathbf{f}_{t,\mathbf{U}} + \mathbf{D}_{t,\mathbf{U}} \mathbf{\nu} \\
\text{vec}(\mathbf{Q}_t) &= \mathbf{f}_{t,\mathbf{Q}} + \mathbf{D}_{t,\mathbf{Q}} \mathbf{\eta} \\
\text{vec}(\mathbf{Z}_t) &= \mathbf{f}_{t,\mathbf{Z}} + \mathbf{D}_{t,\mathbf{Z}} \mathbf{\zeta} \\
\text{vec}(\mathbf{A}_t) &= \mathbf{f}_{t,\mathbf{A}} + \mathbf{D}_{t,\mathbf{A}} \mathbf{\alpha} \\
\text{vec}(\mathbf{R}_t) &= \mathbf{f}_{t,\mathbf{R}} + \mathbf{D}_{t,\mathbf{R}} \mathbf{\rho} \\
\text{vec}(\mathbf{\Lambda}) &= \mathbf{f}_\mathbf{L} + \mathbf{D}_\mathbf{L} \mathbf{\lambda} \\
\text{vec}(\mathbf{\pi}) &= \mathbf{f}_\mathbf{\pi} + \mathbf{D}_\mathbf{\pi} \mathbf{p}
\end{align*}
\]

The internal MARSS model specification (element \$\text{marss}\$ in a fitted marssMLE object output by a MARSS() call) is a list with the \(\mathbf{f}\) ("fixed") and \(\mathbf{D}\) ("free") matrices for each parameter. The output from fitting are the vectors, \(\mathbf{\beta}, \mathbf{\nu},\) etc. The trick is to rewrite the user’s linear multivariate problem into the general form (Equation 3.1). MARSS does this using functions that take more familiar arguments as input and then constructs the \(\mathbf{f}\) and \(\mathbf{D}\) matrices. Because the \(\mathbf{f}\) and \(\mathbf{D}\) can be whatever the user wants (assuming they are the right shape), this allows users to include covariates, trends (linear, sinusoidal, etc) or indicator variables in a variety of ways. It also means that terms like \(1 + b + 2c\) can appear in the parameter matrices.

Although the above form looks unusual, it is equivalent to the commonly seen form but leads to a log-likelihood function where all terms have form \(\mathbf{Mm},\)
where $M$ is a matrix and $m$ is a column vector of only the different estimated values. This makes it easy to do the partial differentiation with respect to $m$ necessary for the EM algorithm and as a result, easy to impose linear constraints and structure on the elements in a parameter matrix (Holmes, 2012).

### 3.2 Maximum-likelihood parameter estimation

#### 3.2.1 EM algorithm

Function `MARSSkem` in the MARSS package provides a maximum-likelihood algorithm for parameter estimation based on an Expectation-Maximization (EM) algorithm (Holmes, 2012). EM algorithms are widely used algorithms that extend maximum-likelihood estimation to cases where there are hidden random variables in a model (Dempster et al., 1977; Harvey, 1989; Harvey and Shephard, 1993; McLachlan and Krishnan, 2008). Expectation-Maximization algorithms for unconstrained MARSS models have been around for many years and algorithms for certain constrained cases have also been published. What makes the EM algorithm in MARSS different is that it is a general constrained algorithm that allows generic linear constraints among matrix elements (thus allows fixed, shared and linear combinations of estimated elements).

The EM algorithm finds the maximum-likelihood estimates of the parameters in a MARSS model using an iterative process. Starting with an initial set of parameters $\hat{\Theta}_1$, an updated parameter set $\hat{\Theta}_2$ is obtained by finding the $\hat{\Theta}_2$ that maximizes the expected value of the likelihood over the distribution of the states ($X$) conditioned on $\hat{\Theta}_1$. This distribution of states is computed via the Kalman smoother (Section 3.3). Mathematically, each iteration of an EM algorithm does this maximization:

$$\hat{\Theta}_2 = \arg\max_\Theta E_{X|\hat{\Theta}_1} [\log L(\Theta | Y = y^T_1, X)]$$  \hspace{1cm} (3.2)

Then using $\hat{\Theta}_2$, the distribution of $X$ conditioned on $\hat{\Theta}_2$ is computed. Then that distribution along with $\hat{\Theta}_2$ in place of $\hat{\Theta}_1$ is used in Equation (3.2) to produce an updated parameter set $\hat{\Theta}_3$. This is repeated until the expected log-likelihood stops increasing (or increases less than some set tolerance level).

Implementing this algorithm is straight-forward, hence its popularity.

1. Set an initial set of parameters, $\hat{\Theta}_1$.
2. E step: using the model for the hidden states ($X$) and $\hat{\Theta}_1$, calculate the expected values of $X$ conditioned on all the data $y^T_1$; this is $xtT$ output by the Kalman smoother (function `MARSSk` in MARSS).

---

1 You can choose these however you wish, however choosing something not too far off from the correct values will make the algorithm go faster.
expected values of any functions of $X$ (or $Y$ if there are missing $Y$ values) that appear in your expected log-likelihood function.

3. **M step:** put those $E(X|Y = y_1^T, \hat{\Theta}_1)$ and $E(g(X)|Y = y_1^T, \hat{\Theta}_1)$ into your expected log-likelihood function in place of $X$ (and $g(X)$) and maximize with respect to $\Theta$. This gives you $\hat{\Theta}_2$.

4. Repeat the E and M steps until the log likelihood stops increasing.

The EM equations used in the MARSS package (function `MARSSkem`) are described in Holmes (2012) and are extensions of those in Shumway and Stoffer (1982) and Ghahramani and Hinton (1996). Our EM algorithm is an extended version because our algorithm is for cases where there are constraints within the parameter matrices (shared values, linear combinations, diagonal structure, block-diagonal structure, ...), where there are fixed values within the parameter matrices, or where there may be 0s on the diagonal of $Q$, $R$ and $\Lambda$.

The EM algorithm is a hill-climbing algorithm and like all hill-climbing algorithms can get stuck on local maxima. See chapter 4 for a discussion on how to implement a Monte-Carlo initial conditions search based on Biernacki et al. (2003) to minimize this problem. EM algorithms are also known to get close to the maximum very quickly but then creep toward the absolute maximum. Once in the vicinity of the maximum, quasi-Newton methods find the absolute maximum much faster, but they can be sensitive to initial conditions and in practice, we have found the EM algorithm to be much faster for large problems.

### 3.3 Kalman filter and smoother

The Kalman filter (Kalman, 1960) is a recursive algorithm that solves for the expected value of the hidden states (the $X$) in a MARSS model (Equation 1.1) at time $t$ conditioned on the data up to time $t$: $E(X|y_1^T)$. The Kalman filter gives the optimal (lowest mean square error) estimate of the unobserved $x$, based on the observed data up to time $t$ for this class of linear dynamical system. The Kalman smoother (Rauch et al., 1965) solves for the expected value of the hidden state(s) conditioned on all the data: $E(X|y_1^T)$. If the errors in the stochastic process are Gaussian, then the estimators from the Kalman filter and smoother are also the maximum-likelihood estimates.

However, even if the the errors are not Gaussian, the estimators are optimal in the sense that they are estimators with the least variability possible. This robustness is one reason the Kalman filter is so powerful—it provides well-behaving estimates of the hidden states for all kinds of multivariate autoregressive processes, not just Gaussian processes. The Kalman filter and smoother are widely used in time-series analysis, and there are many textbooks covering it and its applications. In the interest of giving the reader a single point of reference, we use Shumway and Stoffer (2006) as our primary reference.
The MARSSkf function provides the Kalman filter and smoother output using one of two algorithms (specified by fun.kf). The algorithm in MARSSkfss is that shown in Shumway and Stoffer (2006). This algorithm is not computationally efficient; see Koopman et al. (1999) sec. 4.3) for a more efficient Kalman filter implementation. The Koopman et al. implementation is provided in the functions MARSSkfas using the KFAS R package.

MARSSkfss (and MARSSkfas with a few exceptions) has the following outputs:

- \( xtt1 \) The expected value of \( X_t \) conditioned on the data up to time \( t-1 \).
- \( xtt \) The expected value of \( X_t \) conditioned on the data up to time \( t \).
- \( xttT \) The expected value of \( X_t \) conditioned on all the data from time 1 to \( T \). This is the smoothed state estimate.
- \( Vtt1 \) The variance of \( X_t \) conditioned on the data up to time \( t-1 \). Denoted \( P_{t-1} \) in section 6.2 in Shumway and Stoffer (2006).
- \( Vtt \) The variance of \( X_t \) conditioned on the data up to time \( t \). Denoted \( P_t \) in section 6.2 in Shumway and Stoffer (2006).
- \( VtT \) The variance of \( X_t \) conditioned on all the data from time 1 to \( T \).
- \( VttT \) The lag-one covariance of \( X_t \) and \( X_{t-1} \) conditioned on all the data, 1 to \( T \).
- \( K_t \) The Kalman gain. This is part of the update equations and relates to the amount \( xtt1 \) is updated by the data at time \( t \) to produce \( xtt \). Not output by MARSSkfas.
- \( J \) This is similar to the Kalman gain but is part of the Kalman smoother. See Equation 6.49 in Shumway and Stoffer (2006). Not output by MARSSkfas.
- \( \text{Innov} \) This has the innovations at time \( t \), defined as \( \varepsilon_t \equiv y_t - E(Y_t) \). These are the residuals, the difference between the data and their predicted values. See Equation 6.24 in Shumway and Stoffer (2006). Not output by MARSSkfas.
- \( \Sigma \) This has the \( \Sigma_t \), the variance-covariance matrices for the innovations at time \( t \). This is used for the calculation of confidence intervals, the s.e. on the state estimates and the likelihood. See Equation 6.25 in Shumway and Stoffer (2006) for the \( \Sigma \) calculation. Not output by MARSSkfas.
- \( \logLik \) The log-likelihood of the data conditioned on the model parameters.

### 3.4 The exact likelihood

The likelihood of the data given a set of MARSS parameters is part of the output of the MARSSkfss and MARSSkfas functions. The likelihood computation is based on the innovations form of the likelihood (Schweppe 1965) and uses the output from the Kalman filter:

\[
\log L(\Theta | \text{data}) = -\frac{N}{2\log(2\pi)} - \frac{1}{2} \left( \sum_{t=1}^{T} \log |\Sigma_t| + \sum_{t=1}^{T} (\varepsilon_t)^\top \Sigma_t^{-1} \varepsilon_t \right) \tag{3.3}
\]
where \( N \) is the total number of data points, \( \varepsilon_t \) is the innovations at time \( t \) and \( |\Sigma_t| \) is the determinant of the innovations variance-covariance matrix at time \( t \). This likelihood function is shown in Equation 6.62 in Shumway and Stoffer (2006). However there are a few differences between the log-likelihood output by \texttt{MARSSkf} and \texttt{MARSSkfas} and that described in Shumway and Stoffer (2006).

The standard likelihood calculation (Equation 6.62 in Shumway and Stoffer (2006)) is biased when there are missing values in the data, and the missing data modifications discussed in Section 6.4 in Shumway and Stoffer (2006) do not correct for this bias. Harvey (1989), Section 3.4.7, discusses at length that the standard missing values correction leads to an inexact likelihood when there are missing values. The bias is minor if there are few missing values, but it becomes severe as the number of missing values increases. Many ecological datasets have high fractions of missing values and this leads to a very biased likelihood if one uses the inexact formula. Harvey (1989) provides some non-trivial ways to compute the exact likelihood.

The MARSS package uses instead the exact likelihood correction for missing values that is presented in Section 12.3 in Brockwell and Davis (1991). This solution is straight-forward to implement. The correction involves the following changes to \( \varepsilon_t \) and \( \Sigma_t \) in the Equation 3.3. Suppose the value \( y_i, t \) is missing. First, the corresponding \( i \)-th value of \( \varepsilon_t \) is set to 0. Second, the \( i \)-th diagonal value of \( \Sigma_t \) is set to 1 and the off-diagonal elements on the \( i \)-th column and \( i \)-th row are set to 0.

### 3.5 Parametric and innovations bootstrapping

Bootstrapping can be used to construct frequentist confidence intervals on the parameter estimates (Stoffer and Wall, 1991) and to compute the small-sample AIC corrector for MARSS models (Cavanaugh and Shumway, 1997); the functions \texttt{MARSSparamCIs} and \texttt{MARSSaic} do these computations.

The \texttt{MARSSboot} function provides both parametric and innovations bootstrapping of MARSS models. The innovations bootstrap algorithm by Stoffer and Wall (1991) bootstraps the model residuals (the innovations). This is a semi-parametric bootstrap since it uses, partially, the maximum-likelihood parameter estimates. This algorithm cannot be used if there are missing values in the data. Also for short time series, it gives biased bootstraps because one cannot resample the first few innovations.

\texttt{MARSSboot} also provides a fully parametric bootstrap. This uses the maximum-likelihood MARSS parameters to simulate data from which bootstrap parameter estimates are obtained. Our research (Holmes and Ward, 2010) indicates that this provides unbiased bootstrap parameter estimates, and it works with datasets with missing values. Lastly, \texttt{MARSSboot} can also output parameters sampled from the Hessian matrix.
3.6 Simulation and forecasting

The \texttt{MARSSsimulate} function simulates from a fitted marssMLE object (e.g., output from a \texttt{MARSS()} call). It uses \texttt{rmvnorm} (package mvtnorm) to produce draws of the process and observation errors from multivariate normal distributions for each time step.

3.7 Model selection

The package provides the \texttt{MARSSaic} function for computing AIC, AICc and AICb. The latter is a small-sample corrector for autoregressive state-space models. The bias problem with AIC and AICc for short time-series data has been shown in \cite{Cavanaugh1997} and \cite{Holmes2010}. AIC and AICc tend to select overly complex MARSS models when the time-series data are short. AICb corrects this bias. The algorithm for a non-parametric AICb is given in \cite{Cavanaugh1997}. Their algorithm uses the innovations bootstrap \cite{Stoffer1991}, which means it cannot be used when there are missing data. We added a parametric AICb \cite{Holmes2010}, which uses a parametric bootstrap. This algorithm allows one to compute AICb when there are missing data and it provides unbiased AIC even for short time series. See \cite{Holmes2010} for discussion and testing of parametric AICb for MARSS models.

AICb is comprised of the familiar AIC fit term, $-2\log L$, plus a penalty term that is the mean difference between the log likelihood the data under the bootstrapped maximum-likelihood parameter estimates and the log likelihood of the data under the original maximum-likelihood parameter estimate:

\begin{equation}
\text{AICb} = -2\log L(\hat{\Theta}|y) + 2\left(\frac{1}{N_b} \sum_{i=1}^{N_b} \log \frac{L(\hat{\Theta}^*(i)|y)}{L(\hat{\Theta}|y)}\right) \tag{3.4}
\end{equation}

where $\hat{\Theta}$ is the maximum-likelihood parameter set under the original data $y$, $\hat{\Theta}^*(i)$ is a maximum-likelihood parameter set estimated from the $i$-th bootstrapped data set $y^*(i)$, and $N_b$ is the number of bootstrap data sets. It is important to notice that the likelihood in the AICb equation is $L(\hat{\Theta}^*|y)$ not $L(\hat{\Theta}|y^*)$. In other words, we are taking the average of the likelihood of the original data given the bootstrapped parameter sets.
Fitting models with MARSS
Fitting models: the MARSS() function

From the user perspective, the main package function is `MARSS()`. This fits a MARSS model (Equation 1.1) to a matrix of data. The function call takes the form:

```r
MARSS(data, model=list(), form="marxss")
```

The model argument is a list with names `B`, `U`, `C`, `q`, `Z`, `A`, `d`, `R`, `x0`, `V0`. Elements can be left off to use default values. The form argument tells `MARSS()` how to use the model list elements. The default is `form="marxss"` which is the model in Equation 1.1.

The data must be passed in as a $n \times T$ matrix; that is time goes across columns. A vector is not a matrix, nor is a data frame, nor is a time-series (`ts`) object. A data matrix consisting of three time series ($n = 3$) with six time steps might look like

$$
y = \begin{bmatrix}
1 & 2 & NA & NA & 3.2 & 8 \\
2 & 5 & 3 & NA & 5.1 & 5 \\
1 & NA & 2 & 2.2 & NA & 7
\end{bmatrix}
$$

where `NA` denotes a missing value.

The argument `model` specifies the structure of the MARSS model. It is a list, where the list elements for each model parameter specify the form of that parameter.

The most general way to specify model structure is to use a list matrix. The list matrix allows one to combine fixed and estimated elements in one’s parameter specification. It allows a one-to-one correspondence between how you write the parameter matrix on paper and how you specify it in R. For example, let’s say `Q` and `u` have the following forms in your model:

$$
Q = \begin{bmatrix}
q & 0 & 0 \\
0 & q & 0 \\
0 & 0 & 1
\end{bmatrix}
$$

and

$$
u = \begin{bmatrix}
0.05 \\
u_1 \\
u_2
\end{bmatrix}
$$
So $Q$ is a diagonal matrix with the 3rd variance fixed at 1 and the 1st and 2nd estimated and equal. The 1st element of $u$ is fixed, and the 2nd and 3rd are estimated and different. You can specify this using a list matrix:

$$Q = \text{matrix}(\text{list}("q", 0, 0, 0, "q", 0, 0, 1), 3, 3)$$
$$U = \text{matrix}(\text{list}(0.05, "u1", "u2"), 3, 1)$$

If you print out $Q$ and $U$, you will see they look exactly like $Q$ and $u$ written above. MARSS will keep the fixed values fixed and estimate $q$, $u1$, and $u2$.

List matrices allow the most flexible model structures, but MARSS also has text shortcuts for a number of common model structures. Below, the possible ways to specify each model parameter are shown, using $m = 3$ (the number of hidden state processes) and $n = 3$ (number of observation time series).

### 4.1 $u$, $a$ and $\pi$ model structures

$u$, $a$ and $\pi$ are all row matrices and the options for specifying their structures are the same. $a$ has one special option, "scaling" described below. The allowable structures are shown using $u$ as an example. Note that you should be careful about specifying shared structure in $\pi$ because you need to make sure the structure in $\Lambda$ matches. For example, if you require that all the $\pi$ values are shared (equal) then $\Lambda$ cannot be a diagonal matrix since that would be saying that the $\pi$ values are independent, which they are clearly not if you force them to be equal.

- **$U = \text{matrix}(\text{list}(), m, 1)$**: This is the most general form and allows one to specify fixed and estimated elements in $u$. Each character string in $u$ is the name of one of the $u$ elements to be estimated. For example if $U = \text{matrix}(\text{list}(0.01, "u", "u"), 3, 1)$, then $u$ in the model has the following structure:

$$ \begin{bmatrix} 0.01 \\ u \\ u \end{bmatrix} $$

- **$U = \text{matrix}(\text{c}(), m, 1)$**, where the values in $\text{c}()$ are all character strings; each character string is the name of an element to be estimated. For example if $U = \text{matrix}(\text{c}("u1", "u1", "u2"), 3, 1)$, then $u$ in the model has the following structure:

$$ \begin{bmatrix} u1 \\ u1 \\ u2 \end{bmatrix} $$

with two values being estimated. $U = \text{matrix}(\text{list}("u1", "u1", "u2"), 3, 1)$ has the same effect.

- **$U = "unequal"$** or **$U = "unconstrained"$**: Both of these stings indicate that each element of $u$ is estimated. If $m = 3$, then $u$ would have the form:
U="equal": There is only one value in u:

\[
\begin{bmatrix}
u_1 \\
u_2 \\
u_3
\end{bmatrix}
\]

U=matrix(c(),m,1), where the values in c() all numerical values: u is fixed and has no estimated values. If U=matrix(c(0.01,1,-0.5),3,1), then u in the model is:

\[
\begin{bmatrix}
0.01 \\
1 \\
-0.5
\end{bmatrix}
\]

U=matrix(list(0.01,1,-0.5),3,1) would have the same effect.

U="zero": u is all zero:

\[
\begin{bmatrix}
0 \\
0 \\
0
\end{bmatrix}
\]

The a parameter has a special option, "scaling", which is the default behavior. In this case, a is treated like a scaling parameter. If there is only one y row associated with an x row, then the corresponding a element is 0. If there are more than one y rows associated with an x row, then the first a element is set to 0 and the others are estimated. For example, say m = 2 and n = 4 and Z looks like the following:

\[
Z = \begin{bmatrix}
1 & 0 \\
1 & 0 \\
1 & 0 \\
0 & 1
\end{bmatrix}
\]

Then the 1st-3rd rows of y are associated with the first row of x, and the 4th row of y is associated with the last row of x. Then if a is specified as "scaling", a has the following structure:

\[
\begin{bmatrix}
0 \\
a_1 \\
a_2 \\
0
\end{bmatrix}
\]

4.2 Q, R, Λ model structures

The possible Q, R, and Λ model structures are identical, except that R is n × n while Q and Λ are m × m. All types of structures can be specified using a list
matrix, but there are also text shortcuts for specifying common structures.
The structures are shown using $Q$ as the example.

$Q=\text{matrix(list()},m,m)$: This is the most general way to specify the parameters and allows there to be fixed and estimated elements. Each character string in the list matrix is the name of one of the $Q$ elements to be estimated, and each numerical value is a fixed value. For example if $Q=\text{matrix(list("s2a",0,0,0,"s2a",0,0,0,"s2b"),3,3)}$, then $Q$ has the following structure:

$$
\begin{bmatrix}
\sigma^2_{s2a} & 0 & 0 \\
0 & \sigma^2_{s2a} & 0 \\
0 & 0 & \sigma^2_{s2b}
\end{bmatrix}
$$

Note that $\text{diag(c("s2a","s2a","s2b"))}$ will not have the desired effect of producing a matrix with numeric 0s on the off-diagonals. It will have character 0s and MARSS will interpret “0” as the name of an element of $Q$ to be estimated. Instead, the following two lines can be used:

$Q=\text{matrix(list(0),3,3)}$
$\text{diag(Q)=c("s2a","s2a","s2b")}$

$Q=\text{"diagonal and equal"}$: There is only one process variance value in this case:

$$
\begin{bmatrix}
\sigma^2 & 0 & 0 \\
0 & \sigma^2 & 0 \\
0 & 0 & \sigma^2
\end{bmatrix}
$$

$Q=\text{"diagonal and unequal"}$: There are $m$ process variance values in this case:

$$
\begin{bmatrix}
\sigma^2_1 & 0 & 0 \\
0 & \sigma^2_2 & 0 \\
0 & 0 & \sigma^2_3
\end{bmatrix}
$$

$Q=\text{"unconstrained"}$: There are values on the diagonal and the off-diagonals of $Q$ and the variances and covariances are all different:

$$
\begin{bmatrix}
\sigma^2_1 & \sigma_{1,2} & \sigma_{1,3} \\
\sigma_{1,2} & \sigma^2_2 & \sigma_{2,3} \\
\sigma_{1,3} & \sigma_{2,3} & \sigma^2_3
\end{bmatrix}
$$

There are $m$ process variances and $(m^2 - m)/2$ covariances in this case, so $(m^2 + m)/2$ values to be estimated. Note that variance-covariance matrices are never truly unconstrained since the upper and lower triangles of the matrix must be equal.

$Q=\text{"equalvarcov"}$: There is one process variance and one covariance:

$$
\begin{bmatrix}
\sigma^2 & \beta & \beta \\
\beta & \sigma^2 & \beta \\
\beta & \beta & \sigma^2
\end{bmatrix}
$$
4.4 Z model

\( Q = \text{matrix}(c(), m, m) \), where all values in \( c() \) are character strings: Each element in \( Q \) is estimated and each character string is the name of a value to be estimated. Note if \( m = 1 \), you still need to wrap its value in \text{matrix}() \) so that its class is matrix.

\( Q = \text{matrix}(c(), m, m) \), where all values in \( c() \) are numeric values: Each element in \( Q \) is fixed to the values in the matrix.

\( Q = \text{"identity"} \): The \( Q \) matrix is the identity matrix:

\[
\begin{bmatrix}
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & 1
\end{bmatrix}
\]

\( Q = \text{"zero"} \): The \( Q \) matrix is all zeros:

\[
\begin{bmatrix}
0 & 0 & 0 \\
0 & 0 & 0 \\
0 & 0 & 0
\end{bmatrix}
\]

Be careful when setting \( \Lambda \) model structures. Mis-specifying the structure of \( \Lambda \) can have catastrophic, but difficult to discern, effects on your estimates. See the comments on priors in Chapter 1.

4.3 B model structures

Like the variance-covariance matrices (\( Q, R \) and \( \Lambda \)), \( B \) can be specified with a list matrix to allow you to have both fixed and shared elements in the \( B \) matrix. Character matrices and matrices with fixed values operate the same way as for the variance-covariance matrices. In addition, the same text shortcuts are available: “unconstrained”, “identity”, “diagonal and equal”, “diagonal and unequal”, “equalvarcov”, and “zero”. A fixed \( B \) can be specified with a numeric matrix, but all eigenvalues must fall within the unit circle; meaning \( \text{all(abs(eigen(B)$values)<1)} \) must be true.

4.4 Z model

Like \( B \) and the variance-covariance matrices, \( Z \) can be specified with a list matrix to allow you to have both fixed and estimated elements in \( Z \). If \( Z \) is a square matrix, many of the same text shortcuts are available: “diagonal and equal”, “diagonal and equal”, and “equalvarcov”. If \( Z \) is a design matrix\(^1\), then a special shortcut is available using \text{factor}() \) which allows you to specify which \( y \) rows are associated with which \( x \) rows. See Chapter 5 and the applications chapters for more examples.

\(^1\) a matrix with only 0s and 1s and where the row sums are all equal to 1
Z = factor(c(1,1,1)): All y time series are observing the same (and only) hidden state trajectory x (n = 3 and m = 1):

\[
Z = \begin{bmatrix}
1 \\
1 \\
1
\end{bmatrix}
\]

Z = factor(c(1,2,3)): Each time series in y corresponds to a different hidden state trajectory. This is the default Z model and in this case n = m:

\[
Z = \begin{bmatrix}
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & 1
\end{bmatrix}
\]

Z = factor(c(1,1,2)): The first two time series in y corresponds to one hidden state trajectory and the third y time series corresponds to a different hidden state trajectory. Here n = 3 and m = 2:

\[
Z = \begin{bmatrix}
1 & 0 \\
1 & 0 \\
0 & 1
\end{bmatrix}
\]

The Z model can be specified using either numeric or character factor levels. c(1,1,2) is the same as c("north","north","south")

Z = "identity": This is the default behavior. This means Z is a n x n identity matrix and m = n. If n = 3, it is the same as Z = factor(c(1,2,3)).

Z = matrix(c(), n, m), where the elements in c() are all strings: Passing in a n x m character matrix, means that each character string is a value to be estimated. Be careful that you are specifying an identifiable model when using this option.

Z = matrix(c(), n, m), where the elements in c() are all numeric: Passing in a n x m numeric matrix means that Z is fixed to the values in the matrix. The matrix must be numeric but it does not need to be a design matrix.

Z = matrix(list(), n, m): Passing in a n x m list matrix allows you to combine fixed and estimated values in the Z matrix. Be careful that you are specifying an identifiable model.

4.5 Default model structures

The defaults for the model arguments in form = "marxss" are

Z = "identity" each y in y corresponds to one x in x
B = "identity" no interactions among the x's in x
U = "unequal" the u's in u are all different
Q="diagonal and unequal" process errors are independent but have different variances
R="diagonal and equal" the observations are i.i.d.
A="scaling" $a$ is a set of scaling factors
C="zero" and D="zero" no inputs.
c="zero" and d="zero" no inputs.
pi="unequal" all initial states are different
V0="zero" the initial condition on the states ($x_0$ or $x_1$) is fixed but unknown
$\text{tinitx}=0$ the initial state refers to $t = 0$ instead of $t = 1$. 
5

Short Examples

In this chapter, we work through a series of short examples to illustrate the MARSS package functions. This chapter is oriented towards those who are already somewhat familiar with MARSS models and want to get started quickly. We provide little explanatory text. Those unfamiliar with MARSS models might want to start with the application chapters.

In these examples, we will use the default form="marxss" argument for a MARSS() call. This specifies a MARSS model of the form:

\[ x_t = B_t x_{t-1} + u_t + C_t c_t + G_t w_t, \quad \text{where } w_t \sim \text{MVN}(0, Q_t) \]  
\[ y_t = Z_t x_t + a_t + D_t d_t + H_t v_t, \quad \text{where } v_t \sim \text{MVN}(0, R_t) \]  
\[ x_0 \sim \text{MVN}(\pi, \Lambda) \]  

The \( c \) and \( d \) are inputs (not estimated). In the examples here, we leave off \( c \) and \( d \). We address including inputs only briefly at the end of the chapter. See Chapter 13 for extended examples of including covariates as inputs in a MARSS model. We will also not use \( G_t \) or \( H_t \) in this chapter.

5.1 Fixed and estimated elements in parameter matrices

Suppose one has a MARSS model (Equation 5.1) with the following structure:

\[
\begin{bmatrix} x_{1,t} \\ x_{2,t} \end{bmatrix} = \begin{bmatrix} b_1 & 0.1 \\ b_2 & 2 \end{bmatrix} \begin{bmatrix} x_{1,t-1} \\ x_{2,t-1} \end{bmatrix} + \begin{bmatrix} u_t \\ w_{1,t} \end{bmatrix}, \quad w_t \sim \text{MVN}\left( \begin{bmatrix} 0 \\ q_1 \\ q_2 \end{bmatrix} \right)
\]

\[
\begin{bmatrix} y_{1,t} \\ y_{2,t} \\ y_{3,t} \end{bmatrix} = \begin{bmatrix} z_1 & 0 \\ z_2 & z_2 \\ 0 & 3 \end{bmatrix} \begin{bmatrix} x_{1,t} \\ x_{2,t} \end{bmatrix} + \begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix} + \begin{bmatrix} v_{1,t} \\ v_{2,t} \\ v_{3,t} \end{bmatrix}, \quad v_t \sim \text{MVN}\left( \begin{bmatrix} 0 \\ r \\ 0 \end{bmatrix} \right)
\]

\[ x_0 \sim \text{MVN}\left( \begin{bmatrix} \pi_1 \\ \pi_2 \end{bmatrix}, \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} \right) \]
Notice how this model mixes fixed values, estimated values and shared values.

In MARSS, model structure is specified using a list with the names, Z, A, R, B, U, Q, x0 and V0. Each element is a matrix (class `matrix`) with the same dimensions as the matrix of the same name in the MARSS model. MARSS distinguishes between the estimated and fixed values in a matrix by using a list matrix in which you can have numeric and character elements. Numeric elements are fixed; character elements are names of things to be estimated. The model above would be specified as:

```r
Z = matrix(list("z1","z2",0,0,"z2",3),3,2)
A = matrix(0,3,1)
R = matrix(list(0),3,3); diag(R) = c("r","r",1)
B = matrix(list("b1",0.1,"b2",2),2,2)
U = matrix(c("u","u"),2,1)
Q = matrix(c("q1","q3","q3","q2"),2,2)
x0 = matrix(c("pi1","pi2"),2,1)
V0 = diag(1,2)
model.gen = list(Z=Z,A=A,R=R,B=B,U=U,Q=Q,x0=x0,V0=V0,tinitx=0)
```

Notice that there is a one-to-one correspondence between the model list in R and the model on paper. Fitting the model is then just a matter of passing the data and model list to the MARSS function:

```r
kemfit = MARSS(dat, model=model.gen)
```

If you work often with MARSS models then you will probably know whether prior sensitivity is a problem for your types of MARSS applications. If so, note that the MARSS package is unusual in that it allows you to set $\Lambda = 0$ and treat $x_0$ as an unknown estimated parameter. This eliminates the prior and thus the prior sensitivity problems—at the cost of adding $m$ parameters. Depending on your application, you may need to set the initial conditions at $t = 1$ instead of the default of $t = 0$. If you are unsure, look in the index and read all the sections that talk about troubleshooting priors.

### 5.2 Different numbers of state processes

Here we show a series of short examples using a dataset on Washington harbor seals (`?harborSealWA`), which has five observation time series. The dataset is a little unusual in that it has four missing years from years 2 to 5. This causes some interesting issues with prior specification. Before starting the harbor seal examples, we set up the data, making time go across the columns and removing the year column:

```r
dat = t(harborSealWA)
dat = dat[2:nrow(dat),] #remove the year row
```
5.2.1 One hidden state process for each observation time series

This is the default model for the MARSS() function. In this case, \( n = m \), the observation errors are i.i.d. and the process errors are independent and have different variances. The elements in \( \mathbf{u} \) are all different (meaning, they are not forced to be the same). Mathematically, the MARSS model being fit is:

\[
\begin{bmatrix}
    x_{1,t} \\
    x_{2,t} \\
    x_{3,t} \\
    x_{4,t} \\
    x_{5,t}
\end{bmatrix} =
\begin{bmatrix}
    1 & 0 & 0 & 0 & 0 \\
    0 & 1 & 0 & 0 & 0 \\
    0 & 0 & 1 & 0 & 0 \\
    0 & 0 & 0 & 1 & 0 \\
    0 & 0 & 0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
    x_{1,t-1} \\
    x_{2,t-1} \\
    x_{3,t-1} \\
    x_{4,t-1} \\
    x_{5,t-1}
\end{bmatrix} +
\begin{bmatrix}
    u_1 \\
    u_2 \\
    u_3 \\
    u_4 \\
    u_5
\end{bmatrix} +
\begin{bmatrix}
    w_{1,t} \\
    w_{2,t} \\
    w_{3,t} \\
    w_{4,t} \\
    w_{5,t}
\end{bmatrix}, \quad \mathbf{w}_t \sim \text{MVN}
\begin{bmatrix}
    0 \\
    0 \\
    0 \\
    0 \\
    0
\end{bmatrix},
\begin{bmatrix}
    q_1 & 0 & 0 & 0 & 0 \\
    0 & q_2 & 0 & 0 & 0 \\
    0 & 0 & q_3 & 0 & 0 \\
    0 & 0 & 0 & q_4 & 0 \\
    0 & 0 & 0 & 0 & q_5
\end{bmatrix}
\]

\[
\begin{bmatrix}
    y_{1,t} \\
    y_{2,t} \\
    y_{3,t} \\
    y_{4,t} \\
    y_{5,t}
\end{bmatrix} =
\begin{bmatrix}
    1 & 0 & 0 & 0 & 0 \\
    0 & 1 & 0 & 0 & 0 \\
    0 & 0 & 1 & 0 & 0 \\
    0 & 0 & 0 & 1 & 0 \\
    0 & 0 & 0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
    x_{1,t} \\
    x_{2,t} \\
    x_{3,t} \\
    x_{4,t} \\
    x_{5,t}
\end{bmatrix} +
\begin{bmatrix}
    0 \\
    0 \\
    0 \\
    0 \\
    0
\end{bmatrix} +
\begin{bmatrix}
    v_{1,t} \\
    v_{2,t} \\
    v_{3,t} \\
    v_{4,t} \\
    v_{5,t}
\end{bmatrix}, \quad \mathbf{v}_t \sim \text{MVN}
\begin{bmatrix}
    0 \\
    r \\
    0 \\
    0 \\
    0
\end{bmatrix},
\begin{bmatrix}
    0 & 0 & 0 & 0 & 0 \\
    0 & 0 & 0 & 0 & 0 \\
    0 & 0 & 0 & 0 & 0 \\
    0 & 0 & 0 & 0 & 0 \\
    0 & 0 & 0 & 0 & r
\end{bmatrix}
\]

This is the default model, so you can fit it by simply passing \( \text{dat} \) to MARSS().

\[ \text{kemfit} = \text{MARSS(dat)} \]

Success! abstol and log-log tests passed at 38 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 38 iterations.
Log-likelihood: 19.13428
AIC: -6.268557 AICc: 3.805517

<table>
<thead>
<tr>
<th>Estimate</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
<td>0.00895</td>
</tr>
<tr>
<td>U.X.SJF</td>
<td>0.06839</td>
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<tr>
<td>U.X.SJI</td>
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<tr>
<td>U.X.EBays</td>
<td>0.04179</td>
</tr>
<tr>
<td>U.X.PSnd</td>
<td>0.05226</td>
</tr>
<tr>
<td>U.X.HC</td>
<td>-0.00279</td>
</tr>
<tr>
<td>Q.(X.SJF,X.SJF)</td>
<td>0.03205</td>
</tr>
<tr>
<td>Q.(X.SJI,X.SJI)</td>
<td>0.01098</td>
</tr>
<tr>
<td>Q.(X.EBays,X.EBays)</td>
<td>0.00706</td>
</tr>
<tr>
<td>Q.(X.PSnd,X.PSnd)</td>
<td>0.00414</td>
</tr>
</tbody>
</table>
38  5 Examples

\begin{verbatim}
Q.(X.HC,X.HC)  0.05450
x0.X.SJF      5.98647
x0.X.SJI      6.72487
x0.X.EBays    6.66212
x0.X.PSnd     5.83969
x0.X.HC       6.60482

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

The output warns you that the convergence tolerance is high. You can set it lower by passing in control=list(conv.test.slope.tol=0.1). MARSS() is automatically creating parameter names since you did not tell it the names.
To see exactly where each parameter element appears in its parameter matrix, type summary(kemfit$model).

Though it is not necessary to specify the model for this example since it is the default, here is how you could do so using matrices:

\begin{verbatim}
B=Z=diag(1,5)
U=matrix(c("u1","u2","u3","u4","u5"),5,1)
x0=A=matrix(0,5,1)
R=Q=matrix(list(0),5,5)
\end{verbatim}

Notice that when a matrix has both fixed and estimated elements (like R and Q), a list matrix is used to allow you to specify the fixed elements as numeric and to give the estimated elements character names.

The default MLE method is the EM algorithm (method="kem"). You can also use a quasi-Newton method (BFGS) by setting method="BFGS".

\begin{verbatim}
kemfit.bfgs = MARSS(dat, method="BFGS")
\end{verbatim}

Success! Converged in 99 iterations.
Function MARSSkfas used for likelihood calculation.

MARSS fit is
Estimation method: BFGS
Estimation converged in 99 iterations.
Log-likelihood: 19.13936
AIC: -6.278712  AICc: 3.795362

\begin{verbatim}
Estimate
R.diag       0.00849
U.X.SJF      0.06838
U.X.SJI      0.07152
\end{verbatim}
\end{verbatim}
5.2 Different numbers of state processes

<table>
<thead>
<tr>
<th>Process</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>U.X.EBays</td>
<td>0.04188</td>
</tr>
<tr>
<td>U.X.PSnd</td>
<td>0.05233</td>
</tr>
<tr>
<td>U.X.HC</td>
<td>-0.00271</td>
</tr>
<tr>
<td>Q.(X.SJF,X.SJF)</td>
<td>0.03368</td>
</tr>
<tr>
<td>Q.(X.SJI,X.SJI)</td>
<td>0.01124</td>
</tr>
<tr>
<td>Q.(X.EBays,X.EBays)</td>
<td>0.00722</td>
</tr>
<tr>
<td>Q.(X.PSnd,X.PSnd)</td>
<td>0.00437</td>
</tr>
<tr>
<td>Q.(X.HC,X.HC)</td>
<td>0.05600</td>
</tr>
<tr>
<td>x0.X.SJF</td>
<td>5.98437</td>
</tr>
<tr>
<td>x0.X.SJI</td>
<td>6.72169</td>
</tr>
<tr>
<td>x0.X.EBays</td>
<td>6.65689</td>
</tr>
<tr>
<td>x0.X.PSnd</td>
<td>5.83527</td>
</tr>
<tr>
<td>x0.X.HC</td>
<td>6.60425</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated. Use MARSSparamCIs to compute CIs and bias estimates.

Using the default EM convergence criteria, the EM algorithm stops at a log-likelihood a little lower than the BFGS algorithm does, but the EM algorithm was faster, 6.5 times faster, in this case. If you wanted to use the EM fit as the initial conditions, pass in the inits argument using the $par element (or coef(fit,form="marss")) of the EM fit.

```r
ekemfit.bfgs2 = MARSS(dat, method="BFGS", inits=kemfit$par)
```

The BFGS algorithm now converges in 103 iterations. Output not shown.

We mentioned that the missing years from year 2 to 4 creates an interesting issue with the prior specification. The default behavior of MARSS is to treat the initial state as at \( t = 0 \) instead of \( t = 1 \). Usually this doesn’t make a difference, but for this dataset, if we set the prior at \( t = 1 \), the MLE estimate of \( R \) becomes 0. If we estimate \( x_1 \) as a parameter and let \( R \) go to 0, the likelihood will go to infinity (slowly but surely). This is neither an error nor a pathology, but is probably not what you would like to have happen. Note that the “BFGS” algorithm will not find the maximum in this case; it will stop before \( R \) gets small and the likelihood gets very large. However, the EM algorithm will climb up the peak. You can try it by running the following code. It will report warnings which you can read about in Appendix A.

```r
ekemfit.strange = MARSS(dat, model=list(tinitx=1))
```

5.2.2 Five correlated hidden state processes

This is the same model except that the five hidden states have correlated process errors. Mathematically, this is the model:
\[
\begin{bmatrix}
    x_{1,t} \\
    x_{2,t} \\
    x_{3,t} \\
    x_{4,t} \\
    x_{5,t}
\end{bmatrix}
= \begin{bmatrix}
    x_{1,t-1} \\
    x_{2,t-1} \\
    x_{3,t-1} \\
    x_{4,t-1} \\
    x_{5,t-1}
\end{bmatrix} + \begin{bmatrix}
    u_1 \\
    u_2 \\
    u_3 \\
    u_4 \\
    u_5
\end{bmatrix} + \begin{bmatrix}
    w_{1,t} \\
    w_{2,t} \\
    w_{3,t} \\
    w_{4,t} \\
    w_{5,t}
\end{bmatrix}, \quad \mathbf{w}_t \sim \text{MVN} \left( \begin{bmatrix}
    q_1 & c_1,2 & c_1,3 & c_1,4 & c_1,5 \\
    c_1,2 & q_2 & c_2,3 & c_2,4 & c_2,5 \\
    c_1,3 & c_2,3 & q_3 & c_3,4 & c_3,5 \\
    c_1,4 & c_2,4 & c_3,4 & q_4 & c_4,5 \\
    c_1,5 & c_2,5 & c_3,5 & c_4,5 & q_5
\end{bmatrix} \right)
\]

\[
\begin{bmatrix}
    y_{1,t} \\
    y_{2,t} \\
    y_{3,t} \\
    y_{4,t} \\
    y_{5,t}
\end{bmatrix}
= \begin{bmatrix}
    1 & 0 & 0 & 0 & 0 \\
    0 & 1 & 0 & 0 & 0 \\
    0 & 0 & 1 & 0 & 0 \\
    0 & 0 & 0 & 1 & 0 \\
    0 & 0 & 0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
    x_{1,t} \\
    x_{2,t} \\
    x_{3,t} \\
    x_{4,t} \\
    x_{5,t}
\end{bmatrix}
+ \begin{bmatrix}
    0 \\
    0 \\
    0 \\
    0 \\
    0
\end{bmatrix} + \begin{bmatrix}
    v_{1,t} \\
    v_{2,t} \\
    v_{3,t} \\
    v_{4,t} \\
    v_{5,t}
\end{bmatrix}, \quad \mathbf{v}_t \sim \text{MVN} \left( \begin{bmatrix}
    r & 0 & 0 & 0 & 0 \\
    0 & r & 0 & 0 & 0 \\
    0 & 0 & r & 0 & 0 \\
    0 & 0 & 0 & r & 0 \\
    0 & 0 & 0 & 0 & r
\end{bmatrix} \right)
\]

\mathbf{B} \text{ is not shown in the top equation; it is a } m \times m \text{ identity matrix. To fit, use MARSS()} \text{ with the model argument set (output not shown).}

```r
kemfit = MARSS(dat, model=list(Q="unconstrained"))
```

This shows one of the text shortcuts, "unconstrained", which means estimate all elements in the matrix. This shortcut can be used for all parameter matrices.

### 5.2.3 Five equally correlated hidden state processes

This is the same model except that now there is only one process error variance and one process error covariance. Mathematically, the model is:

\[
\begin{bmatrix}
    x_{1,t} \\
    x_{2,t} \\
    x_{3,t} \\
    x_{4,t} \\
    x_{5,t}
\end{bmatrix}
= \begin{bmatrix}
    x_{1,t-1} \\
    x_{2,t-1} \\
    x_{3,t-1} \\
    x_{4,t-1} \\
    x_{5,t-1}
\end{bmatrix} + \begin{bmatrix}
    u_1 \\
    u_2 \\
    u_3 \\
    u_4 \\
    u_5
\end{bmatrix} + \begin{bmatrix}
    w_{1,t} \\
    w_{2,t} \\
    w_{3,t} \\
    w_{4,t} \\
    w_{5,t}
\end{bmatrix}, \quad \mathbf{w}_t \sim \text{MVN} \left( \begin{bmatrix}
    q & c & c & c & c \\
    c & q & c & c & c \\
    c & c & q & c & c \\
    c & c & c & q & c \\
    c & c & c & c & q
\end{bmatrix} \right)
\]

\[
\begin{bmatrix}
    y_{1,t} \\
    y_{2,t} \\
    y_{3,t} \\
    y_{4,t} \\
    y_{5,t}
\end{bmatrix}
= \begin{bmatrix}
    1 & 0 & 0 & 0 & 0 \\
    0 & 1 & 0 & 0 & 0 \\
    0 & 0 & 1 & 0 & 0 \\
    0 & 0 & 0 & 1 & 0 \\
    0 & 0 & 0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
    x_{1,t} \\
    x_{2,t} \\
    x_{3,t} \\
    x_{4,t} \\
    x_{5,t}
\end{bmatrix}
+ \begin{bmatrix}
    0 \\
    0 \\
    0 \\
    0 \\
    0
\end{bmatrix} + \begin{bmatrix}
    v_{1,t} \\
    v_{2,t} \\
    v_{3,t} \\
    v_{4,t} \\
    v_{5,t}
\end{bmatrix}, \quad \mathbf{v}_t \sim \text{MVN} \left( \begin{bmatrix}
    r & 0 & 0 & 0 & 0 \\
    0 & r & 0 & 0 & 0 \\
    0 & 0 & r & 0 & 0 \\
    0 & 0 & 0 & r & 0 \\
    0 & 0 & 0 & 0 & r
\end{bmatrix} \right)
\]

Again \mathbf{B} \text{ is not shown in the top equation; it is a } m \times m \text{ identity matrix. To fit, use the following code (output not shown):}

```r
kemfit = MARSS(dat, model=list(Q="equalvarcov"))
```

The shortcut "equalvarcov" means one value on the diagonal and one on the off-diagonal. It can be used for all square matrices (\(\mathbf{B, Q, R, and \Lambda}\)).
5.2.4 Five hidden state processes with a “north” and a “south” $u$ and $Q$ elements

Here we fit a model with five independent hidden states where each observation time series is an independent observation of a different hidden trajectory but the hidden trajectories 1-3 share their $u$ and $Q$ elements, while hidden trajectories 4-5 share theirs. This is the model:

$$\begin{bmatrix} x_{1,t} \\ x_{2,t} \\ x_{3,t} \\ x_{4,t} \\ x_{5,t} \end{bmatrix} = \begin{bmatrix} x_{1,t-1} \\ x_{2,t-1} \\ x_{3,t-1} \\ x_{4,t-1} \\ x_{5,t-1} \end{bmatrix} + \begin{bmatrix} u_n \\ u_n \\ u_n \\ u_s \\ u_s \end{bmatrix} + \begin{bmatrix} w_{1,t} \\ w_{2,t} \\ w_{3,t} \\ w_{4,t} \\ w_{5,t} \end{bmatrix}, \quad w_t \sim \text{MVN} \left( \begin{bmatrix} q_n & 0 & 0 & 0 & 0 \\ 0 & q_n & 0 & 0 & 0 \\ 0 & 0 & q_n & 0 & 0 \\ 0 & 0 & 0 & q_s & 0 \\ 0 & 0 & 0 & 0 & q_s \end{bmatrix} \right)$$

To fit use the following code, we specify the `model` argument for $u$ and $Q$ using list matrices. List matrices allow us to combine numeric and character values in a matrix. MARSS will interpret the numeric values as fixed, and the character values as parameters to be estimated. Parameters with the same name are constrained to be identical.

```r
regions=list("N","N","N","S","S")
U=matrix(regions,5,1)
Q=matrix(list(0),5,5); diag(Q)=regions
kemfit = MARSS(dat, model=list(U=U, Q=Q))
```

Only $u$ and $Q$ need to be specified since the other parameters are at their default values.

5.2.5 Fixed observation error variance

Here we fit the same model but with a known observation error variance. This is the model:
5 Examples

\[
\begin{bmatrix}
  x_{1,t} \\
  x_{2,t} \\
  x_{3,t} \\
  x_{4,t} \\
  x_{5,t}
\end{bmatrix}
= \begin{bmatrix}
  x_{1,t-1} \\
  x_{2,t-1} \\
  x_{3,t-1} \\
  x_{4,t-1} \\
  x_{5,t-1}
\end{bmatrix}
+ \begin{bmatrix}
  u_n \\
  u_n \\
  u_n \\
  u_n \\
  u_n
\end{bmatrix}
+ \begin{bmatrix}
  w_{1,t} \\
  w_{2,t} \\
  w_{3,t} \\
  w_{4,t} \\
  w_{5,t}
\end{bmatrix},
\]

\[ w_t \sim \text{MVN} \left( \mathbf{0}, \begin{bmatrix}
  q_n & 0 & 0 & 0 & 0 \\
  0 & q_n & 0 & 0 & 0 \\
  0 & 0 & q_n & 0 & 0 \\
  0 & 0 & 0 & q_s & 0 \\
  0 & 0 & 0 & 0 & q_s
\end{bmatrix} \right) \]

\[
\begin{bmatrix}
  y_{1,t} \\
  y_{2,t} \\
  y_{3,t} \\
  y_{4,t} \\
  y_{5,t}
\end{bmatrix}
= \begin{bmatrix}
  1 & 0 & 0 & 0 & 0 \\
  0 & 1 & 0 & 0 & 0 \\
  0 & 0 & 1 & 0 & 0 \\
  0 & 0 & 0 & 1 & 0 \\
  0 & 0 & 0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
  x_{1,t} \\
  x_{2,t} \\
  x_{3,t} \\
  x_{4,t} \\
  x_{5,t}
\end{bmatrix}
+ \begin{bmatrix}
  v_{1,t} \\
  v_{2,t} \\
  v_{3,t} \\
  v_{4,t} \\
  v_{5,t}
\end{bmatrix},
\]

\[ v_t \sim \text{MVN} \left( \mathbf{0}, \begin{bmatrix}
  0.01 & 0 & 0 & 0 & 0 \\
  0 & 0.01 & 0 & 0 & 0 \\
  0 & 0 & 0.01 & 0 & 0 \\
  0 & 0 & 0 & 0.01 & 0 \\
  0 & 0 & 0 & 0 & 0.01
\end{bmatrix} \right) \]

To fit this model, use the following code (output not shown):

\begin{verbatim}
regions=list("N","N","N","S","S")
U=matrix(regions,5,1)
Q=matrix(list(0),5,5); diag(Q)=regions
R=diag(0.01,5)
kemfit = MARSS(dat, model=list(U=U, Q=Q, R=R))
\end{verbatim}

5.2.6 One hidden state and five i.i.d. observation time series

Instead of five hidden state trajectories, we specify that there is only one and all the observations are of that one trajectory. Mathematically, the model is:

\[ x_t = x_{t-1} + u + w_t, \quad w_t \sim \mathcal{N}(0,q) \]

\[
\begin{bmatrix}
  y_{1,t} \\
  y_{2,t} \\
  y_{3,t} \\
  y_{4,t} \\
  y_{5,t}
\end{bmatrix}
= \begin{bmatrix}
  1 & 0 & 0 & 0 & 0 \\
  0 & 1 & 0 & 0 & 0 \\
  0 & 0 & 1 & 0 & 0 \\
  0 & 0 & 0 & 1 & 0 \\
  0 & 0 & 0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
  x_{t} \\
  a_2 \\
  a_3 \\
  a_4 \\
  a_5
\end{bmatrix}
+ \begin{bmatrix}
  v_{1,t} \\
  v_{2,t} \\
  v_{3,t} \\
  v_{4,t} \\
  v_{5,t}
\end{bmatrix}, \quad v_t \sim \text{MVN} \left( \mathbf{0}, \begin{bmatrix}
  r & 0 & 0 & 0 & 0 \\
  0 & r & 0 & 0 & 0 \\
  0 & 0 & r & 0 & 0 \\
  0 & 0 & 0 & r & 0 \\
  0 & 0 & 0 & 0 & r
\end{bmatrix} \right)
\]

Note the default model for \( R \) is "diagonal and equal" so we can leave this off when specifying the \texttt{model} argument. To fit, use this code (output not shown):

\begin{verbatim}
regions=list("N","N","N","S","S")
U=matrix(regions,5,1)
Q=matrix(list(0),5,5); diag(Q)=regions
R=diag(0.01,5)
kemfit = MARSS(dat, model=list(U=U, Q=Q, R=R))
\end{verbatim}
\begin{verbatim}
Z = factor(c(1,1,1,1,1))
kemfit = MARSS(dat, model=list(Z=Z))

Success! abstol and log-log tests passed at 28 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 28 iterations.
Log-likelihood: 3.593276
AIC: 8.813447  AICc: 11.13603

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A.SJI</td>
<td>0.80153</td>
</tr>
<tr>
<td>A.EBays</td>
<td>0.28245</td>
</tr>
<tr>
<td>A.PSnd</td>
<td>-0.54802</td>
</tr>
<tr>
<td>A.HC</td>
<td>-0.62665</td>
</tr>
<tr>
<td>R.diag</td>
<td>0.04523</td>
</tr>
<tr>
<td>U.U</td>
<td>0.04759</td>
</tr>
<tr>
<td>Q.Q</td>
<td>0.00429</td>
</tr>
<tr>
<td>x0.x0</td>
<td>6.39199</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

You can also pass in Z exactly as it is in the equation: Z=matrix(1,5,2),
but the factor shorthand is handy if you need to assign different observed
time series to different underlying state time series (see next examples). The
default a form is "scaling", which means that the first y row associated with
a given x has a = 0 and the rest are estimated.

5.2.7 One hidden state and five independent observation time
series with different variances

Mathematically, this model is:
\end{verbatim}
\[ x_t = x_{t-1} + u + w_t, \quad w_t \sim N(0, q) \]

\[
\begin{bmatrix}
    y_{1,t} \\
    y_{2,t} \\
    y_{3,t} \\
    y_{4,t} \\
    y_{5,t}
\end{bmatrix}
= \begin{bmatrix}
    1 & 0 \\
    1 & a_2 \\
    1 & a_3 \\
    1 & a_4 \\
    1 & a_5
\end{bmatrix}
\begin{bmatrix}
    x_t \\
    v_{1,t} \\
    v_{2,t} \\
    v_{3,t} \\
    v_{4,t} \\
    v_{5,t}
\end{bmatrix}
+ \begin{bmatrix}
    r_1 & 0 & 0 & 0 & 0 \\
    0 & r_2 & 0 & 0 & 0 \\
    0 & 0 & r_3 & 0 & 0 \\
    0 & 0 & 0 & r_4 & 0 \\
    0 & 0 & 0 & 0 & r_5
\end{bmatrix}
\]

To fit this model:

\[
Z = \text{factor(c(1,1,1,1,1))} \\
R = "\text{diagonal and unequal}" \\
\text{kemfit} = \text{MARSS(dat, model=list(Z=Z, R=R))}
\]

Success! abstol and log-log tests passed at 24 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

\text{MARSS fit is}
\text{Estimation method: kem}
\text{Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001}
\text{Estimation converged in 24 iterations.}
\text{Log-likelihood: 16.66199}
\text{AIC: -9.323982 AICc: -3.944671}

<table>
<thead>
<tr>
<th>Estimate</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.SJI</td>
<td>0.79555</td>
</tr>
<tr>
<td>A.EBays</td>
<td>0.27540</td>
</tr>
<tr>
<td>A.PSnd</td>
<td>-0.53694</td>
</tr>
<tr>
<td>A.HC</td>
<td>-0.60874</td>
</tr>
<tr>
<td>R.(SJF,SJF)</td>
<td>0.03229</td>
</tr>
<tr>
<td>R.(SJI,SJI)</td>
<td>0.03528</td>
</tr>
<tr>
<td>R.(EBays,EBays)</td>
<td>0.01352</td>
</tr>
<tr>
<td>R.(PSnd,PSnd)</td>
<td>0.01082</td>
</tr>
<tr>
<td>R.(HC,HC)</td>
<td>0.19609</td>
</tr>
<tr>
<td>U.U</td>
<td>0.05270</td>
</tr>
<tr>
<td>Q.Q</td>
<td>0.00604</td>
</tr>
<tr>
<td>x0.x0</td>
<td>6.26676</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSParamCIs to compute CIs and bias estimates.
5.2.8 Two hidden state processes

Here we fit a model with two hidden states (north and south) where observation time series 1-3 are for the north and 4-5 are for the south. We make the hidden state processes independent (meaning a diagonal $Q$ matrix) but with the same process variance. We make the observation errors i.i.d. (the default) and the $u$ elements equal. Mathematically, this is the model:

$$
\begin{bmatrix}
  x_{n,t} \\
  x_{s,t}
\end{bmatrix}
= \begin{bmatrix}
  x_{n,t-1} \\
  x_{s,t-1}
\end{bmatrix}
+ \begin{bmatrix} u \\ u \end{bmatrix}
+ \begin{bmatrix} w_{n,t} \\ w_{s,t} \end{bmatrix},
$$

$w_t \sim \text{MVN} \left(0, \begin{bmatrix} q & 0 \\ 0 & q \end{bmatrix} \right)$

$$
\begin{bmatrix}
  y_{1,t} \\
  y_{2,t} \\
  y_{3,t} \\
  y_{4,t} \\
  y_{5,t}
\end{bmatrix}
= \begin{bmatrix}
  1 & 0 \\
  1 & 0 \\
  1 & 0 \\
  0 & 1 \\
  0 & 1
\end{bmatrix}
\begin{bmatrix}
  x_{n,t} \\
  x_{s,t}
\end{bmatrix}
+ \begin{bmatrix} 0 \\ a_2 \\ a_3 \\ 0 \\ a_5 \end{bmatrix}
+ \begin{bmatrix}
  v_{1,t} \\
  v_{2,t} \\
  v_{3,t} \\
  v_{4,t} \\
  v_{5,t}
\end{bmatrix},
$$

$v_t \sim \text{MVN} \left(0, \begin{bmatrix} r & 0 & 0 & 0 & 0 \\ 0 & r & 0 & 0 & 0 \\ 0 & 0 & r & 0 & 0 \\ 0 & 0 & 0 & r & 0 \\ 0 & 0 & 0 & 0 & r \end{bmatrix} \right)$

To fit the model, use the following code (output not shown):

```r
Z = factor(c("N", "N", "N", "S", "S"))
Q = "diagonal and equal"
U = "equal"

kemfit = MARSS(dat, model=list(Z=Z, Q=Q, U=U))
```

You can also pass in $Z$ exactly as it is in the equation as a numeric matrix

```r
Z = matrix(c(1,1,1,0,0,0,0,0,1,1),5,2)
```

The factor notation is a shortcut for making a design matrix (as $Z$ is in these examples. "equal" is a shortcut meaning all elements in a matrix are constrained to be equal. It can be used for all column matrices ($a$, $u$ and $\pi$). "diagonal and equal" can be used as a shortcut for all square matrices ($B, Q, R, \Lambda$).

5.3 Time-varying parameters

Time-varying parameters are specified by passing in an array of matrices (list, numeric or character) where the 3rd dimension of the array is time and must be the same value as the 2nd (time) dimension of the data matrix. No text shortcuts are allowed for time-varying parameters; you need to use the matrix form.

For example, let’s say we wanted a different $u$ for the first half versus second half of the harbor seal time series. We would pass in an array for $u$ as follows:

```r
U1 = matrix("t1",5,1); U2 = matrix("t2",5,1)
Ut = array(U2, dim=c(dim(U1), dim(dat)[2]))
```
\[ TT = \text{dim(dat)}[2] \]
\[ Ut[,,1:\text{floor}(TT/2)] = U1 \]
\[ \text{kemfit.tv=MARSS(dat,model=list(U=Ut,Q="diagonal and equal"\))} \]

You can have some elements in a parameter matrix be time-constant and some be time-varying:

\[ U1 = \text{matrix}(\text{c(rep("t1",4),"hc"),5,1}); U2 = \text{matrix}(\text{c(rep("t2",4),"hc"),5,1}) \]
\[ Ut = \text{array}(U2,\text{dim}=\text{c(dim(U1),dim(dat)[2])}) \]
\[ Ut[,,1:\text{floor}(TT/2)] = U1 \]
\[ \text{kemfit.tv=MARSS(dat,model=list(U=Ut,Q="diagonal and equal"\))} \]

Note that how the time-varying model is specified for MARSS is the same as you would write the time-varying model on paper in matrix math form.

5.4 Including inputs (or covariates)

In MARSS models with covariates, the covariates are often treated as inputs and appear as either the \( c \) or \( d \) in Equation 5.1, depending on the application. However, more generally, \( c \) and \( d \) are simply inputs that are fully-known (no missing values). \( c_t \) is the \( p \times 1 \) vector of inputs at time \( t \) which affect the states and \( d_t \) is a \( q \times 1 \) vector of inputs (potentially the same as \( c_t \)), which affect the observations.

\( C_t \) is an \( m \times p \) matrix of coefficients relating the effects of \( c_t \) to the \( m \times 1 \) state vector \( x_t \), and \( D_t \) is an \( n \times q \) matrix of coefficients relating the effects of \( d_t \) to the \( n \times 1 \) observation vector \( y_t \). The elements of \( C \) and \( D \) can be estimated, and their form is specified much like the other matrices.

With the \text{MARSS()}\ function, one can fit a model with inputs by simply passing in \text{model$C} and/or \text{model$D} in the \text{MARSS()}\ call as a \( p \times T \) or \( q \times T \) matrix, respectively. The form for \( C_t \) and \( D_t \) is similarly specified by passing in \text{model$C} and/or \text{model$D}. If \( C \) and \( D \) are not time-varying, they are passed in as a 2-dimensional matrix. If they are time-varying, they must be passed in as an 3-dimensional array with the 3rd dimension equal to the number of time steps if they are time-varying.

See Chapter 13 for extended examples of including covariates as inputs in a MARSS model. Also note that it is not necessary to have your covariates appear in \( c \) and/or \( d \). That is a common form, however in some MARSS models, covariates will appear in one of the parameter matrices as fixed values.

5.5 Printing and summarizing models and model fits

The package includes print functions for \text{marssMODEL} objects and \text{marssMLE} objects (fitted models).
print(kemfit)
print(kemfit$model)

This will print the basic information on model structure and model fit that you have seen in the previous examples. The package also includes a summary function for models.

summary(kemfit$model)

Output for the summary function is not shown because it is verbose. It prints each matrix with the fixed elements denoted with their values and the free elements denoted by their names. This is very helpful for confirming exactly what model structure you are fitting to the data.

The print function will also print various other types of output such as a vector of the estimated parameters, the estimated states, the state standard errors, etc. You use the `what` argument in the print call to specify the desired output.

print(kemfit, what="par")

will print the `par` element of a marssMLE object. This will only include the estimated elements in a column matrix. To see the entire parameter matrix with both estimated and fixed values, you can use:

print(kemfit, what="Q")

Parameter matrix Q

```
[,1]   [,2]
[1,] 0.007669608   0.000000000
[2,] 0.000000000  0.007669608
```

Type `?print.MARSS` to see a list of the types of output that can be printed with a `print` call. If you want to use the output from `print` instead of printing to the console, then assign the print call to a value:

```r
x=print(kemfit, what="states",silent=TRUE)
```

5.6 Tidy output

The `augment`, `tidy` and `glance` functions from the broom package will provide summaries as a data.frame for use in further analyses and for passing to `ggplot()`. See `?augment.marssMLE` for examples.

```r
require(broom)
head(augment(kemfit))
```

MARSSresiduals.tT reported warnings. See `msg` element of returned residuals object.

```
   .rownames t y .fitted .resids .sigma
1     SJF 1  6.033086 6.215483 -0.182397213 0.1729465
2     SJF 2   NA     NA       NA       NA
```
### 5.7 Confidence intervals on a fitted model

The function `MARSSparamCIs()` is used to compute confidence intervals with a default alpha level of 0.05. The default is to compute approximate confidence intervals using the Hessian matrix (method="hessian"). Confidence intervals can also be computed via parametric (method="parametric") or non-parametric (method="innovations") bootstrapping. Note, if you want confidence intervals on variances, then it is unwise to use the Hessian approximation as it is symmetric and variances are constrained to be positive.

#### 5.7.1 Approximate confidence intervals from a Hessian matrix

The default method for `MARSSparamCIs` computes approximate confidence intervals using an analytically computed Hessian matrix [Harvey (1989) sec 3.4.5]. The call is:
kem.with.hess.CIs = MARSSparamCIs(kemfit)

See ?MARSShessian for a discussion of the Hessian calculations. Use print or just type the marssMLE object name to see the confidence intervals:

    print(kem.with.hess.CIs)

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 22 iterations.
Log-likelihood: 7.949236
AIC: 0.1015284  AICc: 2.424109

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ML.Est</th>
<th>Std.Err</th>
<th>low.CI</th>
<th>up.CI</th>
<th>Est.Bias</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.SJI</td>
<td>0.79786</td>
<td>0.06152</td>
<td>0.67729</td>
<td>0.9184</td>
<td></td>
</tr>
<tr>
<td>A.EBays</td>
<td>0.27743</td>
<td>0.06253</td>
<td>0.15487</td>
<td>0.4000</td>
<td></td>
</tr>
<tr>
<td>A.HC</td>
<td>-0.07035</td>
<td>0.08882</td>
<td>-0.24442</td>
<td>0.1037</td>
<td></td>
</tr>
<tr>
<td>R.diag</td>
<td>0.03406</td>
<td>0.00671</td>
<td>0.02091</td>
<td>0.0472</td>
<td></td>
</tr>
<tr>
<td>U.1</td>
<td>0.04318</td>
<td>0.01438</td>
<td>0.01499</td>
<td>0.0714</td>
<td></td>
</tr>
<tr>
<td>Q.diag</td>
<td>0.00767</td>
<td>0.00460</td>
<td>-0.00135</td>
<td>0.0167</td>
<td></td>
</tr>
<tr>
<td>x0.N</td>
<td>6.17205</td>
<td>0.13776</td>
<td>5.90205</td>
<td>6.4420</td>
<td></td>
</tr>
<tr>
<td>x0.S</td>
<td>6.20615</td>
<td>0.15708</td>
<td>5.89828</td>
<td>6.5140</td>
<td></td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

CIs calculated at alpha = 0.05 via method=hessian

5.7.2 Confidence intervals from a parametric bootstrap

Use method="parametric" to use a parametric bootstrap to compute confidence intervals and bias using a parametric bootstrap.

    kem.w.boot.CIs=MARSSparamCIs(kemfit,method="parametric",nboot=10)
    #nboot should be more like 1000, but set low for example's sake
    print(kem.w.boot.CIs)

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 22 iterations.
Log-likelihood: 7.949236
AIC: 0.1015284  AICc: 2.424109

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ML.Est</th>
<th>Std.Err</th>
<th>low.CI</th>
<th>up.CI</th>
<th>Est.Bias</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.SJI</td>
<td>0.79786</td>
<td>0.08192</td>
<td>0.6333</td>
<td>0.8744</td>
<td>1.46e-02</td>
</tr>
<tr>
<td>A.EBays</td>
<td>0.27743</td>
<td>0.04407</td>
<td>0.2077</td>
<td>0.3370</td>
<td>2.83e-05</td>
</tr>
<tr>
<td>A.HC</td>
<td>-0.07035</td>
<td>0.07749</td>
<td>-0.1461</td>
<td>0.0650</td>
<td>-2.36e-02</td>
</tr>
</tbody>
</table>
5.8 Vectors of just the estimated parameters

Often it is useful to have a vector of the estimated parameters. For example, if you are writing a call to \texttt{optim}, you will need a vector of just the estimated parameters. You can use the function \texttt{coef}:

\begin{verbatim}
parvec = coef(kemfit, type="vector")
\end{verbatim}

\begin{verbatim}
parvec
\end{verbatim}

```r
A.SJI A.EBays A.HC R.diag
0.79786453 0.27743473 -0.07035021 0.03406192
U.1 Q.diag x0.N x0.S
0.04317640 0.00766961 6.17204803 6.20615469
```

5.9 Kalman filter and smoother output

All the standard Kalman filter and smoother output (along with the lag-one covariance smoother output) is available using the \texttt{MARSSkf} function. Read the help file (\texttt{?MARSSkf}) for details and meanings of the names in the output list.

\begin{verbatim}
kf = MARSSkf(kemfit)
names(kf)
\end{verbatim}

```
[1] "xtT"  "VtT"  "Vtt1T"  "x0T"
[5] "V0T"  "x10T"  "V10T"  "x00T"
```
5.10 Degenerate variance estimates

If your data are short relative to the number of parameters you are estimating, then you are liable to find that some of the variance elements are degenerate (equal to zero). Try the following:

```r
dat.short = dat[1:4,1:10]
kem.degen = MARSS(dat.short, control=list(allow.degen=FALSE))
```

Warning! Abstol convergence only. Maxit (=500) reached before log-log convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
WARNING: Abstol convergence only no log-log convergence.
maxit (=500) reached before log-log convergence.
The likelihood and params might not be at the ML values.
Try setting control$maxit higher.
Log-likelihood: 11.67854
AIC: 2.642914 AICc: 63.30958

<table>
<thead>
<tr>
<th>Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
</tr>
<tr>
<td>U.X.SJF</td>
</tr>
<tr>
<td>U.X.SJI</td>
</tr>
<tr>
<td>U.X.EBays</td>
</tr>
<tr>
<td>U.X.PSnd</td>
</tr>
<tr>
<td>Q.(X.SJF,X.SJF)</td>
</tr>
<tr>
<td>Q.(X.SJI,X.SJI)</td>
</tr>
<tr>
<td>Q.(X.EBays,X.EBays)</td>
</tr>
<tr>
<td>Q.(X.PSnd,X.PSnd)</td>
</tr>
<tr>
<td>x0.X.SJF</td>
</tr>
<tr>
<td>x0.X.SJI</td>
</tr>
<tr>
<td>x0.X.EBays</td>
</tr>
</tbody>
</table>
x0.X.PSnd  5.71e+00
Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

Convergence warnings
Warning: the Q.(X.SJI,X.SJI) parameter value has not converged.
Warning: the Q.(X.EBays,X.EBays) parameter value has not converged.
Warning: the Q.(X.PSnd,X.PSnd) parameter value has not converged.
Type MARSSinfo("convergence") for more info on this warning.

This will print a warning that the maximum number of iterations was reached before convergence of some of the Q parameters. It might be that if you just ran a few more iterations the variances will converge. So first try setting control$maxit higher.

kem.degen2 = MARSS(dat.short, control=list(maxit=1000, allow.degen=FALSE), silent=2)

Output not shown, but if you run the code, you will see that some of the Q terms are still not converging. MARSS can detect if a variance is going to zero and it will try zero to see if that has a higher likelihood. Try removing the allow.degen=FALSE which was turning off this feature.

kem.short = MARSS(dat.short)

Warning! Abstol convergence only. Maxit (=500) reached before log-log convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
WARNING: Abstol convergence only no log-log convergence.
maxit (=500) reached before log-log convergence.
The likelihood and params might not be at the ML values.
Try setting control$maxit higher.
Log-likelihood: 11.6907
AIC: 2.6186  AICc: 63.28527

<table>
<thead>
<tr>
<th>Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
</tr>
<tr>
<td>U.X.SJF</td>
</tr>
<tr>
<td>U.X.SJI</td>
</tr>
<tr>
<td>U.X.EBays</td>
</tr>
<tr>
<td>U.X.PSnd</td>
</tr>
<tr>
<td>Q.(X.SJF,X.SJF)</td>
</tr>
<tr>
<td>Q.(X.SJI,X.SJI)</td>
</tr>
</tbody>
</table>
5.10 Degenerate variance estimates

\[ Q.(X.EBays, X.EBays) \approx 0.00e+00 \]
\[ Q.(X.PSnd, X.PSnd) \approx 3.04e-05 \]
\[ x0.X.SJF \approx 5.96e+00 \]
\[ x0.X.SJI \approx 6.73e+00 \]
\[ x0.X.EBays \approx 6.60e+00 \]
\[ x0.X.PSnd \approx 5.71e+00 \]

Initial states \((x0)\) defined at \(t=0\)

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

Convergence warnings
- Warning: the \(Q.(X.SJI, X.SJI)\) parameter value has not converged.
- Warning: the \(Q.(X.PSnd, X.PSnd)\) parameter value has not converged.
- Type `MARSSinfo("convergence")` for more info on this warning.

So three of the four \(Q\) elements are going to zero. This often happens when you do not have enough data to estimate both observation and process variance.

Perhaps we are trying to estimate too many variances. We can try using only one variance value in \(Q\) and one \(u\) value in \(U\):

```r
kem.small = MARSS(dat.short, model=list(Q="diagonal and equal", U="equal"))
```

Success! abstol and log-log tests passed at 164 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 164 iterations.
Log-likelihood: 11.19
AIC: -8.379994  AICC: 0.9533396

<table>
<thead>
<tr>
<th>Estimate</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
<td>0.0191</td>
</tr>
<tr>
<td>U.1</td>
<td>0.1027</td>
</tr>
<tr>
<td>Q.diag</td>
<td>0.0000</td>
</tr>
<tr>
<td>x0.X.SJF</td>
<td>6.0609</td>
</tr>
<tr>
<td>x0.X.SJI</td>
<td>6.7698</td>
</tr>
<tr>
<td>x0.X.EBays</td>
<td>6.5307</td>
</tr>
<tr>
<td>x0.X.PSnd</td>
<td>5.7451</td>
</tr>
</tbody>
</table>

Initial states \((x0)\) defined at \(t=0\)

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.
5.11 Bootstrap parameter estimates

You can easily produce bootstrap parameter estimates from a fitted model using \texttt{MARSSboot()}:\footnote{\texttt{boot.params = MARSSboot(kemfit, nboot=20, output="parameters", sim="parametric")$boot.params}}

```
progress: |||||||||||||||||||||||||||||||||||||||||||||||||||||
```

Use \texttt{silent=TRUE} to stop the progress bar from printing. The function will also produce parameter sets generated using a Hessian matrix (\texttt{sim="hessian").} or a non-parametric bootstrap (\texttt{sim="innovations").}

5.12 Data simulation

5.12.1 Simulated data from a fitted MARSS model

Data can be simulated from \texttt{marssMLE} object using \texttt{MARSSsimulate()}.\footnote{\texttt{sim.data=MARSSsimulate(kemfit, nsim=2, tSteps=100)$sim.data}}

Then you might want to estimate parameters from that simulated data. Above we created two simulated datasets (\texttt{nsim=2}). We will fit to the first one. Here the default settings for \texttt{MARSS()} are used.

```
  kem.sim.1 = MARSS(sim.data[,1])
```

Then we might like to see the likelihood of the second set of simulated data under the model fit to the first set of data. We do that with the Kalman filter function. This function takes a \texttt{marssMLE} object (as output by say the \texttt{MARSS} function), and we have to replace the data in kem.sim.1 with the second set of simulated data.

```
kem.sim.2 = kem.sim.1
kem.sim.2$model$data = sim.data[,2]
MARSSkf( kem.sim.2 )$logLik
```

\begin{verbatim}
[1] -11.31445
\end{verbatim}
5.13 Bootstrap AIC

The function `MARSSaic()` computes a bootstrap AIC for model selection purposes. Use `output="AICbp"` to produce a parameter bootstrap. Use `output="AICbb"` to produce a non-parametric bootstrap AIC. You will need a large number of bootstraps (`nboot`). We use only 10 bootstraps to show you how to compute AICb with the MARSS package, but the AICbp estimate will be terrible with this few bootstraps.

```r
kemfit.with.AICb = MARSSaic(kemfit, output = "AICbp",
   Options = list(nboot = 10, silent=TRUE))
```

#nboot should be more like 1000, but set low here for example sake

```r
print(kemfit.with.AICb)
```

MARSS fit is

Estimation method: kem

Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001

Estimation converged in 22 iterations.

Log-likelihood: 7.949236

AIC: 0.1015284  AICc: 2.424109  AICbp(param): 291.4368

Estimate
A.SJI 0.79786
A.EBays 0.27743
A.HC -0.07035
R.diag 0.03406
U.1 0.04318
Q.diag 0.00767
x0.N 6.17205
x0.S 6.20615

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

5.14 Convergence

MARSS uses two convergence tests. The first is

\[ \log \text{Lik}_{i+1} - \log \text{Lik}_i < \text{tol} \]

This is called `abstol` (meaning absolute tolerance) in the output. The second is called the `conv.test.slope`. This looks at the slope of the log parameter value (or likelihood) versus log iteration number and asks whether that is close to zero (not changing).
If you are having trouble getting the model to converge, then start by addressing the following 1) Are you trying to fit a bad model, e.g. a non-stationary model fit to stationary data or the opposite or a model that specifies independence of errors or states to data that clearly violate that or a model that implies a particular stationary distribution (particular mean and variance) to data that strongly violate that? 2) Do you have confounded parameters, e.g. two parameters that have the same effect (like effectively two intercepts)?, 3) Are you trying to fit a model to 1 data point somewhere, e.g. in a big multivariate dataset with lots of missing values? 4) How many parameters are you trying to estimate per data point? 5) Check your residuals ($\text{residuals(kemfit)}$) for normality. 6) Did you do any data transformations that would cause one of the variances to go to zero? Replacing 0s with a constant will do that. Try replacing them with NAs (missing). Do you have long strings of constant numbers in your data? Binned data often look like that, and that will drive \( Q \) to 0.
Setting and searching initial conditions

The EM algorithm is very robust to initial starting conditions however before final results are accepted, they should be tested using other initial conditions. Other times you will want to pass in specific initial conditions because the MARSS() function cannot find initial conditions on its own, which is the case for certain models with certain $Z$ matrices in particular or you might want to start with initial conditions at the MLEs of another fit. This chapter shows you how to set initial conditions.

The chapter will also cover using a Monte Carlo search over random initial values. This is a brute force method for finding optimal initial conditions. It simply uses random initial conditions and runs the EM algorithm for a number of iterations and selects the initial conditions with the highest log-likelihood after the given number of iterations. In MARSS versions 3.9 and earlier, there was a utility function to perform a Monte Carlo search. However, it is very hard for the function to come up with reasonable random initial conditions for the wide variety of models that MARSS() can fit. In MARSS version 3.10, the MARSSmcinit() function was removed and replaced with this chapter discussing how to do your own Monte Carlo initial conditions search. The original MARSSmcinit() function is included in the R code with this chapter (see footnote).

6.1 Fitting a model with a new set of initial conditions

Fitting a model with a new set of initial conditions is straight-forward with the MARSS() function. Simply pass in the argument `inits`. This is illustrated with an example from Chapter 13.

We will fit a model with covariates to phytoplankton data:

Type `RShowDoc("Chapter_inits.R",package="MARSS")` at the R command line to open a file with all the code for this chapter and see a copy of the old MARSSmcinit function.
fulldat = lakeWAplanktonTrans
years = fulldat[,\"Year\"]>=1965 & fulldat[,\"Year\"]<1975
dat = t(fulldat[years,c("Greens", "Bluegreens")])
the.mean = apply(dat,1,mean,na.rm=TRUE)
the.sigma = sqrt(apply(dat,1,var,na.rm=TRUE))
dat = (dat-the.mean)*(1/the.sigma)

The covariates for this example are temperature and total phosphorous.
covariates = rbind(
    Temp = fulldat[years,"Temp"],
    TP = fulldat[years,"TP"]
)
# demean the covariates
covariates = covariates-the.mean

We will fit a model where algal abundance is a random walk without drift and where the observation errors are explained by the covariates plus independent unexplained noise:
U = x0 = "zero"
Q = "unconstrained"
d = covariates
A = "zero"
D = "unconstrained"
R = "diagonal and equal"
model.list = list(U=U,Q=Q,A=A,R=R,D=D,d=d,x0=x0)
kem = MARSS(dat, model=model.list)

Success! abstol and log-log tests passed at 72 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 72 iterations.
Log-likelihood: -236.5911
AIC: 489.1822 AICc: 489.8582

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
<td>0.0720</td>
</tr>
<tr>
<td>Q.(1,1)</td>
<td>0.9946</td>
</tr>
<tr>
<td>Q.(2,1)</td>
<td>-0.0290</td>
</tr>
<tr>
<td>Q.(2,2)</td>
<td>0.0976</td>
</tr>
<tr>
<td>D.(Greens,Temp)</td>
<td>0.3572</td>
</tr>
<tr>
<td>D.(Bluegreens,Temp)</td>
<td>0.2537</td>
</tr>
</tbody>
</table>
6.1 Fitting a model with a new set of initial conditions

Standard errors have not been calculated. Use MARSSparamCIs to compute CIs and bias estimates.

The \texttt{inits} argument can be set either as a list that is the same as that from the following:

\begin{verbatim}
coef(kem, what="par")
\end{verbatim}

or as a marssMLE object from previous MARSS() call for a model with the same structure. 'same structure' means the model list used for the \texttt{model} argument is the same and the dimensions of the model (number of states and rows of data) are the same.

6.1.1 Specifying initial conditions as a list

The output from \begin{verbatim}coef(kem, what="par")\end{verbatim} is a list with a column vector of the estimated values for each parameter matrix. Here is the value for $D$ and $Q$:

\begin{verbatim}
out=coef(kem, what="par")
out$D
\end{verbatim}

\begin{verbatim}
[,1]
(Greens,Temp)  0.35722321
(Bluegreens,Temp)  0.25372912
(Greens,TP)  -0.02151303
(Bluegreens,TP)  0.03544925
\end{verbatim}

\begin{verbatim}
out$Q
\end{verbatim}

\begin{verbatim}
[,1]
(1,1)  0.99456586
(2,1) -0.02895620
(2,2)  0.09757775
\end{verbatim}

MARSS() gave names to the $D$ and $Q$ estimated values. It is important to look at the output from \begin{verbatim}coef(., what="par")\end{verbatim} before passing in the \texttt{inits} list so that you know where the parameter values fall in the parameter column vector. For example, note that in the $Q$ column vector, the variance for the first row in $x$ (Greens) is first, the (1,1) element, then the covariance, and last value is the variance of the second row in $x$ (Bluegreens) which appears in the (2,2) element of $Q$.

You can pass in \texttt{inits} for any of the parameters in \begin{verbatim}coef(., what="par")\end{verbatim}. You can pass in either a column vector that is the same size as that output by
setting inits

c coef, so for Q, the column vectors must be 3 × 1, or a scalar. If a scalar, then all values in the par column vector will be set to that value—except for Q, R and Λ which will be set to diagonal matrices with the scalar on the diagonal.

Examples

Pass in an initial value for Q that is a diagonal matrix with 1 on the diagonal.

```r
inits=list(Q=1)
kem = MARSS(dat, model=model.list, inits=inits)
```

Success! abstol and log-log tests passed at 134 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 134 iterations.
Log-likelihood: -236.6064
AIC: 489.2127  AICc: 489.8888

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
<td>0.0686</td>
</tr>
<tr>
<td>Q.(1,1)</td>
<td>1.0079</td>
</tr>
<tr>
<td>Q.(2,1)</td>
<td>-0.0296</td>
</tr>
<tr>
<td>Q.(2,2)</td>
<td>0.1004</td>
</tr>
<tr>
<td>D.(Greens,Temp)</td>
<td>0.3312</td>
</tr>
<tr>
<td>D.(Bluegreens,Temp)</td>
<td>0.2531</td>
</tr>
<tr>
<td>D.(Greens,TP)</td>
<td>-0.0294</td>
</tr>
<tr>
<td>D.(Bluegreens,TP)</td>
<td>0.0345</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

#or

```r
inits=list(Q=matrix(c(1,0,1),3,1))
kem = MARSS(dat, model=model.list, inits=inits)
```

Success! abstol and log-log tests passed at 134 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
6.1 Fitting a model with a new set of initial conditions

Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 134 iterations.
Log-likelihood: -236.6064
AIC: 489.2127 AICc: 489.8888

Estimate

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
<td>0.0686</td>
</tr>
<tr>
<td>Q.(1,1)</td>
<td>1.0079</td>
</tr>
<tr>
<td>Q.(2,1)</td>
<td>-0.0296</td>
</tr>
<tr>
<td>Q.(2,2)</td>
<td>0.1004</td>
</tr>
<tr>
<td>D.(Greens,Temp)</td>
<td>0.3312</td>
</tr>
<tr>
<td>D.(Bluegreens,Temp)</td>
<td>0.2531</td>
</tr>
<tr>
<td>D.(Greens,TP)</td>
<td>-0.0294</td>
</tr>
<tr>
<td>D.(Bluegreens,TP)</td>
<td>0.0345</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

Pass in an initial value for \( Q \) that is this non-diagonal matrix:

\[
\begin{pmatrix}
1 & 0.5 \\
0.5 & 0.7
\end{pmatrix}
\]

\( \text{inits=list(Q=matrix(c(1,0.5,0.7),3,1))} \)
\( \text{kem = MARSS(dat, model=model.list, inits=inits)} \)

Pass in an initial value for \( Q \) and \( D \):

\( \text{inits=list(Q=matrix(c(1,0.5,0.7),3,1), D=1)} \)
\( \text{kem = MARSS(dat, model=model.list, inits=inits)} \)

The initial values for \( D \) will be all 1s.

Pass in an initial value for \( D \) set to the value from a previous fit but use default inits for everything else:

\( \text{inits=list(D=coef(kem, what="par")$D)} \)
\( \text{kem = MARSS(dat, model=model.list, inits=inits)} \)

6.1.2 Specifying initial conditions using output from another fit

You can also use a MARSS() fit as an initial condition. The model must be the same structure. This is typically used when you want to use an EM fit as a start for a BFGS fit if you are using BFGS for the final MLE search.

You can pass in the inits as a list using the coef() function.

\( \text{# create the par list from the output} \)
\( \text{inits=coef(kem, what="par")} \)
\( \text{kem.bfgs = MARSS(dat, model=model.list, inits=inits, method="BFGS")} \)
Or you can pass in a MARSS() fit. This is a shortcut for the above call.

```r
# create the par list from the output
kem.bfgs = MARSS(dat, model=model.list, inits=kem, method="BFGS")
```

### 6.2 Searching across initial values using a Monte Carlo routine

The EM algorithm is a hill-climbing algorithm and like all hill-climbing algorithms it can get stuck on local maxima and ridges. There are a number of approaches to doing a pre-search of the initial conditions space, but a brute force random Monte Carlo search appears to work well (Biernacki et al., 2003).

It is slow, but normally sufficient. In our papers on the distributional properties of MARSS parameter estimates, we rarely found that an initial conditions search changed the estimates—except in cases where $Z$ and $B$ are estimated as unconstrained or as the fraction of missing data in the data set became large. Regardless an initial conditions search should be done before reporting final estimates for an analysis$^1$.

The idea behind a Monte Carlo search of initial conditions is simple. One simply randomly generates initial conditions, runs the EM algorithm a few iterations (10-20), and save the log-likelihood at the end of those iterations. The starting initial conditions is selected from that giving the lowest log-likelihood.

The R code included for this chapter includes a function that will do a simple Monte Carlo search using a fitted MARSS model (output from a MARSS() call) and drawing random initial conditions from a uniform distribution or a Wishart distribution for the variance-covariance matrices. The function will not work for all MARSS models but will give you a starting point for setting up your own Monte Carlo search. The function uses a control list to set `numInits`, the number of random initial value draws, `numInitSteps`, the maximum number of EM iterations for each random initial value draw, and `boundsInits`, the bounds for the random distributions. It outputs a list with specifying the initial values that give the lowest log-likelihood.

Here is a simple example of using the function. `numInits` is set low so that the example runs quickly.

```r
dat = t(harborSeal)
dat = dat[c(2,nrow(dat)),]
fit1=MARSS(dat)
```

Success! abstol and log-log tests passed at 59 iterations.

Alert: conv.test.slope.tol is 0.5.

Test with smaller values (<0.1) to ensure convergence.

---

$^1$ It is also a good idea to try `method=BFGS` to see if this changes the estimates.
6.2 Searching across initial values using a Monte Carlo routine

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 59 iterations.
Log-likelihood: 11.68334
AIC: -9.366688 AICc: -5.366688

<table>
<thead>
<tr>
<th>Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
</tr>
<tr>
<td>U.X.CoastalEstuaries</td>
</tr>
<tr>
<td>U.X.Georgia.Strait</td>
</tr>
<tr>
<td>Q.(X.CoastalEstuaries,X.CoastalEstuaries)</td>
</tr>
<tr>
<td>Q.(X.Georgia.Strait,X.Georgia.Strait)</td>
</tr>
<tr>
<td>x0.X.CoastalEstuaries</td>
</tr>
<tr>
<td>x0.X.Georgia.Strait</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

```r
MCinits = MARSSmcinit(fit1, control=list(numInits = 10))

> Starting Monte Carlo Initializations
Progress: |||||||||||||||||||||||||||||||||||||||||||||||||||||||||||||||||||||||

fit2=MARSS(dat, inits=MCinits)

Success! abstol and log-log tests passed at 52 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 52 iterations.
Log-likelihood: 11.68371
AIC: -9.367418 AICc: -5.367418

<table>
<thead>
<tr>
<th>Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
</tr>
<tr>
<td>U.X.CoastalEstuaries</td>
</tr>
<tr>
<td>U.X.Georgia.Strait</td>
</tr>
<tr>
<td>Q.(X.CoastalEstuaries,X.CoastalEstuaries)</td>
</tr>
<tr>
<td>Q.(X.Georgia.Strait,X.Georgia.Strait)</td>
</tr>
<tr>
<td>x0.X.CoastalEstuaries</td>
</tr>
</tbody>
</table>
x0.X.Georgia.Strait 8.40874
Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.
Applications
In this part, we walk you through some longer analyses using MARSS models for a variety of different applications. Most of these are analyses of ecological data, but the same models are used in many other fields. These longer examples will take you through both the conceptual steps (with pencil and paper) and a R step which translates the conceptual model into code.

Set-up

- If you haven’t already, install the MARSS package. See directions on the CRAN webpage (http://cran.r-project.org/) for instructions on installing packages. You will need write permissions for your R program directories to install packages. See the help pages on CRAN for workarounds if you don’t have write permission.
- Type in `library(MARSS)` at the R command line to load the package after you install it.

Tips

- `summary(foo$model)`, where `foo` is a fitted model object, will print detailed information on the structure of the MARSS model that was fit in the call `foo = MARSS(logdata)`. This allows you to double check the model you fit. `print(foo)` will print a ‘English’ version of the model structure along with the parameter estimates.
- When you run `MARSS()`, it will output the number of iterations used. If you reached the maximum, re-run with `control=list(maxit=...)` set higher than the default.
- If you mis-specify the model, `MARSS()` will post an error that should give you an idea of the problem (make sure `silent=FALSE` to see full error reports). Remember, the number of rows in your data is \( n \), time is across the columns, and the length of the vector of factors passed in for `model$Z` must be \( n \) while the number of unique factors must be \( m \), the number of \( x \) hidden state trajectories in your model.
- The missing value indicator is NA.
- Running `MARSS(data)`, with no arguments except your data, will fit a MARSS model with \( m = n \), a diagonal \( Q \) matrix with \( m \) variances, and i.i.d. observation errors.
- Try `MARSSinfo()` at the command line if you get errors or warnings you don’t understand. You might find insight there. Or look at the warnings and errors notes in the appendix of this user guide.
Count-based population viability analysis (PVA) using corrupted data

7.1 Background

Estimates of extinction and quasi-extinction risk are an important risk metric used in the management and conservation of endangered and threatened species. By necessity, these estimates are based on data that contain both variability due to real year-to-year changes in the population growth rate (process errors) and variability in the relationship between the true population size and the actual count (observation errors). Classic approaches to extinction risk assume the data have only process error, i.e. no observation error. In reality, observation error is ubiquitous both because of the sampling variability and also because of year-to-year (and day-to-day) variability in sightability.

In this application, we will fit a univariate (meaning one time series) state-space model to population count data with observation error. We will compute the extinction risk metrics given in Dennis et al. (1991), however instead of using a process-error only model (as is done in the original paper), we use a model with both process and observation error. The risk metrics and their interpretations are the same as in Dennis et al. (1991). The only real difference is how we compute \( \sigma^2 \), the process error variance. However this difference has a large effect on our risk estimates, as you will see.

We use here a density-independent model, a stochastic exponential growth model in log space. This equivalent to a MARSS model with \( B = 1 \). Density-independence is often a reasonable assumption when doing a population viability analysis because we do such calculations for at-risk populations that are either declining or that are well below historical levels (and presumably carrying capacity). In an actual population viability analysis, it is necessary to justify this assumption and if there is reason to doubt the assumption,
one tests for density-dependence (Taper and Dennis, 1994) and does sensitivity analyses using state-space models with density-dependence (Dennis et al., 2006).

The univariate model is written:

\[ x_t = x_{t-1} + u + w_t \quad \text{where } w_t \sim N(0, \sigma^2) \quad (7.1) \]

\[ y_t = x_t + v_t \quad \text{where } v_t \sim N(0, \eta^2) \quad (7.2) \]

where \( y_t \) is the logarithm of the observed population size at time \( t \), \( x_t \) is the unobserved state at time \( t \), \( u \) is the growth rate, and \( \sigma^2 \) and \( \eta^2 \) are the process and observation error variances, respectively. In the R code to follow, \( \sigma^2 \) is denoted \( Q \) and \( \eta^2 \) is denoted \( R \) because the functions we are using are also for multivariate state-space models and those models use \( Q \) and \( R \) for the respective variance-covariance matrices.

### 7.2 Simulated data with process and observation error

We will start by using simulated data to see the difference between data and estimates from a model with process error only versus a model that also includes observation error. For our simulated data, we used a decline of 5\% per year, process variability of 0.02 (typical for small to medium-sized vertebrates), and a observation variability of 0.05 (which is a bit on the high end). We’ll randomly set 10\% of the values as missing. Here is the code:

First, set things up:

```r
sim.u = -0.05 # growth rate
sim.Q = 0.02 # process error variance
sim.R = 0.05 # non-process error variance
nYr = 50 # number of years of data to generate
fracmissing = 0.1 # fraction of years that are missing
init = 7 # log of initial pop abundance
years = seq(1:nYr) # sequence 1 to nYr
x = rep(NA,nYr) # replicate NA nYr times
y = rep(NA,nYr)
```

Then generate the population sizes using Equation 7.1:

```r
x[1]=init
for(t in 2:nYr){
  x[t] = x[t-1] + sim.u + rnorm(1,mean=0,sd=sqrt(sim.Q))
}
```

Lastly, add observation error using Equation 7.2 and then add missing values:

```r
for(t in 1:nYr){
  y[t] = x[t] + rnorm(1,mean=0,sd=sqrt(sim.R))
}
```
missYears = sample(years[2:(nYr-1)],floor(fracmissing*nYr), replace = FALSE)
y[missYears]=NA

Stochastic population trajectories show much variation, so it is best to look at a few simulated data sets at once. In Figure 7.1, nine simulations from the identical parameters are shown.

![Stochastic Population Trajectories](image)

**Fig. 7.1.** Plot of nine simulated population time series with process and observation error. Circles are observation and the dashed line is the true population size.

**Example 7.1 (The effect of parameter values on parameter estimates)**

A good way to get a feel for reasonable $\sigma^2$ values is to generate simulated data and look at the time series. A biologist would have a pretty good idea of what kind of year-to-year population changes are reasonable for their study species. For example for many large mammalian species, the maximum population yearly increase would be around 50% (the population could go from 1000 to
1500 in one year), but some of fish species could easily double or even triple in a really good year. Observed data may bounce around a lot for many different reasons having to do with sightability, sampling error, age-structure, etc., but the underlying population trajectory is constrained by the kinds of year-to-year changes in population size that are biologically possible. $\sigma^2$ describes those true population changes.

You should run the example code several times using different parameter values to get a feel for how different the time series can look based on identical parameter values. You can cut and paste from the pdf into the R command line. Typical vertebrate $\sigma^2$ values are 0.002 to 0.02, and typical $\eta^2$ values are 0.005 to 0.1. A $u$ of -0.01 translates to an average 1% per year decline and a $u$ of -0.1 translates to an average 10% per year decline (approximately).

---

**Example 7.1 code**

```r
par(mfrow=c(3,3))
sim.u = -0.05
sim.Q = 0.02
sim.R = 0.05
nYr= 50
fracmiss = 0.1
init = 7
years = seq(1:nYr)
for(i in 1:9){  
x = rep(NA,nYr)  # vector for ts w/o measurement error
y = rep(NA,nYr)  # vector for ts w/ measurement error
x[1]=init
for(t in 2:nYr){
  x[t] = x[t-1]+ sim.u + rnorm(1, mean=0, sd=sqrt(sim.Q))
}
for(t in 1:nYr){
  y[t]= x[t] + rnorm(1,mean=0,sd=sqrt(sim.R))
}
missYears = sample(years[2:(nYr-1)],floor(fracmiss*nYr),replace = FALSE)
y[missYears]=NA
plot(years, y,
  xlab="",ylab="log abundance",lwd=2,bty="l")
lines(years,x,type="l",lwd=2,lty=2)
title(paste("simulation ",i))
}
legend("topright", c("Observed","True"),
  lty = c(-1, 2), pch = c(1, -1))
```
7.3 Maximum-likelihood parameter estimation

7.3.1 Model with process and observation error

Using the simulated data, we estimate the parameters, \( u, \sigma^2, \) and \( \eta^2 \), and the hidden population sizes. These are the estimates using a model with process and observation variability. The function call is

\[
\text{kem} = \text{MARSS}(\text{data})
\]

where \( \text{data} \) is a vector of logged (base e) counts with missing values denoted by NA. After this call, the maximum-likelihood parameter estimates are shown with \( \text{coef}(\text{kem}) \). There are numerous other outputs from the \( \text{MARSS()} \) function. To get a list of the standard model output available type in \( \text{?print.MARSS} \). Note that \( \text{kem} \) is just a name; the output could have been called \( \text{foo} \). Here’s code to fit to the simulated time series:

\[
\text{kem} = \text{MARSS}(y)
\]

Let’s look at the parameter estimates for the nine simulated time series in Figure 7.1 to get a feel for the variation. The \( \text{MARSS()} \) function was used on each time series to produce parameter estimate for each simulation. The estimates are followed by the mean (over the nine simulations) and the true values:

<table>
<thead>
<tr>
<th>Sim</th>
<th>kem.U</th>
<th>kem.Q</th>
<th>kem.R</th>
</tr>
</thead>
<tbody>
<tr>
<td>sim 1</td>
<td>-0.05617</td>
<td>0.02310128</td>
<td>0.07715297</td>
</tr>
<tr>
<td>sim 2</td>
<td>-0.04806465</td>
<td>0.0000000000</td>
<td>0.07556365</td>
</tr>
<tr>
<td>sim 3</td>
<td>-0.04764285</td>
<td>0.031376280</td>
<td>0.04995952</td>
</tr>
<tr>
<td>sim 4</td>
<td>-0.05811163</td>
<td>0.0000000000</td>
<td>0.05812225</td>
</tr>
<tr>
<td>sim 5</td>
<td>-0.08248527</td>
<td>0.026518617</td>
<td>0.07384139</td>
</tr>
<tr>
<td>sim 6</td>
<td>-0.05123457</td>
<td>0.0000000000</td>
<td>0.07918993</td>
</tr>
<tr>
<td>sim 7</td>
<td>-0.08101421</td>
<td>0.012375455</td>
<td>0.05964581</td>
</tr>
<tr>
<td>sim 8</td>
<td>-0.03909302</td>
<td>0.007472725</td>
<td>0.01955799</td>
</tr>
<tr>
<td>sim 9</td>
<td>-0.05187126</td>
<td>0.010529914</td>
<td>0.05741644</td>
</tr>
<tr>
<td>mean sim</td>
<td>-0.05729874</td>
<td>0.012374891</td>
<td>0.06116111</td>
</tr>
<tr>
<td>true</td>
<td>-0.05000000</td>
<td>0.0200000000</td>
<td>0.05000000</td>
</tr>
</tbody>
</table>

As expected, the estimated parameters do not exactly match the true parameters, but the average should be fairly close (although nine simulations is a small sample size). Also note that although we do not get \( u \) quite right, our estimates are usually negative. Thus our estimates usually indicate declining dynamics. Some of the \( \text{kem.Q} \) estimates may be 0. This means that the maximum-likelihood estimate that the data are generated by is a process with no environment variation and only observation error.

The MARSS model fit also gives an estimate of the true population size with observation error removed. This is in \( \text{kem$states} \). Figure 7.2 shows the estimated true states of the population over time as a solid line. Note that the solid line is considerably closer to the actual true states (dashed line) than the observations. On the other hand with certain datasets, the estimates can be quite wrong as well!
7.3.2 Model with no observation error

We used the MARSS model to estimate the mean population rate $u$ and process variability $\sigma^2$ under the assumption that the count data have observation error. However, the classic approach to this problem, referred to as the “Dennis model” ([Dennis et al., 1991]), uses a model that assumes the data have no observation error (a MAR model); all the variability in the data is assumed to result from process error. This approach works well if the observation error in the data is low, but not so well if the observation error is high. We will next fit the data using the classic approach so that we can compare and contrast parameter estimates from the different methods.

Using the estimation method in [Dennis et al., 1991], our data need to be re-specified as the observed population changes ($\delta.pop$) between censuses along with the time between censuses ($\tau$). We re-specify the data as follows:

```r
den.years = years[!is.na(y)]  # the non missing years
den.y = y[!is.na(y)]  # the non missing counts
```
7.3 Maximum-likelihood parameter estimation

\[ \text{den.n.y} = \text{length(den.years)} \]
\[ \text{delta.pop} = \text{rep(NA, den.n.y-1)} \] # population transitions
\[ \text{tau} = \text{rep(NA, den.n.y-1)} \] # step sizes
for (i in 2:den.n.y){
    \[ \text{delta.pop}[i-1] = \text{den.y}[i] - \text{den.y}[i-1] \]
    \[ \text{tau}[i-1] = \text{den.years}[i] - \text{den.years}[i-1] \]
} # end i loop

Next, we regress the changes in population size between censuses (\text{delta.pop}) on the time between censuses (\text{tau}) while setting the regression intercept to 0. The slope of the resulting regression line is an estimate of \( u \), while the variance of the residuals around the line is an estimate of \( \sigma^2 \). The regression is shown in Figure 7.3. Here is the code to do that regression:

\[ \text{den91} \leftarrow \text{lm(delta.pop} \sim \text{-1 + tau)} \]
# note: the "-1" specifies no intercept
\[ \text{den91.u} = \text{den91$coefficients} \]
\[ \text{den91.Q} = \text{var(resid(den91))} \]
#type ?lm to learn about the linear regression function in R
#form is lm(dependent.var \sim response.var1 + response.var2 + ...)
#type summary(den91) to see other info about our regression fit

Here are the parameter values for the data in Figure 7.2 using the process-error only model:

\[
\begin{array}{ccc}
\text{sim} & \text{den91.U} & \text{den91.Q} \\
1 & -0.05503434 & 0.18996075 \\
2 & -0.03488140 & 0.13338948 \\
3 & -0.04358910 & 0.14493459 \\
4 & -0.06667344 & 0.11338246 \\
5 & -0.08998937 & 0.18480318 \\
6 & -0.04311427 & 0.10186497 \\
7 & -0.07546359 & 0.12511394 \\
8 & -0.04217965 & 0.05024078 \\
9 & -0.06165750 & 0.13467394 \\
\text{mean sim} & -0.05684252 & 0.13092934 \\
\text{true} & -0.05000000 & 0.02000000 \\
\end{array}
\]

Notice that the \( u \) estimates are similar to those from MARSS model, but the \( \sigma^2 \) estimate (\( Q \)) is much larger. That is because this approach treats all the variance as process variance, so any observation variance in the data is lumped into process variance (in fact it appears as an additional variance of twice the observation variance).

Example 7.2 (The variability in parameter estimates)
Fig. 7.3. The regression of $\log(N_{t+\tau}) - \log(N_t)$ against $\tau$. The slope is the estimate of $u$ and the variance of the residuals is the estimate of $\sigma^2$. The regression is constrained to go through (0,0).

In this example, you will look at how variable the parameter estimates are by generating multiple simulated data sets and then estimating parameter values for each. You’ll compare the MARSS estimates to the estimates using a process error only model (i.e. ignoring the observation error).

Run the example code a few times to compare the estimates using a state-space model (`kem`) versus the model with no observation error (`den91`). You can copy and paste the code from the pdf file into R. Next, change the observation variance in the code, `sim.R`, in the data generation step in order to get a feel for the estimation performance as observations are further corrupted. What happens as observation error is increased? Next, decrease the number of years of data, `nYr`, and re-run the parameter estimation. What is the effect of fewer years of data? If you find that the example code takes too long to run, reduce the number of simulations (by reducing `nsim` in the code).
Example 7.2 code

```r
sim.u = -0.05 # growth rate
sim.Q = 0.02 # process error variance
sim.R = 0.05 # non-process error variance
nYr= 50 # number of years of data to generate
fracmiss = 0.1 # fraction of years that are missing
init = 7 # log of initial pop abundance (~1100 individuals)
nsim = 9
years = seq(1:nYr) # col of years
params = matrix(NA, nrow=(nsim+2), ncol=5,
dimnames=list(c(paste("sim",1:nsim),"mean sim","true"),
x.ts = matrix(NA,nrow=nsim,ncol=nYr) # ts w/o measurement error
y.ts = matrix(NA,nrow=nsim,ncol=nYr) # ts w/ measurement error
for(i in 1:nsim){
x.ts[i,1]=init
for(t in 2:nYr){
x.ts[i,t] = x.ts[i,t-1]+sim.u+rnorm(1,mean=0,sd=sqrt(sim.Q))
} # end for
for(t in 1:nYr){
y.ts[i,t] = x.ts[i,t]+rnorm(1,mean=0,sd=sqrt(sim.R))
} # end for
missYears = sample(years[2:(nYr-1)], floor(fracmiss*nYr), replace = FALSE)
y.ts[i,missYears]=NA

#MARSS estimates
kem = MARSS(y.ts[i,], silent=TRUE)
type=vector outputs the estimates as a vector instead of a list
params[i,c(1,3,4)] = coef(kem,type="vector")[c(2,3,1)]

#Dennis et al 1991 estimates
den.years = years[!is.na(y.ts[i,])] # the non missing years
den.yts = y.ts[i,!is.na(y.ts[i,])] # the non missing counts
den.n.yts = length(den.years)
delta.pop = rep(NA, den.n.yts-1) # transitions
tau = rep(NA, den.n.yts-1) # time step lengths
for (t in 2:den.n.yts ){
delta.pop[t-1] = den.yts[t] - den.yts[t-1] # transitions
tau[t-1] = den.years[t]-den.years[t-1] # time step length
} # end i loop
den91 <- lm(delta.pop ~ -1 + tau) # -1 specifies no intercept
params[i,c(2,5)] = c(den91$coefficients, var(resid(den91)))
}
params[nsim+1,]=apply(params[1:nsim,],2,mean)
params[nsim+2,]=c(sim.u,sim.u,sim.Q,sim.R,sim.Q)
```
Here is an example of the output from the Example 7.2 code:

```
print(params,digits=3)
```

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>sim 1</td>
<td>-0.0668</td>
<td>-0.050894</td>
<td>0.03371</td>
<td>0.0400</td>
<td>0.1235</td>
</tr>
<tr>
<td>sim 2</td>
<td>-0.0666</td>
<td>-0.051231</td>
<td>0.04121</td>
<td>0.0287</td>
<td>0.1031</td>
</tr>
<tr>
<td>sim 3</td>
<td>-0.0551</td>
<td>-0.042510</td>
<td>0.02386</td>
<td>0.0570</td>
<td>0.1452</td>
</tr>
<tr>
<td>sim 4</td>
<td>-0.0384</td>
<td>-0.029270</td>
<td>0.03573</td>
<td>0.0338</td>
<td>0.1002</td>
</tr>
<tr>
<td>sim 5</td>
<td>-0.0357</td>
<td>-0.048896</td>
<td>0.01202</td>
<td>0.0650</td>
<td>0.1340</td>
</tr>
<tr>
<td>sim 6</td>
<td>-0.0532</td>
<td>-0.030510</td>
<td>0.00816</td>
<td>0.0345</td>
<td>0.0805</td>
</tr>
<tr>
<td>sim 7</td>
<td>-0.0311</td>
<td>-0.024622</td>
<td>0.00637</td>
<td>0.0713</td>
<td>0.1602</td>
</tr>
<tr>
<td>sim 8</td>
<td>0.0104</td>
<td>0.000885</td>
<td>0.00000</td>
<td>0.0662</td>
<td>0.1285</td>
</tr>
<tr>
<td>sim 9</td>
<td>-0.0405</td>
<td>-0.041409</td>
<td>0.03923</td>
<td>0.0200</td>
<td>0.0972</td>
</tr>
<tr>
<td>mean sim</td>
<td>-0.0419</td>
<td>-0.035384</td>
<td>0.02225</td>
<td>0.0463</td>
<td>0.1192</td>
</tr>
<tr>
<td>true</td>
<td>-0.0500</td>
<td>-0.050000</td>
<td>0.02000</td>
<td>0.0500</td>
<td>0.0200</td>
</tr>
</tbody>
</table>

7.4 Probability of hitting a threshold \( \Pi(x_d, t_e) \)

A common extinction risk metric is ‘the probability that a population will hit a certain threshold \( x_d \) within a certain time frame \( t_e \) — if the observed trends continue’. In practice, the threshold used is not \( N_e = 1 \), which would be true extinction. Often a ‘functional’ extinction threshold will be used (\( N_e >> 1 \)). Other times a threshold representing some fraction of current levels is used. The latter is used because we often have imprecise information about the relationship between the true population size and what we measure in the field; that is, many population counts are index counts. In these cases, one must use ‘fractional declines’ as the threshold. Also, extinction estimates that use an absolute threshold (like 100 individuals) are quite sensitive to error in the estimate of true population size. Here, we are going to use fractional declines as the threshold, specifically \( p_d = 0.1 \) which means a 90% decline.

The probability of hitting a threshold, denoted \( \Pi(x_d, t_e) \), is typically presented as a curve showing the probabilities of hitting the threshold (y-axis) over different time horizons (\( t_e \)) on the x-axis. Extinction probabilities can be computed through Monte Carlo simulations or analytically using Equation 16 in Dennis et al. (1991) (note there is a typo in Equation 16; the last + is supposed to be a −). We will use the latter method:

\[
\Pi(x_d, t_e) = \pi(u) \times \Phi \left( \frac{-x_d + |u| t_e}{\sqrt{\sigma^2 t_e}} \right) + \exp(2x_d|u|/\sigma^2)\Phi \left( \frac{-x_d - |u| t_e}{\sqrt{\sigma^2 t_e}} \right)
\] (7.3)
where \( x_e \) is the threshold and is defined as \( x_e = \log(N_0/N_e) \). \( N_0 \) is the current population estimate and \( N_e \) is the threshold. If we are using fractional declines then \( x_e = \log(N_0/(p d \times N_0)) = - \log(pd) \). \( \pi(u) \) is the probability that the threshold is eventually hit (by \( t_e = \infty \)). \( \pi(u) = 1 \) if \( u \leq 0 \) and \( \pi(u) = \exp(-2ux_d/\sigma^2) \) if \( u > 0 \). \( \Phi() \) is the cumulative probability distribution of the standard normal (mean = 0, sd = 1).

Here is the R code for that computation:

```r
pd = 0.1 # means a 90 percent decline
tyrs = 1:100
xd = -log(pd)
p.ever = ifelse(u<=0,1,exp(-2*u*xd/Q)) #Q=sigma2
for (i in 1:100){
  Pi[i] = p.ever * pnorm((-xd+abs(u)*tyrs[i])/sqrt(Q*tyrs[i])) +
    exp(2*xd*abs(u)/Q)*pnorm((-xd-abs(u)*tyrs[i])/sqrt(Q*tyrs[i]))
}
```

Figure 7.4 shows the estimated probabilities of hitting the 90% decline for the nine 30-year times series simulated with \( u = -0.05 \), \( \sigma^2 = 0.01 \) and \( \eta^2 = 0.05 \). The dashed line shows the estimates using the MARSS parameter estimates and the solid line shows the estimates using a process-error only model (the den91 estimates). The circles are the true probabilities. The difference between the estimates and the true probabilities is due to errors in \( \hat{u} \). Those errors are due largely to process error—not observation error. As we saw earlier, by chance population trajectories with a \( u < 0 \) will increase, even over a 50-year period. In this case, \( \hat{u} \) will be positive when in fact \( u < 0 \).

Looking at the figure, it is obvious that the probability estimates are highly variable. However, look at the first panel. This is the average estimate (over nine simulations). Note that on average (over nine simulations), the estimates are good. If we had averaged over 1000 simulations instead of nine, you would see that the MARSS line falls on the true line. It is an unbiased predictor. While that may seem a small consolation if estimates for individual simulations are all over the map, it is important for correctly specifying our uncertainty about our estimates. Second, rather than focusing on how the estimates and true lines match up, see if there are any types of forecasts that seem better than others. For example, are 20-year predictions better than 50-year and are 100-year forecasts better or worse. In Example 7.3, you will remake this figure with different \( u \). You’ll discover from that forecasts are more certain for populations that are declining faster.

Example 7.3 (The effect of parameter values on risk estimates)

In this example, you will recreate Figure 7.4 using different parameter values. This will give you a feel for how variability in the data and population pro-
cess affect the risk estimates. You’ll need to run the Example 7.2 code before running the Example 7.3 code.

Begin by changing sim.R and rerunning the Example 7.2 code. Now run the Example 7.3 code and generate parameter estimates. When are the estimates using the process-error only model (den91) worse and in what way are they worse? You might imagine that you should always use a model that includes observation error, since in practice observations are never perfect. However, there is a cost to estimating that extra variance parameter and the cost is a more variable $\sigma^2$ (Q) estimate. Play with shortening the time series and decreasing the sim.R values. Are there situations when the ‘cost’ of the extra parameter is greater than the ‘cost’ of ignoring observation error?

Next change the rate of decline in the simulated data. To do this, rerun the Example 7.2 code using a lower sim.u; then run the Example 7.3 code. Do the estimates seem better or worse for rapidly declining populations? Rerun the Example 7.2 code using fewer number of years (nYr smaller) and increase
Run the Example 7.3 code again. The graphs will start to look peculiar. Why do you think it is doing that? Hint: look at the estimated parameters.

Last change the extinction threshold (\( p_d \) in the Example 7.3 code). How does changing the extinction threshold change the extinction probability curves? Do not remake the data, i.e. don’t rerun the Example 7.2 code.
Example 7.3 code

# Needs Example 2 to be run first
par(mfrow=c(3,3))
pd = 0.1; xd = -log(pd)  # decline threshold
te = 100; tyrs = 1:te  # extinction time horizon
for(j in c(10,1:8)){
  real.ex = denn.ex = kal.ex = matrix(nrow=te)
  # MARSS parameter estimates
  u=params[j,1]; Q=params[j,3]
  if(Q==0) Q=1e-4  # just so the extinction calc doesn't choke
  p.ever = ifelse(u<=0,1,exp(-2*u*xd/Q))
  for (i in 1:100){
    if(is.finite(exp(2*xd*abs(u)/Q))){
      sec.part = exp(2*xd*abs(u)/Q)*pnorm((-xd-abs(u)*tyrs[i])/sqrt(Q*tyrs[i]))
    }else sec.part=0
    kal.ex[i]=p.ever*pnorm((-xd+abs(u)*tyrs[i])/sqrt(Q*tyrs[i]))+sec.part
  }  # end i loop

  # Dennis et al 1991 parameter estimates
  u=params[j,2]; Q=params[j,5]
  p.ever = ifelse(u<=0,1,exp(-2*u*xd/Q))
  for (i in 1:100){
    denn.ex[i]=p.ever*pnorm((-xd+abs(u)*tyrs[i])/sqrt(Q*tyrs[i]))+
    exp(2*xd*abs(u)/Q)*pnorm((-xd-abs(u)*tyrs[i])/sqrt(Q*tyrs[i]))
  }  # end i loop

  # True parameter values
  u=sim.u; Q=sim.Q
  p.ever = ifelse(u<=0,1,exp(-2*u*xd/Q))
  for (i in 1:100){
    real.ex[i]=p.ever*pnorm((-xd+abs(u)*tyrs[i])/sqrt(Q*tyrs[i]))+
    exp(2*xd*abs(u)/Q)*pnorm((-xd-abs(u)*tyrs[i])/sqrt(Q*tyrs[i]))
  }  # end i loop

  # plot it
  plot(tyrs, real.ex, xlab="time steps into future",
       ylab="probability of extinction", ylim=c(0,1), bty="l")
  if(j<=8) title(paste("simulation ",j) )
  if(j==10) title("average over sims")
  lines(tyrs,denn.ex,type="l",col="red",lwd=2,lty=1)
  lines(tyrs,kal.ex,type="l",col="green",lwd=2,lty=2)
}
legend("bottomright",c("True","Dennis","KalmanEM"),pch=c(1,-1,-1),
        col=c(1,2,3),lty=c(-1,1,2),lwd=c(-1,2,2),bty="n")
7.5 Certain and uncertain regions

From Example 7.3, you have observed one of the problems with estimates of the probability of hitting thresholds. Looking over the nine simulations, your risk estimates will be on the true line sometimes and other times they are way off. So your estimates are variable and one should not present only the point estimates of the probability of 90% decline. At the minimum, confidence intervals need to be added (next section), but even with confidence intervals, the probability of hitting declines often does not capture our certainty and uncertainty about extinction risk estimates.

From Example 7.3, you might have also noticed that there are some time horizons (10, 20 years) for which the estimate are highly certain (the threshold is never hit), while for other time horizons (30, 50 years) the estimates are all over the map. Put another way, you may be able to say with high confidence that a 90% decline will not occur between years 1 to 20 and that by year 100 it most surely will have occurred. However, between the years 20 and 100, you are very uncertain about the risk. The point is that you can be certain about some forecasts while at the same time being uncertain about other forecasts.

One way to show this is to plot the uncertainty as a function of the forecast, where the forecast is defined in terms of the forecast length (number of years) and forecasted decline (percentage). Uncertainty is defined as how much of the 0-1 range your 95% confidence interval covers. Ellner and Holmes (2008) show such a figure (their Figure 1). Figure 7.5 shows a version of this figure that you can produce with the function `CSEGtmufigure(u= val, N= val, s2p= val)`. For the figure, the values $u = -0.05$ which is a 5% per year decline, $N = 25$ so 25 years between the first and last census, and $s^2_p = 0.01$ are used. The process variability for big mammals is typically in the range of 0.002 to 0.02.

Example 7.4 (Uncertain and certain regions)

Use the Example 7.4 code to re-create Figure 7.5 and get a feel for when risk estimates are more certain and when they are less certain. $N$ are the number of years of data, $u$ is the mean population growth rate, and $s^2_p$ is the process variance.

Example 7.4 code

```r
par(mfrow = c(1, 1))
CSEGtmufigure(N = 50, u = -0.05, s2p = 0.02)
```
Fig. 7.5. This figure shows your region of high uncertainty (dark gray). In this region, the minimum 95% confidence intervals (meaning if you had no observation error) span 80% of the 0 to 1 probability. That is, you are uncertain if the probability of a specified decline is close to 0 or close to 1. The white area shows where your upper 95% CIs does not exceed $P=0.05$. So you are quite sure the probability of a specified decline is less than 0.05. The black area shows where your lower 95% confidence interval is above $P=0.95$. So you are quite sure the probability is greater than $P=0.95$. The light gray is between these two certain/uncertain extremes.

7.6 More risk metrics and some real data

The previous sections have focused on the probability of hitting thresholds because this is an important and common risk metric used in population viability analysis and it appears in IUCN Red List criteria. However, as you have seen, there is high uncertainty associated with such estimates. Part of the problem is that probability is constrained to be 0 to 1, and it is easy to get estimates with confidence intervals that span 0 to 1. Other metrics of risk, $\hat{u}$ and the distribution of the time to hit a threshold (Dennis et al., 1991), do not have this problem and may be more informative. Figure 7.6 shows different risk metrics from (Dennis et al., 1991) on a single plot. This figure is generated by a call to the function `CSEGriskfigure()`.
dat = read.table(datafile, skip=1)
dat = as.matrix(dat)
CSEGriskfigure(dat)

The datafile is the name of the data file, with years in column 1 and population count (logged) in column 2. CSEGriskfigure() has a number of arguments that can be passed in to change the default behavior. The variable te is the forecast length (default is 100 years), threshold is the extinction threshold either as an absolute number, if absolutethresh=TRUE, or as a fraction of current population count, if absolutethresh=FALSE. The default is absolutethresh=FALSE and threshold=0.1. datlogged=TRUE means the data are already logged; this is the default.

**Example 7.5 (Risk figures for different species)**

*Use the Example 7.5 code to re-create Figure 7.6.* The package includes other data for you to run: prairiechicken from the endangered Attwater Prairie Chicken, graywhales from Gerber et al. (1999), and grouse from the Sharp-tailed Grouse (a species of U.S. federal concern) in Washington State. Note for some of these other datasets, the Hessian matrix cannot be inverted and you will need to use CI.method="parametric". If you have other text files of data, you can run those too. The commented lines show how to read in data from a tab-delimited text file with a header line.

---

**Example 7.5 code**

```
#If you have your data in a tab delimited file with a header
#This is how you would read it in using file.choose()
#to call up a directory browser.
#However, the package has the datasets for the examples
#dat=read.table(file.choose(), skip=1)
#dat=as.matrix(dat)
dat = wilddogs
CSEGriskfigure(dat, CI.method="hessian", silent=TRUE)
```

---

**7.7 Confidence intervals**

The figures produced by CSEGriskfigure() have confidence intervals (95% and 75%) on the probabilities in the top right panel. A standard way to
produce these intervals is via parametric bootstrapping. Here are the steps in a parametric bootstrap:

- You estimate $u$, $\sigma^2$ and $\eta^2$
- Then you simulate time series using those estimates and Equations 7.1 and 7.2
- Then you re-estimate your parameters from the simulated data (using say MARSS(simdata))
- Repeat for 1000s of time series simulated using your estimated parameters.

This gives you a large set of bootstrapped parameter estimates.

Fig. 7.6. Risk figure using data for the critically endangered African Wild Dog (data from Ginsberg et al. 1995). This population went extinct after 1992.
- For each bootstrapped parameter set, compute a set of extinction estimates (you use Equation 7.3 and code from Example 7.3).
- The $\alpha\%$ ranges on those bootstrapped extinction estimates gives you your $\alpha$ confidence intervals on your probabilities of hitting thresholds.

The MARSS package provides the function `MARSSparamCIs()` to add bootstrapped confidence intervals to fitted models (type `?MARSSparamCIs` to learn about the function).

In the function `CSEGriskfigure()`, you can set `CI.method = c("hessian", "parametric", "innovations", "none")` to tell it how to compute the confidence intervals. The methods ‘parametric’ and ‘innovations’ specify parametric and non-parametric bootstrapping respectively. Producing parameter estimates by bootstrapping is quite slow. Approximate confidence intervals on the parameters can be generated rapidly using the inverse of a numerically estimated Hessian matrix (method ‘hessian’). This uses an estimate of the variance-covariance matrix of the parameters (the inverse of the Hessian matrix). Using an estimated Hessian matrix to compute confidence intervals is a handy trick that can be used for all sorts of maximum-likelihood parameter estimates.

### 7.8 Discussion

Data with cycles, from age-structure or predator-prey interactions, are difficult to analyze and the EM algorithm used in the MARSS package will give poor estimates for this type of data. The slope method [Holmes, 2001] is more robust to those problems. [Holmes et al., 2007] used the slope method in a large study of data from endangered and threatened species, and [Ellner and Holmes, 2008] showed that the slope estimates are close to the theoretical minimum uncertainty. Especially, when doing a population viability analysis using a time series with fewer than 25 years of data, the slope method is often less biased and (much) less variable because that method is less data-hungry [Holmes, 2004]. However the slope method is not a true maximum-likelihood method and thus constrains the types of further analyses you can do (such as model selection).
8

Combining multi-site data to estimate regional population trends

8.1 Harbor seals in the Puget Sound, WA.

In this application, we will use multivariate state-space models to combine surveys from multiple regions (or sites) into one estimate of the average long-term population growth rate and the year-to-year variability in that growth rate. Note this is not quite the same as estimating the trend; “trend” often means “what population change happened?”, whereas the long-term population growth rate refers to the underlying population dynamics. We will use as our example a dataset from harbor seals in Puget Sound, Washington, USA.

We have five regions (or sites) where harbor seals were censused from 1978-1999 while hauled out of land [Jeffries et al. 2003]. During the period of this dataset, harbor seals were recovering steadily after having been reduced to low levels by hunting prior to protection. The methodologies were consistent throughout the 20 years of the data but we do not know what fraction of the population that each region represents nor do we know the observation-error variance for each region. Given differences between behaviors of animals in different regions and the numbers of haul-outs in each region, the observation errors may be quite different. The regions have had different levels of sampling; the best sampled region has only 4 years missing while the worst has over half the years missing (Figure 8.1).

We will assume that the underlying population process is a stochastic exponential growth process with rates of increase that were not changing through 1978-1999. However, we are not sure if all five regions sample a single “total Puget Sound” population or if there are independent subpopulations. We will estimate the long-term population growth rate using different assumptions about the population structures (one big population versus multiple smaller

Type RShowDoc("Chapter_SealTrend.R",package="MARSS") at the R command line to open a file with all the code for the examples in this chapter.
Combining multi-site and subpopulation data

Fig. 8.1. Plot of the count data from the five harbor seal regions (Jeffries et al. 2003). The numbers on each line denote the different regions: 1) Strait of Juan de Fuca (SJF), 2) San Juan Islands (SJI), 2) Eastern Bays (EBays), 4) Puget Sound (PSnd), and 5) Hood Canal (HC). Each region is an index of the total harbor seal population, but the bias (the difference between the index and the true population size) for each region is unknown.

ones) and observation error structures to see how different assumptions change the trend estimates.

The harbor seal data are included in the MARSS package. The data have time running down the rows and years in the first column. We need time across the columns for the\ MARSS() function, so we will transpose the data:

```r
dat=t(harborSealWA) #Transpose
years = dat[1,] #[1,] means row 1
n = nrow(dat)-1
dat = dat[2:nrow(dat),] #no years
```

The years are in column 1 of dat and the logged data are in the rest of the columns. The number of observation time series (n) is the number of rows in dat minus 1 (for years row). Let’s look at the first few years of data:

```r
print(harborSealWA[1:8,], digits=3)
```
8.2 A single well-mixed population with i.i.d. errors

<table>
<thead>
<tr>
<th>Year</th>
<th>SJF</th>
<th>SJI</th>
<th>EBays</th>
<th>PSnd</th>
<th>HC</th>
</tr>
</thead>
<tbody>
<tr>
<td>[1,]</td>
<td>1978</td>
<td>6.03</td>
<td>6.75</td>
<td>6.63</td>
<td>5.82</td>
</tr>
<tr>
<td>[2,]</td>
<td>1979</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>[3,]</td>
<td>1980</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>[4,]</td>
<td>1981</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>[5,]</td>
<td>1982</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>[6,]</td>
<td>1983</td>
<td>6.78</td>
<td>7.43</td>
<td>7.21</td>
<td>NA</td>
</tr>
<tr>
<td>[7,]</td>
<td>1984</td>
<td>6.93</td>
<td>7.74</td>
<td>7.45</td>
<td>NA</td>
</tr>
<tr>
<td>[8,]</td>
<td>1985</td>
<td>7.16</td>
<td>7.53</td>
<td>7.26</td>
<td>6.60</td>
</tr>
</tbody>
</table>

The NA’s in the data are missing values.

8.1.1 A MARSS model for Puget Sound harbor seals

The first step is to mathematically specify the population structure and how the regions relate to that structure. The general state-space model is

\[ x_t = B x_{t-1} + u + w_t, \quad \text{where } w_t \sim \text{MVN}(0, Q) \]

\[ y_t = Z x_t + a + v_t, \quad \text{where } v_t \sim \text{MVN}(0, R) \]

where all the bolded symbols are matrices. To specify the structure of the population and observations, we will specify what those matrices look like.

8.2 A single well-mixed population with i.i.d. errors

When we are looking at data over a large geographic region, we might make the assumption that the different census regions are measuring a single population if we think animals are moving sufficiently such that the whole area (multiple regions together) is “well-mixed”. We write a model of the total population abundance for this case as:

\[ n_t = \exp(u + w_t) n_{t-1}, \quad (8.1) \]

where \( n_t \) is the total count in year \( t \), \( u \) is the mean population growth rate, and \( w_t \) is the deviation from that average in year \( t \). We then take the log of both sides and write the model in log space:

\[ x_t = x_{t-1} + u + w_t, \quad \text{where } w_t \sim \text{N}(0, q) \quad (8.2) \]

\( x_t = \log n_t \). When there is one effective population, there is one \( x \), therefore \( x_t \) is a 1 \times 1 matrix. There is one population growth rate (\( u \)) and there is one process variance (\( q \)). Thus \( u \) and \( Q \) are 1 \times 1 matrices.
8.2.1 The observation process

We assume that all five regional time series are observations of this one population trajectory but they are scaled up or down relative to that trajectory. In effect, we think that animals are moving around a lot and our regional samples are some fraction of the population. There is year-to-year variation in the fraction in each region, just by chance. Notice that under this analysis, we do not think the regions represent independent subpopulations but rather independent observations of one population. Our model for the data, \( y_t = Zx_t + a + v_t \), is written as:

\[
\begin{pmatrix}
  y_1 \\
  y_2 \\
  y_3 \\
  y_4 \\
  y_5 \\
\end{pmatrix}
= \begin{pmatrix}
  1 \\
  1 \\
  1 \\
  1 \\
  1 \\
\end{pmatrix}
 x_t + \begin{pmatrix}
  a_2 \\
  a_3 \\
  a_4 \\
  a_5 \\
\end{pmatrix} + \begin{pmatrix}
  v_1 \\
  v_2 \\
  v_3 \\
  v_4 \\
  v_5 \\
\end{pmatrix}
\]

(8.3)

Each \( y_i \) is the time series for a different region. The \( a \)'s are the bias between the regional sample and the total population. The \( a \)'s are scaling (or intercept-like) parameters. We allow that each region could have a unique observation variance and that the observation errors are independent between regions. Lastly, we assume that the observations errors on log(counts) are normal and thus the errors on (counts) are log-normal.

For our first analysis, we assume that the observation variance is equal across regions but the errors are independent. This means we estimate one observation variance instead of five. This is a fairly standard assumption for data that come from the uniform survey methodology. We specify independent observation errors with identical variances by specifying that the \( v \)'s come from a multivariate normal distribution with variance-covariance matrix \( R (v \sim MVN(0, R)) \), where

---

1 To get rid of the \( a \)'s, we scale multiple observation time series against each other; thus one \( a \) will be fixed at 0. Estimating the bias between regional indices and the total population is important for getting an estimate of the total population size. The type of time-series analysis that we are doing here (trend analysis) is not useful for estimating \( a \)'s. Instead to get \( a \)'s one would need some type of mark-recapture data. However, for trend estimation, the \( a \)'s are not important. The regional observation variance captures increased variance due to a regional estimate being a smaller sample of the total population.

2 The assumption of normality is not unreasonable since these regional counts are the sum of counts across multiple haul-outs.

3 By the way, this is not a good assumption for these data since the number haul-outs in each region varies and the regional counts are the sums across all haul-outs in a region. We will change this assumption in the next fit and see that the AIC values decline.
8.2 A single well-mixed population with i.i.d. errors

\[
R = \begin{bmatrix}
  r & 0 & 0 & 0 \\
  0 & r & 0 & 0 \\
  0 & 0 & r & 0 \\
  0 & 0 & 0 & r \\
\end{bmatrix}
\tag{8.4}
\]

\(Z\) specifies which observation time series, \(y_{i,1:T}\), is associated with which population trajectory, \(x_{j,1:T}\). \(Z\) is like a look up table with 1 row for each of the \(n\) observation time series and 1 column for each of the \(m\) population trajectories. A 1 in row \(i\) column \(j\) means that observation time series \(i\) is measuring state process \(j\). Otherwise the value in \(Z_{ij} = 0\). Since we have only 1 population trajectory, all the regions must be measuring that one population trajectory. Thus \(Z\) is \(n \times 1\):

\[
Z = \begin{bmatrix}
  1 \\
  1 \\
  1 \\
  1 \\
  1 \\
\end{bmatrix}
\tag{8.5}
\]

8.2.2 Fitting the model

We have specified the mathematical form of our state-space model. The next step is to fit this model with \texttt{MARSS()}. The function call will now look like:

\[
\texttt{kem1 = MARSS(dat, model=list(Z=Z.model, R=R.model) )}
\]

The \texttt{model} list argument tells the \texttt{MARSS()} function the model structure, i.e. the form of \(Z, u, Q,\) etc. For our first analysis, we only need to set the model structure for \(Z\) and \(R\). Since there is only one population, there is only one \(u\) and \(Q\) (they are scalars), so they have no 'structure'.

First we specify the \(Z\) matrix. We need to tell the \texttt{MARSS} function that \(Z\) is a \(5 \times 1\) matrix of 1s (as in Equation 8.3). We can do this two ways. We can pass in \(Z\cdot\text{model}\) as a matrix of ones, \texttt{matrix(1,5,1)}, just like in Equation 8.3 or we can pass in a vector of five factors, \texttt{factor(c(1,1,1,1,1))}. The \(i\)-th factor specifies which population trajectory the \(i\)-th observation time series belongs to. Since there is only one population trajectory in this first analysis, we will have a vector of five 1's: every observation time series is measuring the first, and only, population trajectory.

\[
Z\cdot\text{model} = \texttt{factor(c(1,1,1,1,1))}
\]

Note, the vector (the \texttt{c()} bit) must be wrapped in \texttt{factor()} so that \texttt{MARSS} recognizes what it is. You can use either numeric or character vectors: \(c(1,1,1,1,1)\) is the same as \(c("PS","PS","PS","PS","PS")\).

Next we specify that the \(R\) variance-covariance matrix only has terms on the diagonal (the variances) with the off-diagonal terms (the covariances) equal to zero:
\textit{R.model} = "\textit{diagonal and equal}"

The ‘and equal’ part specifies that the variances are the same value. We will relax this assumption later.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{plot}
\caption{Plot of the estimate of “log total harbor seals in Puget Sound”. The estimate of the total count has been scaled relative to the first time series. The 95\% confidence intervals on the population estimates are the dashed lines. These are not the confidence intervals on the observations, and the observations (the numbers) will not fall between the confidence interval lines.}
\end{figure}

Code \texttt{8.2} shows you how to fit the single population model (Equations \texttt{8.2} and \texttt{8.3}) to the harbor seal data.
8.2 A single well-mixed population with i.i.d. errors

---

**Code 8.2**

```r
#Code to fit the single population model with i.i.d. errors
#Read in data
dat=t(harborSealWA)  #Transpose since MARSS needs time ACROSS columns
years = dat[1,]
n = nrow(dat)-1
dat = dat[2:nrow(dat),]
legendnames = (unlist(dimnames(dat)[1]))

#estimate parameters
Z.model = factor(c(1,1,1,1,1))
R.model = "diagonal and equal"
kem1 = MARSS(dat, model=list(Z=Z.model, R=R.model))

#make figure
matplot(years, t(dat),xlab="",ylab="index of log abundance",
        pch=c("1","2","3","4","5"),ylim=c(5,9),bty="L")
lines(years,kem1$states-1.96*kem1$states.se,type="l",
       lwd=1,lty=2,col="red")
lines(years,kem1$states+1.96*kem1$states.se,type="l",
       lwd=1,lty=2,col="red")
lines(years,kem1$states,type="l",lwd=2)
title("Observations and total population estimate",cex.main=.9)
coef(kem1, type="vector")  #show the estimated parameter elements as a vector
coef(kem1)
kem1$logLik  #show the log-likelihood
kem1$AIC  #show the AIC
```

---

**8.2.3 The MARSS() output**

The output from `MARSS()`, here assigned the name `kem1`, is a list of objects:

```r
names(kem1)
```

The maximum-likelihood estimates of “total harbor seal population” scaled to the first observation data series (Figure 8.2) are in `kem1$states`, and `kem1$states.se` are the standard errors on those estimates. To get 95% confidence intervals, use `kem1$states +/- 1.96*kem1$states.se`. Figure 8.2 shows a plot of `kem1$states` with its 95% confidence intervals over the data. Because `kem1$states` has been scaled relative to the first time series, it is on top of that time series. One of the biases, the `/aa`, cannot be estimated and arbitrarily our algorithm chooses $a_1 = 0$, so the population estimate is scaled to the first observation time series.
The estimated parameters are output with the function `coef(kem1)`. To get the estimate just for $U$, which is the estimated long-term population growth rate, use `coef(kem1)$U`. Multiply by 100 to get the percent increase per year. The estimated process variance is given by `coef(kem2)$Q`.

The log-likelihood of the fitted model is in `kem1$logLik`. We estimated one initial $x (t = 1)$, one process variance, one $u$, four $a$'s, and five observation variances. So $K = 12$ parameters. The AIC of this model is $-2 \times \text{log-like} + 2K$, which we can show by typing `kem1$AIC`.

### 8.3 Single population model with independent and non-identical errors

Here is the estimated $R$ matrix for our first model:

```r
coef(kem1, type="matrix")$R
```

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>[1,]</td>
<td>0.04523437</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.00000000</td>
</tr>
<tr>
<td>[2,]</td>
<td>0.00000000</td>
<td>0.04523437</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.00000000</td>
</tr>
<tr>
<td>[3,]</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.04523437</td>
<td>0.00000000</td>
<td>0.00000000</td>
</tr>
<tr>
<td>[4,]</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.04523437</td>
<td>0.00000000</td>
</tr>
<tr>
<td>[5,]</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.00000000</td>
<td>0.04523437</td>
</tr>
</tbody>
</table>

Notice that the variances along the diagonal are all the same—we estimated one observation variance and it applied to all observation time series. We might be able to improve the fit (at the cost of more parameters) by assuming that the observation variance is different across regions while the errors are still independent. This means we estimate five observation variances instead of one. In this case, $R$ has the form:

$$
R = \begin{pmatrix}
  r_1 & 0 & 0 & 0 & 0 \\
  0 & r_2 & 0 & 0 & 0 \\
  0 & 0 & r_3 & 0 & 0 \\
  0 & 0 & 0 & r_4 & 0 \\
  0 & 0 & 0 & 0 & r_5 \\
\end{pmatrix}
$$

To impose this model, we set the $R$ model to

```r
R.model = "diagonal and unequal"
```

This tells MARSS that all the $r$'s along the diagonal in $R$ are different. To fit this model to the data, call `MARSS()` as:

```r
Z.model = factor(c(1,1,1,1,1))
R.model = "diagonal and unequal"
kem2 = MARSS(dat, model=list(Z=Z.model, R=R.model))
```
We estimated one initial $x$, one process variance, one $u$, four $a$’s, and five observation variances. So $K = 11$ parameters. The AIC for this new model compared to the old model with one observation variance is:

\[
c(kem1$AIC,kem2$AIC)
\]

\>[1] 8.813447 -9.323982

A smaller AIC means a better model. The difference between the one observation variance versus the unique observation variances is $>10$, suggesting that the unique observation variances model is better.

One of the key diagnostics when you are comparing fits from multiple models is whether the model is flexible enough to fit the data. This can be checked by looking for temporal trends in the residuals between the estimated population states (e.g. `kem2$states`) and the data. In Figure 8.3, the residuals for the second analysis are shown. Ideally, these residuals should not have a temporal trend. They should look cloud-like. The fact that the residuals have a strong temporal trend is an indication that our one population model is too restrictive for the data\(^4\). Code 8.3 shows you how to fit the second model and make the diagnostics plot.

\(^4\) When comparing models via AIC, it is important that you only compare models that are flexible enough to fit the data. Fortunately if you neglect to do this, the inadequate models will usually have very high AICs and fall out of the mix anyhow.
Fig. 8.3. Residuals for the model with a single population. The plots of the residuals should not have trends with time, but they do. This is an indication that the single population model is inconsistent with the data. The code to make this plot is given in the script file for this chapter.

**Code 8.3**

```R
#Code to fit the single population model with independent and unequal errors

Z.model = factor(c(1,1,1,1,1))
R.model = "diagonal and unequal"
kem2 = MARSS(dat, model=list(Z=Z.model, R=R.model))
coef(kem2) #the estimated parameter elements
kem2$logLik #log likelihood
c(kem1$AIC,kem2$AIC) #AICs

#plot residuals
plotdat = t(dat)
matrix.of.biases = matrix(coef(kem2, type="matrix")$A,
    nrow=nrow(plotdat),ncol=ncol(plotdat),byrow=T)
xs = matrix(kem2$states,
    nrow=dim(plotdat)[1],ncol=dim(plotdat)[2],byrow=F)
resids = plotdat-matrix.of.biases-xs
par(mfrow=c(2,3))
for(i in 1:n){
    plot(resids[!is.na(resids[,i]),i],ylab="residuals")
    title(legendnames[i])
}
par(mfrow=c(1,1))
```
8.4 Two subpopulations, north and south

For the third analysis, we will change our assumption about the structure of the population. We will assume that there are two subpopulations, north and south, and that regions 1 and 2 (Strait of Juan de Fuca and San Juan Islands) fall in the north subpopulation and regions 3, 4 and 5 fall in the south subpopulation. For this analysis, we will assume that these two subpopulations share their growth parameter, $u$, and process variance, $q$, since they share a similar environment and prey base. However, we postulate that because of fidelity to natal rookeries for breeding, animals do not move much year-to-year between the north and south and the two subpopulations are independent.

We need to write down the state-space model to reflect this population structure. There are two subpopulations, $x_n$ and $x_s$, and they have the same growth rate $u$:

\[
\begin{bmatrix}
    x_n \\
    x_s
\end{bmatrix}_t = \begin{bmatrix}
    x_n \\
    x_s
\end{bmatrix}_{t-1} + \begin{bmatrix}
    u \\
    u
\end{bmatrix} + \begin{bmatrix}
    w_n \\
    w_s
\end{bmatrix}_t
\] (8.7)

We specify that they are independent by specifying that their year-to-year population fluctuations (their process errors) come from a multivariate normal with no covariance:

\[
\begin{bmatrix}
    w_n \\
    w_s
\end{bmatrix}_t \sim \text{MVN} \left( \begin{bmatrix}
    0 \\
    q
\end{bmatrix}, \begin{bmatrix}
    q & 0 \\
    0 & q
\end{bmatrix} \right)
\] (8.8)

For the observation process, we use the $Z$ matrix to associate the regions with their respective $x_n$ and $x_s$ values:

\[
\begin{bmatrix}
    y_1 \\
    y_2 \\
    y_3 \\
    y_4 \\
    y_5
\end{bmatrix}_t = \begin{bmatrix}
    1 & 0 \\
    0 & 1 \\
    0 & 1 \\
    0 & 1 \\
    0 & 1
\end{bmatrix} \begin{bmatrix}
    x_n \\
    x_s
\end{bmatrix}_t + \begin{bmatrix}
    v_1 \\
    v_2 \\
    v_3 \\
    v_4 \\
    v_5
\end{bmatrix}_t
\] (8.9)

8.4.1 Specifying the model elements

We need to change the $Z$ specification to indicate that there are two subpopulations (north and south), and that regions 1 and 2 are in the north subpopulation and regions 3, 4 and 5 are in the south subpopulation. There are a few ways, we can specify this $Z$ matrix for \texttt{MARSS}():

\begin{verbatim}
Z.model = matrix(c(1,1,0,0,0,0,0,1,1,1),5,2)
Z.model = factor(c(1,1,2,2,2))
Z.model = factor(c("N","N","S","S","S"))
\end{verbatim}

Which you choose is a matter of preference as they all specify the same form for $Z$.

We also want to specify that the $u$'s are the same for each subpopulation and that $Q$ is diagonal with equal $q$'s. To do this, we set
This says that there is one u and one q parameter and both subpopulations share it (if we wanted the u’s to be different, we would use U.model="unequal" or leave off the u model since the default behavior is U.model="unequal").

Code 8.4 puts all the pieces together and shows you how to fit the north and south population model and create the residuals plot (Figure 8.4). The residuals look better (more cloud-like) but the Hood Canal residuals are still temporally correlated.

```r
#fit the north and south population model
Z.model = factor(c(1,1,2,2,2))
U.model = "equal"
Q.model = "diagonal and equal"
R.model = "diagonal and unequal"
kem3 = MARSS(dat, model=list(Z=Z.model,
    R=R.model, U=U.model, Q=Q.model))

#plot residuals
plotdat = t(dat)
matrix.of.biases = matrix(coef(kem3,type="matrix")$A,
    nrow=nrow(plotdat),ncol=ncol(plotdat),byrow=T)
par(mfrow=c(2,3))
for(i in 1:n){
    j=c(1,1,2,2,2)
    xs = kem3$states[j[i],]
    resids = plotdat[,i]-matrix.of.biases[,i]-xs
    plot(resids[!is.na(resids)],ylab="residuals")
    title(legendnames[i])
}
par(mfrow=c(1,1))
```

8.5 Other population structures

Now work through a number of different structures and examine how your estimation of the mean population growth rate varies under different assumptions about the structure of the population and the data. You can compare the model fits using AIC (or AICc). For AIC, lower is better and only the relative differences matter. A difference of 10 between two AICs means substantially more support for the model with lower AIC. A difference of 30 or 40 between two AICs is very large.
8.5 Other population structures

8.5.1 Five subpopulations

Analyze the data using a model with five subpopulations, where each of the five census regions is sampling one of the subpopulations. Assume that the subpopulation are independent (diagonal $Q$), however let each subpopulation share the same population parameters, $u$ and $q$. Code 8.5.1 shows how to set the `MARSS()` arguments for this case. You can use `R.model="diagonal and equal"` to make all the observation variances equal.
8.5.1 Code

\[
\begin{align*}
\text{Z.model} &= \text{factor(c(1,2,3,4,5))} \\
\text{U.model} &= \text{"equal"} \\
\text{Q.model} &= \text{"diagonal and equal"} \\
\text{R.model} &= \text{"diagonal and unequal"} \\
\text{kem} &= \text{MARSS(dat, model=list(Z=Z.model,} \\
& \quad \text{U=U.model, Q=Q.model, R=R.model) )}
\end{align*}
\]

8.5.2 Two subpopulations with different population parameters

Analyze the data using a model that assumes that the Strait of Juan de Fuca and San Juan Islands census regions represent a northern Puget Sound subpopulation, while the other three regions represent a southern Puget Sound subpopulation. This time assume that each population trajectory (north and south) has different \( u \) and \( q \) parameters: \( u_n, u_s \) and \( q_n, q_s \). Also assume that each of the five census regions has a different observation variance. Try to write your own code. If you get stuck (or want to check your work, you can open a script file with sample R code by typing \texttt{RShowDoc("Chapter_SealTrend.R",package="MARSS")} at the R command line.

In math form, this model is:

\[
\begin{bmatrix} x_n \\ x_s \end{bmatrix}_t = \begin{bmatrix} x_n \\ x_s \end{bmatrix}_{t-1} + \begin{bmatrix} u_n \\ u_s \end{bmatrix} + \begin{bmatrix} w_n \\ w_s \end{bmatrix}, \begin{bmatrix} w_n \\ w_s \end{bmatrix} \sim \text{MVN}\left(0, \begin{bmatrix} q_n & 0 \\ 0 & q_s \end{bmatrix}\right) (8.10)
\]

\[
\begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ y_4 \\ y_5 \end{bmatrix}_t = \begin{bmatrix} 1 & 0 \\ 1 & 0 \\ 0 & 1 \\ 0 & 1 \\ 0 & 1 \end{bmatrix} \begin{bmatrix} x_n \\ x_s \end{bmatrix}_t + \begin{bmatrix} a_2 \\ 0 \\ a_4 \\ a_5 \end{bmatrix} + \begin{bmatrix} v_1 \\ v_2 \\ v_3 \\ v_4 \\ v_5 \end{bmatrix}_t (8.11)
\]

8.5.3 Hood Canal covaries with the other regions

Analyze the data using a model with two subpopulations with the divisions being Hood Canal versus everywhere else. In math form, this model is:

\[
\begin{bmatrix} x_p \\ x_h \end{bmatrix}_t = \begin{bmatrix} x_p \\ x_h \end{bmatrix}_{t-1} + \begin{bmatrix} u_p \\ u_h \end{bmatrix} + \begin{bmatrix} w_p \\ w_h \end{bmatrix}, \begin{bmatrix} w_p \\ w_h \end{bmatrix} \sim \text{MVN}\left(0, \begin{bmatrix} q & c \\ c & q \end{bmatrix}\right) (8.12)
\]

\[
\begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ y_4 \\ y_5 \end{bmatrix}_t = \begin{bmatrix} 1 & 0 \\ 1 & 0 \\ 1 & 0 \\ 1 & 0 \\ 0 & 1 \end{bmatrix} \begin{bmatrix} x_p \\ x_h \end{bmatrix}_t + \begin{bmatrix} a_2 \\ a_3 \\ a_4 \\ a_5 \end{bmatrix} + \begin{bmatrix} v_1 \\ v_2 \\ v_3 \\ v_4 \\ v_5 \end{bmatrix}_t (8.13)
\]
To specify that $Q$ has one value on the diagonal (one variance) and one value on the off-diagonal (covariance) you can specify `Q.model` two ways:

```r
Q.model = "equalvarcov"
Q.model = matrix(c("q","c","c","q"),2,2)
```

### 8.5.4 Three subpopulations with shared parameter values

Analyze the data using a model with three subpopulations as follows: north (regions 1 and 2), south (regions 3 and 4), Hood Canal (region 5). You can specify that some subpopulations share parameters while others do not. First, let's specify that each population is affected by independent environmental variability, but that the variance of that variability is the same for the two interior populations:

```r
Q.model=matrix(list(0),3,3)
diag(Q.model)=c("coastal","interior","interior")
print(Q.model)
```

Notice that $Q$ is a diagonal matrix (independent year-to-year environmental variability) but the variance of two of the populations is the same. Notice too that the off-diagonal terms are numeric; they do not have quotes. We specified $Q$ using a matrix of class list, so that we could have numeric values (fixed) and character values (estimated parameters).

In a similar way, we specify that the observation errors are independent but that estimates from a plane do not have the same variance as those from a boat:

```r
R.model=matrix(list(0),5,5)
diag(R.model)=c("boat","boat","plane","plane","plane")
```

For the long-term trends, we specify that $x_1$ and $x_2$ share a long-term trend ("puget sound") while $x_3$ is allowed to have a separate trend ("hood canal").

```r
U.model=matrix(c("puget sound","puget sound","hood canal"),3,1)
```

### 8.6 Discussion

There are a number of corners that we cut in order to show you code that runs quickly:

- We ran the code starting from one initial condition. For a real analysis, you should start from a large number of random initial conditions and use the one that gives the highest likelihood. Since the EM algorithm is a "hill-climbing" algorithm, this ensures that it did not get stuck on a local maxima. See Chapter [6](#) for a discussion of initial conditions searchers.
We assume independent observation and process errors. Depending on your system, observation errors may be driven by large-scale environmental factors (temperature, tides, prey locations) that would cause your observation errors to covary across regions. If your observation errors strongly covary between regions and you treat them as independent, this could be bad for your analysis. Unfortunately, separating covariance across observation versus process errors will require much data (to have any power). In practice, the first step is to think hard about what drives sightability for your species and what are the relative levels of process and observation variance. You may be able to subsample your data in a way that will make the observation errors more independent.

The `MARSS()` argument `control` specifies the options for the EM algorithm. We left the default tolerance for the convergence test. You would want to set this lower for a real analysis. You will need to up the `maxit` argument correspondingly.

We used the large-sample approximation for AIC instead of a bootstrap AIC that is designed to correct for small sample size in state-space models. The bootstrap metric, AICb, takes a long time to run. Use the call `MARSSaic(kem, output=c("AICbp"))` to compute AICb. We could have shown AICc, which is the small-sample size corrector for non-state-space models. Type `kem$AICc` to get that.

Finally, in a real (maximum-likelihood) analysis, one needs to be careful not to dredge the data. The temptation is to look at the data and pick a population structure that will fit that data. This can lead to including models in your analysis that have no biological basis. In practice, we spend a lot of time discussing the population structure with biologists working on the species and review all the biological data that might tell us what are reasonable structures. From that, a set of model structures to use are selected. Other times, a particular model structure needs to be used because the population structure is not in question rather it is a matter of using that pre-specified structure and using all the data to get parameter estimates for forecasting.

Some more questions you might ponder

Do different assumptions about whether the observation error variances are all identical versus different affect your estimate of the long-term population growth rate ($\mu$)? You may want to rerun Examples 3-7 with the `R.model` changed. `R.model="diagonal and unequal"` means measurement variances all different versus "diagonal and equal".

Do assumptions about the underlying structure of the population affect your estimates of $\mu$? Structure here means number of subpopulations and which areas are in which subpopulation.

The confidence intervals for the first two analyses are very tight because the estimated process variance, $Q$, was very small. Why do you think process
variance \( (q) \) was forced to be so small? [Hint: We are forcing there to be one and only one true population trajectory and all the observation time series have to fit that one time series. Look at the AICs too.]
Identifying spatial population structure and covariance

9.1 Harbor seals on the U.S. west coast

In this application, we use harbor seal abundance estimates along the west coast to examine large-scale spatial structure. Harbor seals are distributed along the west coast of the U.S. from California to Washington. The populations been surveyed at haul-out sites since the mid-1970s (Figure 9.1) and have been increasing steadily since the 1972 Marine Mammal Protection Act. See ?harborSeal for the data sources.

For management purposes, three stocks are recognized: the CA stock, the OR/WA coastal stock which consists of four regions (Northern/Southern Oregon, Coastal Estuaries, Olympic Peninsula), and the inland WA stock which consists of the regions in the WA inland waters minus Hood Canal (Figure 9.1). Differences exist in the demographics across regions (e.g. pupping dates), however mtDNA analyses and tagging studies support the larger stock structure. Harbor seals are known for strong site fidelity, but at the same time travel large distances to forage.

Our goal is to address the following questions about spatial structure: 1) Does population abundance data support the existing management boundaries, or are there alternative groupings that receive more support?, 2) Do subpopulations (if they exist) experience independent environmental variability or correlated variability? and 3) Does the Hood Canal site represent a distinct subpopulation? To address these questions, we will mathematically formulate different hypotheses about population structure via different MARSS models. We will then compare the data support for different models using model selection criteria, specifically AICc and AIC weights.

Type RShowDoc("Chapter_SeaPopStructure.R",package="MARSS") at the R command line to open a file with all the code for the examples in this chapter.
Fig. 9.1. Map of spatial distribution of harbor seal survey regions in Washington and Oregon. In addition to these nine survey regions, we also have data from the Georgia Strait just north of the San Juan Islands, the California coast and the Channel Islands in Southern California.
9.2 Question 1, How many distinct subpopulations?

9.1.1 MARSS models for a population with spatial structure

The mathematical form of the model we will use is

\[
\begin{align*}
x_t &= x_{t-1} + u_t + w_t \quad \text{where } w_t &\sim \text{MVN}(0, Q) \\
y_t &= Zx_t + a + v_t \quad \text{where } v_t &\sim \text{MVN}(0, R) \\
x_0 &\sim \text{MVN}(\pi, \Lambda)
\end{align*}
\] (9.1)

\(B\) is in front of \(x\) but is left off above since it is the identity matrix\(^1\). We will use \(Z\), \(u\), and \(Q\) to specify different hypotheses about the population structure. The form of \(a\) will be “scaling” in all cases. Aerial survey methodology has been relatively constant across time and space, and we will assume that all the time series from each region has identical and independent observation error variance, which means a diagonal \(R\) matrix with one variance term on the diagonal\(^2\).

Each call to \text{MARSS()}\ will look like

\[
\text{fit} = \text{MARSS(sealData, model=list(}
\quad Z = Z.model, \ \ Q = Q.model, \ \ ...))
\]

where the \(\ldots\) are components of the model list that are the same across all models. We will specify different \(Z.model\) and \(Q.model\) in order to model different population spatial structures.

9.2 Question 1, How many distinct subpopulations?

We will start by evaluating the data support for the following hypotheses about the population structure:

- H1 3 subpopulations defined by stock
- H2 2 subpopulations defined by coastal versus WA inland
- H3 2 subpopulations defined by north and south split in the middle of Oregon
- H4 4 subpopulations defined by N coastal, S coastal, SJF+Georgia Strait, and Puget Sound
- H5 All regions are part of the same panmictic population
- H6 Each of the 11 regions is a subpopulation

We will analyze each of these under the assumption of independent process errors with each subpopulation having different variances or the same variance.

\(^1\) a diagonal matrix with 1s on the diagonal
\(^2\) The sampling regions have different number of sites where animals are counted.

But we are working with log counts. We assume that the distribution of percent errors is the same (the probability of a 10% over-count is the same) and thus that the variances are similar on the log-scale.
9.2.1 Specify the $Z$ matrices

The $Z$ matrices specify the relationship between the survey regions and the subpopulations and allow us to specify the spatial population structures in the hypotheses. Each column of $Z$ corresponds to a different subpopulation and associates regions with particular subpopulations. For example for hypothesis 1, column 1 of the $Z$ matrix is OR/WA Coastal, column 2 is inland WA (ps for Puget Sound) and column 3 is CA. The $Z$ matrix for hypotheses 1, 2, 4, and 5 take the following form:

<table>
<thead>
<tr>
<th></th>
<th>H1</th>
<th>H2</th>
<th>H4</th>
<th>H5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$Z$</td>
<td>$Z$</td>
<td>$Z$</td>
<td>$Z$</td>
</tr>
<tr>
<td>wa.or</td>
<td>1 0 0</td>
<td>1 0</td>
<td>1 0 0 0</td>
<td>1</td>
</tr>
<tr>
<td>ps</td>
<td>1 0 0</td>
<td>1 0</td>
<td>1 0 0 0</td>
<td>1</td>
</tr>
<tr>
<td>ca</td>
<td>0 1 0</td>
<td>0 1</td>
<td>0 1 0 0</td>
<td>1</td>
</tr>
<tr>
<td>pan</td>
<td>0 1 0</td>
<td>0 1</td>
<td>0 1 0 0</td>
<td>1</td>
</tr>
<tr>
<td>Coastal Estuaries</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Olympic Peninsula</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Str. Juan de Fuca</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>San Juan Islands</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Eastern Bays</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Puget Sound</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CA.Maine</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CA.ChannelIslands</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OR North Coast</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OR South Coast</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Georgia Strait</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

To tell MARSS() the form of $Z$, we construct the same matrix in R. For example, for hypotheses 1, we can write:

```r
Z.model=matrix(0,11,3)
Z.model[c(1,2,9,10),1]=1  # which elements in col 1 are 1
Z.model[c(3:6,11),2]=1  # which elements in col 2 are 1
Z.model[7:8,3]=1  # which elements in col 3 are 1

MARSS has a shortcut for making this kind of $Z$ matrix using factor(). The following code specifies the same $Z$ matrix:

```r
Z1=factor(c("wa.or","wa.or",rep("ps",4),"ca","ca","wa.or","wa.or","bc"))
```

Each element in the c() vector is for one of the rows of $Z$ and indicates which column the “1” appears in or which row of your data belongs to which subpopulation. Notice the vector is 11 elements in length; one element for each row of data (in this case survey region).
9.2.2 Specify the \textbf{u} structure

We will assume that subpopulations can have a unique population growth rate. Mathematically, this means that the \textbf{u} matrix in Equation 9.1 looks like this for hypotheses 1 (3 subpopulations):

\[
\begin{pmatrix}
  u_1 \\
  u_2 \\
  u_3
\end{pmatrix}
\]

To specify this, we construct \texttt{U.model} as a character matrix where shared elements have the same character name. For example,

\begin{verbatim}
U.model=matrix(c("u1","u2","u3"),3,1)
\end{verbatim}

for a three subpopulation model. Alternatively, we can use the shortcut \texttt{U.model="unequal"}.

9.2.3 Specify the \textbf{Q} structures

For our first analysis, we fit a model where the subpopulations experience independent process errors. We will use two different types of independent process errors: independent process errors with different variances and independent process errors with identical variance. Independence is specified with a diagonal variance-covariance matrix with 0s on the off-diagonals.

Independent process errors with different variances is a diagonal matrix with different values on the diagonal:

\[
\begin{pmatrix}
  q_1 & 0 & 0 \\
  0 & q_2 & 0 \\
  0 & 0 & q_3
\end{pmatrix}
\]

This matrix has fixed numeric values, the zeros, combined with symbols \(q_1\), \(q_2\) and \(q_3\), representing estimated values. We specified this for MARSS() using a list matrix which combines numeric values (the fixed zeros) with character values (names of the estimated elements). The following produces this and printing it shows that it combines numeric values and character strings in quotes.

\begin{verbatim}
Q.model=matrix(list(0),3,3)
diag(Q.model)=c("q1","q2","q3")
Q.model
\end{verbatim}

\begin{verbatim}
[,1] [,2] [,3]
[1,] "q1" 0 0
[2,] 0 "q2" 0
[3,] 0 0 "q3"
\end{verbatim}
9 Using MARSS models to identify spatial population structure and covariance

We can also use the shortcut `Q.model="diagonal and unequal"`. Independent process errors with identical variance is a diagonal matrix with one value on the diagonal:

\[
\begin{bmatrix}
q & 0 & 0 \\
0 & q & 0 \\
0 & 0 & q \\
\end{bmatrix}
\]

```
Q.model=matrix(list(0),3,3)
diag(Q.model)="q"
Q.model
```

<table>
<thead>
<tr>
<th>[,1]</th>
<th>[,2]</th>
<th>[,3]</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;q&quot;</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>&quot;q&quot;</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>&quot;q&quot;</td>
</tr>
</tbody>
</table>

The shortcut for this form is `Q.model="diagonal and equal"`.

### 9.3 Fit the different models

The dataset `harborSeal` is a 29-year dataset of abundance indices for each of 12 regions between 1975-2004 (Figure 9.2). We start by setting up our data matrix. We will leave off Hood Canal (column 8) for now.

```
years = harborSeal[,1]  # first col is years
# leave off Hood Canal data for now
sealData = t(harborSeal[,c(2:7,9:13)])
```

We will set up our models so we can fit all of them with one loop of code. First the `Z` models.

**H1** stock
```
Z1=factor(c("wa.or","wa.or",rep("ps",4),"ca","ca","wa.or","wa.or","bc"))
```

**H2** coastal+PS
```
Z2=factor(c(rep("coast",2),rep("ps",4),rep("coast",4),"ps"))
```

**H3** N and S
```
Z3=factor(c(rep("N",6),"S","S","S","S","N"))
```

**H4** North Coast, Inland Strait, Puget Sound, South Coast
```
Z4=factor(c("nc","nc","is","is","ps","ps","sc","sc","nc","sc","is"))
```

**H5** panmictic
```
Z5=factor(rep("pan",11))
```

**H6** Site
```
Z6=factor(1:11)  # site
```

```
Z.models=list(Z1,Z2,Z3,Z4,Z5,Z6)
names(Z.models)=
  c("stock","coast+PS","N-S","NC+Strait+PS+SC","panmictic","site")
```
Fig. 9.2. Plot of the harbor seal sites in the harborSeal dataset. Each region is an index of the harbor seal abundance in that region.

Next we set up the Q models.

Q.models=c("diagonal and equal", "diagonal and unequal")

The rest of the model matrices have the same form across all models.

U.model="unequal"
R.model="diagonal and equal"
A.model="scaling"
B.model="identity"
x0.model="unequal"
V0.model="zero"
model.constant=list(
  U=U.model, R=R.model, A=A.model,
  x0=x0.model, V0=V0.model, tinitx=0)

We loop through the models, fit and store the results:

out.tab=NULL
fits=list()
for(i in 1:length(Z.models)){
  for(Q.model in Q.models){
    fit.model = c(list(Z=Z.models[[i]], Q=Q.model), model.constant)
    fit = MARSS(sealData, model=fit.model,
                silent=TRUE, control=list(maxit=1000))
    out=data.frame(H=names(Z.models)[i], Q=Q.model, U=U.model,
                    logLik=fit$logLik, AICc=fit$AICc, num.param=fit$num.params,
                    m=length(unique(Z.models[[i]])),
                    num.iter=fit$numIter, converged=!fit$convergence)
    out.tab=rbind(out.tab,out)
    fits=c(fits,list(fit))
  }
  if(i==5) next #one m for panmictic so only run 1 Q
}

9.4 Summarize the data support

We will use AICc and AIC weights to summarize the data support for the
different hypotheses. First we will sort the fits based on AICc:

min.AICc=order(out.tab$AICc)
out.tab.1=out.tab[min.AICc,]

Next we add the \(\Delta\)AICc values by subtracting the lowest AICc:

out.tab.1=cbind(out.tab.1,
                delta.AICc=out.tab.1$AICc-out.tab.1$AICc[1])

Relative likelihood is defined as \(\exp(-\Delta\text{AICc}/2)\).

out.tab.1=cbind(out.tab.1,
                rel.like=exp(-1*out.tab.1$delta.AICc/2))

The AIC weight for a model is its relative likelihood divided by the sum of all
the relative likelihoods.

out.tab.1=cbind(out.tab.1,
                AIC.weight = out.tab.1$rel.like/sum(out.tab.1$rel.like))

Let’s look at the model weights (out.tab.1):

<table>
<thead>
<tr>
<th>H</th>
<th>Q</th>
<th>delta.AICc</th>
<th>AIC.weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC+Strait+PS+SC</td>
<td>diagonal and equal</td>
<td>0.00</td>
<td>0.886</td>
</tr>
<tr>
<td>NC+Strait+PS+SC</td>
<td>diagonal and unequal</td>
<td>4.15</td>
<td>0.112</td>
</tr>
<tr>
<td>N-S</td>
<td>diagonal and unequal</td>
<td>12.67</td>
<td>0.002</td>
</tr>
<tr>
<td>N-S</td>
<td>diagonal and equal</td>
<td>14.78</td>
<td>0.001</td>
</tr>
<tr>
<td>coast+PS</td>
<td>diagonal and equal</td>
<td>31.23</td>
<td>0.000</td>
</tr>
<tr>
<td>coast+PS</td>
<td>diagonal and unequal</td>
<td>33.36</td>
<td>0.000</td>
</tr>
</tbody>
</table>
It appears that a population structure north and south coast subpopulations and two inland subpopulations is more supported than any of the other population structures—under the assumption of independent process errors. The latter means that good and bad years are not correlated across the subpopulations. The stock structure, supported by genetic information, does not appear to correspond to independent subpopulations and the individual survey regions, which are characterized by differential pupping times, does not appear to correspond to independent subpopulations either.

Figure 9.3 shows the four subpopulation trajectories estimated by the best fit model. The trajectories have been rescaled so that each starts at 0 in 1975 (to facilitate comparison).

![Figure 9.3](image-url)

**Fig. 9.3.** Estimated trajectories for the four subpopulations in the best-fit model. The plots have been rescaled so that each is at 0 at 1975.
9.5 Question 2, Are the subpopulations independent?

The assumption of independent process errors is unrealistic given ocean conditions are correlated across large spatial scales. We will repeat the analysis allowing correlated process errors using two different $\mathbf{Q}$ models. The first correlated $\mathbf{Q}$ model is correlated process errors with the same variance and covariance. For a model with three subpopulations, this $\mathbf{Q}$ would look like:

$$
\begin{bmatrix}
q & c & c \\
c & q & c \\
c & c & q
\end{bmatrix}
$$

We can construct this like so

```r
# identical variances
Q.model = matrix("c", 3, 3)
diag(Q.model) = "q"
```

or use the short-cut `Q.model="equalvarcov"`. The second type of correlated $\mathbf{Q}$ we will use is allows each subpopulation to have a different process variance and covariances. For a model with three subpopulations, this is the following variance-covariance matrix:

$$
\begin{bmatrix}
q_1 & c_{1,2} & c_{1,3} \\
c_{1,2} & q_2 & c_{2,3} \\
c_{1,2} & c_{2,3} & q_3
\end{bmatrix}
$$

Constructing this is tedious in R, but there is a short-cut: `Q.model="unconstrained".` We will re-run all the $\mathbf{Z}$ matrices with these two extra $\mathbf{Q}$ types and add them to our results table.

```r
for(i in 1:length(Z.models)){
  if(i==5) next # don't rerun panmictic
  for(Q.model in c("equalvarcov", "unconstrained")){
    fit.model = c(list(Z=Z.models[[i]], Q=Q.model), model.constant)
    fit = MARSS(sealData, model=fit.model,
                silent=TRUE, control=list(maxit=1000))
    out = data.frame(H=names(Z.models)[i], Q=Q.model, U=U.model,
                     logLik=fit$logLik, AICc=fit$AICc, num.param=fit$num.params,
                     m=length(unique(Z.models[[i]])),
                     num.iter=fit$numIter, converged=fit$convergence)
    out.tab = rbind(out.tab, out)
    fits = c(fits, list(fit))
  }
}
```

Again we sort the models by AICc and compute model weights.
9.5 Question 2, Are the subpopulations independent?

```r
min.AICc = order(out.tab$AICc)
out.tab.2 = out.tab[min.AICc,]
fits = fits[min.AICc]
out.tab.2 = cbind(out.tab.2, delta.AICc = out.tab.2$AICc - out.tab.2$AICc[1])
out.tab.2 = cbind(out.tab.2, rel.like = exp(-1*out.tab.2$delta.AICc/2))
out.tab.2 = cbind(out.tab.2, AIC.weight = out.tab.2$rel.like/sum(out.tab.2$rel.like))
```

Examination of the expanded results table (out.tab.2) shows there is strong support for correlated process errors; top 10 models shown:

<table>
<thead>
<tr>
<th>Model</th>
<th>Q</th>
<th>delta.AICc</th>
<th>AIC.weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC+Strait+PS+SC equalvarcov</td>
<td></td>
<td>0.00</td>
<td>0.976</td>
</tr>
<tr>
<td>site</td>
<td></td>
<td>7.65</td>
<td>0.021</td>
</tr>
<tr>
<td>NC+Strait+PS+SC unconstrained</td>
<td></td>
<td>11.47</td>
<td>0.003</td>
</tr>
<tr>
<td>NC+Strait+PS+SC diagonal and equal</td>
<td></td>
<td>23.39</td>
<td>0.000</td>
</tr>
<tr>
<td>NC+Strait+PS+SC diagonal and unequal</td>
<td></td>
<td>27.53</td>
<td>0.000</td>
</tr>
<tr>
<td>N-S unconstrained</td>
<td></td>
<td>32.61</td>
<td>0.000</td>
</tr>
<tr>
<td>N-S diagonal and unequal</td>
<td></td>
<td>36.06</td>
<td>0.000</td>
</tr>
<tr>
<td>N-S equalvarcov</td>
<td></td>
<td>36.97</td>
<td>0.000</td>
</tr>
<tr>
<td>stock</td>
<td></td>
<td>37.82</td>
<td>0.000</td>
</tr>
<tr>
<td>N-S diagonal and equal</td>
<td></td>
<td>38.16</td>
<td>0.000</td>
</tr>
</tbody>
</table>

The model weight for “equalvarcov”, “unconstrained”, versus “diagonal and equal” is:

```r
c(sum(out.tab.2$AIC.weight[out.tab.2$Q=="equalvarcov"]), sum(out.tab.2$AIC.weight[out.tab.2$Q=="unconstrained"]),
sum(out.tab.2$AIC.weight[out.tab.2$Q=="diagonal and equal"])
)
```

[1] 0.997 0.003 0.000

9.5.1 Looking at the correlation structure in the Q matrix

The 3rd model in the output table is a model with all elements of the process error variance-covariance matrix estimated. Estimating a variance-covariance matrix with so many extra parameters is not supported relative to a constrained variance-covariance matrix with two parameters (compare the AICc for the 1st model and 3rd model) but looking at the full variance-covariance matrix shows some interesting and not surprising patterns.

The Q matrix is recovered from the model fit using this command:

```r
Q.unc = coef(fits[[3]], type="matrix")$Q
```

The diagonal of this matrix shows that each region appears to experience process variability of a similar magnitude:

```r
diag(Q.unc)
```
We can compute the correlation matrix as follows. Row names are added to make the matrix more readable.

\[
h = \text{diag}(1/\sqrt{\text{diag}(Q.\text{unc})})
\]
\[
Q.\text{corr} = h^T \cdot Q.\text{unc} \cdot h
\]
\[
\text{rownames}(Q.\text{corr}) = \text{unique}(Z4)
\]
\[
\text{colnames}(Q.\text{corr}) = \text{unique}(Z4)
\]
\[
Q.\text{corr}
\]

<table>
<thead>
<tr>
<th></th>
<th>nc</th>
<th>is</th>
<th>ps</th>
<th>sc</th>
</tr>
</thead>
<tbody>
<tr>
<td>nc</td>
<td>1.00</td>
<td>0.59</td>
<td>0.64</td>
<td>0.92</td>
</tr>
<tr>
<td>is</td>
<td>0.59</td>
<td>1.00</td>
<td>0.99</td>
<td>0.23</td>
</tr>
<tr>
<td>ps</td>
<td>0.64</td>
<td>0.99</td>
<td>1.00</td>
<td>0.28</td>
</tr>
<tr>
<td>sc</td>
<td>0.92</td>
<td>0.23</td>
<td>0.28</td>
<td>1.00</td>
</tr>
</tbody>
</table>

The correlation matrix indicates that the inland strait (‘is’) subpopulation experiences process errors (good and bad years) that are almost perfectly correlated with the Puget Sound subpopulation though the two have different population growth rates (Figure 9.3). Similarly the north and south coastal subpopulations (‘nc’ and ‘sc’) experience highly correlated process errors, though again population growth rates are much higher to the north. There is much higher correlation between the process errors of the north coastal subpopulation and the nearby inland straits and Puget Sound subpopulations than between the two inland subpopulations and the much farther south coastal subpopulation. These patterns are not ecologically surprising but are not easy to discern looking at the raw count data.

### 9.6 Question 3, Is the Hood Canal independent?

In the initial analysis, the data from Hood Canal were removed. Hood Canal has experienced a series of hypoxic events which has led to large perturbations to the harbor seal prey. We will add the Hood Canal data back in and look at whether treating Hood Canal as separate is supported compared to treating it as part of the Puget Sound subpopulation in the top model.

\[
\text{sealData.hc} = \text{rbind}(\text{sealData}, \text{harborSeal[,8]})
\]
\[
\text{rownames}(\text{sealData.hc})[12] = \text{"Hood.Canal"}
\]

Here are the two Z matrices for a ‘Hood Canal in the Puget Sound’ and ‘Hood Canal separate’ model:

\[
ZH1 = \text{factor(c("nc","nc","is","is","ps","ps","sc","sc","nc","sc","is","ps"))}
\]
\[
ZH2 = \text{factor(c("nc","nc","is","is","ps","ps","sc","sc","nc","sc","is","sc","hc"))}
\]
Z.models.hc=list(ZH1, ZH2)
names(Z.models.hc)=c("hood.in.ps","hood.separate")

We will test three different \( Q \) matrices: a matrix with one variance and one covariance, an unconstrained variance-covariance matrix and a variance-covariance where the Hood Canal subpopulation has independent process errors.

\[
Q3=\begin{bmatrix}
\text{"offdiag"} & \text{"offdiag"} & \text{"offdiag"} & \text{"offdiag"} & 0 \\
\text{"offdiag"} & \text{"q"} & \text{"offdiag"} & \text{"offdiag"} & 0 \\
\text{"offdiag"} & \text{"offdiag"} & \text{"q"} & \text{"offdiag"} & 0 \\
\text{"offdiag"} & \text{"offdiag"} & \text{"offdiag"} & \text{"q"} & 0 \\
0 & 0 & 0 & 0 & \text{"q.hc"}
\end{bmatrix}
\]

As before, we loop through the model and create a results table:

\[
\text{out.tab.hc}=\text{NULL} \\
\text{fits.hc}=\text{list()} \\
\text{for}(i \text{ in } 1:\text{length(Z.models.hc)}){ \\
  \text{for}(j \text{ in } 1:\text{length(Q.models)}){ \\
    \text{if}(i==1 & j==3) \text{ next #Q3 is only for Hood Separate model} \\
    Q.model=Q.models[j] \\
    fit.model = c(list(Z=Z.models.hc[[i]], Q=Q.model), model.constant) \\
    fit = \text{MARSS(sealData.hc, model=fit.model, silent=\text{TRUE}, control=list(maxit=1000))} \\
    \text{out}=\text{data.frame(H=names(Z.models.hc)[[i]], Q=names(Q.models)[j], U=U.model, logLik=fit$logLik, AICc=fit$AICc, num.param=fit$num.params, m=length(unique(Z.models.hc)[[i]])), num.iter=fit$numIter, converged=fit$convergence) \\
    \text{out.tab.hc}=\text{rbind(out.tab.hc, out)} \\
    \text{fits.hc}=\text{c(fits.hc, list(fit))} \\
  } \\
}\}

We sort the results by AICc and compute the \( \Delta \text{AICc} \).

\[
\text{min.AICc}=\text{order(out.tab.hc$AICc)} \\
\text{out.tab.hc}=\text{out.tab.hc[min.AICc,]}
\]
The results table \((\text{out.tab hc})\) indicates strong support for treating Hood Canal as a separate subpopulation but not support for completely independent process errors.

<table>
<thead>
<tr>
<th>H</th>
<th>Q</th>
<th>delta.AICc</th>
<th>AICc.weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>hood.separate</td>
<td>equalvarcov</td>
<td>0.00</td>
<td>0.988</td>
</tr>
<tr>
<td>hood.separate hood.independent</td>
<td></td>
<td>8.74</td>
<td>0.012</td>
</tr>
<tr>
<td>hood.in.ps</td>
<td>equalvarcov</td>
<td>23.53</td>
<td>0.000</td>
</tr>
<tr>
<td>hood.separate unconstrained</td>
<td></td>
<td>30.65</td>
<td>0.000</td>
</tr>
<tr>
<td>hood.in.ps unconstrained</td>
<td>unconstrained</td>
<td>36.66</td>
<td>0.000</td>
</tr>
</tbody>
</table>

### 9.7 Discussion

In this chapter, we used model selection and AICc model weights to explore the temporal correlation structure in the harbor seal abundance data from the U.S. west coast. We used the term ‘subpopulation’, however it should be kept in mind that we are actually looking at the data support for different spatial patterns of temporal correlation in the process errors. Treating region A and B as a ‘subpopulation’ in this context means that we are asking if the counts from A and B can be treated as observations of the same underlying stochastic trajectory.

Metapopulation structure refers to a case where a larger population is composed on a collection of smaller temporally independent subpopulations. Metapopulation structure buffers the variability seen in the larger population and has important consequences for the viability of a population. We tested for temporal independence using diagonal versus non-diagonal \(Q\) matrices. Although the west coast harbor seal population appears to divided into ‘subpopulations’ that experience different population growth rates, there is strong temporal correlation in the year-to-year variability experienced in these subpopulations. This suggests that this harbor seal population does not function as a true metapopulation with independent subpopulations but rather as a collection of subpopulations that are temporal correlated.
10

Dynamic factor analysis (DFA)

10.1 Overview

In this chapter, we use MARSS to do dynamic factor analysis (DFA), which allows us to look for a set of common underlying trends among a relatively large set of time series (Harvey [1989] sec. 8.5). See also Zuur et al. [2003], which shows a number of examples of DFA applied to fisheries catch data and densities of zoobenthos. We will walk through some examples to show you the math behind DFA, and then in Section 10.4, we will show a short-cut for doing a DFA with MARSS using form="dfa".

DFA is conceptually different than what we have been doing in the previous applications. Here we are trying to explain temporal variation in a set of \( n \) observed time series using linear combinations of a set of \( m \) hidden random walks, where \( m \ll n \). A DFA model is a type of MARSS model with the following structure:

\[
x_t = x_{t-1} + \mathbf{w}_t \quad \text{where} \quad \mathbf{w}_t \sim \text{MVN}(0, \mathbf{Q}) \\
y_t = Zx_t + \mathbf{a} + \mathbf{v}_t \quad \text{where} \quad \mathbf{v}_t \sim \text{MVN}(0, \mathbf{R}) \\
x_0 \sim \text{MVN}(\pi, \Lambda)
\]  

(10.1)

The general idea is that the observations \( \mathbf{y} \) are modeled as a linear combination of hidden trends \( \mathbf{x} \) and factor loadings \( Z \) plus some offsets \( \mathbf{a} \). The DFA model in Equation 10.1 and the standard MARSS model in Equation 1.1 are equivalent—we have simply set the matrix \( B \) equal to an \( m \times m \) identity matrix\(^1\) and the vector \( \mathbf{u} = \mathbf{0} \).

---

\(^1\) a diagonal matrix with 1’s on the diagonal
10.1.1 Writing out a DFA model in MARSS form

Imagine a case where we had a data set with six observed time series \((n = 6)\) and we want to fit a model with three hidden trends \((m = 3)\). If we write out our DFA model in MARSS matrix form (ignoring the error structures and initial conditions for now), it would look like this:

\[
\begin{bmatrix}
  x_1 \\
  x_2 \\
  x_3
\end{bmatrix}_t =
\begin{bmatrix}
  1 & 0 & 0 \\
  0 & 1 & 0 \\
  0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
  x_1 \\
  x_2 \\
  x_3
\end{bmatrix}_t - 1 +
\begin{bmatrix}
  w_1 \\
  w_2 \\
  w_3
\end{bmatrix}_t
\]

\[
\begin{bmatrix}
  y_1 \\
  y_2 \\
  y_3 \\
  y_4 \\
  y_5 \\
  y_6
\end{bmatrix}_t =
\begin{bmatrix}
  z_{11} & z_{12} & z_{13} \\
  z_{21} & z_{22} & z_{23} \\
  z_{31} & z_{32} & z_{33} \\
  z_{41} & z_{42} & z_{43} \\
  z_{51} & z_{52} & z_{53} \\
  z_{61} & z_{62} & z_{63}
\end{bmatrix}
\begin{bmatrix}
  x_1 \\
  x_2 \\
  x_3
\end{bmatrix}_t +
\begin{bmatrix}
  a_1 \\
  a_2 \\
  a_3 \\
  a_4 \\
  a_5 \\
  a_6
\end{bmatrix}_t +
\begin{bmatrix}
  v_1 \\
  v_2 \\
  v_3 \\
  v_4 \\
  v_5 \\
  v_6
\end{bmatrix}_t.
\]

(10.2)

The process errors of the hidden trends would be

\[
\begin{bmatrix}
  w_1 \\
  w_2 \\
  w_3
\end{bmatrix}_t \sim MVN \left( \begin{bmatrix} 0 \\
  0 \\
  0 \end{bmatrix}, \begin{bmatrix} q_{11} & q_{12} & q_{13} \\
  q_{12} & q_{22} & q_{23} \\
  q_{13} & q_{23} & q_{33} \end{bmatrix} \right),
\]

(10.3)

and the observation errors would be

\[
\begin{bmatrix}
  v_1 \\
  v_2 \\
  v_3 \\
  v_4 \\
  v_5 \\
  v_6
\end{bmatrix}_t \sim MVN \left( \begin{bmatrix} 0 \\
  0 \\
  0 \\
  0 \\
  0 \\
  0 \end{bmatrix}, \begin{bmatrix} r_{11} & r_{12} & r_{13} & r_{14} & r_{15} & r_{16} \\
  r_{12} & r_{22} & r_{23} & r_{24} & r_{25} & r_{26} \\
  r_{13} & r_{23} & r_{33} & r_{34} & r_{35} & r_{36} \\
  r_{14} & r_{24} & r_{34} & r_{44} & r_{45} & r_{46} \\
  r_{15} & r_{25} & r_{35} & r_{45} & r_{55} & r_{56} \\
  r_{16} & r_{26} & r_{36} & r_{46} & r_{56} & r_{66} \end{bmatrix} \right).
\]

(10.4)

10.1.2 Constraints to ensure identifiability

If \(Z, a,\) and \(Q\) in Equation 10.1 are not constrained, then the DFA model above is unidentifiable (Harvey, 1989, sec 4.4). Harvey (1989, sec. 8.5.1) suggests the following parameter constraints to make the model identifiable:

- in the first \(m - 1\) rows of \(Z,\) the \(z\)-value in the \(j\)-th column and \(i\)-th row is set to zero if \(j > i;\)
- \(a\) is constrained so that the first \(m\) values are set to zero; and
- \(Q\) is set equal to the identity matrix \((I_m)\).
However, found that with Harvey’s second constraint, the EM algorithm is not particularly robust, and it takes a long time to converge. Zuur et al. found that the EM estimates are much better behaved if you instead constrain each of the time series in $x$ to have a mean of zero across $t = 1$ to $T$. To do so, they replaced the estimates of the hidden states, $x_t^T$, coming out of the Kalman smoother with $x_t^T - \bar{x}$ for $t = 1$ to $T$, where $\bar{x}$ is the mean of $x_t$ across $t$. With this approach, you estimate all of the $a$ elements, which represent the average level of $y_t$ relative to $Z(x_t - \bar{x})$. We found that demeaning the $x_t^T$ in this way can cause the EM algorithm to have errors (decline in log-likelihood). Instead, we demean our data, and fix all elements of $a$ to zero.

Using these constraints, the DFA model in Equation 10.2 becomes

$$
\begin{bmatrix}
x_1 \\
x_2 \\
x_3
\end{bmatrix}_t =
\begin{bmatrix}
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
x_1 \\
x_2 \\
x_3
\end{bmatrix}_{t-1} +
\begin{bmatrix}
0 \\
0 \\
0
\end{bmatrix} +
\begin{bmatrix}
w_1 \\
w_2 \\
w_3
\end{bmatrix}_t
$$

(10.5)

The process errors of the hidden trends in Equation 10.3 would then become

$$
\begin{bmatrix}
w_1 \\
w_2 \\
w_3
\end{bmatrix}_t \sim \text{MVN}\left(\begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix}, \begin{bmatrix} 5 & 0 & 0 \\ 0 & 5 & 0 \\ 0 & 0 & 5 \end{bmatrix}\right),
$$

(10.6)

but the observation errors in Equation 10.4 would stay the same, such that

$$
\begin{bmatrix}
v_1 \\
v_2 \\
v_3 \\
v_4 \\
v_5 \\
v_6
\end{bmatrix}_t \sim \text{MVN}\left(\begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix}, \begin{bmatrix} r_{11} & r_{12} & r_{13} & r_{14} & r_{15} & r_{16} \\ r_{12} & r_{22} & r_{23} & r_{24} & r_{25} & r_{26} \\ r_{13} & r_{23} & r_{33} & r_{34} & r_{35} & r_{36} \\ r_{14} & r_{24} & r_{34} & r_{44} & r_{45} & r_{46} \\ r_{15} & r_{25} & r_{35} & r_{45} & r_{55} & r_{56} \\ r_{16} & r_{26} & r_{36} & r_{46} & r_{56} & r_{66} \end{bmatrix}\right),
$$

(10.7)

To complete our model, we still need the final form for the initial conditions of the state. Following Zuur et al. (2003), we set the initial state vector ($x_0$) to have zero mean and a diagonal variance-covariance matrix with large variances, such that

$$
x_0 \sim \text{MVN}\left(\begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \end{bmatrix}, \begin{bmatrix} 5 & 0 & 0 \\ 0 & 5 & 0 \\ 0 & 0 & 5 \end{bmatrix}\right).
$$

(10.8)
10.2 The data

We will analyze some of the Lake Washington plankton data included in the MARSS package. This dataset includes 33 years of monthly counts for 13 plankton species along with data on water temperature, total phosphorous (TP), and pH. First, we load the data and then extract a subset of columns corresponding to the phytoplankton species only. For the purpose of speeding up model fitting times and to limit our analysis to years with no missing covariate data, we will only examine 10 years of data (1980-1989).

\[
data(lakeWAplankton)
\]
# we want lakeWAplanktonTrans, which has been log-transformed # and the 0s replaced with NAs
plankdat = lakeWAplanktonTrans
years = plankdat[,"Year"]>=1980 & plankdat[,"Year"]<1990
phytos = c("Cryptomonas", "Diatoms", "Greens", "Unicells", "Other.algae")

\[
dat.spp.1980 = plankdat[years,phytos]
\]

Next, we transpose the data and calculate the number of time series and their length.

# transpose data so time goes across columns
dat.spp.1980 = t(dat.spp.1980)
N.ts = nrow(dat.spp.1980)
TT = ncol(dat.spp.1980)

It is normal in this type of analysis to standardize each time series by first subtracting its mean and then dividing by its standard deviation (i.e., create a \(z\)-score \(y^*_t\) with mean = 0 and standard deviation = 1), such that

\[
y^*_t = \Sigma^{-1}(y_t - \bar{y}),
\]

\(\Sigma\) is a diagonal matrix with the standard deviations of each time series along the diagonal, and \(\bar{y}\) is a vector of the means. In R, this can be done as follows

\[
Sigma = sqrt(apply(dat.spp.1980, 1, var, na.rm=TRUE))
y.bar = apply(dat.spp.1980, 1, mean, na.rm=TRUE)
dat.z = (dat.spp.1980 - y.bar) * (1/Sigma)
rownames(dat.z) = rownames(dat.spp.1980)
\]

MARSS also has a helper function to \(z\)-score data:

\[
dat.z = zscore(dat.spp.1980)
\]

Figure 10.1 shows time series of Lake Washington phytoplankton data following \(z\)-score transformation.
Fig. 10.1. Time series of Lake Washington phytoplankton data following z-score transformation.

10.3 Setting up the model in MARSS

As we have seen in other cases, setting up the model structure for MARSS requires that the parameter matrices have a one-to-one correspondence to the model as you would write it on paper (i.e., Equations 10.5 through 10.8). If a parameter matrix has a combination of fixed and estimated values, then you specify that using \texttt{matrix(list(), nrow, ncol)}. This is a matrix of class \texttt{list()} and allows you to combine numeric and character values in a single matrix. MARSS recognizes the numeric values as fixed values and the character values as estimated values.

This is how we set up \textbf{Z} for MARSS, assuming a model with 5 observed time series and 3 hidden trends:

\begin{verbatim}
Z.vals = list(
    "z11", 0 , 0 ,
    "z21","z22", 0 ,
    "z31","z32","z33",
    "z41","z42","z43",
\end{verbatim}
"z51","z52","z53")

\[ Z = \text{matrix}(Z.\text{vals}, \text{nrow}=N.\text{ts}, \text{ncol}=3, \text{byrow}=\text{TRUE}) \]

When specifying the list values, spacing and carriage returns were added to help show the correspondence with the \(Z\) matrix in Equation [10.3] If you print \(Z\) (at the R command line), you will see that it is a matrix with character values (the estimated elements) and numeric values (the fixed 0's).

\[
\begin{array}{ccc}
[1,] & \text{"z11"} & 0 & 0 \\
[2,] & \text{"z21"} & \text{"z22"} & 0 \\
[3,] & \text{"z31"} & \text{"z32"} & \text{"z33"} \\
[4,] & \text{"z41"} & \text{"z42"} & \text{"z43"} \\
[5,] & \text{"z51"} & \text{"z52"} & \text{"z53"} \\
\end{array}
\]

Notice that the 0's do not have quotes around them. If they did, it would mean the "0" is a character value and would be interpreted as the name of a parameter to be estimated rather than a fixed numeric value.

The \(Q\) and \(B\) matrices are both set equal to the identity matrix using \(\text{diag()}\).

\[
Q = B = \text{diag}(1,3)
\]

For our first analysis, we will assume that each time series of phytoplankton has a different observation variance, but that there is no covariance among time series. Thus, \(R\) should be a diagonal matrix that looks like:

\[
\begin{bmatrix}
 r_{11} & 0 & 0 & 0 & 0 \\
 0 & r_{22} & 0 & 0 & 0 \\
 0 & 0 & r_{33} & 0 & 0 \\
 0 & 0 & 0 & r_{44} & 0 \\
 0 & 0 & 0 & 0 & r_{55}
\end{bmatrix},
\]

and each of the \(r_{ij}\) elements is a different parameter to be estimated. We can also specify this \(R\) structure using a list matrix as follows:

\[
R.\text{vals} = \text{list}(\\n \text{"r11"},0,0,0,0, \\
0,\text{"r22"},0,0,0, \\
0,0,\text{"r33"},0,0, \\
0,0,0,\text{"r44"},0, \\
0,0,0,0,\text{"r55"})
\]

\[
R = \text{matrix}(R.\text{vals}, \text{nrow}=N.\text{ts}, \text{ncol}=N.\text{ts}, \text{byrow}=\text{TRUE})
\]
You can print `R` at the R command line to see what it looks like:

```r
print(R)
[1,] "r11" 0 0 0 0
[2,] 0 "r22" 0 0 0
[3,] 0 0 "r33" 0 0
[4,] 0 0 0 "r44" 0
[5,] 0 0 0 0 "r55"
```

This form of variance-covariance matrix is commonly used, and therefore MARSS has a built-in shorthand for this structure. Alternatively, we could simply type:

```r
R = "diagonal and unequal"
```

Type `?MARSS` at the R command line to see a list of the shorthand options for each parameter vector/matrix.

The parameter vectors $\pi$ (termed $x_0$ in MARSS), $a$ and $u$ are each set to be a column vector of zeros. Any of the following can be used:

```r
x0 = U = matrix(0, nrow=3, ncol=1)
A = matrix(0, nrow=6, ncol=1)
x0 = U = A = "zero"
```

The $\Lambda$ matrix (termed $V_0$ in MARSS) is a diagonal matrix with 5’s along the diagonal:

```r
V0 = diag(5,3)
```

Finally, we make a list of the model parameters to pass to the `MARSS()` function and set the control list:

```r
dfa.model = list(Z=Z, A="zero", R=R, B=B, U=U, Q=Q, x0=x0, V0=V0)
cntl.list = list(maxit=50)
```

For the examples in this chapter, we have set the maximum iterations to 50 to speed up model fitting. Note, however, that the parameter estimates will not have converged to their maximum likelihood values, which would likely take 100s, if not 1000+, iterations.

### 10.3.1 Fitting the model

We can now pass the DFA model list to `MARSS()` to estimate the $Z$ matrix and underlying hidden states ($x$). The output is not shown because it is voluminous, but the model fits are plotted in Figure 10.2. The warnings regarding non-convergence are due to setting `maxit` to 50.

```r
kemz.3 = MARSS(dat.z, model=dfa.model, control=cntl.list)
```
Warning! Reached maxit before parameters converged. Maxit was 50. Neither abstol nor log-log convergence tests were passed.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
WARNING: maxit reached at 50 iter before convergence. Neither abstol nor log-log convergence test were passed. The likelihood and params are not at the ML values. Try setting control$maxit higher.
Log-likelihood: -782.202
AIC: 1598.404  AICc: 1599.463

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Z.z11</td>
<td>0.4163</td>
</tr>
<tr>
<td>Z.z21</td>
<td>0.5364</td>
</tr>
<tr>
<td>Z.z31</td>
<td>0.2780</td>
</tr>
<tr>
<td>Z.z41</td>
<td>0.5179</td>
</tr>
<tr>
<td>Z.z51</td>
<td>0.1611</td>
</tr>
<tr>
<td>Z.z22</td>
<td>0.6757</td>
</tr>
<tr>
<td>Z.z32</td>
<td>-0.2381</td>
</tr>
<tr>
<td>Z.z42</td>
<td>-0.2381</td>
</tr>
<tr>
<td>Z.z52</td>
<td>-0.2230</td>
</tr>
<tr>
<td>Z.z33</td>
<td>0.2305</td>
</tr>
<tr>
<td>Z.z43</td>
<td>-0.1225</td>
</tr>
<tr>
<td>Z.z53</td>
<td>0.3887</td>
</tr>
<tr>
<td>R.(Cryptomonas,Cryptomonas)</td>
<td>0.6705</td>
</tr>
<tr>
<td>R.(Diatoms,Diatoms)</td>
<td>0.0882</td>
</tr>
<tr>
<td>R.(Greens,Greens)</td>
<td>0.7201</td>
</tr>
<tr>
<td>R.(Unicells,Unicells)</td>
<td>0.1865</td>
</tr>
<tr>
<td>R.(Other.algae,Other.algae)</td>
<td>0.5441</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0
Standard errors have not been calculated.
Use MARSSParamCIs to compute CIs and bias estimates.

Convergence warnings
10 warnings. First 10 shown. Type cat(object$errors) to see the full list.
Warning: the Z.z51 parameter value has not converged.
Warning: the Z.z32 parameter value has not converged.
Warning: the Z.z52 parameter value has not converged.
Warning: the Z.z33 parameter value has not converged.
Warning: the Z.z43 parameter value has not converged.
Warning: the R.(Diatoms,Diatoms) parameter value has not converged.
Warning: the R.(Greens,Greens) parameter value has not converged.
Warning: the R.(Other.algae,Other.algae) parameter value has not converged.
Warning: the logLik parameter value has not converged.
Type MARSSinfo("convergence") for more info on this warning.

Fig. 10.2. Plots of Lake Washington phytoplankton data with model fits (dark lines) from a model with 3 trends and a diagonal and unequal variance-covariance matrix for the observation errors. This model was run to convergence so is different than that shown in the text which uses \( \text{maxit}=50 \).

10.4 Using model selection to determine the number of trends

Following \cite{Zuur et al. (2003)}, we use model selection criteria (specifically AICc) to determine the number of underlying trends that have the highest data support. Our first model had three underlying trends \((m = 3)\). Let's compare this to a model with two underlying trends. The forms for parameter matrix \( R \) and vector \( a \) will stay the same but we need to change the other parameter vectors and matrices because \( m \) is different.
After showing you the matrix math behind a DFA model, we will now use the `form` argument for a MARSS call to specify that we want to fit a DFA model. Type `?MARSS.dfa` to learn about the `MARSS()` call with `form="dfa"`. This will set up the $Z$ matrix and the other parameters for you. Specify how many trends you want by passing in `model=list(m=x)`. You can also pass in different forms for the $R$ matrix in the usual way.

Here is how to fit two trends using `form="dfa"`:

```r
model.list = list(m=2, R="diagonal and unequal")
kemz.2 = MARSS(dat.spp.1980, model=model.list,
  z.score=TRUE, form="dfa", control=cntl.list)
if(!saved.res) {
  model.list = list(m=2, R="diagonal and unequal")
kemz.2 = MARSS(dat.spp.1980, model=model.list,
  z.score=TRUE, form="dfa", control=big.maxit.cntl.list)
}
```

and compare its AICc value to that from the 3-trend model.

```r
print(cbind(model=c("3 trends", "2 trends"),
  AICc=round(c(kemz.3$AICc, kemz.2$AICc))),
  quote=FALSE)
```

```r
   model  AICc
[1,] 3 trends 1589
[2,] 2 trends 1608
```

It looks like a model with 3 trends has much more support from the data because its AICc value is more than 10 units less than that for the 2-trend model.

### 10.4.1 Comparing many model structures

Now let’s examine a larger suite of possible models. We will test from one to four underlying trends ($m = 1$ to 4) and four different structures for the $R$ matrix:

1. same variances & no covariance ("diagonal and equal");
2. different variances & no covariance ("diagonal and unequal");
3. same variances & same covariance ("equalvarcov"); and
4. different variances & covariances ("unconstrained").

The following code builds our model matrices; you could also write out each matrix as we did in the first example, but this allows us to build and run all of the models together. **NOTE:** the following piece of code will take a very long time to run!
10.5 Using varimax rotation to determine the loadings and trends

As Harvey [1989] p. 450, sec. 8.5.1) discusses, there are multiple equivalent solutions to the dynamic factor loadings. We arbitrarily constrained Z in such a way to choose only one of these solutions, but fortunately the different solutions are equivalent, and they can be related to each other by a rotation matrix H. Let H be any $m \times m$ non-singular matrix. The following are then equivalent solutions:

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

(10.9)

and
Table 10.1. Model selection results.

<table>
<thead>
<tr>
<th>R</th>
<th>m</th>
<th>logLik</th>
<th>delta.AICc</th>
<th>Ak.wt</th>
<th>Ak.wt.cum</th>
</tr>
</thead>
<tbody>
<tr>
<td>unconstrained</td>
<td>3</td>
<td>-762.5</td>
<td>0.0</td>
<td>0.39</td>
<td>0.39</td>
</tr>
<tr>
<td>unconstrained</td>
<td>2</td>
<td>-765.9</td>
<td>0.1</td>
<td>0.37</td>
<td>0.76</td>
</tr>
<tr>
<td>unconstrained</td>
<td>4</td>
<td>-761.5</td>
<td>2.3</td>
<td>0.12</td>
<td>0.89</td>
</tr>
<tr>
<td>unconstrained</td>
<td>1</td>
<td>-772.4</td>
<td>4.4</td>
<td>0.04</td>
<td>0.93</td>
</tr>
<tr>
<td>diagonal and unequal</td>
<td>4</td>
<td>-774.2</td>
<td>5.9</td>
<td>0.02</td>
<td>0.95</td>
</tr>
<tr>
<td>equalvarcov</td>
<td>2</td>
<td>-782.7</td>
<td>6.1</td>
<td>0.02</td>
<td>0.97</td>
</tr>
<tr>
<td>diagonal and unequal</td>
<td>3</td>
<td>-777.1</td>
<td>7.5</td>
<td>0.01</td>
<td>0.98</td>
</tr>
<tr>
<td>diagonal and equal</td>
<td>4</td>
<td>-779.3</td>
<td>7.7</td>
<td>0.01</td>
<td>0.99</td>
</tr>
<tr>
<td>diagonal and equal</td>
<td>3</td>
<td>-781.8</td>
<td>8.4</td>
<td>0.01</td>
<td>0.99</td>
</tr>
<tr>
<td>equalvarcov</td>
<td>4</td>
<td>-779.0</td>
<td>9.1</td>
<td>0.00</td>
<td>1.00</td>
</tr>
<tr>
<td>equalvarcov</td>
<td>3</td>
<td>-781.4</td>
<td>9.9</td>
<td>0.00</td>
<td>1.00</td>
</tr>
<tr>
<td>diagonal and unequal</td>
<td>2</td>
<td>-786.6</td>
<td>20.2</td>
<td>0.00</td>
<td>1.00</td>
</tr>
<tr>
<td>equalvarcov</td>
<td>1</td>
<td>-799.9</td>
<td>32.3</td>
<td>0.00</td>
<td>1.00</td>
</tr>
<tr>
<td>diagonal and equal</td>
<td>2</td>
<td>-798.4</td>
<td>35.4</td>
<td>0.00</td>
<td>1.00</td>
</tr>
<tr>
<td>diagonal and unequal</td>
<td>1</td>
<td>-798.4</td>
<td>35.4</td>
<td>0.00</td>
<td>1.00</td>
</tr>
<tr>
<td>diagonal and equal</td>
<td>1</td>
<td>-813.5</td>
<td>57.4</td>
<td>0.00</td>
<td>1.00</td>
</tr>
</tbody>
</table>

\[
y_t = ZH^{-1}x_t^\dagger + a + v_t
\]

\[
x_t^\dagger = x_{t-1}^\dagger + w_t^\dagger
\]

\[
x_t^\dagger = Hx_t; w_t^\dagger = Hw_t
\]

\(x^\dagger\) are the rotated trends.

There are many ways of doing factor rotations, but a common approach is the varimax rotation which seeks a rotation matrix \(H\) that creates the largest difference between loadings. For example, let’s say there are three trends in our model. In our estimated \(Z\) matrix, let’s say row 3 is \((0.2, 0.2, 0.2)\). That would mean that data series 3 is equally described by trends 1, 2, and 3. If instead row 3 was \((0.8, 0.1, 0.1)\), this would make interpretation easier because we could say that data time series 3 was mostly described by trend 1. The varimax rotation finds the \(H\) matrix that makes the \(Z\) rows more like \((0.8, 0.1, 0.1)\) and less like \((0.2, 0.2, 0.2)\).

The varimax rotation is easy to compute because \(R\) has a built in function that returns \(H^{-1}\). We will illustrate the use of the varimax rotation with the 2-state model with \(R\) unconstrained. We will fit this with a large maxit.

```r
big.maxit.cntl.list = list(minit=200, maxit=5000, allow.degen=FALSE)
model.list = list(m=2, R="unconstrained")
the.fit = MARSS(dat.z, model=model.list, form="dfa", control=big.maxit.cntl.list)
```

Next, we retrieve the matrix used for varimax rotation.

```r
# get the inverse of the rotation matrix
Z.est = coef(the.fit, type="matrix")$Z
```
The rotation matrix that varimax returns $H^{-1}$ rather than $H$. If $Z$ has one column, there is only one $Z$; there is only a rotation matrix if $Z$ has more than one column. We use $H^{-1}$ to rotate the factor loadings and $H$ to rotate the trends as in Equation 10.10.

```
# rotate factor loadings
Z.rot = Z.est %*% H.inv
# rotate trends
trends.rot = solve(H.inv) %*% the.fit$states
```

Rotated factor loadings for the model are shown in Figure 10.3. Oddly, some taxa appear to have no loadings on some trends (e.g., diatoms on trend 1). The reason is that, merely for display purposes, we chose to plot only those loadings that are greater than 0.05, and it turns out that after varimax rotation, several loadings are close to 0.

Recall that we set $\text{Var}(w_t) = Q = I_m$ in order to make our DFA model identifiable. Does the variance in the process errors also change following varimax rotation? Interestingly, no. Because $H$ is a non-singular, orthogonal matrix, $\text{Var}(Hw_t) = H\text{Var}(w_t)H^\top = HI_mH^\top = I_m$.

10.6 Examining model fits

Now that we have done the appropriate factor and trends rotations, we should examine some plots of model fits. To do so, we will create a function `getDFAfits()` to extract the model fits and estimated $(1-\alpha)\%$ confidence intervals. Note, the function `residuals()` and `broom::augment()` will also return this information.

```
fit.b = getDFAfits(the.fit)
```

First, it looks like this model captures some of the high frequency variation (i.e., seasonality) in the time series (see Figure 10.5). Second, some of the time series had much better overall fits than others (e.g., compare Cryptomonas and Unicells). Given the obvious seasonal patterns in the phytoplankton data, it would be worthwhile to first detrend the data and then repeat the model fitting exercise to see (1) how many trends would be favored, and (2) the shape of those trends.

Alternatively, you can use the `broom::augment()` function to obtain a summary of the estimates and use `ggplot()` to plot:

```
Type `RShowDoc("Chapter_DFA.R",package="MARSS")` at the R command line to open a file with all the code for this chapter and search for the function name.
```
#plot the factor loadings
spp = rownames(dat.z)
minZ = 0.05
m=dim(trends.rot)[1]
ylims = c(-1.1*max(abs(Z.rot)), 1.1*max(abs(Z.rot)))
par(mfrow=c(ceiling(m/2),2), mar=c(3,4,1.5,0.5), oma=c(0.4,1,1,1))
for(i in 1:m) {
  plot(c(1:N.ts)[abs(Z.rot[,i])>minZ], as.vector(Z.rot[abs(Z.rot[,i])>minZ,i]),
       type="h", lwd=2, xlab="", ylab="", xaxt="n", ylim=ylims, xlim=c(0,N.ts+1))
  for(j in 1:N.ts) {
    if(Z.rot[j,i] > minZ) {text(j, -0.05, spp[j], srt=90, adj=1, cex=0.9)}
    if(Z.rot[j,i] < -minZ) {text(j, 0.05, spp[j], srt=90, adj=0, cex=0.9)}
    abline(h=0, lwd=1, col="gray")
  } # end j loop
  mtext(paste("Factor loadings on trend",i,sep=" "),side=3, line=.5)
} # end i loop

Fig. 10.3. Plot of the factor loadings (following varimax rotation) from the 2-state model fit to the phytoplankton data.
Fig. 10.4. Plot of the unobserved trends (following varimax rotation) from the 2-state model fit to the phytoplankton data.

Fig. 10.5. Plot of the 2-state model fits to the phytoplankton data.
10 Dynamic factor analysis

```r
require(broom)
require(ggplot2)
theme_set(theme_bw())
d <- augment(the.fit, interval="confidence")
ggplot(data = d) +
  geom_line(aes(t, .fitted)) +
  geom_point(aes(t, y)) +
  geom_ribbon(aes(x=t, ymin=.conf.low, ymax=.conf.up), linetype=2, alpha=0.2) +
  facet_grid(~.rownames) +
  xlab("Time Step") +
  ylab("Count")
```

![Plot of the 2-state model fits to the phytoplankton data using ggplot.](image)

**Fig. 10.6.** Plot of the 2-state model fits to the phytoplankton data using ggplot.

### 10.7 Adding covariates

It is standard to add covariates to the analysis so that one removes known important drivers. The DFA with covariates is written:

\[
x_t = x_{t-1} + w_t \quad \text{where} \quad w_t \sim \text{MVN}(0, Q)
\]
\[
y_t = Zx_t + a + Dd_t + v_t \quad \text{where} \quad v_t \sim \text{MVN}(0, R)
\]
\[
x_0 \sim \text{MVN}(\pi, \Lambda)
\]  

(10.11)
where the $q \times 1$ vector $d_t$ contains the covariate(s) at time $t$, and the $n \times q$ matrix $D$ contains the effect(s) of the covariate(s) on the observations. Using `form="dfa"` and `covariates=<covariate name(s)>`, we can easily add covariates to our DFA, but this means that the covariates are input, not data, and there can be no missing values (see Chapter 13 for how to include covariates with missing values).

The Lake Washington dataset has two environmental covariates that we might expect to have effects on phytoplankton growth, and hence, abundance: temperature (Temp) and total phosphorous (TP).

```
temp = t(plankdat[years,"Temp",drop=FALSE])
TP = t(plankdat[years,"TP",drop=FALSE])
```

We will now fit three different models that each add covariate effects (i.e., Temp, TP, Temp & TP) to our 2-state model with R “unconstrained”.

```
model.list=list(m=2, R="unconstrained")
kemz.temp = MARSS(dat.spp.1980, model=model.list, z.score=TRUE,
form="dfa", control=cntl.list, covariates=temp)
kemz.TP = MARSS(dat.spp.1980, model=model.list, z.score=TRUE,
form="dfa", control=cntl.list, covariates=TP)
kemz.both = MARSS(dat.spp.1980, model=model.list, z.score=TRUE,
form="dfa", control=cntl.list, covariates=rbind(temp,TP))
```

Next we can compare whether the addition of the covariates improves the model fit (effectively less residual error while accounting for the additional parameters). *NOTE:* The following results were obtained by letting the EM algorithm run for a very long time, so your results may differ.

```
print(cbind(model=c("no covars", "Temp", "TP", "Temp & TP"),
AICc=round(c(the.fit$AICc, kemz.temp$AICc, kemz.TP$AICc,
   kemz.both$AICc))), quote=FALSE)
```

```
   model  AICc
[1,] no covars 1582
[2,] Temp      1518
[3,] TP        1568
[4,] Temp & TP 1522
```

This suggests that adding temperature or phosphorus to the model, either alone or in combination with one another, improves overall model fit. If we were interested in assessing the “best” model structure that includes covariates, however, we should examine all combinations of trends and structures for R. The model fits for the temperature-only model are shown in Fig 10.7 and they appear much better than the model without any covariates.
Fig. 10.7. Plot of the fits from the temperature-only model to the phytoplankton data.

10.8 Discussion

We analyzed the phytoplankton data alone. You can try analyzing the zooplankton data (type `head(plankdat)` to see the names). You can also try analyzing the phytoplankton and zooplankton together. You can also try different assumptions concerning the structure of $R$; we just tried unconstrained, diagonal and unequal, and diagonal and equal. To see all the R code behind the figures, type `RShowDoc("Chapter_DFA.R", package="MARSS")`. This opens a file with all the code. Copy and paste the code into a new file, and then you can edit that code. DFA models often take an unusually long time to converge. In a real DFA, you will want to make sure to try different initial starting values (see chapter 6), and force the algorithm to run a long time by using `minit=x` and `maxit=(x+c)`, where $x$ and $c$ are something like 200 and 5000, respectively.
Analyzing noisy animal tracking data

11.1 A simple random walk model of animal movement

A simple random walk model of movement with drift (directional movement) but no correlation is

\[ x_{1,t} = x_{1,t-1} + u_1 + w_{1,t}, \quad w_{1,t} \sim N(0, \sigma^2_1) \]  \hspace{1cm} (11.1)

\[ x_{2,t} = x_{2,t-1} + u_2 + w_{2,t}, \quad w_{2,t} \sim N(0, \sigma^2_2) \]  \hspace{1cm} (11.2)

where \( x_{1,t} \) is the location at time \( t \) along one axis (here, longitude) and \( x_{2,t} \) is for another, generally orthogonal, axis (here, latitude). The parameter \( u_1 \) is the rate of longitudinal movement and \( u_2 \) is the rate of latitudinal movement.

We add errors to our observations of location:

\[ y_{1,t} = x_{1,t} + v_{1,t}, \quad v_{1,t} \sim N(0, \eta^2_1) \]  \hspace{1cm} (11.3)

\[ y_{2,t} = x_{2,t} + v_{2,t}, \quad v_{2,t} \sim N(0, \eta^2_2), \]  \hspace{1cm} (11.4)

This model is comprised of two separate univariate state-space models. Note that \( y_1 \) depends only on \( x_1 \) and \( y_2 \) depends only on \( x_2 \). There are no actual interactions between these two univariate models. However, we can write the model down in the form of a multivariate model using diagonal variance-covariance matrices and a diagonal design (\( Z \)) matrix. Because the variance-covariance matrices and \( Z \) are diagonal, the \( x_1:y_1 \) and \( x_2:y_2 \) processes will be independent as intended. Here are Equations 11.2 and 11.4 written as a MARSS model (in matrix form):

Type `RShowDoc("Chapter_AnimalTracking.R",package="MARSS")` at the R command line to open a file with all the code for the examples in this chapter.
Analyzing animal tracking data

\[ \begin{bmatrix} x_{1,t} \\ x_{2,t} \end{bmatrix} = \begin{bmatrix} x_{1,t-1} \\ x_{2,t-1} \end{bmatrix} + \begin{bmatrix} \mu_1 \\ \mu_2 \end{bmatrix} + \begin{bmatrix} w_{1,t} \\ w_{2,t} \end{bmatrix}, \quad w_t \sim \text{MVN} \left( 0, \begin{bmatrix} \sigma_1^2 & 0 \\ 0 & \sigma_2^2 \end{bmatrix} \right) \] (11.5)

\[ \begin{bmatrix} y_{1,t} \\ y_{2,t} \end{bmatrix} = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} \begin{bmatrix} x_{1,t} \\ x_{2,t} \end{bmatrix} + \begin{bmatrix} v_{1,t} \\ v_{2,t} \end{bmatrix}, \quad v_t \sim \text{MVN} \left( 0, \begin{bmatrix} \eta_1^2 & 0 \\ 0 & \eta_2^2 \end{bmatrix} \right) \] (11.6)

The variance-covariance matrix for \( w_t \) is a diagonal matrix with unequal variances, \( \sigma_1^2 \) and \( \sigma_2^2 \). The variance-covariance matrix for \( v_t \) is a diagonal matrix with unequal variances, \( \eta_1^2 \) and \( \eta_2^2 \). We can write this succinctly as

\[ x_t = x_{t-1} + u + w_t, \quad w_t \sim \text{MVN}(0, Q) \] (11.7)

\[ y_t = x_t + v_t, \quad v_t \sim \text{MVN}(0, R). \] (11.8)

11.2 Loggerhead sea turtle tracking data

Loggerhead sea turtles (\textit{Caretta caretta}) are listed as threatened under the United States Endangered Species Act of 1973. Over the last ten years, a number of state and local agencies have been deploying ARGOS tags on loggerhead turtles on the east coast of the United States. We have data on eight individuals over that period. In this chapter, we use some turtle data from the WhaleNet Archive of STOP Data, however we have corrupted this data severely by adding random errors in order to create a “bad tag” problem. We corrupted latitude and longitude data by errors (Figure 11.1) and it would appear that our sea turtles are becoming land turtles (at least part of the time). We will the MARSS model to estimate true positions and speeds from the corrupted data.

Our noisy data are in \textit{loggerheadNoisy}. They consist of daily readings of location (longitude/latitude). If data are missing for a day, then the entries for latitude and longitude for that day should be NA. However, to make the code in this chapter run quickly, we have interpolated all missing values in the original, uncorrupted, dataset (\textit{loggerhead}). The first six lines of the corrupted data look like so

\texttt{loggerheadNoisy[1:6,]}

\begin{verbatim}
   turtle month day year lon lat
1  BigMama  5  28 2001 -81.45989 31.70337
2  BigMama  5  29 2001 -80.88292 32.18865
3  BigMama  5  30 2001 -81.27393 31.67568
4  BigMama  5  31 2001 -81.59317 31.83092
5  BigMama  6  1  2001 -81.35969 32.12685
6  BigMama  6  2  2001 -81.15644 31.89568
\end{verbatim}

The file has data for eight turtles:
We will analyze the position data for “Big Mama”. We put the data for “Big Mama” into matrix \texttt{dat}. \texttt{dat} is transposed because we need time across the columns.

\begin{verbatim}
turtlename="BigMama"
theTurtle=which(loggerheadNoisy$turtle==turtlename)
dat = loggerheadNoisy[theTurtle, 5:6]
dat = t(dat) #transpose
\end{verbatim}

Figure 11.1 shows the corrupted location data for Big Mama. The figure code uses the \texttt{maps} R package. You will need to install this R package in order to run the example code.

\begin{verbatim}
#load the map package; you have to install it first library(maps)
# Read in our noisy data (no missing values)
pdat = loggerheadNoisy #for plotting
turtlename="BigMama"
theTurtle=which(loggerheadNoisy$turtle==turtlename)
par(mai = c(0,0,0,0),mfrow=c(1,1))
map('state', region = c('florida', 'georgia', 'south carolina',
'north carolina', 'virginia', 'delaware','new jersey','maryland'),
xlim=c(-85,-70))
points(pdat$lon[theTurtle], pdat$lat[theTurtle],
col="blue",pch=21, cex=0.7)
\end{verbatim}

11.3 Estimate locations from bad tag data

We will begin by specifying the structure of the MARSS model and then use \texttt{MARSS()} to fit that model to the data. There are two state processes (one for latitude and the other for longitude), and there is one observation time series for each state process. As we saw in Equation 11.6, \( Z \) is the an identity matrix (a diagonal matrix with 1s on the diagonal). We could specify this structure as \texttt{Z.model="identity"} or \texttt{Z.model=\texttt{factor(c(1,2))}}). Although technically, this is unnecessary as this is the default form for \( Z \).

We will assume that the errors are independent and that there are different drift rates (\( u \)), process variances (\( \sigma^2 \)) and observation variances for latitude and longitude (\( \eta^2 \)).
Fig. 11.1. Plot of the tag data from the turtle Big Mama. Errors in the location data make it seem that Big Mama has been moving overland.

```r
Z.model="identity"
U.model="unequal"
Q.model="diagonal and unequal"
R.model="diagonal and unequal"
```

Fit the model to the data:

```r
kem = MARSS(dat, model=list(Z = Z.model,
                    Q = Q.model, R = R.model, U = U.model))
```

We can create a plot comparing the estimated and actual locations (Figure 11.2). The real locations (from which `loggerheadNoisy` was produced by adding noise) are in `loggerhead` and plotted with crosses. There are only a few data points for the real data because in the real tag data, there are many missing days.

```r
#Code to plot estimated turtle track against observations
#The estimates
pred.lon = kem$states[1,]
pred.lat = kem$states[2,]
```
```
par(mai = c(0,0,0,0),mfrow=c(1,1))
library(maps)
pdat=loggerheadNoisy
turtlename="BigMama"
map('state', region = c('florida', 'georgia', 'south carolina',
  'north carolina', 'virginia', 'delaware','new jersey','maryland'),
  xlim=c(-85,-70))
points(pdat$lon[theTurtle], pdat$lat[theTurtle],
  col="blue",pch=21, cex=0.7)
lines(pred.lon, pred.lat,col="red", lwd=2)
goodturtles = loggerhead
gooddat = goodturtles[which(goodturtles$turtle==turtlename),5:6]
points(gooddat[,1], gooddat[,2],col="black", lwd=2, pch=3,cex=1.1)
legend("bottomright",c("bad locations", "estimated true location",
  "good location data"),pch=c(1,-1,3),lty=c(-1,1,-1),
  col=c("blue","red","black"), bty=FALSE)
```

![Plot of the estimated track of the turtle Big Mama versus the good location data (before we corrupted it with noise).](image)
11.4 Estimate speeds for each turtle

Turtle biologists designated one of these loggerheads “Big Mama,” presumably for her size and speed. For each of the eight turtles, estimate the average miles traveled per day. To calculate the distance traveled by a turtle each day, you use the estimate (from \texttt{MARSS()}) of the lat/lon location of turtle at day $t$ and at day $t-1$. To calculate distance traveled in miles from lat/lon start and finish locations, we will use the function \texttt{GCDF}:

\begin{verbatim}
GCDF <- function(lon1, lon2, lat1, lat2, degrees=TRUE, units="miles") {
  temp = ifelse(degrees==FALSE,
    acos(sin(lat1)*sin(lat2)+cos(lat1)*cos(lat2)*cos(lon2-lon1)),
    acos(sin(lat1/57.2958)*sin(lat2/57.2958)+cos(lat1/57.2958)*cos(lat2/57.2958))
  r=3963.0 # (statute miles), default
  if("units"=="nm") r=3437.74677 # (nautical miles)
  if("units"=="km") r=6378.7 # (kilometers)
  return (r * temp)
}
\end{verbatim}

We can now compute the distance traveled each day by passing in lat/lon estimates from day $i-1$ and day $i$:

\begin{verbatim}
distance\[i-1\]=GCDF(pred.lon\[i-1\],pred.lon\[i\],
                   pred.lat\[i-1\],pred.lat\[i\])
\end{verbatim}

\texttt{pred.lon} and \texttt{pred.lat} are the predicted longitudes and latitudes from \texttt{MARSS()}: rows one and two in \texttt{kem$states}. To calculate the distances for all days, we put this through a \texttt{for} loop:

\begin{verbatim}
distance = array(NA, dim=c(dim(dat)[2]-1,1))
for(i in 2:dim(dat)[2])
  distance\[i-1\]=GCDF(pred.lon\[i-1\],pred.lon\[i\],
                        pred.lat\[i-1\],pred.lat\[i\])
\end{verbatim}

The command \texttt{mean(distance)} gives us the average distance per day. We can also make a histogram of the distances traveled per day (Figure [11.3]). We can compare the histogram of daily distances to what we would get if we had not accounted for measurement error (Figure [11.4]). We can also compare the mean miles per day:

\begin{verbatim}
#accounting for observation error
mean(distance)
[1] 15.53858

#assuming the data have no observation error
mean(distance.noerr)
[1] 34.80579
\end{verbatim}
11.5 Using specialized packages to analyze tag data

If you have real tag data to analyze, you should use a state-space modeling package that is customized for fitting MARSS models to tracking data. The MARSS package does not have all the bells and whistles that you would want for analyzing tracking data, particularly tracking data in the marine environment. Examples are the Ukfsst and kfttrack R packages:
# Compare to the distance traveled per day if you used the raw data

distance.noerr = array(NA, dim=c(dim(dat)[2]-1,1))
for(i in 2:dim(dat)[2])
  distance.noerr[i-1]=GCDF(dat[1,i-1],dat[1,i],dat[2,i-1],dat[2,i])
hist(distance.noerr) #make a histogram of distance traveled per day

![Histogram of distance.noerr](image)

**Fig. 11.4.** Histogram of the miles traveled per day for Big Mama with estimates that account for measurement error in the data.

UKFSST [https://github.com/positioning/kalmanfilter/wiki/ArticleUkfsst](https://github.com/positioning/kalmanfilter/wiki/ArticleUkfsst)

KFTRACK [https://github.com/positioning/kalmanfilter/wiki/ArticleKftrack](https://github.com/positioning/kalmanfilter/wiki/ArticleKftrack)

`kftrack` is a full-featured toolbox for analyzing tag data with extended Kalman filtering. It incorporates a number of extensions that are important for analyzing track data: barriers to movement such as coastlines and non-Gaussian movement distributions. With `kftrack`, you can use the real tag data which has big gaps, i.e. days with no location. `MARSS()` will struggle with these data because it will estimate states for all the unseen days; `kftrack` only fits to the seen days.

To use `kftrack` to fit the turtle data, type

```r
library(kftrack) # must be installed from github
loggerhead = loggerhead
```
turtlename = "BigMama"
dat = loggerhead[which(loggerhead$turtle == turtlename),2:6]
model = kftrack(dat, fix.first=FALSE, fix.last=FALSE,
                 var.struct="uniform")
12 Detection of outliers and structural breaks

12.1 Overview

This chapter is based on a short example shown on pages 147-148 in Koopman et al. (1999) using a 100-year record of river flow on the Nile River. The methods are based on Harvey et al. (1998) which is in turn based on techniques in Harvey and Koopman (1992) and Koopman (1993). The Nile dataset is included in R. Figure 12.1 shows the data.

12.2 Different models for the Nile flow levels

We begin by fitting different flow models to the data and compare these models with AIC. After that, we will use the model residuals to look for outliers and structural breaks.

12.2.1 Flat level model

We will start by modeling these data as a simple average river flow with variability around this level.

\[ y_t = a + v_t \text{ where } v_t \sim N(0,r) \]  \hspace{1cm} (12.1)

where \( y_t \) is the river flow volume at year \( t \) and \( x \) is some constant average flow level (notice it has no \( t \) subscript).

To fit this model with MARSS, we will explicitly show all the MARSS parameters.

Type `RShowDoc("Chapter_StructuralBreaks.R",package="MARSS")` at the R command line to open a file with all the code for the examples in this chapter.
Fig. 12.1. The Nile River flow volume 1871 to 1970 (included dataset in R).

\[
x_t = 1 \times x_{t-1} + 0 + w_t \text{ where } w_t \sim N(0,0) \]
\[
y_t = 0 \times x_t + a + v_t \text{ where } v_t \sim N(0,r) \]
\[
x_0 = 0
\]

(12.2)

MARSS includes the state process \( x_t \) but we are setting \( Z \) to zero so that does not appear in our observation model. We need to fix all the state parameters to zero so that the algorithm doesn’t “chase its tail” trying to fit \( x_t \) to the data.

An equivalent way to write this model is to use \( x_t \) as the average flow level and make it be a constant level by setting \( q = 0 \). The average flow appears as the \( x_0 \) parameter. In MARSS form, the model is:

\[
x_t = 1 \times x_{t-1} + 0 + w_t \text{ where } w_t \sim N(0,0) \]
\[
y_t = 1 \times x_t + 0 + v_t \text{ where } v_t \sim N(0,r) \]
\[
x_0 = a
\]

(12.3)
We will use this latter format since we will be building on this form. The model is specified as a list as follows and we denote this model “0”:

```r
mod.nile.0 = list(
  Z=matrix(1), A=matrix(0), R=matrix("r"),
  B=matrix(1), U=matrix(0), Q=matrix(0),
  x0=matrix("a")
)
```

We then fit the model with `MARSS()`:

```r
# The data is in a ts format, and we need a matrix
dat = t(as.matrix(Nile))
# Now we fit the model
kem.0 = MARSS(dat, model=mod.nile.0)
```

Success! algorithm run for 15 iterations. abstol and log-log tests passed. Alert: conv.test.slope.tol is 0.5. Test with smaller values (<0.1) to ensure convergence.

MARSS fit is

Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Algorithm ran 15 (=minit) iterations and convergence was reached.
Log-likelihood: -654.5157
AIC: 1313.031 AICc: 1313.155

Estimate
R.r 28352
x0.a 919
Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

### 12.2.2 Linear trend in flow model

Figure 12.2 shows the fit for the flat average river flow model. Looking at the data, we might expect that a declining average river flow would be better. In MARSS form, that model would be:

\[
\begin{align*}
  x_t &= 1 \times x_{t-1} + u + w_t \quad \text{where } w_t \sim N(0,0) \\
  y_t &= 1 \times x_t + 0 + v_t \quad \text{where } v_t \sim N(0,r) \\
  x_0 &= a
\end{align*}
\]  

(12.4)

where \( u \) is now the average per-year decline in river flow volume. The model is specified as a list as follows and we denote this model “1”:
mod.nile.1 = list(
  Z=matrix(1), A=matrix(0), R=matrix("r"),
  B=matrix(1), U=matrix("u"), Q=matrix(0),
  x0=matrix("a") )

We then fit the model with MARSS():

kem.1 = MARSS(dat, model=mod.nile.1, silent=TRUE)
glance(kem.1)

MARSSresiduals.tT reported warnings. See msg element of returned residuals object. coef.de
1 0.2081979 22644.48 3 -642.3159 1290.632 1290.882 0 0

Figure 12.2 shows the fits for the two models with deterministic models (flat and declining) for mean river flow along with their AICc values (smaller AICc is better). The AICc for the model with a declining river flow is lower by over 20 (which is a lot).

12.2.3 Stochastic level model

Looking at the flow levels, we might suspect that a model that allows the average flow to change would model the data better and we might suspect that there have been sudden, and anomalous, changes in the river flow level. We will now model the average river flow at year $t$ as a random walk, specifically an autoregressive process which means that average river flow is year $t$ is a function of average river flow in year $t-1$.

$$x_t = x_{t-1} + w_t \text{ where } w_t \sim N(0, q)$$
$$y_t = x_t + v_t \text{ where } v_t \sim N(0, r) \quad (12.5)$$
$$x_0 = \pi$$

As before, $y_t$ is the river flow volume at year $t$. With all the MARSS parameters shown, the model is:

$$x_t = 1 \times x_{t-1} + 0 + w_t \text{ where } w_t \sim N(0, q)$$
$$y_t = 1 \times x_t + 0 + v_t \text{ where } v_t \sim N(0, r) \quad (12.6)$$
$$x_0 = \pi$$

Thus, $Z = 1$, $a = 0$, $R = r$, $B = 1$, $u = 0$, $Q = q$, and $x_0 = \pi$. The model is then specified as:

mod.nile.2 = list(
  Z=matrix(1), A=matrix(0), R=matrix("r"),
  B=matrix(1), U=matrix("u"), Q=matrix("q"),
  x0=matrix("pi") )
We could also use the text shortcuts to specify the model. Because \( R \) and \( Q \) are \( 1 \times 1 \) matrices, “unconstrained”, “diagonal and unequal”, “diagonal and equal” and “equalvarcov” will all lead to a \( 1 \times 1 \) matrix with one estimated element. For \( a \) and \( u \), the following shortcut could be used:

\[
A=U=\text{"zero"}
\]

Because \( x_0 \) is \( 1 \times 1 \), it could be specified as “unequal”, “equal” or “unconstrained”.

We fit the model with the \texttt{MARSS()} function. We are using the “BFGS” algorithm to polish off the estimates, since it will get the maximum faster than the default EM algorithm as long as we start it close to the maximum.

\[
kem.2em = \text{MARSS}(\text{dat}, \text{model}=\text{mod.nile.2}, \text{silent}=\text{TRUE})
\]

\[
kem.2 = \text{MARSS}(\text{dat}, \text{model}=\text{mod.nile.2},
\text{inits}=\text{kem.2em}\$\text{par}, \text{method}=\text{"BFGS"}, \text{silent}=\text{TRUE})
\]

\[
tidy(kem.2, \text{conf.int}=\text{FALSE})
\]

<table>
<thead>
<tr>
<th>term</th>
<th>estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 R.r</td>
<td>15336.530</td>
</tr>
<tr>
<td>2 Q.q</td>
<td>1218.137</td>
</tr>
<tr>
<td>3 x0.pi</td>
<td>1111.591</td>
</tr>
</tbody>
</table>

This is the same model fit in \cite{Koopman1999} p. 148 except that we estimate \( x_1 \) as parameter rather than specifying \( x_1 \) via a diffuse prior. As a result, the log-likelihood value and \( R \) and \( Q \) are a little different than in \cite{Koopman1999}.

### 12.3 Observation and state residuals

Figure 12.2 shows the MARSS fits to the data. From these model fits, auxiliary residuals can be computed which contain information about whether the data and models fits at time \( t \) differ more than you would expect given the model and the model fits at time \( t-1 \). In this section, we follow the example shown on page 147-148 in \cite{Koopman1999} and use these residuals to look for outliers and sudden flow level changes. Using auxiliary residuals this way follows mainly from \cite{Harvey1992}, but see also \cite{Koopman1993}, \cite{deJong1998} and \cite{Penzer2001} for discussions of using auxiliary residuals for detection of outliers and structural breaks.

The MARSS function will output the expected values of \( x_t \) conditioned on the maximum-likelihood values of \( q, r, \) and \( x_1 \) on the data (\( y \) from \( t = 1 \) to \( T \)). In time-series literature, these are called the smoothed state estimates and they are output by the Kalman filter-smoother. We will call these smoothed estimates \( \hat{x}_{t|T} \) (and are \( x_{tT} \) in the MARSS output). The time value after the \( | \) in the subscript indicates the data on which the estimate was conditioned (in
this case, 1 to $T$). From these, we can compute the model predicted value of $y_t$, denoted or $\hat{y}_{i\mid T}$. This is the predicted value of $y_t$ conditioned on $\tilde{x}_{i\mid T}$.

$$\hat{x}_{i\mid T} = E(X_i\mid \hat{\theta}, y_1^T)$$
$$\hat{y}_{i\mid T} = E(Y_i\mid \hat{\theta}, \tilde{x}_{i\mid T})$$

$$= \tilde{x}_{i\mid T} + E(w_i\mid \hat{\theta}, y_1^T) = \tilde{x}_{i\mid T}$$

(12.7)

where $\hat{\theta}$ are the maximum-likelihood estimates of the parameters. The $\hat{y}_{i\mid T}$ equation comes directly from equation (12.5). This expectation is not conditioned on the data $y_1^T$, directly. It is conditioned on $\tilde{x}_{i\mid T}$, which is conditioned on $y_1^T$. 

**Fig. 12.2.** The Nile River flow volume with the model estimated flow rates (solid lines). The bottom model is a stochastic level model, and the 2 standard deviations for the level are also shown. The other two models are deterministic level models so the state is not stochastic and does not have a standard deviation.
12.3 Observation and state residuals

12.3.1 Using observation residuals to detect outliers

The standardized smoothed observation residuals\(^1\) are the difference between the data at time \(t\) and the model fit at time \(t\) conditioned on all the data standardized by the observation variance:

\[
\hat{v}_t = y_t - \hat{y}_t|f \\
e_t = \frac{1}{\sqrt{\text{var}(\hat{v}_t)}} \hat{v}_t
\] (12.8)

These residuals should have (asymptotically) a \(t\)-distribution (Kohn and Ansley [1989], sec. 3) and by looking at the residuals, we can identify potential outlier data points—or more accurately, we can identify data points that do not fit the model (Equation 12.5). The call `residuals()` will compute these residuals for a marssMLE object (output by a MARSS call). It returns the standardized residuals (also called auxiliary residuals) as a \(n+m\times T\) matrix. The first \(n\) rows are the estimated \(v_t\) standardized observation residuals and the next \(m\) rows are the estimated \(w_t\) standardized state residuals (discussed below).

\[
\text{resids.0} = \text{residuals(kem.0)}$std.residuals
\]

MARSSresiduals.tT reported warnings. See msg element of returned residuals object.

\[
\text{resids.1} = \text{residuals(kem.1)}$std.residuals
\]

MARSSresiduals.tT reported warnings. See msg element of returned residuals object.

\[
\text{resids.2} = \text{residuals(kem.2)}$std.residuals
\]

MARSSresiduals.tT reported warnings. See msg element of returned residuals object.

Figure [12.3] shows the observation residuals for the three models developed above. We immediately see that model 0 (flat level) and model 1 (linear declining level) have problems because the residuals are all positive for the first part of the time series and then all negative. The residuals should not be temporally correlated like that. Model 2 with a stochastic level shows well-behaving residuals with low temporal correlation between \(t\) and \(t-1\). Looking at the residuals for model 2, we see that there are a number of years with flow levels that appear to be outliers (are beyond the dashed level lines).

\(^1\) also called smoothations in the literature to distinguish them from innovations, which are \(y_t - E(Y_t|\hat{y}_{t-1})\). Notice that for innovations the expectation is conditioned on the data up to time \(t-1\) while for smoothations, we condition on all the data.
12.3.2 Detecting sudden level changes

The standardized smoothed state residuals \( f_t \) below are the difference between the estimated state at time \( t \) and the estimated state at time \( t - 1 \) conditioned on all the data standardized by the standard deviation:

\[
\hat{w}_t = \hat{x}_{t|T} - \hat{x}_{t-1|T}
\]

\[
f_t = \frac{1}{\sqrt{\text{var}(\hat{w}_t)}} \hat{w}_t
\]  

(12.9)

These state residuals do not show simple changes in the average level; \( x_t \) is clearly changing in Figure [12.2] bottom panel. Instead we are looking for “breaks” or sudden changes in the level. The bottom panel of Figure [12.4] shows the standardized state residuals \( (f_t) \). This shows, as we can see by eye, the average flow level in the Nile appears to have suddenly changed around the turn of the century when the first Aswan dam was built. The top panel shows the standardized observation residuals for comparison.
Fig. 12.4. Top panel, the standardized observation residuals. Bottom panel, the standardized state residuals. This replicates Figure 12 in Koopman et al. (1999).
13.1 Covariates as inputs

A MARSS model with covariate effects in both the process and observation components is written as:

\[
\begin{align*}
    x_t &= B_t x_{t-1} + u_t + C_t c_t + w_t, \quad \text{where } w_t \sim \text{MVN}(0, Q_t) \\
    y_t &= Z_t x_t + a_t + D_t d_t + v_t, \quad \text{where } v_t \sim \text{MVN}(0, R_t)
\end{align*}
\]

where \( c_t \) is the \( p \times 1 \) vector of covariates (e.g., temperature, rainfall) which affect the states and \( d_t \) is a \( q \times 1 \) vector of covariates (potentially the same as \( c_t \)), which affect the observations. \( C_t \) is an \( m \times p \) matrix of coefficients relating the effects of \( c_t \) to the \( m \times 1 \) state vector \( x_t \), and \( D_t \) is an \( n \times q \) matrix of coefficients relating the effects of \( d_t \) to the \( n \times 1 \) observation vector \( y_t \).

With the MARSS() function, one can fit this model by passing in \( \text{model}\$c \) and/or \( \text{model}\$d \) in the MARSS() call as a \( p \times T \) or \( q \times T \) matrix, respectively. The form for \( C_t \) and \( D_t \) is similarly specified by passing in \( \text{model}\$C \) and/or \( \text{model}\$D \). Because \( C \) and \( D \) are matrices, they must be passed in as an 3-dimensional array with the 3rd dimension equal to the number of time steps if they are time-varying. If they are time-constant, then they can be specified as 2-dimensional matrices.

13.2 Examples using plankton data

Here we show some examples using the Lake Washington plankton data set and covariates in that dataset. We use the 10 years of data from 1965-1974.

---

Type `RShowDoc("Chapter_Covariates.R", package="MARSS")` at the R command line to open a file with all the code for the examples in this chapter.
(Figure 13.1), a decade with particularly high green and blue-green algae levels. We use the transformed plankton dataset which has 0s replaced with NAs. Below, we set up the data and z-score the data. The original data were already z-scored, but we changed the mean when we sub-sampled the years so need to z-score again.

```r
fulldat = lakeWApTankonTrans
years = fulldat[, "Year"] >= 1965 & fulldat[, "Year"] < 1975
dat = t(fulldat[years, c("Greens", "Bluegreens")])
the.mean = apply(dat, 1, mean, na.rm = TRUE)
the.sigma = sqrt(apply(dat, 1, var, na.rm = TRUE))
dat = (dat - the.mean) * (1/the.sigma)
```

Next we set up the covariate data, temperature and total phosphorous. We z-score the covariates to standardize and remove the mean.

```r
covariates = rbind(
  Temp = fulldat[years, "Temp"],
  TP = fulldat[years, "TP"])
# z.score the covariates
# covariates = zscore(covariates)
```

### 13.3 Observation-error only model

We can estimate the effect of the covariates using a process-error only model, an observation-error only model, or a model with both types of error. An observation-error only model is a multivariate regression, and we will start here so you see the relationship of MARSS model to more familiar linear regression models.

#### 13.3.1 Multivariate linear regression

In a standard multivariate linear regression, we only have an observation model with independent errors (i.e., the state process does not appear in the model):

\[
y_t = a + Dd_t + v_t, \text{ where } v_t \sim MVN(0, R)
\]  \hspace{1cm} (13.2)

The elements in \( a \) are the intercepts and those in \( D \) are the slopes (effects). We have dropped the \( t \) subscript on \( a \) and \( D \) because these will be modeled as time-constant. Writing this out for the two plankton and the two covariates we get:

\[
\begin{bmatrix}
y_g \\
y_{bg}
\end{bmatrix}_t = \begin{bmatrix} a_1 \\ a_2 \end{bmatrix} + \begin{bmatrix} \beta_{g, temp} & \beta_{g, tp} \\ \beta_{bg, temp} & \beta_{bg, tp} \end{bmatrix} \begin{bmatrix} temp \\ tp \end{bmatrix}_{t-1} + \begin{bmatrix} v_1 \\ v_2 \end{bmatrix}
\]  \hspace{1cm} (13.3)

Let's fit this model with MARSS. The \( x \) part of the model is irrelevant so we want to fix the parameters in that part of the model. We won't set \( B = 0 \)
or $Z = 0$ since that might cause numerical issues for the Kalman filter. Instead we fix them as identity matrices and fix $x_0 = 0$ so that $x_t = 0$ for all $t$.

$Q = U = x0 = "zero"$; $B = Z = "identity"

$d = \text{covariates}$

$A = "zero"

$D = "unconstrained"

$y = \text{dat} \# \text{to show relationship between dat & the equation}$

model.list = list(B=B, U=U, Q=Q, Z=Z, A=A, D=D, d=d, x0=x0)

kem = MARSS(y, model=model.list)

Success! algorithm run for 15 iterations. abstol and log-log tests passed. Alert: conv.test.slope.tol is 0.5. Test with smaller values (<0.1) to ensure convergence.

MARSS fit is

Estimation method: kem

Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001

Algorithm ran 15 (=minit) iterations and convergence was reached.
Log-likelihood: -276.4287
AIC: 562.8573 AICc: 563.1351

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>R.diag</td>
<td>0.706</td>
</tr>
<tr>
<td>D. (Greens, Temp)</td>
<td>0.367</td>
</tr>
<tr>
<td>D. (Bluegreens, Temp)</td>
<td>0.392</td>
</tr>
<tr>
<td>D. (Greens, TP)</td>
<td>0.058</td>
</tr>
<tr>
<td>D. (Bluegreens, TP)</td>
<td>0.535</td>
</tr>
</tbody>
</table>

Initial states \((x_0)\) defined at \(t=0\)

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

We set \(A=\text{"zero"}\) because the data and covariates have been demeaned. Of course, one can do multiple regression in R using, say, \(\text{lm()}\), and that would be much, much faster. The EM algorithm is over-kill here, but it is shown so that you see how a standard multivariate linear regression model is written as a MARSS model in matrix form.

### 13.3.2 Multivariate linear regression with autocorrelated errors

We can add a twist to the standard multivariate linear regression model, and instead of having temporally \(i.i.d\). errors in the observation process, we’ll assume autoregressive errors. There is still no state process in our model, but we will use the state part of a MARSS model to model our errors. Mathematically, this can be written as

\[
x_t = B x_{t-1} + w_t, \quad \text{where} \quad w_t \sim \text{MVN}(0, Q)
\]

\[
y_t = D_d + x_t
\]

(13.4)

Here, the \(x_t\) are the errors for the observation model; they are modeled as an autoregressive process via the \(x\) equation. We drop the \(v_t\) (set \(R = 0\)) because the \(x_t\) in the \(y\) equation are now the observation errors. As usual, we have left the intercepts \((a\text{ and }u)\) off since the data and covariates are all demeaned.

Here’s how we fit this model in MARSS:

\[Q = \text{"unconstrained"}\]
\[B = \text{"diagonal and unequal"}\]
\[A = U = x0 = \text{"zero"}\]
\[R = \text{"diagonal and equal"}\]
\[d = \text{covariates}\]
\[D = \text{"unconstrained"}\]
\[y = \text{dat}\]

\[
\text{model.list = list(B=B, U=U, Q=Q, Z=Z, A=A, R=R, D=D, d=d, x0=x0)}
\]

\[
\text{control.list = list(maxit=1500)}
\]

\[
\text{kem = MARSS(y, model=model.list, control=control.list)}
\]
Success! abstol and log-log tests passed at 79 iterations.  
Alert: conv.test.slope.tol is 0.5.  
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 79 iterations.
Log-likelihood: -209.3408  
AIC: 438.6816  AICc: 439.7243

<table>
<thead>
<tr>
<th>Estimate</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.(diag)</td>
<td>0.0428</td>
</tr>
<tr>
<td>B.(X.Greens,X.Greens)</td>
<td>0.2479</td>
</tr>
<tr>
<td>B.(X.Bluegreens,X.Bluegreens)</td>
<td>0.9136</td>
</tr>
<tr>
<td>Q.(1,1)</td>
<td>0.7639</td>
</tr>
<tr>
<td>Q.(2,1)</td>
<td>-0.0285</td>
</tr>
<tr>
<td>Q.(2,2)</td>
<td>0.1265</td>
</tr>
<tr>
<td>D.(Greens,Temp)</td>
<td>0.3777</td>
</tr>
<tr>
<td>D.(Bluegreens,Temp)</td>
<td>0.2621</td>
</tr>
<tr>
<td>D.(Greens,TP)</td>
<td>0.0459</td>
</tr>
<tr>
<td>D.(Bluegreens,TP)</td>
<td>0.0675</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.  
Use MARSSparamCIs to compute CIs and bias estimates.

You can try setting B to identity and MARSS will fit a model with non-mean-reverting autoregressive errors to the data. It is not done here since it turns out that that is not a very good model and it takes a long time to fit. If you try it, you’ll see that Q gets small meaning that the x part is being removed from the model.

13.4 Process-error only model

Now let’s model the data as an autoregressive process observed without error, and incorporate the covariates into the process model. Note that this is much different from typical linear regression models. The x part represents our model of the data (in this case plankton species). How is this different from the autoregressive observation errors? Well, we are modeling our data as autoregressive so data at t − 1 affects the data at t. Population abundances are inherently autoregressive so this model is a bit closer to the underlying mechanism generating the data. Here is our new process model for plankton abundance. x is the plankton abundance.
\( x_t = x_{t-1} + Cc_t + w_t, \) where \( w_t \sim \text{MVN}(0, Q) \) \hspace{1cm} (13.5)

We can fit this as follows:

\[
R = A = U = "zero"; \quad B = Z = "identity" \\
Q = "equalvarcov" \\
C = "unconstrained" \\
x = \text{dat} \quad \text{to show the relation between dat & the equations} \\
\text{model.list = list(B=B, U=U, Q=Q, Z=Z, A=A, R=R, C=C, c=covariates)} \\
kem = \text{MARSS(x, model=model.list)}
\]

Success! algorithm run for 15 iterations. abstol and log-log tests passed. Alert: conv.test.slope.tol is 0.5.

Test with smaller values (<0.1) to ensure convergence.

MARSS fit is

Estimation method: kem

Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001

Algorithm ran 15 (=minit) iterations and convergence was reached.

Log-likelihood: -285.0732

AIC: 586.1465 \quad \text{AICc: 586.8225}

\[
\begin{array}{l}
\text{Estimate} \\
Q.\text{diag} & 0.7269 \\
Q.\text{offdiag} & -0.0210 \\
x0.\text{X.Greens} & -0.5189 \\
x0.\text{X.Bluegreens} & -0.2431 \\
C.(\text{X.Greens,Temp}) & -0.0434 \\
C.(\text{X.Bluegreens,Temp}) & 0.0988 \\
C.(\text{X.Greens,TP}) & -0.0589 \\
C.(\text{X.Bluegreens,TP}) & 0.0104 \\
\end{array}
\]

Initial states (x0) defined at t=0

Standard errors have not been calculated.

Use MARSSparamCIs to compute CIs and bias estimates.

Now, it looks like temperature has a strong negative effect on algae? Also our log-likelihood dropped a lot. Well, the data do not look at all like a random walk model (i.e., where \( B = 1 \)), which we can see from the plot of the data (Figure 13.1). The data are fluctuating about some mean so let’s switch to a better autoregressive model—a mean-reverting model. To do this, we will allow the diagonal elements of \( B \) to be something other than 1.

\[
\text{model.list}$B = "\text{diagonal and unequal}" \\
kem = \text{MARSS(dat, model=model.list)}
\]

Success! algorithm run for 15 iterations. abstol and log-log tests passed. Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Algorithm ran 15 (=minit) iterations and convergence was reached.
Log-likelihood: -236.6106
AIC: 493.2211  AICc: 494.2638

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>B.(X.Greens,X.Greens)</td>
<td>0.1981</td>
</tr>
<tr>
<td>B.(X.Bluegreens,X.Bluegreens)</td>
<td>0.7672</td>
</tr>
<tr>
<td>Q.diag</td>
<td>0.4899</td>
</tr>
<tr>
<td>Q.offdiag</td>
<td>-0.0221</td>
</tr>
<tr>
<td>x0.X.Greens</td>
<td>-1.2915</td>
</tr>
<tr>
<td>x0.X.Bluegreens</td>
<td>-0.4179</td>
</tr>
<tr>
<td>C.(X.Greens,Temp)</td>
<td>0.2844</td>
</tr>
<tr>
<td>C.(X.Bluegreens,Temp)</td>
<td>0.1655</td>
</tr>
<tr>
<td>C.(X.Greens,TP)</td>
<td>0.0332</td>
</tr>
<tr>
<td>C.(X.Bluegreens,TP)</td>
<td>0.1340</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

Notice that the log-likelihood goes up quite a bit, which means that the mean-reverting model fits the data much better.

With this model, we are estimating $x_0^0$ (the starting value for the Kalman filter). If we set `model$tinitx=1`, to use $x_0^0$ as the starting value instead, we will get an error message that $R$ diagonals are equal to 0 and we need to fix $x_0$. This is a restriction of the (default) EM algorithm having to do with the update equation for $x_0^0$. We cannot use BFGS unless we set $Q$ to be either unconstrained or diagonal because the way $Q$ is being estimated using the BFGS algorithm to ensure that the matrix stays positive-definite (via a Cholesky transformation) does not allow any constraints on $Q$.

### 13.5 Both process- & observation-error model

The MARSS package is really designed for state-space models where you have errors ($v$ and $w$) in both the process and observation models. For example,

$$
\begin{align*}
    x_t &= Bx_{t-1} + C_t c_t + w_t, \text{ where } w_t \sim \text{MVN}(0,Q) \\
    y_t &= x_t + v_t, \text{ where } v_t \sim \text{MVN}(0,R)
\end{align*}
$$

(13.6)

$x$ is the true algae abundances and $y$ is the observation of the $x$'s.
Let’s say we knew that the observation variance on the algae measurements was about 0.16 and we wanted to include that known value in the model. To do that, we can simply add $R$ to the model list from the process-error only model in the last example.

```r
model.list$R = diag(0.16,2)
kem = MARSS(dat, model=model.list)
```

Success! abstol and log-log tests passed at 27 iterations. Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 27 iterations.
Log-likelihood: -241.5301
AIC: 503.0603  AICc: 504.1029

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>B.(X.Greens,X.Greens)</td>
<td>0.31497</td>
</tr>
<tr>
<td>B.(X.Bluegreens,X.Bluegreens)</td>
<td>0.76205</td>
</tr>
<tr>
<td>Q.diag</td>
<td>0.33374</td>
</tr>
<tr>
<td>Q.offdiag</td>
<td>-0.00331</td>
</tr>
<tr>
<td>x0.X.Greens</td>
<td>-0.90020</td>
</tr>
<tr>
<td>x0.X.Bluegreens</td>
<td>-0.40473</td>
</tr>
<tr>
<td>C.(X.Greens,Temp)</td>
<td>0.23448</td>
</tr>
<tr>
<td>C.(X.Bluegreens,Temp)</td>
<td>0.16960</td>
</tr>
<tr>
<td>C.(X.Greens,TP)</td>
<td>0.02423</td>
</tr>
<tr>
<td>C.(X.Bluegreens,TP)</td>
<td>0.14120</td>
</tr>
</tbody>
</table>
Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

Note, our estimates of the effect of temperature and total phosphorous are not that different than what you get from a simple multiple regression (our first example). This might be because the autoregressive component is small, meaning the estimated diagonals on the $B$ matrix are small.

13.6 Including seasonal effects in MARSS models

Time-series data are often collected at intervals with some implicit seasonality. For example, quarterly earnings for a business, monthly rainfall totals, or
hourly air temperatures. In those cases, it is often helpful to extract any recurring seasonal patterns that might otherwise mask some of the other temporal dynamics we are interested in examining.

Here we show a few approaches for including seasonal effects using the Lake Washington plankton data, which were collected monthly. The following examples will use all five phytoplankton species from Lake Washington. First, let’s set up the data.

```r
years = fulldat[, "Year"]>=1965 & fulldat[, "Year"]<1975
phytos = c("Diatoms", "Greens", "Bluegreens", "Unicells", "Other.algae")
dat = t(fulldat[years, phytos])
# z.score data again because we changed the mean when we subsampled
dat = zscore(dat)
# number of time periods/samples
TT = ncol(dat)
```

### 13.6.1 Seasonal effects as fixed factors

One common approach for estimating seasonal effects is to treat each one as a fixed factor. This adds an estimated parameter for each season (e.g., 24 hours per day, 4 quarters per year). The plankton data are collected monthly, so we will treat each month as a fixed factor. To fit a model with fixed month effects, we create a $12 \times T$ covariate matrix $c$ with one row for each month (Jan, Feb, ...) and one column for each time point. We put a 1 in the January row for each column corresponding to a January time point, a 1 in the February row for each column corresponding to a February time point, and so on. All other values of $c$ equal 0. The following code will create such a $c$ matrix.

```r
# number of "seasons" (e.g., 12 months per year)
period = 12
# first "season" (e.g., Jan = 1, July = 7)
per.1st = 1
# create factors for seasons
c.in = diag(period)
for(i in 2:ceiling(TT/period)) {c.in = cbind(c.in, diag(period))}
# trim c.in to correct start & length
c.in = c.in[, (1:TT)+{per.1st-1}]
# better row names
rownames(c.in) = month.abb
```

Next we need to set up the form of the $C$ matrix which defines any constraints we want to set on the month effects. $C$ is a $5 \times 12$ matrix. Five taxon and 12 month effects. If we wanted each taxon to have the same month effect, a common month effect across all taxon, then we have the same value in each $C$ column$^1$:

$^1$ *month.abb* is a R constant that gives month abbreviations in text.
C = matrix(month.abb,5,12,byrow=TRUE)

[1,] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"
[2,] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"
[3,] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"
[4,] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"
[5,] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"

Notice, that C only has 12 values in it, the 12 common month effects. However, for this example, we will let each taxon have a different month effect thus allowing different seasonality for each taxon. For this model, we want each value in C to be unique:

C = "unconstrained"

Now C has 5 x 12 = 60 separate effects.

Then we set up the form for the rest of the model parameters. We make the following assumptions:

# Each taxon has unique density-dependence
B = "diagonal and unequal"
# Independent process errors
Q = "diagonal and unequal"
# We have demeaned the data & are fitting a mean-reverting model
# by estimating a diagonal B, thus
U = "zero"
# Each obs time series is associated with only one process
Z = "identity"
# The data are demeaned & fluctuate around a mean
A = "zero"
# Observation errors are independent, but they
# have similar variance due to similar collection methods
R = "diagonal and equal"
# No covariate effects in the obs equation
D = "zero"
d = "zero"

Now we can set up the model list for MARSS and fit the model (results are not shown since they are verbose with 60 different month effects).
13.6 Including seasonal effects in MARSS models

```r
model.list = list(B=B, U=U, Q=Q, Z=Z, A=A, R=R, C=C, c=c.in, D=D, d=d)
seas.mod.1 = MARSS(dat, model=model.list, control=list(maxit=1500))
```

# Get the estimated seasonal effects
# rows are taxa, cols are seasonal effects
seas.1 = coef(seas.mod.1, type="matrix")$C
rownames(seas.1) = phytos
colnames(seas.1) = month.abb

The top panel in Figure 13.2 shows the estimated seasonal effects for this model. Note that if we had set U="unequal", we would need to set one of the columns of C to zero because the model would be under-determined (infinite number of solutions). If we subtracted the mean January abundance off each time series, we could set the January column in C to 0 and get rid of 5 estimated effects.

### 13.6.2 Seasonal effects as a polynomial

The fixed factor approach required estimating 60 effects. Another approach is to model the month effect as a 3rd-order (or higher) polynomial: \(a + b \times m + c \times m^2 + d \times m^3\) where \(m\) is the month number. This approach has less flexibility but requires only 20 estimated parameters (i.e., 4 regression parameters times 5 taxa). To do so, we create a \(4 \times T\) covariate matrix \(c\) with the rows corresponding to 1, \(m\), \(m^2\), and \(m^3\), and the columns again corresponding to the time points. Here is how to set up this matrix:

```r
# number of "seasons" (e.g., 12 months per year)
period = 12
# first "season" (e.g., Jan = 1, July = 7)
per.1st = 1
# order of polynomial
poly.order = 3
# create polynomials of months
month.cov = matrix(1,1,period)
for(i in 1:poly.order) {month.cov = rbind(month.cov,(1:12)^i)}
# our c matrix is month.cov replicated once for each year
c.m.poly = matrix(month.cov, poly.order+1, TT+period, byrow=FALSE)
# trim c.in to correct start & length
c.m.poly = c.m.poly[,(1:TT)+(per.1st-1)]
# Everything else remains the same as in the previous example
model.list = list(B=B, U=U, Q=Q, Z=Z, A=A, R=R, C=C, c=c.m.poly, D=D, d=d)
seas.mod.2 = MARSS(dat, model=model.list, control=list(maxit=1500))
```

The effect of month \(m\) for taxon \(i\) is \(a_i + b_i \times m + c_i \times m^2 + d_i \times m^3\), where \(a_i\), \(b_i\), \(c_i\) and \(d_i\) are in the \(i\)-th row of \(C\). We can now calculate the matrix of seasonal effects as follows, where each row is a taxon and each column is a month:
\[ C.2 = \text{coef(seas.mod.2,type="matrix")$C} \]
\[ \text{seas.2 = C.2 \times \% \% \text{month.cov}} \]
\[ \text{rownames(seas.2) = phytos} \]
\[ \text{colnames(seas.2) = month.abb} \]

The middle panel in Figure 13.2 shows the estimated seasonal effects for this polynomial model.

### 13.6.3 Seasonal effects as a Fourier series

The factor approach required estimating 60 effects, and the 3rd order polynomial model was an improvement at only 20 parameters. A third option is to use a discrete Fourier series, which is combination of sine and cosine waves; it would require only 10 parameters. Specifically, the effect of month \( m \) on taxon \( i \) is \( a_i \times \cos(2\pi m/p) + b_i \times \sin(2\pi m/p) \), where \( p \) is the period (e.g., 12 months, 4 quarters), and \( a_i \) and \( b_i \) are contained in the \( i \)-th row of \( C \).

We begin by defining the \( 2 \times T \) seasonal covariate matrix \( c \) as a combination of 1 cosine and 1 sine wave:

\[ \begin{align*}
\text{cos.t} &= \cos(2 \times \pi \times \text{seq(TT) / period}) \\
\text{sin.t} &= \sin(2 \times \pi \times \text{seq(TT) / period}) \\
\text{c.Four} &= \text{rbind(cos.t,sin.t)}
\end{align*} \]

Everything else remains the same and we can fit this model as follows:

\[ \text{model.list = list(B=B,U=U,Q=Q,Z=Z,A=A,R=R,C=C,Four=D,D=d)} \]
\[ \text{seas.mod.3 = MARSS(dat, model=model.list, control=list(maxit=1500))} \]

We make our seasonal effect matrix as follows:

\[ \begin{align*}
\text{C.3 = coef(seas.mod.3, type="matrix")$C} \\
\# The time series of net seasonal effects \\
\text{seas.3 = C.3 \times \% \% \text{c.Four[,1:period]}} \\
\text{rownames(seas.3) = phytos} \\
\text{colnames(seas.3) = month.abb}
\end{align*} \]

The bottom panel in Figure 13.2 shows the estimated seasonal effects for this seasonal-effects model based on a discrete Fourier series.

Rather than rely on our eyes to judge model fits, we should formally assess which of the 3 approaches offers the most parsimonious fit to the data. Here is a table of AICc values for the 3 models:

\[ \text{data.frame(Model=c("Fixed", "Cubic", "Fourier"), AICc=round(c(seas.mod.1$AICc, seas.mod.2$AICc, seas.mod.3$AICc),1))} \]

<table>
<thead>
<tr>
<th>Model</th>
<th>AICc</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Fixed</td>
<td>1188.4</td>
</tr>
<tr>
<td>2 Cubic</td>
<td>1144.9</td>
</tr>
<tr>
<td>3 Fourier</td>
<td>1127.4</td>
</tr>
</tbody>
</table>
The model selection results indicate that the model with monthly seasonal effects estimated via the discrete Fourier sequence is the most parsimonious of the 3 models. Its AICc value is much lower than either the polynomial or fixed-effects models.

13.7 Model diagnostics

We will examine some basic model diagnostics for these three approaches by looking at plots of the model residuals and their autocorrelation functions (ACFs) for all five taxa using the following code:

```r
for(i in 1:3) {
  dev.new()
  modn = paste("seas.mod",i,sep=".")
  for(j in 1:5) {
    # Code for plotting residuals and ACFs
  }
}
```
Figures 13.3–13.5 shows these diagnostics for the three models. The model residuals for all taxa and models appear to show significant negative autocorrelation at lag=1, suggesting that a model with seasonal effects is inadequate to capture all of the systematic variation in phytoplankton abundance.

MARSSresiduals.tT reported warnings. See msg element of returned residuals object.

Fig. 13.3. Model residuals and their ACF for the model with fixed monthly effects.

13.8 Covariates with missing values or observation error

The specific formulation of Equation 13.1 creates restrictions on the assumptions regarding the covariate data. You have to assume that your covariate
data has no error, which is probably not true. You cannot have missing values in your covariate data, again unlikely. You cannot combine instrument time series; for example, if you have two temperature recorders with different error rates and biases. Also, what if you have one noisy temperature recorder in the first part of your time series and then you switch to a much better recorder in the second half of your time series? All these problems require pre-analysis massaging of the covariate data, leaving out noisy and gappy covariate data, and making what can feel like arbitrary choices about which covariate time series to include.

To circumvent these potential problems and allow more flexibility in how we incorporate covariate data, one can instead treat the covariates as components of an auto-regressive process by including them in both the process and observation models. Beginning with the process equation, we can write

\[
\text{residuals(seas.mod.2)$model.residuals[i,]
\]

\[
\text{data has no error, which is probably not true. You cannot have missing values in your covariate data, again unlikely. You cannot combine instrument time series; for example, if you have two temperature recorders with different error rates and biases. Also, what if you have one noisy temperature recorder in the first part of your time series and then you switch to a much better recorder in the second half of your time series? All these problems require pre-analysis massaging of the covariate data, leaving out noisy and gappy covariate data, and making what can feel like arbitrary choices about which covariate time series to include.}

To circumvent these potential problems and allow more flexibility in how we incorporate covariate data, one can instead treat the covariates as components of an auto-regressive process by including them in both the process and observation models. Beginning with the process equation, we can write
MARSSresiduals.tT reported warnings. See msg element of returned residuals object. MARSSresiduals.tT reported warnings. See msg element of returned residuals object.

Fig. 13.5. Model residuals and their ACF for the model with monthly effects estimated using a Fourier transform.

\[
\begin{bmatrix} x^{(v)}_t \\ x^{(c)}_t \end{bmatrix} = \begin{bmatrix} B^{(v)} & C \\ 0 & B^{(c)} \end{bmatrix} \begin{bmatrix} x^{(v)}_t \\ x^{(c)}_{t-1} \end{bmatrix} + \begin{bmatrix} u^{(v)}_t \\ u^{(c)}_t \end{bmatrix} + w_t,
\]

\[w_t \sim \text{MVN} \left( 0, \begin{bmatrix} Q^{(v)} & 0 \\ 0 & Q^{(c)} \end{bmatrix} \right) \] (13.7)

The elements with superscript \((v)\) are for the \(k\) variate states and those with superscript \((c)\) are for the \(q\) covariate states. The dimension of \(x^{(c)}\) is \(q \times 1\) and \(q\) is not necessarily equal to \(p\), the number of covariate observation time series in your dataset. Imagine, for example, that you have two temperature sensors and you are combining these data. Then you have two covariate observation time series \((p = 2)\) but only one underlying covariate state time series \((q = 1)\).

The matrix \(C\) is dimension \(k \times q\), and \(B^{(c)}\) and \(Q^{(c)}\) are dimension \(q \times q\). The dimension\(^2\) of \(x^{(v)}\) is \(k \times 1\), and \(B^{(v)}\) and \(Q^{(v)}\) are dimension \(k \times k\).

\(^2\) The dimension of \(x\) is always denoted \(m\). If your process model includes only variates, then \(k = m\), but now your process model includes \(k\) variates and \(q\) covariate states so \(m = k + q\).
Next, we can write the observation equation in an analogous manner, such that
\[
\begin{align*}
\begin{bmatrix} y^{(v)} \\ y^{(c)} \end{bmatrix}_t &= \begin{bmatrix} Z^{(v)} & D \\ 0 & Z^{(c)} \end{bmatrix} \begin{bmatrix} x^{(v)} \\ x^{(c)} \end{bmatrix}_t + \begin{bmatrix} a^{(v)} \\ a^{(c)} \end{bmatrix} + v_t,
\end{align*}
\]
\[v_t \sim \text{MVN} \left( 0, \begin{bmatrix} R^{(v)} & 0 \\ 0 & R^{(c)} \end{bmatrix} \right) \tag{13.8}
\]

The dimension of \(y^{(c)}\) is \(p \times 1\), where \(p\) is the number of covariate observation time series in your dataset. The dimension of \(y^{(v)}\) is \(l \times 1\), where \(l\) is the number of variate observation time series in your dataset. The total dimension of \(y\) is \(l + p\). The matrix \(D\) is dimension \(l \times q\), \(Z^{(c)}\) is dimension \(p \times q\), and \(R^{(c)}\) are dimension \(p \times p\). The dimension of \(Z^{(v)}\) is dimension \(l \times k\), and \(R^{(v)}\) are dimension \(l \times l\).

The \(D\) matrix would presumably have a number of all zero rows in it, as would the \(C\) matrix. The covariates that affect the states would often be different than the covariates that affect the observations. For example, mean annual temperature would affect population growth rates for many species while having little or no effect on observability, and turbidity might strongly affect observability in many types of aquatic, say, surveys but have little affect on population growth rate.

Our MARSS model with covariates now looks on the surface like a regular MARSS model:
\[
\begin{align*}
x_t &= Bx_{t-1} + u + w_t, \text{ where } w_t \sim \text{MVN}(0, Q) \\
y_t &= Zx_t + a + v_t, \text{ where } v_t \sim \text{MVN}(0, R) 
\end{align*}
\tag{13.9}
\]
with the \(x_t\), \(y_t\) and parameter matrices redefined as in Equations 13.7 and 13.8.

\[
\begin{align*}
x &= \begin{bmatrix} x^{(v)} \\ x^{(c)} \end{bmatrix}, & B &= \begin{bmatrix} B^{(v)} & C \\ 0 & B^{(c)} \end{bmatrix}, & u &= \begin{bmatrix} u^{(v)} \\ u^{(c)} \end{bmatrix}, & Q &= \begin{bmatrix} Q^{(v)} & 0 \\ 0 & Q^{(c)} \end{bmatrix} \\
y &= \begin{bmatrix} y^{(v)} \\ y^{(c)} \end{bmatrix}, & Z &= \begin{bmatrix} Z^{(v)} & D \\ 0 & Z^{(c)} \end{bmatrix}, & a &= \begin{bmatrix} a^{(v)} \\ a^{(c)} \end{bmatrix}, & R &= \begin{bmatrix} R^{(v)} & 0 \\ 0 & R^{(c)} \end{bmatrix} 
\end{align*}
\tag{13.10}
\]

Note \(Q\) and \(R\) are written as block diagonal matrices, but you could allow covariances if that made sense. \(u\) and \(a\) are column vectors here. We can fit the model (Equation 13.9) as usual using the\texttt{MARSS()} function.

The log-likelihood that is returned by MARSS will include the log-likelihood of the covariates under the covariate state model. If you want only the log-likelihood of the non-covariate data, you will need to subtract off the log-likelihood of the covariate model:
\[
\begin{align*}
x_t^{(c)} &= B^{(c)}x_{t-1}^{(c)} + u^{(c)} + w_t, \text{ where } w_t \sim \text{MVN}(0, Q^{(c)}) \\
y_t^{(c)} &= Z^{(c)}x_t^{(c)} + a^{(c)} + v_t, \text{ where } v_t \sim \text{MVN}(0, R^{(c)}) 
\end{align*}
\tag{13.11}
\]
An easy way to get this log-likelihood for the covariate data only is use the augmented model (Equation 13.9 with terms defined as in Equation 13.10) but pass in missing values for the non-covariate data. The following code shows how to do this.

```r
y.aug = rbind(data, covariates)
fit.aug = MARSS(y.aug, model=model.aug)
```

`fit.aug` is the MLE object that can be passed to `MARSSkf()`. You need to make a version of this MLE object with the non-covariate data filled with NAs so that you can compute the logLik without the covariates. This needs to be done in the `marss` element since that is what is used by MARSSkf(). Below is code to do this.

```r
fit.cov = fit.aug; fit.cov$marss$data[1:dim(data)[1],] = NA
extra.LL = MARSSkf(fit.cov)$logLik
```

Note that when you fit the augmented model, the estimates of $C$ and $B^{(c)}$ are affected by the non-covariate data since the model for both the non-covariate and covariate data are estimated simultaneously and are not independent (since the covariate states affect the non-covariates states). If you want the covariate model to be unaffected by the non-covariate data, you can fit the covariate model separately and use the estimates for $B^{(c)}$ and $Q^{(c)}$ as fixed values in your augmented model.
Estimation of species interaction strengths with and without covariates

14.1 Background

Multivariate autoregressive models (commonly termed MAR models) have been developed as a tool for analyzing community dynamics from time series data (Ives 1995, Ives et al. 1999, 2003, Hampton et al. 2013). These models are based on a process model for log abundances ($x_t$) of the form

$$x_t = Bx_{t-1} + u + w_t, \quad w_t \sim \text{MVN}(0, Q)$$

(14.1)

$B$ is the interaction matrix; self interaction strengths (density-dependence) are on the diagonal and inter-specific interaction strengths are on the off-diagonals such that $B_{ij}$ is the ‘effect’ of species $j$ on species $i$. This model has a stochastic equilibrium—it fluctuates around mean, $(I - B)^{-1}u$.

The term $u$ determines the mean level but once the system is at equilibrium, it does not affect the fluctuations relative to the mean. To see this, compare two models with $b = 0.5$ and $u = 1$ versus $u = 0$. The mean for the first is $1/(1 - 0.5) = 2$ and for the second is 0. If we start both 1 above the mean, the next $x$ is the same distance from the mean: $x_2 = 0.5(2 + 1) + 1 = 2.5$ and $x_2 = 0.5(0 + 1) + 0 = 0.5$. So both end up at 0.5 above the mean. So once the system is at equilibrium, it is ‘scale invariant’, where $u$ is the scaling term.

The way that Ives et al. (2003) write their process model (their Equation 10) is $X_t = A + BX_{t-1} + E_t$. The $A$ in Ives’s equation is the $u$ appearing in Equation 14.1 and the $E_t$ is our $w_t$.

Often the models include environmental covariates, but we will leave off covariates for the moment and address them at the end of the chapter. If we

Type `RShowDoc("Chapter_SpeciesInteractions.R", package="MARSS")` at the R command line to open a file with all the code for the examples in this chapter.
add a measurement process\(^1\), we have a MARSS model:

\[
y_t = Z x_t + a + v_t \text{ where } v_t \sim \text{MVN}(0, R)
\] (14.2)

Typically, we have one time series per species and thus we assume that \(m = n\) and \(Z\) is an \(m \times m\) identity matrix (when \(m = n\), \(a\) is set to 0). However, it is certainly possible to have multiple time series per species (for example data taken at multiple sites).

In this chapter, we will estimate the \(B\) matrix of species interactions for a simple wolf-moose system and for a four-species freshwater plankton system.

### 14.2 Two-species example using wolves and moose

Population dynamics of wolves and moose on Isle Royale, Michigan make an interesting case study of a two-species predator-prey interactions. These populations have been studied intensively since 1958\(^2\). Unlike other populations of gray wolves, the Isle Royale population has a diet dominated by one prey item, moose. The only predator of moose on Isle Royale is the gray wolf, as this population is not hunted.

We will use the wolf and moose winter census data from Isle Royale to learn how to fit community dynamics models to time-series data. The long-term January (wolf) and February (moose) population estimates are provided at [http://www.isleroyalewolf.org](http://www.isleroyalewolf.org).

The mathematical form of the process model for the wolf-moose population dynamics is

\[
\begin{bmatrix}
x_w \\
x_m
\end{bmatrix}_t =
\begin{bmatrix}
b_{w \rightarrow w} & b_{m \rightarrow w} \\
b_{w \rightarrow m} & b_{m \rightarrow m}
\end{bmatrix}
\begin{bmatrix}
x_w \\
x_m
\end{bmatrix}_{t-1} +
\begin{bmatrix}
\mu_w \\
\mu_m
\end{bmatrix} +
\begin{bmatrix}
w_w \\
w_m
\end{bmatrix},
\end{bmatrix}
\sim \text{MVN} \left( 0, \begin{bmatrix} q_w & 0 \\ 0 & q_m \end{bmatrix} \right)
\] (14.3)

where \(w\) denotes wolf and \(m\) denotes moose. \(w \rightarrow w\) is effect of wolf on wolf (density-dependence) and \(w \rightarrow m\) is effect of wolf on moose (predation effect on moose).

#### 14.2.1 Load in and plot the data

We will use 1960 to 2011. We will hold out 1959 as we will need that year when we look at the effect of covariates.

\(^1\) You can fit a MAR model with no observation error by setting \(R = 0\), but a conditional least-squares algorithm is vastly faster than EM or BFGS for the \(R = 0\) case (assuming no missing data).

\(^2\) There are many publications from this long-term study site; see [http://www.isleroyalewolf.org/wolfhome/tech_pubs.html](http://www.isleroyalewolf.org/wolfhome/tech_pubs.html) and the review here [http://www.isleroyalewolf.org/data/data/home.html](http://www.isleroyalewolf.org/data/data/home.html).
14.2 Two-species example using wolves and moose

\[
\text{yr1960to2011} = \text{isleRoyal[,"Year"]}\geq1960 \& \text{isleRoyal[,"Year"]}\leq2011 \\
\text{royale.dat} = \log(t(\text{isleRoyal}[\text{yr1960to2011}, c("Wolf", "Moose")]))
\]

\[
x = \text{isleRoyal[,"Year"]} \\
y = \log(\text{isleRoyal[,c("Wolf", "Moose")])} \\
\text{matplot}(x, y, \\
   \quad \text{ylab=\"log count\", xlab=\"Year\", type=\"l",} \\
   \quad \text{lwd=3, bty=\"L\", col=\"black\")} \\
\text{legend(\"topright\", c("Wolf", "Moose"), lty=c(1,2), bty=\"n\")}
\]

\[
\text{Fig. 14.1. Plot of the Isle Royale wolf and moose data.}
\]

14.2.2 Fit the model to the wolf-moose data

The naive way to fit the model is to use Equations 14.2 and 14.1 “as is”:

\[
\text{royale.model.0}=\text{list}(B=\text{"unconstrained"}, Q=\text{"diagonal and unequal"}, R=\text{"diagonal and unequal"}, U=\text{"unequal"}) \\
\text{kem.0=MARSS(royale.dat, model=royale.model.0)}
\]
If you try this, you will notice that it does not converge but stops when it reaches maxit and prints a number of warnings about non-convergence. The problem is that when you try to estimate $B$ and $u$, they are often confounded. This a well-known problem, and you will need to find a way to fix $u$ at some value. If you are willing to assume that the process is at equilibrium (i.e. not recovering to equilibrium from a big perturbation), then you can simply demean the data and set $u$ to 0. It is also common to standardize the variance by dividing by the square root of the variance of the data. This is called z-scoring the data.

```r
# if missing values are in the data, they should be NAs
z.royale.dat = zscore(royale.dat)
```

We can fit the model to the z-scored data, but we still have convergence issues.

```r
royale.model.1 = list(Z="identity", B="unconstrained",
        Q="diagonal and unequal", R="diagonal and unequal",
        U="zero", tinitx=1)
cntl.list = list(allow.degen=FALSE, maxit=200)
kem.1 = MARSS(z.royale.dat, model=royale.model.1, control=cntl.list)
```

Warning! Reached maxit before parameters converged. Maxit was 200. Neither abstol nor log-log convergence tests were passed.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
WARNING: maxit reached at 200 iter before convergence. Neither abstol nor log-log convergence test were passed. The likelihood and params are not at the ML values. Try setting control$maxit higher.
Log-likelihood: -75.56383
AIC: 171.1277  AICc: 173.4933

<table>
<thead>
<tr>
<th>Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>R.(Wolf,Wolf)</td>
</tr>
<tr>
<td>0.001433</td>
</tr>
<tr>
<td>R.(Moose,Moose)</td>
</tr>
<tr>
<td>0.000362</td>
</tr>
<tr>
<td>B.(1,1)</td>
</tr>
<tr>
<td>0.768031</td>
</tr>
<tr>
<td>B.(2,1)</td>
</tr>
<tr>
<td>-0.178990</td>
</tr>
<tr>
<td>B.(1,2)</td>
</tr>
<tr>
<td>0.078335</td>
</tr>
<tr>
<td>B.(2,2)</td>
</tr>
<tr>
<td>0.827922</td>
</tr>
<tr>
<td>Q.(X.Wolf,X.Wolf)</td>
</tr>
<tr>
<td>0.455214</td>
</tr>
<tr>
<td>Q.(X.Moose,X.Moose)</td>
</tr>
<tr>
<td>0.178661</td>
</tr>
<tr>
<td>x0.X.Wolf</td>
</tr>
<tr>
<td>0.002926</td>
</tr>
<tr>
<td>x0.X.Moose</td>
</tr>
<tr>
<td>-1.192645</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=1
Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

Convergence warnings
Warning: the R.(Wolf,Wolf) parameter value has not converged.
Warning: the R.(Moose,Moose) parameter value has not converged.
Warning: the x0.X.Wolf parameter value has not converged.
Warning: the logLik parameter value has not converged.
Type MARSSinfo("convergence") for more info on this warning.

It looks like $R$ is going to zero, meaning that the maximum-likelihood model is a process error only model. That is not too surprising given that the data look more like a random walk than white noise. We will set $R$ manually to zero and assume that the census is complete (they count all individuals):

```r
royale.model.2=list(Z="identity", B="unconstrained",
Q="diagonal and unequal", R="zero", U="zero")
kem.2=MARSS(z.royale.dat, model=royale.model.2)
```

Success! abstol and log-log tests passed at 16 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 16 iterations.
Log-likelihood: -81.53121
AIC: 179.0624 AICc: 180.5782

<table>
<thead>
<tr>
<th>Estimate</th>
</tr>
</thead>
<tbody>
<tr>
<td>B.(1,1)</td>
</tr>
<tr>
<td>B.(2,1)</td>
</tr>
<tr>
<td>B.(1,2)</td>
</tr>
<tr>
<td>B.(2,2)</td>
</tr>
<tr>
<td>Q.(X.Wolf,X.Wolf)</td>
</tr>
<tr>
<td>Q.(X.Moose,X.Moose)</td>
</tr>
<tr>
<td>x0.X.Wolf</td>
</tr>
<tr>
<td>x0.X.Moose</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.
14.2.3 Look at the estimated interactions

The estimated $B$ elements are $\text{coef(kem.2)}$.$B$.

```r
data <- data.frame(  
  wolf.B=coef(kem.2, type="matrix")$B,  
  rownames(wolf.B)=colnames(wolf.B)=rownames(royale.dat)  
)

print(wolf.B, digits=2)
```

Wolf Moose  
Wolf 0.77 0.078  
Moose -0.18 0.828

The `coef()` function returns the estimated parameters of the fitted object, but in this case we want to see the estimates in matrix form. Thus we use `type="matrix"`. Element row=$i$, col=$j$ in $B$ is the effect of species $j$ on species $i$, so $B_{2,1}$ is the effect of wolves on moose and $B_{1,2}$ is the effect of moose on wolves. The $B$ matrix suggests that wolves have a negative effect on moose and that moose have a positive effect on wolves—as one would expect. The diagonals are interpreted differently than the off-diagonals since the diagonals are $(b_{ii} - 1)$ so subtract off 1 from the diagonals to get the effect of species $i$ on itself. If the species are density-independent, then $B_{i,i}$ would equal 1. Smaller $B_{i,i}$ means more density dependence.

14.2.4 Adding covariates

It is well-known that moose numbers are strongly affected by winter and summer climate. The Isle Royale data set provided with MARSS has climate data from climate stations in Northeastern Minnesota, near Isle Royale\(^3\). The covariate data include January-February, July-September and April-May average temperature and precipitation. Also included are three-year running means of these data, where the number for year $x$ is the average of years $x-1$, $x$ and $x+1$. We will include these covariates in the analysis to see how they change our interaction estimates. We have to adjust our covariates because the census numbers are from winter in year $x$ and we want the climate data from the previous year to affect this winter’s moose count. As usual, we will need to demean our covariate data so that we can set $u$ equal to zero. We will standardize the variance also so that we can more easily compare the effects across different covariates.

The mathematical form of our new process model for the wolf-moose population dynamics is

$$
\begin{bmatrix}
  x_w \\
  x_m \\
\end{bmatrix}_t = 
B \begin{bmatrix}
  x_w \\
  x_m \\
\end{bmatrix}_{t-1} 
+ \begin{bmatrix}
  0 & 0 & 0 \\
  C_{21} & C_{22} & C_{23} \\
\end{bmatrix} \begin{bmatrix}
  \text{win temp} \\
  \text{win precip} \\
  \text{sum temp} \\
\end{bmatrix}_{t-1} 
+ \begin{bmatrix}
  w_w \\
  w_m \\
\end{bmatrix}_t
$$

\(^3\) From the Western Regional Climate Center. See the help file for this dataset for references (?isleRoyal).
The $C_{21}$, $C_{22}$, etc. terms are the effect of winter temperature, winter precipitation, previous summer temperature and previous summer precipitation on winter moose numbers. Since climate is known to mainly affect the moose, we set the climate effects to 0 for wolves (top row of $C$).

First we prepare the covariate data and select the winter temperature and precipitation data and the summer temperature data. We need to use the previous year’s climate data with this winter’s abundance data, so 1959 to 2010.

```r
clim.variables = c("jan.feb.ave.temp","jan.feb.ave.precip","july.sept.ave.temp")
yr1959to2010 = isleRoyal[,"Year"]>=1959 & isleRoyal[,"Year"]<=2010
clim.dat= t(isleRoyal[yr1959to2010, clim.variables])
z.score.clim.dat=zscore(clim.dat)
```

A plot of the covariate data against each other indicates that there is not much correlation between winter temperature and precipitation (Figure 14.2, which is good for analysis purposes, but warm winters are somewhat correlated with warm summers. The latter will make it harder to interpret the effect of winter versus summer temperature although the correlation is not too strong fortunately.

Next we prepare the list with the structure of all the model matrices. We give descriptive names to the $C$ elements so we can remember what each $C$ element means.

```r
royale.model.3=list(Z="identity", B="unconstrained",
Q="diagonal and unequal", R="zero", U="zero",
C=matrix(list(0,"Moose win temp",0,"Moose win precip",
0,"Moose sum temp"),2,3),
c=z.score.clim.dat)
```

Then we fit the model with covariates.

```r
kem.3=MARSS(z.royale.dat, model=royale.model.3)
```

Success! abstol and log-log tests passed at 16 iterations. Alert: conv.test.slope.tol is 0.5. Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 16 iterations.
Log-likelihood: -80.65261
AIC: 183.3052  AICc: 186.1748

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>B.(1,1)</td>
<td>0.7670</td>
</tr>
<tr>
<td>B.(2,1)</td>
<td>-0.1638</td>
</tr>
</tbody>
</table>
Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

The results suggest what is already known about this system: cold winters and heavy snow are bad for moose as are hot summers.

Fig. 14.2. Pairs plot of the covariate data for Isle Royale with correlations in the lower panel. The R code that produced this plot was cor.fun=function(x, y)text(0.5,0.5,format(cor(x,y),digits=2),cex=2) pairs(t(z.score.clim.dat),lower.panel=cor.fun).
14.3 Some settings to improve performance when estimating \( B \)

14.2.5 Change the model and data

You can explore the sensitivity of the \( B \) estimates when the measurement error is increased by adding white noise to the data:

\[
\text{bad.data} = \text{z.royale.dat} + \text{matrix(rnorm(100,0,sqrt(.2)),2,50)}
\]

\[
\text{kem.bad} = \text{MARSS(bad.data, model=model)}
\]

You can change the model by changing the constraints on \( R \) and \( Q \).

14.3 Some settings to improve performance when estimating \( B \)

In the default MARSS model, the value of \( E[x_0|y_0] \) (what \( x_0 \) denotes when \( tinitx=0 \) in the model list) is estimated. If we are estimating the \( B \) matrix, it is better to set \( tinitx=1 \) so that we are estimating \( E[x_1|y_0] \) instead\(^4\). The model will fit either way, but setting \( tinitx=1 \) in the model list will speed up and stabilize the fitting. It doesn’t make much of a difference for the wolf-moose dataset but can have a large effect for larger models. The reason is that the likelihood surface for \( E[x_1|y_0] \) is better behaved when \( B \) is small. For example, if \( B \) equal to 0, there is little information about \( E[x_0|y_0] \) so the algorithm goes in circles trying to estimate it while there is good information \( E[x_1|y_0] \) from \( y_1 \). We could use a prior on the initial \( x \) but this requires its variance-covariance structure, which depends on the unknown \( B \) and specifying a variance-covariance structure that conflicts with \( B \) will change your \( B \) estimates.

For the wolf-moose model, we set \( R = 0 \). The EM algorithm (default) cannot estimate \( x_0 \) when \( tinitx=1 \) therefore to fit the model with \( tinitx=1 \) we need to use \text{method}="BFGS". This is only for the case when \( R = 0 \)\(^5\).

\[
\text{royale.model.4} = \text{list(B="unconstrained", U="zero", Q="diagonal and unequal", Z="identity", R="zero", tinitx=1)}
\]

\[
\text{kem.4} = \text{MARSS(z.royale.dat, model=royale.model.4)}
\]

The other setting we may want to change is \text{allow.degen} in the control list. This sets the diagonals of \( Q \) or \( R \) to zero if they are heading towards zero. When the initial \( x \) is at \( t = 1 \), this can have non-intuitive (not wrong but puzzling; see Appendix A) consequences if \( R \) is going to zero. So, we will set \text{control$allow.degen=}F\text{ALSE} \text{and manually set } R \text{ to 0 if needed.}

\(^4\) If there are many missing values at \( t = 1 \), we might still have problems and have to adjust accordingly.

\(^5\) because the update for \( (x_0^0)_{j+1} \) is \( x_1^T \) but when \( R = 0 \) and \( V_0^0 = 0 \), \( x_1^T \) will equal \( (x_0^0)_{j} \), i.e. whatever value you started with. Thus the estimate of \( x_0^0 \) never changes.
14.4 Analysis a four-species plankton community

Ives et al. (2003) presented weekly data on the biomass of two species of phytoplankton and two species of zooplankton in two lakes, one with low planktivory and one with high planktivory. They used these data to estimate the interaction terms for the four species. Here we will reanalyze data and compare our results.

Ives et al. (2003) explain the data as: “The data consist of weekly samples of zooplankton and phytoplankton, which for the analyses were divided into two zooplankton groups (Daphnia and non-Daphnia) and two phytoplankton groups (large and small phytoplankton). Daphnia are large, effective herbivores, and small phytoplankton are particularly vulnerable to herbivory, so we anticipate strong interactions between Daphnia and small phytoplankton groups.” Figure 14.3 shows the data. What you can see from the figure is that the data are only collected in the summer.

14.4.1 Load in the plankton data

```r
# only use the plankton, daphnia, & non-daphnia
plank.spp = c("Large Phyto","Small Phyto","Daphnia","Non-daphnia")
plank.dat = ivesDataByWeek[,plank.spp]
#The data are not logged
plank.dat = log(plank.dat)
#Transpose to get time going across the columns
plank.dat = t(plank.dat)
#make a demeaned version
d.plank.dat = (plank.dat-apply(plank.dat,1,mean,na.rm=TRUE))
```

We will demean the data so we can set u to 0. We do not standardize by the variance, however because we are going to fix the R variance later as Ives et al. did.

14.4.2 Specify a MARSS model for the plankton data

We will start by fitting a model with the following assumptions:

- All phytoplankton share the same process variance.
- All zooplankton share the same process variance.
- Phytoplankton and zooplankton have different measurement variances
- Measurement errors are independent.
- Process errors are independent.

```r
Q=matrix(list(0),4,4); diag(Q)=c("Phyto","Phyto","Zoo","Zoo")
R=matrix(list(0),4,4); diag(R)=c("Phyto","Phyto","Zoo","Zoo")
plank.model.0=list(
    B="unconstrained", U="zero", Q=Q,
```
14.4 Analysis a four-species plankton community

Fig. 14.3. Plot of the de-meaned plankton data. Zooplankton are the thicker lines. Phytoplankton are the thinner lines.

\[ Z=\text{"identity"}, \ A=\text{"zero"}, \ R=R, \ x0=\text{"unequal"}, \ tinitx=1 \]

Why did we set \(U=\text{"zero"}\)? Equation [14.1] is a stationary model; it fluctuates about a mean. The \(u\) in Equation [14.1] is a scaling term that just affects the mean level—once the system is at equilibrium. If we assume that the mean of \(y\) (the mean of our data) is a good estimate of the mean of the system (the \(x\)), then we can set \(u\) equal to zero (and \(a\)). The initial states (\(x\)) are set at \(t = 1\) instead of \(t = 0\), which improves estimation for large systems.

14.4.3 Fit the plankton model and look at the estimated \(B\) matrix

The call to fit the model is:

\( \text{kem.plank.0 = MARSS(d.plank.dat, model=plank.model.0) } \)

Now we can print the \(B\) matrix, with a little clean up.

\( \text{#Cleaning up the B matrix for printing} \)

\( \text{B.0 = coef(kem.plank.0, type=\"matrix\")$B[1:4,1:4] } \)

\( \text{rownames(B.0) = colnames(B.0) = c("LP","SP","D","ND")} \)

\( \text{print(B.0,digits=2)} \)

<table>
<thead>
<tr>
<th></th>
<th>LP</th>
<th>SP</th>
<th>D</th>
<th>ND</th>
</tr>
</thead>
<tbody>
<tr>
<td>LP</td>
<td>0.77</td>
<td>0.29</td>
<td>-0.0182</td>
<td>0.131</td>
</tr>
<tr>
<td>SP</td>
<td>0.19</td>
<td>0.51</td>
<td>0.0052</td>
<td>-0.045</td>
</tr>
<tr>
<td>D</td>
<td>-0.43</td>
<td>2.29</td>
<td>0.4916</td>
<td>0.389</td>
</tr>
<tr>
<td>ND</td>
<td>-0.33</td>
<td>1.35</td>
<td>-0.2180</td>
<td>0.831</td>
</tr>
</tbody>
</table>
LP stands for large phytoplankton, SP for small phytoplankton, D for Daphnia and ND for non-Daphnia.

We can compare this to the Ives et al. estimates (in their Table 2, bottom right) and see quite a few differences:

<table>
<thead>
<tr>
<th></th>
<th>LP</th>
<th>SP</th>
<th>D</th>
<th>ND</th>
</tr>
</thead>
<tbody>
<tr>
<td>LP</td>
<td>0.48</td>
<td>-0.39</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>SP</td>
<td>--</td>
<td>0.25</td>
<td>-0.17</td>
<td>-0.11</td>
</tr>
<tr>
<td>D</td>
<td>--</td>
<td>--</td>
<td>0.74</td>
<td>0.00</td>
</tr>
<tr>
<td>ND</td>
<td>--</td>
<td>0.10</td>
<td>0.00</td>
<td>0.60</td>
</tr>
</tbody>
</table>

First, thing you will notice is that the Ives et al. matrix is missing values. The matrix they show is after a model selection step to determine which interactions had little data support and thus could be set to zero. Also, they fixed apriori the interactions between Daphnia and non-Daphnia at zero because they do not prey on each other. The second thing you will notice is that the estimates are not particularly similar. Next we will try some other ways of fitting the data that are closer to the way that Ives et al. fitted the data.

By the way, if you are curious what would happen if we removed all those NAs, you can run the following code.

```r
test.dat = d.plank.dat[, !is.na(d.plank.dat[1, ])]
test = MARSS(test.dat, model = plank.model.0)
```

Removing all the NAs would mean that the end of summer 1 is connected to the beginning of summer 2. This adds some steep steps in the Daphnia time series where Daphnia ended the summer high and started the next summer low.

### 14.4.4 Look at different ways to fit the model

We will try a series of changes to get closer to the way Ives et al. fit the data, and you will see how different assumptions change (or do not change) our species interaction estimates.

First, we change $Q$ to be unconstrained. Making $Q$ diagonal in model 0 meant that we were assuming that whatever environmental factor is driving variation in phytoplankton numbers is uncorrelated with the environmental factor driving zooplankton variation. That is probably not true since they are all in the same lake. This case takes awhile to run.

```r
plank.model.1 = MARSS(d.plank.dat, model = plank.model.0)

plank.model.1$Q = "unconstrained"
kem.plank.1 = MARSS(d.plank.dat, model = plank.model.1)
```

Notice that the $Q$ specification changed to "unconstrained". Everything else stays the same as in model 0. The code now runs longer, and the $B$ estimates are not particularly closer to Ives et al.
Next, we will set some of the interactions to zero as in Table 2 in Ives et al. (2003). In their table, certain interactions were fixed at 0 (denoted with 0s), and some were made 0 after fitting (the blanks). We will fix all to zero. To do this, we need to write out the $B$ matrix as a list matrix so that we can have estimated and fixed values (the 0s) in the $B$ specification.

$$B.2 = \text{matrix(list(0),4,4)} \quad \# \text{set up the list matrix}$$
$$\text{diag(B.2)=c("B11","B22","B33","B44")} \quad \# \text{give names to diagonals}$$
$$\text{and names to the estimated non-diagonals}$$
$$\text{print(B.2)}$$

As you can see, the $B$ matrix now has elements that will be estimated (the names in quotes) and fixed values (the numbers with no quotes). When preparing your list matrix, make sure your fixed values do not have quotes around them. If they do, they are strings (class character) not numbers (class numeric), and MARSS will interpret a string as the name of something to be estimated. If you use the same name for an element, then MARSS will force those elements to be shared (have the same value).

#model 2
$$\text{plank.model.2=plank.model.1}$$
$$\text{plank.model.2$B = B.2}$$
$$\text{kem.plank.2= MARSS(d.plank.dat, model=plank.model.2)}$$

Now we are getting closer to the Ives et al. estimates:
Ives et al. did not estimate $R$. Instead they used a fixed observation variance of 0.04 for phytoplankton and 0.16 for zooplankton. We fit the model with their fixed $R$ as follows:

```r
#model 3
plank.model.3 = plank.model.2
plank.model.3$R = diag(c(.04,.04,.16,.16))
kem.plank.3 = MARSS(d.plank.dat, model=plank.model.3)
```

As you can see from Table 14.1, we are getting closer to the Ives et al. estimates, but we are still a bit off. Now we need to add the environmental covariates: phosphorous and fish biomass.

### 14.4.5 Adding covariates

A standard way that you will see covariate data added to a MARSS model is the following:

$$
\begin{align*}
\mathbf{x}_t &= \mathbf{Bx}_{t-1} + \mathbf{u} + \mathbf{Cc}_t + \mathbf{w}_t, \text{ where } \mathbf{w}_t \sim \text{MVN}(\mathbf{0}, \mathbf{Q}) \\
\mathbf{y}_t &= \mathbf{Zx}_t + \mathbf{a} + \mathbf{Dd}_t + \mathbf{v}_t, \text{ where } \mathbf{v}_t \sim \text{MVN}(\mathbf{0}, \mathbf{R})
\end{align*}
$$

$\mathbf{c}_t$ and $\mathbf{d}_t$ are covariate data, like temperature. At time $t$ and $\mathbf{C}$ is a matrix with the (linear) effects of $\mathbf{c}_t$ on $\mathbf{x}_t$, and $\mathbf{D}$ is a matrix with the (linear) effects of $\mathbf{d}_t$ on $\mathbf{y}_t$.

[14.5] Ives et al. (2003) only include covariates in their process model, and their process model (their Equation 27) is written $\mathbf{X}_t = \mathbf{A} + \mathbf{BX}_{t-1} + \mathbf{CU}_t + \mathbf{E}_t$. In our Equation 14.5, $\mathbf{U}_t = \mathbf{c}_t$, and $\mathbf{C}$ is a $m \times p$ matrix, where $p$ is the number of covariates in $\mathbf{c}_t$. We will set their $\mathbf{A}$ (our $\mathbf{u}$) to zero by demeaning the $\mathbf{y}$ and implicitly assuming that the mean of the $\mathbf{y}$ is a good estimate of the mean of the $\mathbf{x}$’s. Thus the model where covariates only affect the underlying process is

$$
\begin{align*}
\mathbf{x}_t &= \mathbf{Bx}_{t-1} + \mathbf{Cc}_t + \mathbf{w}_t, \text{ where } \mathbf{w}_t \sim \text{MVN}(\mathbf{0}, \mathbf{Q}) \\
\mathbf{y}_t &= \mathbf{x}_t + \mathbf{v}_t, \text{ where } \mathbf{v}_t \sim \text{MVN}(\mathbf{0}, \mathbf{R})
\end{align*}
$$

To fit this model, we first need to prepare the covariate data. We will just use the phosphorous data.

```r
#transpose to make time go across columns
#drop=FALSE so that R doesn't change our matrix to a vector
phos = t(log(ivesDataByWeek[, "Phosph", drop=FALSE]))
d.phos = (phos-apply(phos,1,mean,na.rm=TRUE))
```

Why log the covariate data? It is what Ives et al. did, so we follow their method. However, in general, you want to think about what relationship you want to assume between the covariates and their effects. For example, log (or

---

6 You can compare this to the estimated observation variances by looking at `coef(kem.plank.2)$R`
square-root) transformations mean that extremes have less impact relative to their untransformed value and that a small absolute change, say from 0.01 to 0.0001 in the untransformed value, can mean large difference in the effects since \(\log(0.0001) < \log(0.01)\).

Phosphorous is assumed to only affect phytoplankton so the other terms in \(C\), corresponding to the zooplankton, are set to 0. The \(C\) matrix is defined as follows:

\[
C = \begin{bmatrix}
C_{LP,phos} \\
C_{SP,phos} \\
0 \\
0 
\end{bmatrix}
\] (14.7)

To add \(C\) and \(c\) to our latest model, we add \(C\) and \(c\) to the model list used in the MARSS call:

```r
plank.model.4=plank.model.3
plank.model.4$C=matrix(list("C11","C21",0,0),4,1)
plank.model.4$c=d.phos
```

Then we fit the model as usual:

```r
kem.plank.4= MARSS(d.plank.dat, model=plank.model.4)
```

Temperature and phosphorous have and estimated positive effect on phytoplankton:

```r
[,1] LP -- SP -- D -- ND --
```

14.4.6 Including a covariate observation model

The difficulty with the standard approach to including covariates (Equation 14.5) is that it limits what kind of covariate data you can use and how you model that covariate data. You have to assume that your covariate data has no error, which is probably not true. Assuming that your covariate has no error reduces the reported uncertainty in your covariate effect because you did not include uncertainty in those values. The standard approach also does not allow missing values in your covariate data, which is why we did not include the fish covariate data in the last model. Also you cannot combine instrument time series; for example, if you have two temperature recorders with different error rates and biases. Also, what if you have one noisy temperature recorder in the first part of your time series and then you switch to a much better recorder in the second half of your time series? All these problems require pre-analysis massaging of the covariate data, leaving out noisy and gappy covariate data, and making what can feel like arbitrary choices about which covariate time
series to include. This is especially worrisome when the covariates are then incorporated into the model as if they are known without error.

Instead one can include an observation and process model for the covariates just like for the non-covariate data. Now the covariates are included in \( y \), and are modeled with their own state process(es) in \( x \). A MARSS model with a covariate observation and process model is shown below. The elements with superscript \( (v) \) are for the variates and those with superscript \( (c) \) are for the covariates. The superscripts just help us keep straight which of the state processes and parameters correspond to abundances and which correspond to the environmental covariates.

\[
\begin{bmatrix}
    x^{(v)}_t \\
    x^{(c)}_t
\end{bmatrix} = \begin{bmatrix}
    B^{(v)} & C \\
    0 & B^{(c)}
\end{bmatrix} \begin{bmatrix}
    x^{(v)}_{t-1} \\
    x^{(c)}_{t-1}
\end{bmatrix} + \begin{bmatrix}
    u^{(v)}_t \\
    u^{(c)}_t
\end{bmatrix} + w_t, \quad w_t \sim \text{MVN}\left(0, \begin{bmatrix}
    Q^{(v)} & 0 \\
    0 & Q^{(c)}
\end{bmatrix}\right)
\]

\[
\begin{bmatrix}
    y^{(v)}_t \\
    y^{(c)}_t
\end{bmatrix} = \begin{bmatrix}
    Z^{(v)} & 0 \\
    0 & Z^{(c)}
\end{bmatrix} \begin{bmatrix}
    x^{(v)}_t \\
    x^{(c)}_t
\end{bmatrix} + \begin{bmatrix}
    a^{(v)}_t \\
    a^{(c)}_t
\end{bmatrix} + v_t, \quad v_t \sim \text{MVN}\left(0, \begin{bmatrix}
    R^{(v)} & 0 \\
    0 & R^{(c)}
\end{bmatrix}\right)
\]

(14.8)

Note that when you fit your covariate and non-covariate data jointly as in Equation 14.8, your non-covariate data affect the estimates of the covariate models. When you maximize the likelihood, you do so conditioned on all the data. The likelihood that is output is the likelihood of the non-covariate and covariate data. Depending on your system, you might not want the covariate model affected by the non-covariate data. In this case, you can fit the covariate model separately:

\[
\begin{align*}
    x^{(c)}_t &= B^{(c)} x^{(c)}_{t-1} + u^{(c)}_t + w_t, \quad w_t \sim \text{MVN}(0, Q^{(c)}) \\
    y^{(c)}_t &= Z^{(c)} x^{(c)}_t + a^{(c)}_t, \quad v_t \sim \text{MVN}(0, R^{(c)})
\end{align*}
\]

(14.9)

At this point, you have another choice. Do you want the estimated covariates states, the \( x^{(c)} \), to be affected by the non-covariate data? For example, you have temperature data. You can estimates true temperature for the temperature only from the temperature data or you can decide that the non-covariate data has information about the true temperature, because the non-covariate states are affected by the true temperature. If you want the covariate states to only be affected by the covariate data, then use Equation 14.5 with \( c_t \) set to your estimates of \( x^{(c)} \) from Equation 14.9. Or if you want the non-covariate data to affect the estimates of the covariate states, use Equation 14.8 with the parameters estimated from Equation 14.9.

### 14.4.7 The MARSS model with covariates following Ives et al.

Ives et al. used Equation 14.5 for phosphorous and Equation 14.8 for fish biomass. Phosphorus was treated as observed with no error since it was experimentally manipulated and there were no missing values. Fish biomass
was treated as having observation error and was modeled as a autoregressive
process with unknown parameters as in Equation 14.8.

Their MARSS model takes the form:

\[
x_t = \mathbf{B} x_{t-1} + \mathbf{C} \mathbf{c}_t + \mathbf{w}_t, \quad \text{where } \mathbf{w}_t \sim \text{MVN}(0, \mathbf{Q})
\]

\[
y_t = x_t + \mathbf{v}_t, \quad \text{where } \mathbf{v}_t \sim \text{MVN}(0, \mathbf{R}) \tag{14.10}
\]

where \(x\) and \(y\) are redefined as

\[
\begin{bmatrix}
\text{large phyto} \\
\text{small phyto} \\
\text{Daphnia} \\
\text{Non-Daphnia zooplank} \\
\text{fish biomass}
\end{bmatrix}
\tag{14.11}
\]

The covariate fish biomass appears in \(x\) because it will be modeled, and its interaction terms (Ives et al.'s \(\mathbf{C}\) terms) appear in \(\mathbf{B}\). Phosphorous appears in the \(\mathbf{c}\) terms because it is treated as a known additive term and its interaction terms appear in \(\mathbf{C}\). Recall that we set \(u\) to 0 by demeaning the plankton data, so it does not appear above. The \(\mathbf{Z}\) matrix does not appear in front of the \(x\), since there is a one-to-one correspondence the \(x\)'s and \(y\)'s, and thus \(\mathbf{Z}\) is the identity matrix.

The \(\mathbf{B}\) matrix is

\[
\mathbf{B} = \begin{bmatrix}
\mathbf{B}^{(v)} & \mathbf{C} \\
0 & \mathbf{B}^{(c)}
\end{bmatrix} = \begin{bmatrix}
b_{LP} & b_{LP,SP} & 0 & 0 & 0 \\
0 & b_{SP} & b_{SP,D} & b_{SP,ND} & 0 \\
0 & 0 & b_{D} & 0 & c_{D,fish} \\
0 & b_{ND,SP} & 0 & b_{ND,ND} & c_{ND,fish} \\
0 & 0 & 0 & 0 & b_{fish}
\end{bmatrix} \tag{14.12}
\]

The \(\mathbf{B}\) elements have some interactions fixed at 0 as in our last model fit. The \(c\)'s are the interactions between the fish and the species. We will estimate a \(\mathbf{B}\) term for fish since Ives et al. did, but this is an odd thing to do for the fish data since these data were interpolated from two samples per season.

The \(\mathbf{Q}\) matrix is the same as that in our last model fit, with the addition of an element for the variance for the fish biomass:

\[
\mathbf{Q} = \begin{bmatrix}
\mathbf{Q}^{(v)} & 0 \\
0 & \mathbf{Q}^{(c)}
\end{bmatrix} = \begin{bmatrix}
q_{LP} & q_{LP,SP} & q_{LP,D} & q_{LP,ND} & 0 \\
q_{LP,SP} & q_{SP} & q_{SP,D} & q_{SP,ND} & 0 \\
q_{LP,D} & q_{SP,D} & q_{D} & q_{D,ND} & 0 \\
q_{LP,ND} & q_{SP,ND} & q_{ND} & q_{ND} & 0 \\
0 & 0 & 0 & 0 & q_{fish}
\end{bmatrix} \tag{14.13}
\]

Again it is odd to estimate a variance term for data interpolated from two points, but we follow Ives et al. here.

Ives et al. set the observation variance for the logged fish biomass data to 0.36 (page 320 in Ives et al. (2003)). The observation variances for the plankton data was set as in our previous model.
14.4.8 Setting the model structure for the model with fish covariate data

First we need to add the logged fish biomass to our data matrix.

```r
# transpose to make time go across columns
# drop = FALSE so that R doesn't change our matrix to a vector
fish = t(log(ivesDataByWeek[, "Fish biomass", drop = FALSE]))
d.fish = (fish - apply(fish, 1, mean, na.rm = TRUE))
# plank.dat.w.fish = rbind(plank.dat, fish)
d.plank.dat.w.fish = rbind(d.plank.dat, d.fish)
```

Next make the B matrix. Some elements are estimated and others are fixed at 0.

```r
B = matrix(list(0), 5, 5)
diag(B) = list("B11", "B12", "B22", "B33", "B44", "Bfish")
B[4, 2] = "B42";
B[1:4, 5] = list(0, 0, "C32", "C42")
print(B)

[1,] "B11" "B12" 0 0 0
[2,] 0 "B22" "B23" "B24" 0
[3,] 0 0 "B33" 0 "C32"
[4,] 0 "B42" 0 "B44" "C42"
[5,] 0 0 0 0 "Bfish"
```

Now we have a B matrix that looks like that in Equation 14.12.

We need to add an extra row to C for the fish biomass row in x:

```r
C = matrix(list("C11", "C21", 0, 0, 0), 5, 1)
```

Then we set up the R matrix.

```r
R = matrix(list(0), 5, 5)
diag(R) = list(0.04, 0.04, 0.16, 0.16, 0.36)
```

Last, we need to set up the Q matrix:

```r
Q = matrix(list(0), 5, 5);
Q[1:4, 1:4] = paste(rep(1:4, times = 4), rep(1:4, each = 4), sep = "")
Q[5, 5] = "fish"
```
14.4 Analysis a four-species plankton community

\[
Q[\text{lower.tri}(Q)] = t(Q)[\text{lower.tri}(Q)]
\]

\[
\text{print}(Q)
\]

\[
\begin{bmatrix}
[1,] & [2,] & [3,] & [4,] & [5,] \\
[1,] & "11" & "12" & "13" & "14" & 0 \\
[2,] & "12" & "22" & "23" & "24" & 0 \\
[3,] & "13" & "23" & "33" & "34" & 0 \\
[4,] & "14" & "24" & "34" & "44" & 0 \\
[5,] & 0 & 0 & 0 & 0 & "fish"
\end{bmatrix}
\]

14.4.9 Fit the model with covariates

The model is the same as the previous model with updated process parameters and updated \( \mathbf{R} \). We will pass in the updated data matrix with the fish biomass added:

\[
\text{plank.model.5} = \text{plank.model.4}
\]

\[
\text{plank.model.5}$B$=B
\]

\[
\text{plank.model.5}$C$=C
\]

\[
\text{plank.model.5}$Q$=Q
\]

\[
\text{plank.model.5}$R$=R
\]

\[
\text{kem.plank.5} = \text{MARSS}(d.plank.dat.w.fish, \text{model}=\text{plank.model.5})
\]

This is the new \( \mathbf{B} \) matrix using covariates.

\[
\begin{array}{llllll}
\text{LP} & \text{SP} & \text{D} & \text{ND} \\
\text{LP} & 0.61 & -0.465 & -- & -- \\
\text{SP} & -- & 0.333 & -0.019 & -0.048 \\
\text{D} & -- & -- & 0.896 & -- \\
\text{ND} & -- & 0.044 & -- & 0.675
\end{array}
\]

Now we are getting close to Ives et al.’s estimates. Compare model 5 in Table 14.1 to the first column.

NOTE! When you include your covariates in your state model (the \( \mathbf{x} \) part), the reported log-likelihood is for the variate plus the covariate data. If you want just the log-likelihood for the variates, then you replace the covariate data with NAs and run the Kalman filter with your estimated model:

\[
\text{tmp} = \text{kem.plank.5}
\]

\[
\text{tmp}$\text{marss}$data[5,] = \text{NA}
\]

\[
\text{LL.variates} = \text{MARSSkf}(\text{tmp})\$\text{logLik}
\]

\text{MARSSkf} is the Kalman filter function and it needs a fitted model as output by a \text{MARSS} call. We set up a temporary fitted model, \text{tmp}, equal to our fitted model and then set the covariate data in that to NAs. Note we need to do this for the marssMODEL object used by \text{MARSSkf}, which will be in \text{$\text{marss}}. We then pass that temporary fitted model to \text{MARSSkf} to get the log-likelihood of just the variates.
Table 14.1. The parameter estimates under the different plankton models. Models 0 to 3 do not include covariates, so the C elements are blank. Bij is the effect of species i on species j. 1=large phytoplankton, 2=small phytoplankton, 3=Daphnia, 4=non-Daphnia zooplankton. The Ives et al. (2003) estimates are from their table 2 for the low planktivory lake with the observation model.

<table>
<thead>
<tr>
<th></th>
<th>Ives et al.</th>
<th>Model 0</th>
<th>Model 1</th>
<th>Model 2</th>
<th>Model 3</th>
<th>Model 4</th>
<th>Model 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>B11</td>
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<td>0.77</td>
<td>0.50</td>
<td>0.65</td>
<td>0.62</td>
<td>0.61</td>
<td>0.61</td>
</tr>
<tr>
<td>B22</td>
<td>0.25</td>
<td>0.51</td>
<td>0.90</td>
<td>0.54</td>
<td>0.51</td>
<td>0.33</td>
<td>0.33</td>
</tr>
<tr>
<td>B33</td>
<td>0.74</td>
<td>0.49</td>
<td>0.64</td>
<td>0.83</td>
<td>0.89</td>
<td>0.89</td>
<td>0.90</td>
</tr>
<tr>
<td>B44</td>
<td>0.60</td>
<td>0.83</td>
<td>0.81</td>
<td>0.60</td>
<td>0.67</td>
<td>0.66</td>
<td>0.67</td>
</tr>
<tr>
<td>B12</td>
<td>-0.39</td>
<td>0.29</td>
<td>0.06</td>
<td>-0.33</td>
<td>-0.32</td>
<td>-0.46</td>
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</tr>
<tr>
<td>B23</td>
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<td>0.01</td>
<td>0.07</td>
<td>0.00</td>
<td>-0.02</td>
<td>-0.02</td>
<td>-0.02</td>
</tr>
<tr>
<td>B24</td>
<td>-0.11</td>
<td>-0.04</td>
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<td>0.02</td>
<td>-0.05</td>
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</tr>
<tr>
<td>B42</td>
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<td>0.37</td>
<td>0.13</td>
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<td>0.05</td>
<td>0.04</td>
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<td></td>
<td></td>
<td></td>
<td>0.14</td>
<td>0.14</td>
<td></td>
</tr>
<tr>
<td>C21</td>
<td>0.25</td>
<td></td>
<td></td>
<td></td>
<td>0.16</td>
<td>0.16</td>
<td></td>
</tr>
<tr>
<td>C32</td>
<td>-0.14</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>-0.04</td>
<td></td>
</tr>
<tr>
<td>C42</td>
<td>-0.04</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>-0.01</td>
</tr>
</tbody>
</table>

14.4.10 Discussion

The estimates for our last model are fairly close to the Ives et al. estimates, but still a bit different. There are two big difference between our last model and the Ives et al. analysis. Ives et al. had data from three lakes and the estimate of $Q$ used the data from all lakes.

Combining data, whether it be from different areas or years, can be done in a MARSS model as follows. Let $y_1$ be the first data set (say from site 1) and $y_2$ be the second data set (say from site 2). Then a MARSS model with shared parameters values across datasets would be

$$x_i^+ = B^+ x_{i-1}^+ + u^+ w_r, \text{ where } w_r \sim \text{MVN}(0, Q^+)$$

$$y_i^+ = Z^+ x_i^+ + a^+ + v_r, \text{ where } v_r \sim \text{MVN}(0, R^+)$$

where the + matrices are stacked matrices from the different sites (1 and 2):

$$[x_{1,t}^+, x_{2,t}^+] = \begin{bmatrix} B & 0 \\ 0 & B \end{bmatrix} [x_{1,t-1}^+, x_{2,t-1}^+] + \begin{bmatrix} u^+ \\ u^+ \end{bmatrix} + w_r, w_r \sim \text{MVN}\left(0, \begin{bmatrix} Q & q \\ q & Q \end{bmatrix} \right)$$

$$[y_{1,t}^+, y_{2,t}^+] = \begin{bmatrix} Z & 0 \\ 0 & Z \end{bmatrix} [x_{1,t}^+, x_{2,t}^+] + \begin{bmatrix} a^+ \\ a^+ \end{bmatrix} + v_r, v_r \sim \text{MVN}\left(0, \begin{bmatrix} R & 0 \\ 0 & R \end{bmatrix} \right)$$

The $q$ in the process variance allows that the environmental variability might be correlated between datasets, i.e. if they are replicate plots that are nearby, say. If you did not want all the parameters shared, then you replace the $B$ in $B^+$ with $B_1$ and $B_2$, say.
The second big difference is that Ives et al. did not demean their data, but estimated \( u \). We could have done that too, but with all the NAs in the data (during winter), estimating \( u \) is not robust and takes a long time. You can try the analysis on the data that has not been demeaned and set \( U="unequal" \). The results are not particularly different, but it takes a long, long,...long time to converge.

You can also try using the actual fish data instead of the interpolated data. Fish biomass was estimated at the end and start of the season, so only the values at the start and finish of strings of fish numbers are the real data. The others are interpolated. You can fill in those interpolated values with NAs (missing values) and rerun. The results are not appreciably different, but the effect of fish drops a bit as you might expect when you have less fish information. You don’t see it here, but your estimated confidence in the fish effects would also drop since this estimate is based on less fish data.

14.5 Stability metrics from estimated interaction matrices

The previous sections focused on estimation of the \( B \) and \( C \) matrices. The estimated \( B \) matrix gives a picture of the species interactions, but it also can be used to compute metrics of the intrinsic community stability (Ives et al., 2003). Here we illustrate how to compute these metrics; the reader should see Ives et al. (2003) for details on the meaning of each.

For the examples here, we will use the estimated \( B \) and \( Q \) matrices from our model 5:

\[
B = \text{coef(kem.plank.5,type="matrix")$B[1:4,1:4]}
\]
\[
Q = \text{coef(kem.plank.5,type="matrix")$Q[1:4,1:4]}
\]

14.5.1 Return rate metrics

Return rate metrics measure how rapidly the system returns to the stationary distribution of species abundances after it is perturbed away from the stationary distribution. With a deterministic (\( Q = 0 \)) MARSS community model, the equilibrium is a point or stable limit cycle. In a stochastic model (\( Q \neq 0 \)), the equilibrium is stochastic and is a stationary distribution. Rate of return to the stochastic equilibrium is the rate at which the distribution converges to the stationary distribution after a perturbation away from this stationary distribution. The more rapid the convergence, the more stable the system.

The rate of return of the mean of the stationary distribution is governed by the dominant eigenvalue of \( B \). In R, we can compute this as:

\[
\text{max(eigen(B)$values)}
\]

[1] 0.8964988
The rate of return of the variance of the stationary distribution is governed by the dominant eigenvalue of $B \otimes B$:

$$\max(eigen(kronecker(B,B))\$values)$$

[1] 0.8037101

### 14.5.2 Variance metrics

These metrics measure the variance of the stationary distribution of species abundances (with variance due to environmental drivers removed) relative to the process error variance. The system is considered more stable when the stationary distribution variance is low relative to the process error variance.

To compute variance metrics, we need to first compute the variance-covariance matrix for the stationary distribution, $V_\infty$:

```r
m <- nrow(B)
vecV <- solve(diag(m*m)-kronecker(B,B))%*%as.vector(Q)
V_inf <- matrix(vecV,nrow=m,ncol=m)
```

A measure of the proportion of the “volume” of the stationary distribution due to species interactions is given by the square of the determinant of the $B$ matrix (Eqn. 24 in Ives et al. (2003)):

$$\text{abs(det}(B)\text{)^2}$$

[1] 0.01559078

To compare stability across systems of different sizes, you scale by the number of species:

$$\text{abs(det}(B)\text{)^2}/(\text{nrow}(B))$$

[1] 0.3533596

### 14.5.3 Reactivity metrics

Reactivity measure how the system responds to a perturbation. A highly reactive system tends to move farther away from a stable equilibrium immediately after a perturbation, even though the system will eventually return to the equilibrium. High reactivity occurs when species interactions greatly amplify the environmental variance to produce a stationary distribution with high variances in the abundance of individual species.

Both metrics of reactivity of estimates of the average expected change in distance from the mean of the stationary distribution. The first uses estimates of $Q$ and $V_\infty$.

$$-\text{sum(diag}(Q)/\text{sum(diag}(V_inf))$$

[1] -0.346845
Estimation of $Q$ is prone to high uncertainty. Another metric that uses only $B$ is the worst-case reactivity. This is given by

$$\max(eigen(t(B)^%*%B)$values)-1$$

[1] -0.1957795

14.6 Further information

MAR modeling and models have been used estimate species interaction strengths, stability metrics, and environmental drivers for a variety of freshwater plankton systems ([Ives 1995], [Ives et al. 1999, 2003], [Hampton et al. 2008, 2006], [Hampton and Schindler 2006], [Klug and Cottingham 2001]). They have been used to gain much insight into the dynamics of ecological communities and how environmental drivers affect the system. See [Hampton et al. 2013] for a review of the literature using MAR models to understand plankton dynamics.
Combining data from multiple time series

15.1 Overview

In this section, we consider the case where multiple time series exist and we want to use all the datasets to estimate a common underlying state-process or common underlying parameters. In ecological applications, this situation arises when 1) there are time series of observations from the same species as the original time series (e.g. aerial and land based surveys of the same species) or 2) there are time series collected in the same survey, but represent observations of multiple species (e.g. we might be doing a fisheries trawl survey that collects multiple species in each trawl). Why should we consider using other time series? In the first scenario, where methodology differs between time series of the same species, observation error may be survey-specific. These time series may represent observations of multiple populations, or these may represent multiple observations of the same population. In the second scenario, each species should be treated as a separate process (given its own state vector), but because the survey methodology is the same across species, it might be reasonable to assume a shared observation error variance. If these species have a similar response to environmental stochasticity, it might be possible to also assume a shared process variance.

In both of the above examples, MARSS models offer a way to link multiple time series. If parameters are allowed to be shared among the state processes (trend parameters, process variances) or observation processes (observation variances), parameter estimates will be more precise than if we treated each time series as independent. By improving estimates of variance parameters,
we will also be better able to discriminate between process and observation error variances.

In this chapter, we will show examples of using MARSS models to analyze multiple time series on the same species but either different populations or different survey methods. The multivariate first-order autoregressive state process is written as usual as:

\[ x_t = Bx_{t-1} + u + w_t \quad \text{where} \quad w_t \sim \text{MVN}(0, Q) \] (15.1)

The true population sizes at time \( t \) are represented by the state \( x_t \), whose dimensions are equal to the number of state processes \( m \). The \( m \times m \) matrix \( B \) allows interaction between processes (density dependence and competition, for instance), \( u \) is a vector describing the mean trend, and the correlation of the process deviations is determined by the structure of the matrix \( Q \).

The multivariate observation error model is expressed as,

\[ y_t = Zx_t + a + v_t \quad \text{where} \quad v_t \sim \text{MVN}(0, R) \] (15.2)

where \( y_t \) is a vector of observations at time \( t \), \( Z \) is a design matrix of 0s and 1s, \( a \) is a vector of bias adjustments, and the correlation structure of observation matrices is specified with the matrix \( R \). Including \( Z \) and \( a \) will not be required for every model, but is useful when some processes are observed multiple times.

### 15.2 Salmon spawner surveys

In our first application combining multiple time series, we will analyze a dataset on Chinook salmon \((Oncorhynchus tshawytscha)\). This data comes from the Okanogan River in Washington state, a major tributary of the Columbia River (with headwaters in British Columbia). As an index of the abundance of spawning adults, biologists have conducted redd surveys during summer months (redds are nests or collection of rocks on stream bottoms where females deposit eggs). Our data are aerial surveys of redds on the Okanogan River conducted 1956-2008 and ground surveys of redds from 1990-2008.

#### 15.2.1 Read in and plot the raw data

We will be using the logged counts.

```r
head(okanaganRedds)
```

```
Year aerial ground
[1,] 1956  37   NA
[2,] 1957  53   NA
[3,] 1958  94   NA
```
Year is in the first column, and the counts (in normal space) are in columns 2:3. Missing observations are represented by NAs.

```r
logRedds = log(t(okanaganRedds)[2:3,])
```

Fig. 15.1. The two time series look to be pretty close to one another in the years where there is overlap.
15.2.2 Test hypotheses about whether the data can be combined

Do these surveys represent observations of the same underlying process? We can evaluate data support for this question by testing a few relatively simple models. Using the logged data, we will start with a simple model that assumes the underlying population process is univariate (there is one underlying population trajectory) and each survey is an independent observation of this population process. Mathematically, the model is:

\[
x_t = x_{t-1} + u + w_t, \text{ where } w_t \sim N(0, q)
\]

\[
\begin{bmatrix}
y_{aer} \\
y_{gnd}
\end{bmatrix}_t = \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} x_t \\ x_{t-1} \end{bmatrix} + \begin{bmatrix} u \\ v_{aer} \\ v_{gnd} \end{bmatrix}_t, \text{ where } v_t \sim \text{MVN}\left(0, \begin{bmatrix} r & 0 \\ 0 & r \end{bmatrix}\right)
\] (15.3)

The \(a\) structure means that the \(a\) for one of the \(y\)'s is fixed at 0 and the other \(a\) is estimated relative to that fixed \(a\). In MARSS, this is the “scaling” structure for \(a\). We specify this model as follows. Since \(x\) is univariate, \(Q\) and \(u\) are just scalars (single numbers), and we can leave them off in our specification.

Fit the single state model, where the time series are assumed to be from the same population.

```r
model1 = list()
model1$R = "diagonal and equal"
model1$Z = matrix(1, 2, 1)
model1$A = "scaling"
kem1 = MARSS(logRedds, model=model1)
```

The AIC and AICc values for this model are

We can modify the above model to let the observation error variances to be unique:

```r
model2 = model1 # model2 is based on model1
model2$R = "diagonal and unequal"
model2$Z = matrix(1, 2, 1)
model2$A = "scaling"
kem2 = MARSS(logRedds, model=model2)
```

It is possible that these surveys are measuring different population processes. For our third model, we will fit a model with two different population process with the same process parameters. For simplicity, we will keep the trend and variance parameters the same. Mathematically, the model we are fitting is:

\[
\begin{bmatrix}
x_1 \\
x_2
\end{bmatrix}_t = \begin{bmatrix} x_1 \\ x_{2, t-1} \end{bmatrix} + \begin{bmatrix} u \\ u \end{bmatrix}_t + \begin{bmatrix} w_t \end{bmatrix}, \text{ where } w_t \sim \text{MVN}(0, \begin{bmatrix} q & 0 \\ 0 & q \end{bmatrix})
\]

\[
\begin{bmatrix}
y_{aer} \\
y_{gnd}
\end{bmatrix}_t = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix}_t + \begin{bmatrix} 0 \\ 0 \end{bmatrix} + \begin{bmatrix} v_{aer} \\ v_{gnd} \end{bmatrix}_t, \text{ where } v_t \sim \text{MVN}\left(0, \begin{bmatrix} r & 0 \\ 0 & r \end{bmatrix}\right)
\] (15.4)

We specify this in MARSS as

```r
model3 = list()
model3$R = "diagonal and equal"
model3$Z = matrix(1, 2, 1)
model3$A = "scaling"
kem3 = MARSS(logRedds, model=model3)
```
model3=list()
model3$Q="diagonal and equal"
model3$R="diagonal and equal"
model3$U="equal"
model3$Z="identity"
model3$A="zero"
kem3 = MARSS(logRedds, model=model3)

Based on AICc, it appears that the best model is also the simplest one, with one state vector (model1).

c(mod1=kem1$AICc, mod2=kem2$AICc, mod3=kem3$AICc)

mod1 mod2 mod3
133.9804 136.2164 174.1392

This suggests that the two different surveys are not only measuring the same underlying process, but have the same observation error variance. Finally, we will make a plot of the model-predicted states (with +/- 2 s.e.s) and the log-transformed data (Figure 15.2).

15.3 American kestrel abundance indices

In this example, we evaluate uncertainty in the structure of process variability (environmental stochasticity) using breeding bird surveys data. In this analysis, we use three time series of American kestrel (*Falco sparverius*) abundance from adjacent Canadian provinces along a longitudinal gradient (British Columbia, Alberta, Saskatchewan). The data were collected annually, and corrected for changes in observer coverage and detectability.

15.3.1 Read in and look at the data

Figure 15.3 shows the data. The data are already log transformed.

```r
cirdat = t(kestrel[,2:4])
head(kestrel)
```

<table>
<thead>
<tr>
<th>Year</th>
<th>British.Columbia</th>
<th>Alberta</th>
<th>Saskatchewan</th>
</tr>
</thead>
<tbody>
<tr>
<td>[1,]</td>
<td>1969</td>
<td>0.754</td>
<td>0.460</td>
</tr>
<tr>
<td>[2,]</td>
<td>1970</td>
<td>0.673</td>
<td>0.899</td>
</tr>
<tr>
<td>[3,]</td>
<td>1971</td>
<td>0.734</td>
<td>1.133</td>
</tr>
<tr>
<td>[4,]</td>
<td>1972</td>
<td>0.589</td>
<td>0.528</td>
</tr>
<tr>
<td>[5,]</td>
<td>1973</td>
<td>1.405</td>
<td>0.789</td>
</tr>
<tr>
<td>[6,]</td>
<td>1974</td>
<td>0.624</td>
<td>0.528</td>
</tr>
</tbody>
</table>
Fig. 15.2. The data support the hypothesis that the two redd-count time series are observations of the same population. The points are the data and the thick black line is the estimated underlying state.

We know that the surveys use the same design, so we will force observation error to be shared. Our uncertainty lies in whether these time series are sampling the same population, and how environmental stochasticity varies by subpopulation (if there are subpopulations). Our first model has one population trajectory (meaning there is one panmictic BC/AB/SK population) and each of these three surveys is an observation of this single population with equal observation variances. Mathematically, the model is:

\[
x_t = x_{t-1} + u + w_t, \text{ where } w_t \sim N(0, q)
\]

\[
\begin{bmatrix}
y_{BC} \\
v_{AB} \\
v_{SK}
\end{bmatrix}_t = 
\begin{bmatrix}
1 & 0 \\
a_2 & 1 \\
a_3 & 1
\end{bmatrix}
x_t +
\begin{bmatrix}
v_{BC} \\
v_{AB} \\
v_{SK}
\end{bmatrix}_t,
\text{ where } v_t \sim \text{MVN}
\begin{bmatrix}
0 & 0 \\
r & 0 \\
0 & r
\end{bmatrix}.
\]

(15.5)

In MARSS, we denote the model:

```r
model.b1=list()
model.b1$R="diagonal and equal"
```
We do not need to specify the structure of $Q$ and $u$ since they are scalar and have no structure.

We will compare this to a model where we assume that there is a separate population for British Columbia, Alberta, and Saskatchewan but they have the same process parameters (trend and process variance). Mathematically, this model is:

$$
\begin{bmatrix}
    x_{BC} \\
    x_{AB} \\
    x_{SK}
\end{bmatrix}_t =
\begin{bmatrix}
    x_{BC} \\
    x_{AB} \\
    x_{SK}
\end{bmatrix}_{t-1} +
\begin{bmatrix}
    u \\
    u \\
    u
\end{bmatrix} + w_t, \text{ where } w_t \sim \text{MVN}(0, \begin{bmatrix}
    q & 0 & 0 \\
    0 & q & 0 \\
    0 & 0 & q
\end{bmatrix})
$$

$$
\begin{bmatrix}
    y_{BC} \\
    y_{AB} \\
    y_{SK}
\end{bmatrix}_t =
\begin{bmatrix}
    1 & 0 & 0 \\
    0 & 1 & 0 \\
    0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
    x_{BC} \\
    x_{AB} \\
    x_{SK}
\end{bmatrix}_t +
\begin{bmatrix}
    v_{BC} \\
    v_{AB} \\
    v_{SK}
\end{bmatrix}_t, \text{ where } v_t \sim \text{MVN} \left( 0, \begin{bmatrix}
    r & 0 & 0 \\
    0 & r & 0 \\
    0 & 0 & r
\end{bmatrix} \right)
$$

(15.6)

This is specified as:
Because these populations are surveyed over a relatively large geographic area, it is reasonable to expect that environmental variation may differ between provinces. For our third model, we will fit a model with separate processes that are allowed to have unequal process parameters.

For our last model, we will consider lumping Alberta/Saskatchewan, because the time series suggest similar trends. Mathematically, this model is:

\[
\begin{bmatrix}
  x_{BC} \\
  x_{AB-SK}
\end{bmatrix}_t = \begin{bmatrix}
  x_{BC} \\
  x_{AB-SK}
\end{bmatrix}_{t-1} + \begin{bmatrix}
  u \\
  u
\end{bmatrix} + w_t, \text{ where } w_t \sim \text{MVN} \left( \begin{bmatrix} 0 \\ 0 \end{bmatrix}, \begin{bmatrix} q & 0 \\ 0 & q \end{bmatrix} \right)
\]

\[
\begin{bmatrix}
  y_{BC} \\
  y_{AB} \\
  y_{SK}
\end{bmatrix}_t = \begin{bmatrix}
  1 & 0 \\
  0 & 1 \\
  0 & 1
\end{bmatrix} \begin{bmatrix}
  x_{BC} \\
  x_{AB-SK}
\end{bmatrix}_t + \begin{bmatrix}
  0 \\
  0 \\
  a_3
\end{bmatrix} + \begin{bmatrix}
  v_{BC} \\
  v_{AB} \\
  v_{SK}
\end{bmatrix}_t, \text{ where } v_t \sim \text{MVN} \left( \begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix}, \begin{bmatrix} r & 0 & 0 \\ 0 & r & 0 \\ 0 & 0 & r \end{bmatrix} \right)
\]  

(15.7)

This model is specified as

\[
\text{mod}_{b4} = \text{list()}
\]

\[
\text{mod}_{b4}$Q$="diagonal and unequal"
\]

\[
\text{mod}_{b4}$R$="diagonal and equal"
\]

\[
\text{mod}_{b4}$Z$=\text{factor(c("BC","AB-SK","AB-SK"))}
\]

\[
\text{mod}_{b4}$A$="scaling"
\]

\[
\text{mod}_{b4}$U$="unequal"
\]

\[
\text{kem}_{b4} = \text{MARSS(birddat, model=mod}_{b4})
\]

The AICc values for the four models are

\[
\text{c(mod1=kem}_{b1}$AICc, mod2=kem}_{b2}$AICc, mod3=kem}_{b3}$AICc, mod4=kem}_{b4}$AICc)
\]

<table>
<thead>
<tr>
<th>mod1</th>
<th>mod2</th>
<th>mod3</th>
<th>mod4</th>
</tr>
</thead>
<tbody>
<tr>
<td>20.90670</td>
<td>22.96714</td>
<td>23.75125</td>
<td>14.76889</td>
</tr>
</tbody>
</table>

The last model is superior to the others, and lowers the AICc value compared to model 1 by 8 units. Figure [15.4] shows the fits for this model.
Fig. 15.4. Plot model 4 fits to the kestrel data.
16

Univariate dynamic linear models (DLMs)

16.1 Overview of dynamic linear models

In this chapter, we will use MARSS to analyze dynamic linear models (DLMs), wherein the parameters in a regression model are treated as time-varying. DLMs are used commonly in econometrics, but have received less attention in the ecological literature (c.f. Lamon III et al., 1998; Scheuerell and Williams, 2005). Our treatment of DLMs is rather cursory—we direct the reader to excellent textbooks by Pole et al. (1994) and Petris et al. (2009) for more in-depth treatments of DLMs. The former focuses on Bayesian estimation whereas the latter addresses both likelihood-based and Bayesian estimation methods.

We begin our description of DLMs with a static regression model, wherein the \( i \)-th observation is a linear function of an intercept, predictor variable(s), and a random error term. For example, if we had one predictor variable \((F)\), we could write the model as

\[
y_i = \alpha + \beta F_i + v_i,
\]

where the \( \alpha \) is the intercept, \( \beta \) is the regression slope, \( F_i \) is the predictor variable matched to the \( i \)-th observation \((y_i)\), and \( v_i \sim N(0,\sigma) \). It is important to note here that there is no implicit ordering of the index \( i \). That is, we could shuffle any/all of the \((y_i,F_i)\) pairs in our dataset with no effect on our ability to estimate the model parameters. We can write the model in Equation 16.1 using vector notation, such that

\[
\text{Type } \text{RSpatialDoc("Chapter_UnivariateDLM.R",package="MARSS") at the R command line to open a file with all the code for the examples in this chapter.}
\begin{align*}
y_i &= \begin{bmatrix} 1 \ F_i \end{bmatrix} \begin{bmatrix} \alpha \\ \beta \end{bmatrix} + v_i \\
&= F_i^\top \theta + v_i,
\end{align*}

(16.2)

and \( F_i^\top = \begin{bmatrix} \alpha \\ \beta \end{bmatrix} \) and \( \theta = \begin{bmatrix} \alpha \ \beta \end{bmatrix}^\top \).

In a DLM, however, the regression parameters are dynamic in that they evolve over time. For a single observation at time \( t \), we can write

\begin{equation}
y_t = F_t^\top \theta_t + v_t,
\end{equation}

(16.3)

where \( F_t \) is a column vector of regression variables at time \( t \), \( \theta_t \) is a column vector of regression parameters at time \( t \) and \( v_t \sim N(0, r) \). While seemingly identical, this formulation presents two features that distinguish it from Equation (16.2). First, the observed data are explicitly time ordered (i.e., \( y = \{y_1, y_2, y_3, \ldots, y_T\} \)), which means we expect them to contain implicit information. Second, the relationship between the observed datum and the predictor variables are unique at every time \( t \) (i.e., \( \theta = \{\theta_1, \theta_2, \theta_3, \ldots, \theta_T\} \)).

However, closer examination of Equation (16.3) reveals an apparent complication for parameter estimation. With only one datum at each time step \( t \), we could, at best, estimate only one regression parameter, and even then, the 1:1 correspondence between data and parameters would preclude any estimation of parameter uncertainty. To address this shortcoming, we return to the time ordering of model parameters. Rather than assume the regression parameters are independent from one time step to another, we instead model them as an autoregressive process where

\begin{equation}
\theta_t = G_t \theta_{t-1} + w_t,
\end{equation}

(16.4)

\( G_t \) is the parameter “evolution” matrix, and \( w_t \) is a vector of process errors, such that \( w_t \sim \text{MVN}(0, Q) \). The elements of \( G_t \) may be known and fixed \textit{a priori}, or unknown and estimated from the data. Although we allow for \( G_t \) to be time-varying, we will typically assume that it is time invariant.

The idea is that the evolution matrix \( G_t \) deterministically maps the parameter space from one time step to the next, so the parameters at time \( t \) are temporally related to those before and after. However, the process is stochastic and the mapping includes stochastic error, which leads to a degradation of information over time. If the diagonal elements of \( Q \) are relatively large, then the parameters can vary widely from \( t \) to \( t+1 \). If \( Q = 0 \), then \( \theta_1 = \theta_2 = \theta_T \) and we are back to the static model in Equation (16.1).

\textbf{16.2 Example of a univariate DLM}

Let’s consider an example from the literature. Scheuerell and Williams (2005) used a DLM to examine the relationship between marine survival of Chinook
salmon and an index of ocean upwelling strength along the west coast of the USA. Upwelling brings cool, nutrient-rich waters from the deep ocean to shallower coastal areas. Scheuerell and Williams hypothesized that stronger upwelling in April should create better growing conditions for phytoplankton, which would then translate into more zooplankton. In turn, juvenile salmon (“smolts”) entering the ocean in May and June should find better foraging opportunities. Thus, for smolts entering the ocean in year $t$,

$$survival_t = \alpha_t + \beta_t F_t + \nu_t \text{ with } \nu_t \sim N(0, r),$$

(16.5)

and $F_t$ is the coastal upwelling index\(^1\) for the month of April in year $t$.

Both the intercept and slope are time varying, so

$$\alpha_t = \alpha_{t-1} + w_t^{(1)} \text{ with } w_t^{(1)} \sim N(0, q_1); \text{ and }$$

(16.6)

$$\beta_t = \beta_{t-1} + w_t^{(2)} \text{ with } w_t^{(2)} \sim N(0, q_2).$$

(16.7)

If we define $\theta_t = [\alpha_t \beta_t]^\top$, $G_t = I$ for all $t$, $w_t = [w_t^{(1)} w_t^{(2)}]^\top$, and $Q = \begin{bmatrix} q_1 & 0 \\ 0 & q_2 \end{bmatrix}$, we get Equation (16.4). If we define $y_t = survival_t$ and $F_t = [1 F_t]^\top$, we can write out the full univariate DLM as a state-space model with the following form:

$$\theta_t = G_t \theta_{t-1} + w_t \sim MVN(0, Q);$$

$$y_t = F_t^\top \theta_t + \nu_t \text{ with } \nu_t \sim N(0, r);$$

(16.8)

$$\theta_0 \sim MVN(\pi_0, \Lambda_0).$$

Equation (16.8) is equivalent to our standard MARSS model:

$$x_t = B_t x_{t-1} + u_t + C_t c_t + w_t \sim MVN(0, Q_t);$$

$$y_t = Z_t x_t + a_t + D_t d_t + \nu_t \sim MVN(0, R_t);$$

$$x_0 \sim MVN(\pi, \Lambda);$$

(16.9)

where $x_t = \theta_t$, $B_t = G_t$, $u_t = C_t = c_t = 0$, $y_t = y_t$ (i.e., $y_t$ is $1 \times 1$), $Z_t = F_t^\top$, $a_t = D_t = d_t = 0$, and $R_t = r$ (i.e., $R_t$ is $1 \times 1$).

16.2.1 Fitting a univariate DLM with MARSS

Now let’s go ahead and analyze the DLM specified in Equations (16.5)–(16.8). We begin by getting the data set, which has 3 columns for 1) the year the salmon smolts migrated to the ocean ($year$), 2) logit-transformed survival\(^2\) ($logit.s$), and 3) the coastal upwelling index for April ($CUI.apr$). There are 42 years of data (1964–2005).

\(^1\) cubic meters of seawater per second per 100 m of coastline

\(^2\) Survival in the original context was defined as the proportion of juveniles that survive to adulthood. Thus, we use the logit function, defined as $logit(p) = loge(p/(1 - p))$, to map survival from the open interval $(0,1)$ onto the interval $(-\infty, \infty)$, which allows us to meet our assumption of normally distributed observation errors.
data(SalmonSurvCUI)
years = SalmonSurvCUI[,1]
TT = length(years)
# response data: logit(survival)
dat = matrix(SalmonSurvCUI[,2],nrow=1)

As we have seen in other chapters, standardizing our covariate(s) to have zero-mean and unit-variance can be helpful in model fitting and interpretation. In this case, it is a good idea because the variance of \( \text{CUI}.\text{apr} \) is orders of magnitude greater than \( \text{survival} \).

\[
\text{CUI} = \text{SalmonSurvCUI}[,"\text{CUI.apr"}]
\]
\[
\text{CUI.z} = \text{zscore(CUI)}
\]

Plots of logit-transformed survival and the z-scored April upwelling index are shown in Figure 16.1.

\[
\text{Fig. 16.1.} \text{ Time series of logit-transformed marine survival estimates for Snake River spring/summer Chinook salmon (top) and z-scores of the coastal upwelling index at 45N 125W (bottom). The x-axis indicates the year that the salmon smolts entered the ocean.}
\]

Next, we need to set up the appropriate matrices and vectors for MARSS. Let’s begin with those for the process equation because they are straightforward.

\[
\text{# for process eqn}
\]
\[
B = \text{diag}(m) \quad # 2x2; \text{Identity}
\]
16.2 Example of a univariate DLM

\[ U = \text{matrix}(0, nrow=m, ncol=1) \quad \# \text{2x1; both elements = 0} \]
\[ Q = \text{matrix} \left( \text{list}(0), m, m \right) \quad \# \text{2x2; all 0 for now} \]
\[ \text{diag}(Q) = c("q1","q2") \quad \# \text{2x2; diag = (q1,q2)} \]

Defining the correct form for the observation model is a little more tricky, however, because of how we model the effect(s) of explanatory variables. In a DLM, we need to use \( Z_t \) (instead of \( d_t \)) as the matrix of known regressors/drivers that affect \( y_t \), and \( x_t \) (instead of \( D_t \)) as the regression parameters. Therefore, we need to set \( Z_t \) equal to an \( n \times m \times T \) array, where \( n \) is the number of response variables (= 1; \( y_t \) is univariate), \( m \) is the number of regression parameters (= intercept + slope = 2), and \( T \) is the length of the time series (= 42).

\[ # \text{for observation eqn} \]
\[ Z = \text{array}(\text{NA}, c(1,m,TT)) \quad \# \text{N\timesM\timesT; empty for now} \]
\[ Z[1,1,] = \text{rep}(1,TT) \quad \# \text{N\times1; 1's for intercept} \]
\[ Z[1,2,] = \text{CUI.z} \quad \# \text{N\times1; regr variable} \]
\[ A = \text{matrix}(0) \quad \# \text{1\times1; scalar = 0} \]
\[ R = \text{matrix}("r") \quad \# \text{1\times1; scalar = r} \]

Lastly, we need to define our lists of initial starting values and model matrices/vectors.

\[ # \text{only need starting values for regr parameters} \]
\[ \text{inits.list} = \text{list}(x0=\text{matrix}(c(0, 0), nrow=m)) \]
\[ \# \text{list of model matrices & vectors} \]
\[ \text{mod.list} = \text{list}(B=B, U=U, Q=Q, Z=Z, A=A, R=R) \]

And now we can fit our DLM with MARSS.

\[ \text{dlm1 = MARSS(dat, inits=inits.list, model=mod.list)} \]

Success! abstol and log-log tests passed at 115 iterations. Alert: conv.test.slope.tol is 0.5. Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 115 iterations.
Log-likelihood: -40.03813
AIC: 90.07627   AICc: 91.74293

\[ \begin{array}{l}
\text{Estimate} \\
R.r \quad 0.15708 \\
Q.q1 \quad 0.11264 \\
Q.q2 \quad 0.00564 \\
x0.X1 \quad -3.34023 \\
\end{array} \]
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\[ x_{0,X2} = -0.05388 \]

Initial states \((x_0)\) defined at \(t=0\)

Standard errors have not been calculated.

Use MARSSparamCIs to compute CIs and bias estimates.

Notice that the MARSS output does not list any estimates of the regression parameters themselves. Why not? Remember that in a DLM the matrix of states \((x)\) contains the estimates of the regression parameters \((\theta)\). Therefore, we need to look in \(\text{dlm1}\$\text{states}\) for the MLEs of the regression parameters, and in \(\text{dlm1}\$\text{states.se}\) for their standard errors.

Time series of the estimated intercept and slope are shown in Figure 16.2. It appears as though the intercept is much more dynamic than the slope, as indicated by a much larger estimate of process variance for the former \((Q_{11})\). In fact, although the effect of April upwelling appears to be increasing over time, it doesn’t really become important as an explanatory variable until about 1990 when the approximate 95% confidence interval for the slope no longer overlaps zero.

![Time series of estimated mean states](image)

**Fig. 16.2.** Time series of estimated mean states (thick lines) for the intercept (top) and slope (bottom) parameters from the univariate DLM specified by Equations 16.5–16.8. Thin lines denote the mean ± 2 standard deviations.

16.3 Forecasting with a univariate DLM

Scheuerell and Williams (2005) were interested in how well upwelling could be used to forecast expected survival of salmon. Let’s look at how well our
16.3 Forecasting with a univariate DLM

To do so, we need the predictive distributions for the regression parameters and observation. Beginning with our definition for the distribution of the parameters at time $t = 0$, $\theta_0 \sim \text{MVN}(\pi_0, \Lambda_0)$ in Equation 16.8, we write

$$\theta_{t-1|y_{1:t-1}} \sim \text{MVN}(\pi_{t-1}, \Lambda_{t-1})$$

(16.10)

to indicate the distribution of $\theta$ at time $t - 1$ conditioned on the observed data through time $t - 1$ (i.e., $y_{1:t-1}$). Then, we can write the one-step ahead predictive distribution for $\theta_t$ given $y_{1:t-1}$ as

$$\theta_t|y_{1:t-1} \sim \text{MVN}(\eta_t, \Phi_t),$$

where

$$\eta_t = G_t \pi_{t-1},$$

and

$$\Phi_t = G_t \Lambda_{t-1} G_t^\top + Q.$$

(16.11)

Consequently, the one-step ahead predictive distribution for the observation at time $t$ given $y_{1:t-1}$ is

$$y_t|y_{1:t-1} \sim \text{N}(\zeta_t, \Psi_t),$$

where

$$\zeta_t = F_t \eta_t,$$

and

$$\Psi_t = F_t \Phi_t F_t^\top + R.$$

(16.12)

16.3.1 Forecasting a univariate DLM with MARSS

Working from Equation 16.12, we can now use MARSS to compute the expected value of the forecast at time $t$ ($E[y_t|y_{1:t-1}] = \zeta_t$), and its variance ($\text{var}[y_t|y_{1:t-1}] = \Psi_t$). For the expectation, we need $F_t \eta_t$. Recall that $F_t$ is our $1 \times m$ matrix of explanatory variables at time $t$ ($F_t$ is called $Z_t$ in MARSS notation). The one-step ahead forecsts of the parameters at time $t$ ($\eta_t$) are calculated as part of the Kalman filter algorithm—they are termed $\hat{x}_{t-1}$ in MARSS notation and stored as $\texttt{xtt1}$ in the list produced by the \texttt{MARSSkfss()} function.

```r
# get list of Kalman filter output
kf.out = MARSSkfss(dlm1)
# forecasts of regr parameters; 2xT matrix
eta = kf.out$xt$xtt1
# ts of E(forecasts)
fore.mean = vector()
for(t in 1:TT) {
  fore.mean[t] = Z[,,t] %*% eta[,t,drop=F]
}
```

For the variance of the forecasts, we need $F_t \Phi_t F_t^\top + R$. As with the mean, $F_t \equiv Z_t$. The variances of the one-step ahead forecasts of the parameters at time $t$ ($\Phi_t$) are also calculated as part of the Kalman filter algorithm—they
are stored as Vtt1 in the list produced by the \texttt{MARSSkfss()} function. Lastly, the observation variance $R$ is part of the standard MARSS output.

```r
# variance of regr parameters; 1x2xT array
Phi = kf.out$Vtt1
# obs variance; 1x1 matrix
R.est = coef(dlm1, type="matrix")$R
# ts of Var(forecasts)
fore.var = vector()
for(t in 1:TT) {
  tZ = matrix(Z[,,t],m,1) # transpose of Z
  fore.var[t] = Z[,,t] %*% Phi[,,t] %*% tZ + R.est
}
```

Plots of the model mean forecasts with their estimated uncertainty are shown in Figure 16.3. Nearly all of the observed values fell within the approximate prediction interval. Notice that we have a forecasted value for the first year of the time series (1964), which may seem at odds with our notion of forecasting at time $t$ based on data available only through time $t-1$. In this case, however, MARSS is actually estimating the states at $t=0$ ($\theta_0$), which allows us to compute a forecast for the first time point.

![Fig. 16.3. Time series of logit-transformed survival data (blue dots) and model mean one-step ahead forecasts (thick line). Thin lines denote the approximate 95% prediction intervals.](image)

Although our model forecasts look reasonable in logit-space, it is worthwhile to examine how well they look when the survival data and forecasts are back-transformed onto the interval [0,1] (Figure 16.4). In that case, the accuracy does not seem to be affected, but the precision appears much worse, especially during the early and late portions of the time series when survival is changing rapidly.
16.3 Forecasting with a univariate DLM

16.3.2 DLM forecast diagnostics

As with other time series models, evaluation of a DLM should include model diagnostics. In a forecasting context, we are often interested in the forecast errors, which are simply the observed data minus the forecasts \( e_t = y_t - \zeta_t \). In particular, the following assumptions should hold true for \( e_t \):

1. \( e_t \sim N(0, \sigma^2) \);
2. \( \text{cov}(e_t, e_{t-k}) = 0 \).

In the literature on state-space models, the set of \( e_t \) are commonly referred to as “innovations”. \texttt{MARSS()} calculates the innovations as part of the Kalman filter algorithm—they are stored as \texttt{Innov} in the list produced by the \texttt{MARSSkfss()} function.

```r
# forecast errors
innov = kf.out$Innov
```

Let’s see if our innovations meet the model assumptions. Beginning with (1), we can use a Q-Q plot to see whether the innovations are normally distributed with a mean of zero. We’ll use the \texttt{qqnorm()} function to plot the quantiles of the innovations on the \( y \)-axis versus the theoretical quantiles from a Normal distribution on the \( x \)-axis. If the 2 distributions are similar, the points should fall on the line defined by \( y = x \).

```r
# Q-Q plot of innovations
qqnorm(t(innov), main="", pch=16, col="blue")
# add y=x line for easier interpretation
qline(t(innov))
```

The Q-Q plot (Figure [16.5]) indicates that the innovations appear to be more-or-less normally distributed (i.e., most points fall on the line). Furthermore, it looks like the mean of the innovations is about 0, but we should...
use a more reliable test than simple visual inspection. We can formally test whether the mean of the innovations is significantly different from 0 by using a one-sample $t$-test, based on a null hypothesis of $E[e_t] = 0$. To do so, we will use the function `t.test()` and base our inference on a significance value of $\alpha = 0.05$.

```r
# p-value for t-test of H0: E(innov) = 0
t.test(t(innov), mu=0)$p.value
```

[1] 0.4840901

The $p$-value $> 0.05$ so we cannot reject the null hypothesis that $E[e_t] = 0$.

Moving on to assumption (2), we can use the sample autocorrelation function (ACF) to examine whether the innovations covary with a time-lagged version of themselves. Using the `acf()` function, we can compute and plot the correlations of $e_t$ and $e_{t-k}$ for various values of $k$. Assumption (2) will be met if none of the correlation coefficients exceed the 95% confidence intervals defined by $\pm z_{0.975}/\sqrt{n}$.

```r
# plot ACF of innovations
acf(t(innov), lag.max=10)
```

The ACF plot (Figure 16.6) shows no significant autocorrelation in the innovations at lags 1–10, so it appears that both of our model assumptions have been met.
Fig. 16.6. Autocorrelation plot of the forecast errors (innovations) for the DLM specified in Equations 16.5–16.8. Horizontal blue lines define the upper and lower 95% confidence intervals.
Multivariate linear regression

This chapter shows how to write regression models with multivariate responses and multivariate explanatory variables in MARSS form. R has many excellent packages for multiple linear regression. We will be showing how to use the MARSS() function to fit these models, but note that R’s standard linear regression functions would be much better choices in most cases. The purpose of this chapter is to show the relationship between multivariate linear regression and the MARSS equation.

In a classic linear regression, the response variable \( y \) is univariate and there may one to multiple explanatory variables \((d_1, d_2, \ldots)\) plus an optional intercept \((\alpha)\):

\[
y_t = \alpha + \sum_k \beta_k d_k + e_t, \quad \text{where } e_t \sim N(0, \sigma^2)
\]  

Here the subscript, \( t \) is used since we are working with time-series data. Explanatory variables are normally denoted \( x \) in linear regression however \( x \) is not used here since \( x \) is already used in MARSS models to denote the hidden process trajectory. Instead \( d \) is used when the explanatory variables appear in the \( y \) part of the equation (and \( c \) if they appear in the \( x \) part).

This chapter will start with classical linear regression where the explanatory variables are treated as inputs that are known without error and where we are trying to explain the variation in \( y \) with our explanatory variables. We will extend this to the case of autocorrelated errors.

17.1 Univariate linear regression

A vanilla linear regression where our data are time ordered but we treat them as independent can be written

Type `RShowDoc("Chapter_MLR.R",package="MARSS")` at the R command line to open a file with all the code for the examples in this chapter.
y_t = \alpha + \beta_1 d_{1,t} + \beta_2 d_{2,t} + \epsilon_t, \quad (17.2)

where the \( d \) are our explanatory variables. This model can be written in many different ways in as a MARSS equation. Here we use a specific form where the i.i.d. component of the errors is \( v_t \) in the \( y \) part of the MARSS equation and the autocorrelated errors appear as \( x_t \) in the \( y \) equation. Specifying the MARSS model this way allows us to use the EM-algorithm to fit the model which will prove to be important.

\[
y_t = \alpha + \begin{bmatrix} \beta_1 \\ \beta_2 \\ \vdots \\ \beta_j \end{bmatrix} \begin{bmatrix} d_{1,t} \\ d_{2,t} \\ \vdots \\ d_{j,t} \end{bmatrix} + v_t, v_t \sim N(0, r) \quad (17.3)
\]

\[
x_t = bx_{t-1} + w_t, w_t \sim N(0, q)
\]

The \( v_t \) are the i.i.d. errors and the \( x_t \) are the AR(1) errors.

17.1.1 Univariate response using the Longley dataset: example 1

We will start by using an example from Chapter 6 in Linear Models in R [Faraway, 2004]. This example uses the built-in R dataset “longley” which has the number of people employed from 1947 to 1962 and a number of predictors. For this example we will regress the number of people employed against gross National product and population size (following Faraway).

Mathematically, the model we are fitting is

\[
Employed_t = \alpha + \begin{bmatrix} \beta_{GnP} \\ \beta_{Pop} \end{bmatrix} \begin{bmatrix} GNP_t \\ Pop_t \end{bmatrix} + v_t, v_t \sim N(0, r) \quad (17.4)
\]

\( x \) does not appear in the vanilla linear regression since we do not have autocorrelated errors (yet). We are trying to estimate \( \alpha \) (intercept), \( \beta_{GnP} \) and \( \beta_{Pop} \).

A full multivariate MARSS model looks like

\[
y_t = Zx_t + a + Dd_t + v_t, \text{ where } v_t \sim \text{MVN}(0, R)
\]

\[
x_t = Bx_{t-1} + u + Cc_t + w_t, \text{ where } w_t \sim \text{MVN}(0, Q) \quad (17.5)
\]

We need to specify the parameters in Equation 17.5 such that we get Equation 17.4.

First, we load the data and set up \( y \), the response variable number of people employed, as a matrix with time going across the columns.

\[
data(longley)
Employed = matrix(longley$Employed, nrow=1)
\]

Second create a list to hold our model specification.
Fig. 17.1. Employment time series from the Longley dataset.

```r
longley.model=list()

Set the u, Q and x0 parameters to 0. We will also set a and C to 0 and B and Z to identity although this is not necessary since these are the defaults.

```r
longley.model$U=longley.model$Q="zero"
longley.model$C="zero"
longley.model$B=longley.model$Z="identity"
longley.model$x0="zero"
longley.model$tinitx=0
```  

We will estimate R, the variance of the i.i.d. errors (residuals).

```r
longley.model$R=matrix("r")
```  

The D matrix has the two \( \beta \) (slope) parameters for GNP and Population and \( a \) has the intercept.\(^1\)

\(^1\) A better way to fit the model is to put the intercept into D by adding a row of 1s to d and putting the intercept parameter on the first row of D. This reduces by one the number of matrices being estimated by the EM algorithm. It’s not done here just so the equations look more like standard linear regression equations.
Last we set up our explanatory variables. This is the \( d \) matrix and we need each explanatory variable in a row and time across the columns.

\[
\text{longley.model}\$d = \text{rbind(longley}\$\text{GNP, longley}\$\text{Population)}
\]

Now we can fit the model:

\[
\text{mod1=MARSS(Employed, model=longley.model)}
\]

and look at the estimates.

\[
\text{coef(mod1, type="vector")}
\]

\text{method="BFGS"} can also be used and gives similar results.

We can compare the fit to that from \text{lm()} and see that we get the same estimates:

\[
\text{mod1.lm=lm(Employed \sim GNP + Population, data=longley)}
\]

\[
\text{coef(mod1.lm)}
\]

\begin{verbatim}
(Intercept)   GNP  Population
88.93879831  0.06317244 -0.40974292
\end{verbatim}

### 17.1.2 Univariate response using auto-correlated errors: example 1

As Faraway (2004) discusses, the errors in this dataset are temporally correlated. We can model the errors as an AR(1) process to account for this. This changes our model to

\[
\begin{aligned}
\text{Employed}_t &= \alpha + \begin{bmatrix} \beta_{\text{GNP}} & \beta_{\text{Pop}} \end{bmatrix} \begin{bmatrix} \text{GNP}_t \\ \text{Pop}_t \end{bmatrix} + x_t, v_t \sim N(0, r) \\
x_t &= bx_{t-1} + w_t, w_t \sim N(0, q) \\
x_0 &= 0
\end{aligned}
\] (17.6)

We assume the AR(1) errors have mean 0 so \( u = 0 \) in the \( x_t \) equation. Setting \( u \) to anything else would make the mean of our errors equal to \( u/(1 - b) \) for \(-1 < b < 1\). This would lead to two mean levels in our model, \( \alpha \) and \( u/(1 - b) \), and we would not be able to estimate both. Notice that the model is somewhat confounded since if \( b = 0 \) then \( x_t \) is i.i.d. errors same as \( v_t \). In this case, either \( q \) or \( r \) would be redundant. It is thus possible that either \( r \) or \( q \) will go to zero.

To fit the model with autoregressive errors, we add the \( x \) parameters to our model list. We estimate \( b \) and \( q \).

\[
\text{longley.ar1=longley.model}
\]

\[
\text{longley.ar1}\$B=\text{matrix("b")}
\]

\[
\text{longley.ar1}\$Q=\text{matrix("q")}
\]
Now we can fit the model as before

\[ \text{mod2} = \text{MARSS(employed, model=longley.ar1)} \]

however, this is a difficult model to fit and takes a long, long time to converge. The default \text{maxit} used in the call above is not nearly enough iterations. Using \text{method="BFGS"} helps a little but not much. We can improve behavior by using the fit of the model with i.i.d. errors as initial conditions for \( D \) and \( a \).

\[
\text{inits=list(A=coef(mod1)$A, D=coef(mod1)$D)}
\]

\[
\text{mod2=MARSS(employed, model=longley.ar1, inits=inits, control=list(maxit=1000))}
\]

\[
\text{ests.marss} = c(b=coef(mod2)$b, alpha=coef(mod2)$A, GNP=coef(mod2)$D[1], Population=coef(mod2)$D[2], logLik=logLik(mod2))
\]

We can the fit the same model using \text{gls()} (\text{nlme} package). The \( b \) term is called \( \Phi \) in the \text{gls()} call and is somewhat difficult to recover although it is printed by \text{summary()}.

\[
\text{library(nlme)}
\]

\[
\text{mod2.gls=gls(employed ~ GNP + Population, correlation=corAR1(), data=longley, method="ML")}
\]

\[
\text{mod2.gls.phi = coef(mod2.gls$modelStruct[[1]], unconstrained=FALSE)}
\]

\[
\text{ests.gls} = c(b=mod2.gls.phi, alpha=coef(mod2.gls)[1], GNP=coef(mod2.gls)[2], Population=coef(mod2.gls)[3], logLik=logLik(mod2.gls))
\]

Note we need to set \text{method="ML"} to maximize the likelihood because the default is to maximize the restricted maximum-likelihood (\text{method="REML"}) and that gives a different answer from the MARSS() function since MARSS() is maximizing the likelihood.

Both functions return similar values though \text{gls()} is much faster and the EM algorithm has not quite converged even with 1000 iterations.

\[
\text{rbind(MARSS = ests.marss, GLS = ests.gls)}
\]

<table>
<thead>
<tr>
<th></th>
<th>b</th>
<th>alpha</th>
<th>GNP</th>
<th>Population</th>
<th>logLik</th>
</tr>
</thead>
<tbody>
<tr>
<td>MARSS</td>
<td>0.3509377</td>
<td>95.36017</td>
<td>0.06748567</td>
<td>-0.4784003</td>
<td>-10.53742</td>
</tr>
<tr>
<td>GLS</td>
<td>0.3651196</td>
<td>96.09369</td>
<td>0.06822305</td>
<td>-0.4871554</td>
<td>-10.47396</td>
</tr>
</tbody>
</table>

17.1.3 Univariate response using the Longley dataset: example 2

The full Longley dataset is often used to test the performance of numerical methods for fitting linear regression models because it has severe collinearity problems (Figure 17.2). We can compare the EM and BFGS algorithms for the full dataset and see how fitting a MARSS model with the BFGS algorithm leads to estimates far from the maximum-likelihood values for this problem.
We can fit a regression of Employed to all the Longley explanatory variables using the following code. The mathematical model is the same as in Equation 17.4 except that instead of two explanatory variables with have all seven shown in Figure 17.2.

```r
eVar.names = colnames(longley)[-7]
eVar = t(longley[,eVar.names])
longley.model=list()
longley.model$U=longley.model$Q="zero"
longley.model$C="zero"
longley.model$B=longley.model$Z="identity"
longley.model$A=matrix("intercept")
longley.model$R=matrix("r")
longley.model$D=matrix(eVar.names,nrow=1)
longley.model$d = eVar
longley.model$x0="zero"
longley.model$tinitx=0
```

And then we fit as usual. We will fit with the EM-algorithm (the default) and compare to BFGS.
mod3.em=MARSS(Employed, model=longley.model)
mod3.bfgs=MARSS(Employed, model=longley.model, method="BFGS")

Here are the EM estimates with the log-likelihood.

par.names = c("A.intercept", paste("D",eVar.names,sep="."))
c(coef(mod3.em, type="vector")[,par.names], logLik=mod3.em$logLik)

A.intercept D.GNP.deflator D.GNP D.Unemployed
-3.482259e+03 1.506187e-02 -3.581918e-02 -2.020230e-02
D.Armed.Forces D.Population D.Year logLik
-1.033227e-02 -5.110410e-02 1.829152e+00 9.066497e-01

Compared to the BFGS estimates:

c(coef(mod3.bfgs, type="vector")[,par.names], logLik=mod3.bfgs$logLik)

A.intercept D.GNP.deflator D.GNP D.Unemployed
-14.062105281 -0.052705201 0.070642032 -0.004298481
D.Armed.Forces D.Population D.Year logLik
-0.005744197 -0.412771919 0.055610015 -6.996818702

And compared to the estimates from the lm() function:

mod3.lm=lm(Employed ~ 1 + GNP.deflator + GNP + Unemployed + Armed.Forces + Population + Year, data=longley)
c(coef(mod3.lm),logLik=logLik(mod3.lm))

(Intercept) GNP.deflator GNP Unemployed Armed.Forces Population Year logLik
-3.482259e+03 1.506187e-02 -3.581918e-02 -2.020230e-02 -1.033227e-02 -5.110410e-02 1.829151e+00 9.066497e-01

As you can see the BFGS algorithm struggles with the ridge-like likelihood caused by the collinearity in the explanatory variables.

We can also compare the performance of the model with AR(1) errors. This is Equation 17.6 but with all seven explanatory variables. We set up the MARSS model\(^2\) for a linear regression with correlated errors as before with the addition of \(b\) (called Phi in gls()) and \(q\).

longley.correrr.model=longley.model
longley.correrr.model$B=matrix(\"b\")
longley.correrr.model$Q=matrix(\"q\")

We fit as usual and compare the EM-algorithm (the default) to fits using BFGS. We will use the estimate from the model with i.i.d. errors as initial conditions.

\(^2\) Notice that \(x_0\) is set at 0. The model is having a hard time fitting \(x_0\) because the time series is short. Estimating \(x_0\) or using a diffuse prior by setting \(V_0\) big, leads to poor estimates. Since this is just the error term, we set \(x_0 = 0\) since the mean of the errors is assumed to be 0.
Here are the EM estimates with the log-likelihood. We only show $\phi$ (the $b$ term in the AR(1) error equation) and the log-likelihood.

\[
c(coef(mod4.em, type="vector")["B.b"], logLik=mod4.em$logLik)
\]

```
B.b  logLik
-0.7737457 4.5374547
```

Compared to the BFGS estimates:

\[
c(coef(mod4.bfgs, type="vector")["B.b"], logLik=mod4.bfgs$logLik)
\]

```
B.b  logLik
0.8368905 0.9066497
```

And compared to the estimates from the gls() function:

\[
mod4.gls=gls(Employed ~ 1 + GNP.deflator + GNP + Unemployed + Armed.Forces + Population + Year, correlation=corAR1(), data=longley, method="ML")
mod4.gls.phi = coef(mod4.gls$modelStruct[[1]], unconstrained=FALSE)
c(mod4.gls.phi, logLik=logLik(mod4.gls))
\]

```
Phi  logLik
-0.7288687 4.3865475
```

Again we see that the BFGS algorithm struggles with the ridge-like likelihood caused by the collinearity in the explanatory variables.

17.2 Multivariate response example using longitudinal data

We will illustrate linear regression with a multivariate response using longitudinal data from a sleep study on 18 subjects from the `lme4` R package. These are data on reaction time of subjects after 0 to 9 days of being restricted to 3 hours of sleep.

We load the data from the `lme4` package:

```
data(sleepstudy, package="lme4")
```

We set up the data into the form required for the MARSS() function.

```
#number of subjects
nsub = length(unique(sleepstudy$Subject))
days=length(sleepstudy$Days)/nsub
```
17.2 Multivariate response example using longitudinal data

Days of sleep deprivation
Average reaction time (ms)

Fig. 17.3. Plot of the sleep study data (package lme4).

#each subject is a row with day across the columns
dat = matrix(sleepstudy$Reaction, nsub, ndays,byrow=TRUE)
rownames(dat)=paste("sub",unique(sleepstudy$Subject),sep=".")
#the day number 0 to 9 is the explanatory variable
exp.var=matrix(sleepstudy$Days, 1, ndays,byrow=TRUE)

Let’s start with a simple regression where each subject has a separate intercept (reaction time at day 0) but the slope (increase in reaction time with each successive day) is the same across the 18 subjects. Mathematically the model is

\[
\begin{align*}
[r_{1}]_{t} & = [\alpha_{1}]_{t} + [\beta] day_{t} + [v_{1}]_{t} \\
[r_{2}]_{t} & = [\alpha_{2}]_{t} + [\beta] day_{t} + [v_{2}]_{t} \\
& \ldots \\
[r_{18}]_{t} & = [\alpha_{18}]_{t} + [\beta] day_{t} + [v_{18}]_{t}
\end{align*}
\]

(17.7)

\[
[v_{1}]_{t} \sim N\left(0, \begin{bmatrix} r & 0 & \ldots & 0 \\
0 & r & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & 0 & r \end{bmatrix}\right)
\]
The response time of subject $i$ is a subject specific intercept ($\alpha_i$) plus an effect of day at time $t$ (so 0, 1, 2, etc) that doesn’t vary by subject and error that is i.i.d. across subject and day.

We specify and fit this model as follows

```r
sleep.model=list(
  A="unequal", B="zero", x0="zero", U="zero",
  D=matrix("b1", nsub, 1), d=exp.var, tinitx=0, Q="zero")
```

```r
sleep.mod1 = MARSS(dat, model=sleep.model)
```

This is the same as the following with `lm()`:

```r
sleep.lm1 = lm(Reaction ~ -1 + Subject + Days, data=sleepstudy)
```

Now let’s allow each subject to have different slopes (increase in reaction time with each successive day) across subjects. This model is

\[
\begin{bmatrix}
resp_1 \\
resp_2 \\
\vdots \\
resp_{18}
\end{bmatrix}_t = \begin{bmatrix}
\alpha_1 \\
\alpha_2 \\
\vdots \\
\alpha_{18}
\end{bmatrix}_t + \begin{bmatrix}
\beta_1 \\
\beta_2 \\
\vdots \\
\beta_{18}
\end{bmatrix}_t \text{day}_t + \begin{bmatrix}
v_1 \\
v_2 \\
\vdots \\
v_{18}
\end{bmatrix}_t
\]

\[
\begin{bmatrix}
v_1 \\
v_2 \\
\vdots \\
v_{18}
\end{bmatrix}_t \sim N\left(0, \begin{bmatrix}
0 & 0 & \ldots & 0 \\
0 & 0 & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & 0
\end{bmatrix}\right)
\]

(17.8)

We specify and fit this model as

```r
sleep.model=list(
  A="unequal", B="zero", x0="zero", U="zero",
  D="unequal", d=exp.var, tinitx=0, Q="zero")
sleep.mod2 = MARSS(dat, model=sleep.model, silent=TRUE)
```

This is the same as the following with `lm()`:

```r
sleep.lm2 = lm(Reaction ~ 0 + Subject + Days:Subject, data = sleepstudy)
```

We can repeat the above but allow the residual variance to differ across subjects by setting `R="diagonal and unequal"`. This model is

\[
\begin{bmatrix}
resp_1 \\
resp_2 \\
\vdots \\
resp_{18}
\end{bmatrix}_t = \begin{bmatrix}
\alpha_1 \\
\alpha_2 \\
\vdots \\
\alpha_{18}
\end{bmatrix}_t + \begin{bmatrix}
\beta_1 \\
\beta_2 \\
\vdots \\
\beta_{18}
\end{bmatrix}_t \text{day}_t + \begin{bmatrix}
v_1 \\
v_2 \\
\vdots \\
v_{18}
\end{bmatrix}_t
\]

\[
\begin{bmatrix}
v_1 \\
v_2 \\
\vdots \\
v_{18}
\end{bmatrix}_t \sim N\left(0, \begin{bmatrix}
r_1 & 0 & \ldots & 0 \\
0 & r_2 & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & r_{18}
\end{bmatrix}\right)
\]

(17.9)
17.2 Multivariate response example using longitudinal data

```r
sleep.model = list(A = "unequal", B = "zero", x0 = "zero", U = "zero", D = "unequal", d = exp.var, tinitx = 0, Q = "zero", R = "diagonal and unequal")
sleep.mod3 = MARSS(dat, model = sleep.model, silent = TRUE)
```

Or we can allow AR(1) errors across subjects and allow each subject to have its own AR(1) parameters for this error. This model is

\[
\begin{bmatrix}
resp_1 \\
resp_2 \\
\vdots \\
resp_{18}
\end{bmatrix}
= \begin{bmatrix}
\alpha_1 \\
\alpha_2 \\
\vdots \\
\alpha_{18}
\end{bmatrix}
+ \begin{bmatrix}
\beta_1 \\
\beta_2 \\
\vdots \\
\beta_{18}
\end{bmatrix} \begin{bmatrix}
day_1 \\
day_2 \\
\vdots \\
\vdots \\
day_{18}
\end{bmatrix}
+ \begin{bmatrix}
v_1 \\
v_2 \\
\vdots \\
v_{18}
\end{bmatrix}
\sim N\left(\begin{bmatrix}0 \\ 0 \\ \vdots \\ 0\end{bmatrix}, \begin{bmatrix}r_{11} & 0 & \cdots & 0 \\
0 & r_{22} & \cdots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \cdots & r_{18}\end{bmatrix}\right)
\]

\[
\begin{bmatrix}
x_1 \\
x_2 \\
\vdots \\
x_{18}
\end{bmatrix}
= \begin{bmatrix}
b_1 & 0 & \cdots & 0 \\
0 & b_2 & \cdots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \cdots & b_{18}
\end{bmatrix}
\begin{bmatrix}
x_1 \\
x_2 \\
\vdots \\
x_{18}
\end{bmatrix}_{t-1}
+ \begin{bmatrix}
w_1 \\
w_2 \\
\vdots \\
w_{18}
\end{bmatrix}
\sim N\left(\begin{bmatrix}0 \\ 0 \\ \vdots \\ 0\end{bmatrix}, \begin{bmatrix}q_{11} & 0 & \cdots & 0 \\
0 & q_{22} & \cdots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \cdots & q_{18}\end{bmatrix}\right)
\]

(17.10)

We fit this model as

```r
inits = list(A = coef(sleep.mod3)$A, D = coef(sleep.mod3)$D)
# estimate a separate intercept for each but slope is the same
sleep.model = list(A = "unequal", B = "diagonal and unequal", x0 = "zero", U = "zero", D = "unequal", d = exp.var, tinitx = 0, Q = "diagonal and unequal", R = "diagonal and unequal")
sleep.mod4 = MARSS(dat, model = sleep.model, inits = inits, silent = TRUE)
```

It is not obvious how to specify these last two models using gls() or if it is possible.

We can also allow each subject to have his/her own error process but specify that the parameters of these (b, q and r) are the same across subjects. We do this by using "diagonal and equal". Mathematically this model is
Multivariate linear regression

\[
\begin{bmatrix}
\text{resp}_1 \\
\text{resp}_2 \\
\vdots \\
\text{resp}_{18}
\end{bmatrix}
= \begin{bmatrix}
\alpha_1 \\
\alpha_2 \\
\vdots \\
\alpha_{18}
\end{bmatrix} + \begin{bmatrix}
\beta_1 \\
\beta_2 \\
\vdots \\
\beta_{18}
\end{bmatrix} \text{day}_t + \begin{bmatrix}
v_1 \\
v_2 \\
\vdots \\
v_{18}
\end{bmatrix} + \begin{bmatrix}
x_1 \\
x_2 \\
\vdots \\
x_{18}
\end{bmatrix} \sim N\left(0, \begin{bmatrix}
r & 0 & \ldots & 0 \\
0 & r & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & r
\end{bmatrix}\right)
\]

\[
\begin{bmatrix}
x_1 \\
x_2 \\
\vdots \\
x_{18}
\end{bmatrix}
= \begin{bmatrix}
b & 0 & \ldots & 0 \\
0 & b & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & b
\end{bmatrix} \begin{bmatrix}
x_1 \\
x_2 \\
\vdots \\
x_{18}
\end{bmatrix}_{t-1} + \begin{bmatrix}
w_1 \\
w_2 \\
\vdots \\
w_{18}
\end{bmatrix} \sim N\left(0, \begin{bmatrix}
q & 0 & \ldots & 0 \\
0 & q & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & q
\end{bmatrix}\right)
\]

We specify and fit this model as

```r
inits=list(A=coef(sleep.mod3)$A,D=coef(sleep.mod3)$D)
#estimate a separate intercept for each but slope is the same
sleep.model=list(
  A="unequal",B="diagonal and equal", x0="zero", U="zero",
  D="unequal", d=exp.var, tinitx=0, Q="diagonal and equal",
  R="diagonal and equal")
sleep.mod5 = MARSS(dat, model=sleep.model, inits=inits, silent=TRUE)
```

This is fairly close to this model fit with gls().

```r
sleep.mod5.gls=gls(Reaction ~ 0 + Subject + Days:Subject, data=sleepstudy,
  correlation=corAR1(form= ~ 1|Subject), method="ML")
```

The way the variance-covariance structure is modeled is a little different but it’s the same idea.

### 17.3 Discussion

The purpose of the is chapter is to illustrate how linear regression models with multivariate explanatory variable or response variables can be written in MARSS form and fit with the MARSS() function\(^3\). This is to help one understand the relationship between the MARSS model form and the more familiar multivariate linear model forms. Obviously R has many, many packages for linear regression and generalized linear regression (non-Gaussian errors). While

\(^3\) with caveat that one must always be careful when the likelihood surface has prominent ridges which will occur with collinear explanatory variables.
Table 17.1. Parameter estimates of different versions of the model where each subject has a separate intercept (response time on normal sleep) and different slope by day (increase in response time with each day of sleep deprivation). The model types are discussed in the text.

<table>
<thead>
<tr>
<th></th>
<th>lm</th>
<th>mod2</th>
<th>em</th>
<th>mod3</th>
<th>em</th>
<th>mod4</th>
<th>em</th>
<th>mod5</th>
<th>em</th>
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<tr>
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<td>0.08</td>
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</tbody>
</table>
the MARSS package can fit a variety of linear regression models with Gaussian errors, that is not what it is designed to do. The MARSS package is designed for fitting models that cannot be fit with typical linear regression: multivariate autoregressive state-space models with inputs (explanatory variables).
Lag-p models with MARSS

18.1 Background

Most of the chapters in the User Guide are ‘lag-1’ in the autoregressive part of the model. This means that $x_t$ in the process model only depends on $x_{t-1}$ and not $x_{t-2}$ (lag-2) or more generally $x_{t-p}$ (lag-p). A lag-p model can be written in state-space form as a MARSS lag-1 model, aka a MARSS(1) model (see section 11.3.2 in [Tsay (2010)]). Writing lag-p models in this form allows one to take advantage of the fitting algorithms for MARSS(1) models. There are a number of ways to do the conversion to a MARSS(1) form. We use Hamilton’s form (section 1 in [Hamilton (1994)]) because it can be fit with an EM algorithm while the other forms (Harvey’s and Akaike’s) cannot.

This chapter shows how to convert and fit the following using the MARSS(1) form:

- **AR(p)** A univariate autoregressive model where $x_t$ is a function of $x_{t-p}$ (and the prior lags usually too). No observation error.
- **MAR(p)** The same as AR(p) but the $x$ term is multivariate not univariate.
- **ARSS(p)** The same as AR(p) but with a observation model and observation error. The observations may be multivariate ($y$ can be multivariate) but the $x$ term is univariate.
- **MARSS(p)** The same as ARSS(p) but the $x$ term is multivariate not univariate.

Note that only ARSS(p) and MARSS(p) assume observation error in the data. AR(p) and MAR(p) will be rewritten in the state-space form with a $y$ component to facilitate statistical analysis but the data themselves are considered error free.

Type `RShowDoc("Chapter_MARp.R", package="MARSS")` at the R command line to open a file with all the code for the examples in this chapter.
Note there are many R packages for fitting AR(p) (and ARMA(p,q) for that matter) models. If you are only interested in univariate data with no observation error in the data then you probably want to look into the \texttt{arima()} function included in base R and into R packages that specialize in fitting ARMA models to univariate data. The forecast package in R is a good place to start but others can be found on the CRAN task view: Time Series Analysis.

18.2 MAR(2) models

A MAR(2) model is a lag-2 MAR model, aka a multivariate autoregressive process with no observation process (no SS part). A MAR(2) model is written

\[
x_t' = B_1 x_{t-1} + B_2 x_{t-2} + u_t + w_t, \quad \text{where } w_t \sim \text{MVN}(0, Q) \quad (18.1)
\]

We rewrite this as MARSS(1) by defining \( x^t = \begin{bmatrix} x_t' \\ x_{t-1}' \end{bmatrix} \):

\[
\begin{bmatrix}
  x_t' \\
  x_{t-1}'
\end{bmatrix} =
\begin{bmatrix}
  B_1 & B_2 \\
  I_m & 0
\end{bmatrix}
\begin{bmatrix}
  x_{t-1}' \\
  x_{t-2}'
\end{bmatrix}
+ \begin{bmatrix}
  u_t \\
  0
\end{bmatrix}
+ \begin{bmatrix}
  w_t \\
  0
\end{bmatrix}
\sim \text{MVN} \left( \begin{bmatrix} 0 \\ 0 \end{bmatrix}, \begin{bmatrix} 0 & 0 \\ 0 & 0 \end{bmatrix} \right)
\quad (18.2)
\]

Our observations are of \( x_t \) only, so our observation model is

\[
y_t = \begin{bmatrix} 1 & 0 \end{bmatrix} \begin{bmatrix} x_t' \\ x_{t-1}' \end{bmatrix}
\quad (18.3)
\]

18.2.1 Example of AR(2): univariate data

Here is an example of fitting a univariate AR(2) model written in MARSS(1) form. First, let’s generate some simulated AR(2) data from this AR(2) process:

\[
x_t = -1.5x_{t-1} - 0.75x_{t-2} + w_t, \quad \text{where } w_t \sim N(0, 1) \quad (18.4)
\]

\begin{verbatim}
TT=50
true.2=c(r=0,b1=-1.5,b2=-0.75,q=1)
temp=arima.sim(n=TT,list(ar=true.2[2:3]),sd=sqrt(true.2[4]))
sim.ar2=matrix(temp,nrow=1)
\end{verbatim}

Next, we set up the model list for an AR(2) model written in MARSS(1) form (refer to Equation 18.2 and 18.3):

\begin{verbatim}
Z=matrix(c(1,0),1,2)
B=matrix(list("b1",1,"b2",0),2,2)
U=matrix(0,2,1)
Q=matrix(list("q",0,0,0),2,2)
\end{verbatim}
\[ A= \text{matrix}(0,1,1) \]
\[ R= \text{matrix}(0,1,1) \]
\[ \mu= \text{matrix}(\text{sim.ar2}[2:1],2,1) \]
\[ V= \text{matrix}(0,2,2) \]
\[ \text{model.list.2} = \text{list}(Z=Z,B=B,U=U,Q=Q,A=A,R=R,x0=\mu,V0=V,tinitx=0) \]

Notice that we do not estimate \( \mu \). We will fit our model to the data (\( y \)) starting at \( t = 3 \). Because \( R = 0 \), this means \( E[\mathbf{x}_t|y_t] = \mathbf{x}_t \), and \( \mathbf{x}_{t0} = \begin{bmatrix} y_2 \\ y_1 \end{bmatrix} \).

Note \( E[\mathbf{x}_t|y_{t-1}] = \mathbf{x}_{t-1} \neq y_t \) so we do not use \( \mathbf{x}_{t0} \) as our initial \( \mathbf{x} \).

Then we can then estimate the \( b_1 \) and \( b_2 \) parameters for the AR(2) process.

\[ \text{ar2=MARSS(sim.ar2[3:TT],model=model.list.2)} \]

Success! abstol and log-log tests passed at 16 iterations.

Alert: \( \text{conv.test.slope.tol is 0.5} \).

Test with smaller values (<0.1) to ensure convergence.

\underline{MARSS fit is}

\underline{Estimation method: kem}

Convergence test: \( \text{conv.test.slope.tol = 0.5, abstol = 0.001} \)

Estimation converged in 16 iterations.

Log-likelihood: -63.02523

AIC: 132.0505     AICc: 132.5959

  Estimate
  B.b1  -1.582
  B.b2  -0.777
  Q.q   0.809

Initial states (x0) defined at t=0

Standard errors have not been calculated.

Use \text{MARSSparamCIs} to compute CIs and bias estimates.

Comparison to the true values shows the estimates are close:

\[ \text{print(cbind(true=true.2[2:4],estimates=coef(ar2,type="vector")))} \]

true estimates
b1  -1.50  -1.5816137
b2  -0.75  -0.7767462
q   1.00   0.8091055

Missing values in the data are fine. Let’s make half the data missing being careful that the first data point does not get categorized as missing:

\[ \text{gappy.data=sim.ar2[3:TT]} \]
\[ \text{gappy.data[floor(runif(TT/2,2,TT))]=} \text{NA} \]
\[ \text{ar2.gappy=MARSS(gappy.data,model=model.list.2)} \]
And the estimates are still close:

```r
cbind(true=true.2[2:4],
    estimates.no.miss=coef(ar2,type="vector"),
    estimates.w.miss=coef(ar2.gappy,type="vector"))
```

<table>
<thead>
<tr>
<th></th>
<th>true estimates.no.miss</th>
<th>estimates.w.miss</th>
</tr>
</thead>
<tbody>
<tr>
<td>b1</td>
<td>-1.50</td>
<td>-1.5549387</td>
</tr>
<tr>
<td>b2</td>
<td>-0.75</td>
<td>-0.7463820</td>
</tr>
<tr>
<td>q</td>
<td>1.00</td>
<td>0.8868251</td>
</tr>
</tbody>
</table>

By the way, there are much better and faster functions in R for fitting univariate AR models (no observation error). The MARSS package is really for fitting to multivariate data with observation error not AR-p models. For example, here is how you would fit the AR(2) model using the base `arima()` function:

```r
arima(gappy.data,order=c(2,0,0),include.mean=FALSE)
```

Call: `arima(x = gappy.data, order = c(2, 0, 0), include.mean = FALSE)`

Coefficients:

<table>
<thead>
<tr>
<th></th>
<th>ar1</th>
<th>ar2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>-1.5674</td>
<td>-0.7494</td>
</tr>
</tbody>
</table>

s.e. 0.1033 0.1015

sigma^2 estimated as 0.9428: log likelihood = -51.81, aic = 109.62

The estimates will be different because `arima` sets \( x_0 \) as coming from the stationary distribution. That is a non-linear constraint that MARSS cannot handle.

The assumption that \( x_0 \) comes from the stationary distribution is fine if the initial \( x \) indeed comes from the stationary distribution, but if the initial \( x \) is well outside the stationary distribution the the estimates will be incorrect.

\( TT=50 \)

```r
true.2=c(r=0,b1=-1.5,b2=-0.75,q=1)
sim.ar2.ns=rep(NA,TT)
for(i in 3:TT)
    sim.ar2.ns[i]=true.2[2]*sim.ar2.ns[i-1]+true.2[3]*sim.ar2.ns[i-2]+rnorm(1,0,sqrt(true.2[4]))
model.list.3 = model.list.2
model.list.3$x0 = matrix(sim.ar2.ns[2:1],2,1)
ar3.marss=MARSS(sim.ar2.ns[3:TT],model=model.list.3,silent=TRUE)
ar3.arima=arima(sim.ar2.ns[3:TT],order=c(2,0,0),include.mean=FALSE)
print(cbind(true=true.2[2:4],
    estimates.marss=coef(ar3.marss,type="vector"),
    estimates.arima=c(coef(ar3.arima,type="vector"),ar3.arima$sigma2)))
```

<table>
<thead>
<tr>
<th></th>
<th>true estimates.marss estimates.arima</th>
</tr>
</thead>
<tbody>
<tr>
<td>b1</td>
<td>-1.5816137</td>
</tr>
<tr>
<td>b2</td>
<td>-0.7767462</td>
</tr>
<tr>
<td>q</td>
<td>0.8091055</td>
</tr>
</tbody>
</table>
18.2.2 Example of MAR(2): multivariate data

Here we show an example of fitting a MAR(2) model. Let’s make some simulated data of two realizations of the same AR(2) process:

```r
tT=50
true.2=c(r=0,b1=-1.5,b2=-0.75,q=1)
temp1=arima.sim(n=tT,list(ar=true.2[c("b1","b2")]),sd=sqrt(true.2["q"]))
temp2=arima.sim(n=tT,list(ar=true.2[c("b1","b2")]),sd=sqrt(true.2["q"]))
sim.mar2=rbind(temp1,temp2)
```

Although these are independent time series, we want to fit with a MAR(2) model to allow us to use both datasets together to estimate the AR(2) parameters. We need to set up the model list for the multivariate model (Equation 18.2 and 18.3):

```r
Z=matrix(c(1,0,0,0,0,0,0,0),2,4)
B1=matrix(list(0),2,2); diag(B1)="b1"
B2=matrix(list(0),2,2); diag(B2)="b2"
B=matrix(list(0),4,4)
U=matrix(0,4,1)
Q=matrix(list(0),4,4)
Q[1,1]="q"; Q[2,2]="q"
A=matrix(0,2,1)
R=matrix(0,2,2)
pi=matrix(c(sim.mar2[,2],sim.mar2[,1]),4,1)
V=matrix(0,4,4)
model.list.2m=list(Z=Z,B=B,U=U,Q=Q,A=A,R=R,x0=pi,V0=V,tinitx=1)
```

Notice the form of the Z matrix:

```
[1,] 1 0 0 0
[2,] 0 1 0 0
```

It is a $2 \times 2$ identity matrix followed by a $2 \times 2$ all-zero matrix. The B matrix is composed of B1 and B2 which are diagonal matrices with b1 and b2 respectively on the diagonal.

```
[1,] "b1" 0 "b2" 0
```
We fit the model as usual:

\[ \text{mar2} = \text{MARSS}(\text{sim.mar2}[2:TT], \text{model}=\text{model.list.2m}) \]

Then we can compare how using two time series improves the fit versus using only one alone:

\[ \text{model.list.2$x0=matrix(sim.mar2[1,2:1],2,1)} \]
\[ \text{mar2a} = \text{MARSS}(\text{sim.mar2}[1,2:TT], \text{model}=\text{model.list.2}) \]
\[ \text{model.list.2$x0=matrix(sim.mar2[2,2:1],2,1)} \]
\[ \text{mar2b} = \text{MARSS}(\text{sim.mar2}[2,2:TT], \text{model}=\text{model.list.2}) \]

<table>
<thead>
<tr>
<th></th>
<th>true</th>
<th>est.mar2</th>
<th>est.mar2a</th>
<th>est.mar2b</th>
</tr>
</thead>
<tbody>
<tr>
<td>b1</td>
<td>-1.50</td>
<td>-1.4206301</td>
<td>-0.7209367</td>
<td>-1.3506409</td>
</tr>
<tr>
<td>b2</td>
<td>-0.75</td>
<td>-0.7642604</td>
<td>-0.3954671</td>
<td>-0.6953739</td>
</tr>
<tr>
<td>q</td>
<td>1.00</td>
<td>0.8986820</td>
<td>3.2098084</td>
<td>1.5552943</td>
</tr>
</tbody>
</table>

18.3 MAR(p) models

A MAR(p) model is similar to a MAR(2) except it has lags up to time \( p \):

\[
\mathbf{x}_t' = \mathbf{B}_1 \mathbf{x}_{t-1}' + \mathbf{B}_2 \mathbf{x}_{t-2}' + \cdots + \mathbf{B}_p \mathbf{x}_{t-p}' + \mathbf{u}' + \mathbf{w}_t', \quad \text{where} \quad \mathbf{w}_t' \sim \text{MVN}(0, \mathbf{Q}')
\]

where

\[
\begin{bmatrix}
\mathbf{x}_t' \\
\mathbf{x}_{t-1}' \\
\vdots \\
\mathbf{x}_{t-p}'
\end{bmatrix} = 
\begin{bmatrix}
\mathbf{B}_1 & \mathbf{B}_2 & \cdots & \mathbf{B}_p \\
\mathbf{I}_m & 0 & \cdots & 0 \\
0 & \mathbf{I}_m & \cdots & 0 \\
0 & 0 & \ddots & \vdots \\
0 & 0 & \cdots & \mathbf{I}_m \\
0 & 0 & \cdots & 0
\end{bmatrix}
\begin{bmatrix}
\mathbf{u}' \\
0 \\
\vdots \\
0 \\
0 \\
0 \\
\end{bmatrix},
\]

\( \mathbf{Q}' = \begin{bmatrix} Q' & 0 & \cdots & 0 \\ 0 & 0 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & 0 \end{bmatrix} \)

Here's an example of fitting a univariate AR(3) in MARSS(1) form. We need more data to estimate an AR(3), so use 100 time steps.

\( TT=100 \)
\( \text{true.3} = c(r=0,b1=-1.5,b2=-0.75,b3=.05,q=1) \)
\( \text{temp3=arima.sim(n=TT,list(ar=true.3[c("b1","b2","b3")]),sd=sqrt(true.3["q")])} \)
\( \text{sim.ar3=matrix(temp3,nrow=1)} \)

We set up the model list for the AR(3) in MARSS(1) form as follows:

\( Z=\text{matrix}(c(1,0,0),1,3) \)
\( B=\text{matrix}(\text{list("b1"},1,0,"b2",0,1,"b3",0,0),3,3) \)
\( U=\text{matrix}(0,3,1) \)
We can easily fit MAR(p) processes observed with error using MARSS(p) models, but the difficulty is specifying the initial state condition. \( \pi \equiv x_1 \) and thus involves \( x_1, x_0, \ldots \). However, we do not know the variance covariance structure for these consecutive \( x \). Specifying \( \Lambda = 0 \) and estimating \( \pi \) often causes the EM algorithm to run into numerical problems. But if we have an abundance of data, fixing \( \pi \) might not overly affect the \( B \) and \( Q \) estimates.

Here is an example where we set \( \pi \) to the mean of the data and set \( \Lambda \) to zero. Why not set \( \Lambda \) equal to a diagonal matrix with large values on the diagonal to approximate a vague prior? The temporally consecutive initial states are definitely not independent. A diagonal matrix would imply independence which will conflict with the process model and means our model would be fundamentally inconsistent with the data (and that usually has bad consequences for estimation).

Create some simulated data:

```r
t=1000 #set long
tue.2ss=c(r=.5,b1=-1.5,b2=-0.75,q=.1)
temp=arima.sim(n=tT,list(ar=tue.2ss[c("b1","b2")]),sd=sqrt(tue.2ss["q"]))
sim.ar=matrix(temp,nrow=1)
noise=rnorm(tT-1,0,sqrt(tue.2ss["r"]))
noisy.data=sim.ar[2:tT]+noise
```

Set up the model list for the model in MARSS(1) form:
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\[ Z = \text{matrix}(c(1,0),1,2) \]
\[ B = \text{matrix}(\text{list}("b1",1,"b2",0),2,2) \]
\[ U = \text{matrix}(0,2,1) \]
\[ Q = \text{matrix}(\text{list}("q",0,0,0),2,2) \]
\[ A = \text{matrix}(0,1,1) \]
\[ R = \text{matrix}("r") \]
\[ V = \text{matrix}(0,2,2) \]
\[ \pi = \text{matrix}(\text{mean(noisy.data)},2,1) \]
\[ \text{model.list.2ss} = \text{list}(Z=Z,B=B,U=U,Q=Q,A=A,R=R,x0=\pi,V0=V,tinitx=0) \]

Fit as usual:

\[ \text{ar2ss} = \text{MARSS(noisy.data, model=model.list.2ss)} \]

Success! abstol and log-log tests passed at 101 iterations.
Alert: conv.test.slope.tol is 0.5.
Test with smaller values (<0.1) to ensure convergence.

MARSS fit is
Estimation method: kem
Convergence test: conv.test.slope.tol = 0.5, abstol = 0.001
Estimation converged in 101 iterations.
Log-likelihood: -1368.796
AIC: 2745.592  AICc: 2745.632

<table>
<thead>
<tr>
<th>Estimate</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>R.r</td>
<td>0.477</td>
</tr>
<tr>
<td>B.b1</td>
<td>-1.414</td>
</tr>
<tr>
<td>B.b2</td>
<td>-0.685</td>
</tr>
<tr>
<td>Q.q</td>
<td>0.140</td>
</tr>
</tbody>
</table>

Initial states (x0) defined at t=0

Standard errors have not been calculated.
Use MARSSparamCIs to compute CIs and bias estimates.

We can compare the results to modeling the data as if there is no observation error, and we see that that assumption leads to poor B estimates:

\[ \text{model.list.2ss.bad} = \text{model.list.2ss} \]
\[ \# \text{set R to zero in this model} \]
\[ \text{model.list.2ss.bad}$R = \text{matrix}(0) \]

Fit using the model with R set to 0:

\[ \text{ar2ss2} = \text{MARSS(noisy.data, model=model.list.2ss.bad)} \]

Compare results
The middle column are the estimates assuming that the data have no observation error and the right column are our estimates with the observation error estimated. Clearly, assuming no observation error when it is present has negative consequences for the $B$ and $Q$ estimates.

By the way, there is a straight-forward way to deal with the measurement error if you are working with univariate ARMA models and you are only interested in the AR parameters (the $b$'s). Inclusion of measurement error leads to additional MA components up to lag $p$ (Staudenmayer and Buonaccorsi, 2005). This means that if you are fitting a AR(p) model with measurement error, you can fit a ARMA(p,p) and the measurement error will be absorbed in the $p$ MA components. For the example above, we could estimate the AR parameters for our AR(2) data with measurement error by fitting a ARMA(p,p) model. Here’s how we could do that using R’s `arima()` function:

```r
arima(noisy.data,order=c(2,0,2),include.mean=FALSE)
```

Call:
`arima(x = noisy.data, order = c(2, 0, 2), include.mean = FALSE)`

Coefficients:
```
      ar1      ar2      ma1      ma2
-1.4448  -0.6961   0.9504   0.3428
```

s.e. 0.0593 0.0427 0.0686 0.0482

`sigma^2` estimated as 0.9069: log likelihood = -1368.99, aic = 2747.99

Accounting for the measurement error definitely improves the estimates for the AR component.

### 18.5 Discussion

Although both MARSS(1) and ARMA(p,p) approaches can be used to deal with AR(p) processes (univariate data) observed with error, our simulations suggest that the MARSS(1) approach is less biased and more precise (Figure 18.1) and that the EM algorithm is working better for this problem. The performance of different approaches depends greatly on the underlying model.
Fig. 18.1. Comparison of the AR parameter estimates using different approaches to model ARSS(2) data (univariate AR(2) data observed with error). Results are from 200 simulations of AR(2) data with 100 time steps. Results are shown for the $b_1$ and $b_2$ parameters of the AR process fit with 1) AR(2) model with no correction for measurement error, 2) MARSS(1) model fit via the EM optimization, 3) MARSS(1) model fit via BFGS optimization, 4) ARMA(2,2) model fit with the R `arima` function, and 5) AR(2) model fit 2nd differencing with the R `arima` function. The “x” shows the mean of the simulations and the bar in the boxplot is the median. The true values are shown with the dashed horizontal line. The $\sigma^2$ for the AR process was 0.1 and the $\sigma^2$ for the measurement error was 0.5. The $b_1$ parameters was -1.5, and $b_2$ was -0.75.

We chose AR parameters where both ARMA(p,p) and MARSS(1) approaches work. If we used, for example, $b_1 = 0.8$ and $b_2 = -0.2$, the ARMA(2,2) gives $b_1$ estimates close to 0 (i.e. wrong) while the MARSS(1) EM approach gives estimates close to the truth (though rather variable). One would want to also check REML approaches for fitting the ARMA(p,p) models since REML has been found to be less biased than ML estimation for this class (Cheang and Reinsel 2000; Ives et al. 2010). Ives et al. 2010 has R code for REML estimation of ARMA(p,q) models in their appendix.
For multivariate data observed with error, especially multivariate data without a one-to-one relationship to the underlying autoregressive process, an explicit MARSS model will need to be used rather than an ARMA(p,p) model. The time steps required for good parameter estimates are likely to be large; in our simulations, we used 100 for a AR(3) and 1000 for a ARSS(2). Thorough simulation testing should be conducted to determine if the data available are sufficient to allow estimation of the $B$ terms at multiple lags.
Part IV

Appendices
A

Package MARSS: Warnings and errors

The following are brief descriptions of the warning and error message you may see and what they mean (or might mean). More warning information can be found by typing `MARSSinfo()` at the command line.

**B update is outside the unit circle**

If you are estimating $B$, then if the absolute value of all the eigenvalues of $B$ are less than 1, the system is stationary (meaning the $X$'s have some multivariate distribution that does not change over time). In this case, we say that $B$ is within the unit circle. A pure univariate random walk for example would have $B=1$ and it is not stationary. The distribution of $X$ for the pure random walk has a variance that increases with time. If on the other hand $|B|<1$, you have an Ornstein-Uhlenbeck process and is stationary, with a stationary variance of $Q/(1-B^2)$ (note $B$ is a scalar here because in this example $X$ is univariate). If any of the eigenvalues (real part) are greater than 1, then the system will “explode”—it rapidly diverges.

In the EM algorithm, there is nothing to force $B$ to be on or within the unit circle (real part of the eigenvalues less than or equal to 1). It is possible at one of the EM iterations the $B$ update will be outside the unit circle. The problem is that if you get too far outside the unit circle, the algorithm becomes numerically unstable since small errors are magnified by the “explosive” $B$ term. If you see the ‘B outside the unit circle’ warning, it is fine as long as it is temporary and the log-likelihood does not start decreasing (you will see a separate warning if that happens).

If you do see $B$ outside the unit circle and the log-likelihood decreases, then it probably means that you have poorly specified the model somehow. An easy way to do this is to poorly specify the initial conditions, $\pi$ and $\Lambda$. If, say, you try to specify a vague prior on $x_0$ (or $x_1$) with $\pi$ equal to zero and $\Lambda$ equal to a diagonal matrix with a large variance on the diagonal, you will likely run into trouble if $B$ has off-diagonal terms. The reason is that by
specifying that \( \Lambda \) is diagonal, you specified that the individual \( X \)'s in \( X_0 \) are independent, yet if \( B \) has off-diagonal terms, the stationary distribution of \( X_1 \) is NOT independent. If you force the diagonal terms on \( \Lambda \) to be big enough, you can force the maximum-likelihood estimate of \( B \) to be outside the unit circle since this is the only way to account for \( X_0 \) independent and \( X_1 \) highly correlated.

The problem is that you will not know the stationary distribution of the \( X \)'s (from which \( X_0 \) was presumably drawn) without knowing the parameters you are trying to estimate. One approach is the estimate both \( \pi \) and \( \Lambda \) by setting \( x_0=\"unconstrained\" \) and \( V_0=\"unconstrained\" \) in the model specification. Estimating both \( \pi \) and \( \Lambda \) cannot be done robustly for all MARSS models, and in general, one probably wants to specify the model in such a way as to fix one or both of these. Another, more robust approach, is to treat \( x_1 \) as fixed but unknown (instead of \( x_0 \)). You do this by setting \( \text{model}$tinitx=1 \), so that \( \pi \) refers to \( t = 1 \) not \( t = 0 \). Then estimate \( \pi \) and fix \( \Lambda = 0 \). This eliminates \( \Lambda \) from the model and often eliminates the problems with prior specification—as the expense of \( m \) more parameters. Note, when you set \( \Lambda = 0 \), \( \Lambda \) is truly eliminated from the model by MARSS; the likelihood function is different, so do not expect \( \Lambda = 0 \) and \( \Lambda \sim 0 \) to have the same likelihood under all conditions.

**Warning! Reached maxit before parameters converged**

The maximum number of EM iterations is set by \( \text{control$maxit} \). If you get this warning, it means that one of the parameters or log-likelihood had not yet reached the convergence stopping criteria before \( maxit \) was reached. There are many situations where you might want to set \( \text{control$maxit} \) lower than the value needed to reach convergence. For example, if you are using the EM algorithm to produce initial values for a different algorithm (like a Bayesian MCMC algorithm or a Newton method) then you can set \( maxit \) low, say 20 or 50.

**Stopped at iter=xx in MARSSkem() because numerical errors were generated in MARSSkf**

This means the Kalman filter/smoother algorithm became unstable and most likely one of the variances became ill-conditioned. When that happens the inverses of those matrices are poor, and you will start to get negative values on the diagonals of your variance-covariance matrices. Once that happens, the inverse of that var-covariance matrix produces an error. If you get this error, turn on tracing with \( \text{control$trace}=1 \). This will store the error messages so you can see what is going on. It may be that you have specified the model in such a way that some of the variances are being forced very close to 0, which makes the var-covariance matrix ill-conditioned. The output from the MARSS call will be the parameter values just before the error occurred.
Warning: the xyz parameter value has not converged

The algorithm checks whether the log-likelihood and each individual parameter has converged. If a parameter has not converged, you can try upping `control$maxit` and see if it converges. If you set, `maxit` high, but the parameter is still not converging, then it suggests that one of the variance parameters is so small that the EM update steps for that parameter are tiny. For example, as $Q$ goes to zero, the update steps for $u$ go to zero. As $\Lambda$ goes to zero, the update steps for $\pi$ go to zero. The first thing to do is to reflect on whether you are inadvertently specifying the model in such a way that one of the variances is forced to zero. For example, if the total variance in $X$ is 0.1 and you fix $R = 0.2$ then $Q$ must go to zero. The second thing to do is to try using a Newton algorithm, using your last EM values as the initial conditions for the Newton algorithm. The initial values are set using the `inits` argument for the `MARSS()` function.

MARSSkem: The soln became unstable and logLik DROPPED

This is a more serious error as in the EM algorithm, the log-likelihood should never drop. The first thing to do is check if you have specified a bizarre model or data, inadvertently. Plot the data you are trying to fit. Often, this error arises when a user has inadvertently scrambled their data order during a demeaning or variance-standardization step. Second, check the model you are trying to fit. Use `test=MARSS(data, model=xyz, fit=FALSE)` and then `summary(test$model)`. This shows you what MARSS() thinks your model is. You may be trying to fit an illogical model.

If those checks looks good, then pass `control$trace=1` into the `MARSS()` call. This will report a fuller set of warnings. Look if the error “B is outside the unit circle” appears. If so, you are probably specifying a strange B matrix. Are you forcing the B matrix to be outside the unit circle (eigenvalues $> 1$)? If so, you need to rethink your B matrix contraints. If you do not see that error, look at `test$iter.record$logLik`. If the log-likelihood is steadily dropping (at each iteration) or drops by large amounts (much larger than the machine precision), that is bad and means that the EM algorithm did not work. If however the log-likelihood is just fluctuating by small amounts about some steady value, that is ok as it means that the values converged but the parameters are such that there are slight numerical fluctuations. Try passing `control$safe=TRUE` in the `MARSS()` call. This can sometimes help as it inserts a call to the Kalman filter after each individual parameter update.
Stopped at iter=xx in MARSSkem: solution became unstable. R (or Q) update is not positive definite

First check if you have specified an illegally constrained variance-covariance matrix. For example, if the variances (diagonal) are constrained to be equal, you cannot specify the covariances (off-diagonals) as unequal. Or if you specify that some of the covariances are equal, you cannot specify the variances as all unequal. These are illegal constraints on a variance-covariance matrix from a statistical perspective (nothing to do with MARSS specifically).

This could also be due to numerical instability as $B$ leaves the unit circle or one of the variance matrix becomes ill-conditioned. Try turning on tracing with `control$trace=1` and turn on safe with `control$safe=TRUE`. This will print out the error warnings at each parameter update step. Then consider whether you have inadvertently specified the model in such a way as to force this behavior in the $B$ parameter.

You might also get this error if you inadvertently specified an improper structure for $R$ or $Q$. For example, if you used $R=\text{diag}(c(1,1,"r"))$ with the intent of specifying a diagonal matrix with fixed variance 1 at $R[1,1]$ and $R[2,2]$ and an estimated $R[3,3]$, you would have actually specified a character matrix with "0" on the off-diagonals and $c("1","1","r")$ on the diagonal. `MARSS()` interprets all elements in quotes as names of parameters to be estimated. Thus it will estimate one off-diagonal covariance and two diagonal variances. That happens to put illegal constraints on estimation of a variance-covariance matrix having nothing to do with `MARSS()` but with estimation of variance-covariance matrices in general.

iter=xx MARSSkf: logLik computation is becoming unstable. Condition num. of Sigma[t=1] = Inf and of R = Inf.

This means, generally, that $V_0$ is very small, say 0, and $R$ is very small and very close to zero.

Warning: setting diagonal to 0 blocked at iter=X. logLik was lower in attempt to set 0 diagonals on X

This is a warning not an error. What is happening is that one of the variances (in $Q$ or $R$) is getting small and the EM algorithm is attempting to set the value to 0 (because `control$degen.allow=TRUE`). But when it tried to do this, the new likelihood with the variance equal to 0 was lower and the variance was not set to 0.

A model with a variance minuscule and a model with the same variance equal to 0 are not the same model. In the first, a stochastic process with
small variance exists but in the second, the analogous process is deterministic. And in the first case, you can get a situation where the likelihood term \( L(x|\text{mean}=\mu, \text{sigma}=0) \) appears. That term will be infinite when \( x=\mu \). So in the model with variance minuscule, you will get very large likelihood values as the variance term gets smaller and smaller. In the analogous model with that variance set to 0, that likelihood term does not appear so the likelihood does not go to infinity.

This is not an error nor pathological behavior; the models are fundamentally different. Nonetheless, this will pose a dilemma when you want to chose the best model based on maximum likelihood. The model with minuscule variance will have infinite likelihood but the same behavior as the one with variance 0. In our experience, this dilemma arises when one has a lot of missing data near the beginning of the time series and is affected by how you specify the prior on the initial state. Try setting the prior at \( t=0 \) versus \( t=1 \). Try using a diffuse prior. You absolutely want to compare estimates using the BFGS and EM algorithms in this case, because the different algorithms differ in their ability to find the maximum in this strange case. Neither is uniformly better or worse. It seems to depend on which variance (\( Q \) or \( R \)) is going to zero.

**Warning: kf returned error at iter=\( X \) in attempt to set 0 diagonals for \( X \)**

This is a warning that the EM algorithm tried to set one of the diagonals of element \( X \) to 0 because allow.degen is TRUE and element \( X \) is going to zero. However when this was tried, the Kalman filter returned an error. Typically, this happens when both \( R \) and \( Q \) elements are both trying to be set to 0. If the maximum-likelihood estimate is that both \( R \) and \( Q \) are zero, it probably means that your MARSS model is not a very good description of the data.

**Warning: At iter=\( X \) attempt to set 0 diagonals for \( R \) blocked for elements where corresponding rows of \( A \) or \( Z \) are not fixed.**

You have `control$degen.allow=TRUE` and one of the \( R \) diagonal elements is getting very small. MARSS attempts to set these \( R \) elements to 0, but if row \( i \) of \( R \) is 0, then the corresponding \( i \) rows of \( A \) and \( Z \) must be fixed. This is for the EM algorithm. It might work with the BFGS algorithm, or might spit out garbage without telling you. Always be a suspect, when the EM and BFGS behavior is different. That is a good sign that something is wrong with how your model describes the data. It’s not a problem with the algorithms per se; rather for certain pathological models, the algorithms behave differently from each other.
Stopped at \texttt{iter=}X in MARSSkem. XYZ is not invertible.

There are a series of checks in MARSS that check if matrix inversions are possible before doing the inversion. These errors crop up most often when $Q$ or $R$ are getting very small. At some point, they can get so small that inversions become unstable. If this error is given, then the output will be the last parameter estimates before the error. Try setting \texttt{control$allow.degen=}FALSE. Sometimes the error occurs when a diagonal element of $Q$ or $R$ is being set to 0. You will also have to set \texttt{control$maxit} to something smaller because the EM algorithm will not stop since the problematic diagonal element will walk slowly and inexorably to 0.
B

Package MARSS: Object structures

Model objects: class marssMODEL

Objects of class ‘marssMODEL’ specify Multivariate Autoregressive State Space (MARSS) models. The model component of an ML estimation object (class marssMLE; see below) is a marssMODEL object. These objects have the following components:

- **data**: An optional matrix (not dataframe), in which each row is a time series (time across columns).
- **fixed**: A list with 8 matrices $Z$, $A$, $R$, $B$, $U$, $Q$, $x0$, $V0$, specifying which elements of each parameter are fixed.
- **free**: A list with 8 matrices $Z$, $A$, $R$, $B$, $U$, $Q$, $x0$, $V0$, specifying which elements of each parameter are to be estimated.
- **M**: An array of dim $n \times n \times T$ (an $n \times n$ missing values matrix for each time point). Each matrix is diagonal with 0 at the $i,i$ value if the $i$-th value of $y$ is missing, and 1 otherwise.
- **miss.value**: Deprecated. Replace missing values with NAs before passing to MARSS.

The matrices in **fixed** and **free** work as pairs to specify the fixed and free elements for each model parameter. The dimensions for **fixed** and **free** matrices are as follows, where $n$ is the number of observation time series and $m$ is the number of state processes:

- $Z$: $n \times m$
- $B$: $m \times m$
- $U$: $m \times 1$
- $Q$: $m \times m$
- $A$: $n \times 1$
- $R$: $n \times n$
- $x0$: $m \times 1$
- $V0$: $m \times m$
Use `is.marssMODEL()` to check whether an marssMODEL object is correctly specified. The MARSS package includes an `as.marxssMODEL()` method to convert objects of class popWrap (see next section) to objects of class marssMODEL.

**MARSSInputs**

All the user inputs to a MARSS() call are put into a list and then passed to a function called MARSS.form() (where form is the text specified by the form argument, e.g. MARSS.marss()). This function is used to create the marssMODEL object and then MARSScheckinputs() is called to error check the other arguments.

- **data** A matrix (not dataframe) of observations (rows) × time (columns).
- **model** The specification is form dependent. For the default marxss form, the inputs are a list with up to 14 elements Z, A, R, B, U, Q, x0, V0, C, c, D, d, tinitx, diffuse
- **inits** A list with up to 10 matrices Z, A, R, B, U, Q, x0, V0, C, D specifying initial values for parameters. Dimensions are given in the class ‘marssMODEL’ section.
- **miss.value** Deprecated. Specifies missing value representation (default is NA).
- **form** The form to use to interpret the ‘model’ argument and create the marssMODEL object.
- **control** List of estimation options. These are method dependent.

**ML estimation objects: class marssMLE**

Objects of class marssMLE specify maximum-likelihood estimation for a MARSS model, both before and after fitting. A minimal marssMLE object contains components model, start and control, which must be present for estimation by functions like MARSSkem().

- **model** MARSS model specification (an object of class ‘marssMODEL’).
- **start** List with 7 matrices A, R, B, U, Q, x0, V0, specifying initial values for parameters. Dimensions are given in the class marssMODEL section.
- **control** A list specifying estimation options. For method="kem", these are
  - **minit** Minimum number of iterations in the maximization algorithm.
  - **maxit** Maximum number of iterations in the maximization algorithm.
  - **abstol** Optional tolerance for log-likelihood change. If log-likelihood decreases less than this amount relative to the previous iteration, the EM algorithm exits.
trace A positive integer. If not zero, a record will be created of each variable the maximization iterations. The information recorded depends on the maximization method.

safe If TRUE, \texttt{MARSSkem()} will rerun \texttt{MARSSkf()} after each individual parameter update rather than only after all parameters are updated.

silent Suppresses printing of progress bar and convergence information.

\texttt{MARSSkem()} appends the following components to the \texttt{marssMLE}’ object:

method A string specifying the estimation method (‘kem’ for estimation by \texttt{MARSSkem()}).

par A list with 8 matrices of estimated parameter values \(Z\), \(A\), \(R\), \(B\), \(U\), \(Q\), \(x_0\), \(V_0\). If there are fixed elements in the matrices, the corresponding elements in \$par are set to the fixed values.

kf A list containing Kalman filter/smoother output. See Chapter 2

numIter Number of iterations required for convergence.

convergence Convergence status.

logLik the exact Log-likelihood. See Section 3.4

errors any error messages

iter.record record of the parameter values at each iteration (if \texttt{control$trace=1})

Several functions append additional components to the ‘marssMLE’ object passed in. These include:

\texttt{MARSSaic} Appends AIC, AICc, AICbb, AICbp, depending on the AIC flavors requested.

\texttt{MARSShessian} Appends Hessian, gradient, parMean and parSigma.

\texttt{MARSSparamCIs} Appends par.se, par.bias, par.upCI and par.lowCI.
Model specification in the core functions

Most users will not directly work with the core functions nor build marssMODEL objects from scratch. Instead, they will interact with the core functions via the function `MARSS()` described in Chapter 4. With the `MARSS()` function, the user specifies the model structure with matrices or text strings (“diagonal”, “unconstrained”, etc.) and `MARSS()` builds the marssMODEL object. However, a basic understanding of the structure of marssMODEL objects is useful if one wants to interact directly with the core functions.

C.1 The fixed and free components of the model parameters

In a marssMODEL object, each parameter is written in vec form and specified by the equation of the form \( f + D\beta \) as in Equation 77 in Holmes (2012). \( f \) is the fixed matrix, \( D \) is the free matrix and \( \beta \) is the column vector of parameters. In a marssMODEL object, the fixed list has the \( f \) for each parameter matrix, the free list has the \( D \) matrix for each parameter matrix, and par in the marssMLE object has the \( \beta \) column vector of estimated parameters.

C.2 Examples

C.2.1 \( Q \) is a diagonal matrix with one variance value

In this case, there is only one value on the diagonal and the off-diagonals are 0. Thus there is only one estimated parameter and the fixed values are all 0.

\[
Q = \begin{bmatrix}
\alpha & 0 & 0 \\
0 & \alpha & 0 \\
0 & 0 & \alpha
\end{bmatrix}
\]

`fixed$Q` is
262  C Model specification in the core functions

\[ f = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} \]

par$Q$ is

\[ \beta = ["alpha"] \]

and free$Q$ is

\[ D = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \\ 0 \end{bmatrix} \]

Notice that $f + D\beta$ is the vec of $Q$.

C.2.2 $Q$ is a diagonal matrix with unique variance values

\[ Q = \begin{bmatrix} \alpha_1 & 0 & 0 \\ 0 & \alpha_2 & 0 \\ 0 & 0 & \alpha_3 \end{bmatrix} \]

The fixed matrix is the same with all 0s, but the par and free matrices change. par$Q$ is

\[ \beta = ["alpha1", "alpha2", "alpha3"] \]

and free$Q$ is

\[ D = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 1 \end{bmatrix} \]
C.2.3 Q has one variance and one covariance

\[
Q = \begin{bmatrix}
\alpha & \beta & \beta \\
\beta & \alpha & \beta \\
\beta & \beta & \alpha
\end{bmatrix}
\]

The fixed vector is still the same, all zero. \texttt{par}\$Q is

\[
\beta = \begin{bmatrix}
"alpha" \\
"beta"
\end{bmatrix}
\]

and \texttt{free}\$Q is

\[
D = \begin{bmatrix}
1 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 1
\end{bmatrix}
\]

C.2.4 Q is unconstrained

Since \( Q \) is a variance-covariance matrix, it must be symmetric across the diagonal.

\[
Q = \begin{bmatrix}
\alpha_1 & \beta_1 & \beta_2 \\
\beta_1 & \alpha_2 & \beta_3 \\
\beta_2 & \beta_3 & \alpha_3
\end{bmatrix}
\]

There are no fixed values in \( Q \) so \( f \) is still all zero. \texttt{par}\$Q is

\[
\beta = \begin{bmatrix}
"alpha1" \\
"beta1" \\
"beta2" \\
"alpha2" \\
"beta3" \\
"alpha3"
\end{bmatrix}
\]

and \texttt{free}\$Q is

\[
D = \begin{bmatrix}
1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1
\end{bmatrix}
\]
C.2.5 Q is fixed

For example,

\[ Q = \begin{bmatrix}
0.1 & 0 & 0 \\
0 & 0.1 & 0 \\
0 & 0 & 0.1
\end{bmatrix} \]

The fixed $Q$ matrix is

\[ \begin{bmatrix}
0.1 \\
0 \\
0 \\
0 \\
0.1 \\
0 \\
0 \\
0 \\
0.1
\end{bmatrix} \]

There are no estimated parameters so the free matrix is $9 \times 0$ and the par matrix is $0 \times 1$.

C.3 Limits on the model forms that can be fit

The main limitation is that one must specify a model that has only one solution. The core functions will allow you to attempt to fit an improper model (one with multiple solutions). If you do this accidentally, it may or may not be obvious that you have a problem. The estimation functions may chug along happily and return some solution. Careful thought about your model structure and the structure of the estimated parameter matrices will help you determine if your model is under-constrained and unsolvable.
Textbooks and articles that use MARSS modeling for population modeling

**Textbooks Describing the Estimation of Process and Non-process Variance**

There are many textbooks on Kalman filtering and estimation of state-space models. The following are a sample of books on state-space modeling that we have found especially helpful.


**Maximum-likelihood papers**

This is just a sample of the papers from the population modeling literature.


Bayesian papers

This is a sample of the papers from the population modeling and animal tracking literature.


References


