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An introduction to R for dynamic models in biology Last compile: April 17, 2019

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Preface

These notes for computer labs accompany the textbook *Dynamic Models in Biology* (Princeton University Press 2006), but they can also be used as a standalone introduction to R for simulating dynamic models of biological systems. They are based in part on course materials by former TAs Colleen Webb, Jonathan Rowell and Daniel Fink (then at Cornell University), Lou Gross (University of Tennessee) and Paul Fackler (NC State University), and on the book *Getting Started with Matlab* by Rudra Pratap (Oxford University Press). We also have drawn on the documentation built into R.

The current home for these notes is www.cam.cornell.edu/~dmb/DMBsupplements.html, a web page for the textbook that we maintain ourselves. If that fails, an up-to-date link should be in the book's listing at the publisher (www.pupress.princeton.edu). Parallel notes and script files for MATLAB are also available at those sites. This document was originally written at a Windows PC and may sometimes refer to Windowsspecific aspects of R. We will be happy to make changes as these are brought to our attention.

Sections 1-7 are an autotutorial introduction to basic R programming (we generally cover them in two or three 2-hour lab sessions, depending on how much previous experience students have had). Those sections contain many sample calculations. It is important to do them yourselves – *type them in at your keyboard and see what happens on your screen* – to get the feel of working in R . Exercises in the middle of a section should be done *immediately* when you get to them, and make sure that you have them right before moving on. Exercises at the ends of sections may be more appropriate as homework exercises. He marks the end of an exercise.

Subsequent sections link to the textbook in fairly obvious ways. For example, section 8 on matrix computations goes with Chapter 2 on matrix models for structured populations, and section 11 goes with the Markov Chains section of Chapter 3. Some exercises in these sections are "warmups" for exercises in the textbook, such as simple examples of agent-based models as a warmup for agent-based simulation of infectious disease dynamics.

What is R?

R is an object-oriented scripting language that combines

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aov, anova	Analysis of variance or deviance
lm, glm	Linear and generalized linear models
gam, gamm	Generalized additive models and mixed models (in mgcv package)
nls	Fit nonlinear models by least-squares (in nls package)
lmer,glmer,nlmer	Linear and nonlinear mixed-effects models (in lme4 package)
nonparametric regression	Numerous functions in libraries including stats (smoothing
	splines, loess, kernel), mgcv, fields, KernSmooth, logspline, sm
boot	Package: functions for bootstrap estimates of precision and signif-
	icance
multiv	Package: multivariate analysis
survival	Package: survival analysis
tree	Package: tree-based regression

Table 1: A few of the functions and add-on packages in **R** for statistical modeling and data analysis. There are **many** more, but you will have to learn about them somewhere else.

- the S statistical programming language developed by John Chambers (Chambers and Hastie 1988, Chambers 1998, Venables and Ripley 2000).
- a user interface with a few basic menus and extensive help facilities.
- an enormous set of functions for dynamic and statistical modeling.
- graphics functions for visualizing data and model output.

R is an open-source project available at www.cran.r-project.org for Windows, OS X and several flavors of Linux. Originally a research project in statistical computing (Ihaka and Gentlemen 1996) it is now managed by an international development team and is widely used by statisticians and biologists. R mostly follows version 3 of the S language, but some packages use version 4 features. *These notes refer only to version 3 of* S. We also limit ourselves to graphics functions in the base graphics package, rather than the more advanced packages grid, lattice, ggplot2, and do not use the Hadleyverse (google it).

RStudio provides a GUI for R that's similar to the one for MATLAB. It's available free at rstudio.org/download/desktop. Many people like it. We don't.

Using the help system is an essential R skill. If you know that the plot() function plots things, but you don't know exactly how to make a particular kind of plot, type ?plot at the R command prompt. If plot doesn't do what you want, try ??plot to get a list of other functions that have "plot" as part of their name maybe one of them will do what you need. Once you get the habit of using it, and get familiar with how R help pages are structured, the Help system is enormously helpful.

Also, there is tons of information on the Web at message-board sites like StackOverflow. If you get an error message that you don't understand, a useful strategy is to copy-paste that message from the console into the search bar in your favorite web browser, and add [R] in front of the message before you do the search. Web searches can also find answers to lots of "how do I..." questions.

Statistics in R

Some of the important functions and libraries (collections of functions) for data analysis and statistical modeling are summarized in Table 1. The book by Venables and Ripley (2002) gives a good practical overview, and a list of available libraries and their contents is available at CRAN (www.cran.r-project.org, click on Package sources). For the most part, we are not concerned here with that side of R.

1 Interactive calculations

Launching R opens the **console** window. This has a few basic menus at the top, whose names and content are OS-dependent; check them out on your own. The console window is also where you enter commands for R to execute *interactively*, meaning that the command is executed and the result is displayed as soon as you hit the Enter key. For example, at the command prompt >, type in 2+2 and hit Enter; you will see

```
> 2+2
[1] 4
```

To do anything complicated, the results from calculations have to be stored in variables. For example, type a=2+2; a at the prompt and you see

```
> a=2+2; a [1] 4
```

The variable a has been created, and assigned the value 4. The semicolon allows two or more commands to be typed on a single line; the second of these (a by itself) tells R to print out the value of a. By default, a variable created this way is a vector (an ordered list of numbers); in this case a is a vector length 1, which acts just like a number.

Variable names in R must begin with a letter, and followed by alphanumeric characters (i.e., numbers and letters). Long names can be broken up using a period, as in very.long.variable.number.3, or by the underscore character as in make_iteration_matrix. However, you cannot use blank space as a separator in variable names; new population size=3 will get you an error message. It is currently fashionable to use "camel case", as in newPopulationSize=3 - variable names start with a lower-case letter, and upper case is used to mark a separation. R is case sensitive: Abc and abc are **not** the same variable.

Exercise 1.1 Here are some variable names that cannot be used in R; explain why: cell maximum size; 4min; site#7 . \clubsuit

Calculations are done with variables as if they were numbers. R uses +, -, *, /, and ^ for addition, subtraction, multiplication, division and exponentiation, respectively. For example enter

```
> x=5; y=2; z1=x*y; z2=x/y; z3=x^y; z2; z3
```

and you should see

[1] 2.5

[1] 25

Even though the variable values for x, y were not displayed, R "remembers" that values have been assigned to them. Type > x; y to display the values.

If you mis-enter a command, it can be edited instead of starting again from scratch. The \(\gamma\) key recalls previous commands to the prompt. For example, you can bring back the next-to-last command and edit it to

```
> x=5 y=2 z1=x*y z2=x/y z3=x^y z2 z3
```

so that commands are not separated by a semicolon. Then press Enter, and you will get an error message.

You can do several operations in one calculation, such as

```
> A=3; C=(A+2*sqrt(A))/(A+5*sqrt(A)); C
[1] 0.5543706
```

5

Table 2: Some of the built-in mathematical functions in **R**. You can get a more complete list from the Help system: ?Arithmetic for simple, ?log for logarithmic, ?sin for trigonometric, and ?Special for special functions.

abs(x)	absolute value
cos(x), $sin(x)$, $tan(x)$	cosine, sine, tangent of angle x in radians
exp(x)	exponential function
log(x)	natural (base-e) logarithm
log 10(x)	common (base-10) logarithm
sqrt(x)	square root

The parentheses are specifying the order of operations. The command

$$> C=A+2*sqrt(A)/A+5*sqrt(A)$$

gets a different result – the same as

$$> C=A + 2*(sqrt(A)/A) + 5*sqrt(A).$$

The default order of operations is: (1) Exponentiation, (2) multiplication and division, (3) addition and subtraction.

```
> b = 12-4/2<sup>3</sup> gives 12 - 4/8 = 12 - 0.5 = 11.5

> b = (12-4)/2<sup>3</sup> gives 8/8 = 1

> b = -1<sup>2</sup> gives -(1<sup>2</sup>) = -1

> b = (-1)<sup>2</sup> gives 1
```

In complicated expressions it's best to use parentheses to specify explicitly what you want, such as $> b = 12 - (4/(2^3))$ or at least $> b = 12 - 4/(2^3)$.

R also has many **built-in mathematical functions** that operate on variables (see Table 2).

Exercise 1.2: Have R compute the values of

(a)
$$\frac{2^7}{2^7-1}$$
 and compare it with $\left(1-\frac{1}{2^7}\right)^{-1}$

- (b) $\sin(0.25) + \cos(2\pi/7)$ (pi is pre-defined in R to equal the mathematical constant π).
- (c) $\frac{2^7}{2^7-1} + \sin(0.25)$, using cut-and-paste to assemble parts of your past commands.

Exercise 1.3: Do an Apropos on sin via the Help menu, to see what it does. Next do ??sin

and see what that does (answer: ??sin pulls up all help pages that include 'sin' anywhere in their title or text. Apropos just searches function names for 'sin'.) Note: on a Mac you need to do Apropos at the command line, as in apropos('sin').

Exercise 1.4 Use the Help system to find out what the hist function does, by typing ?hist at the command prompt. Prove that you have succeeded by doing the following: use the command y=rnorm(5000) to generate a vector of 5000 random numbers with a Normal distribution, and then use hist to plot a histogram of the values in y with about 20 bins. Why did we say "about 20" rather than "exactly 20"? Find the answer in the Help page for hist.

2 First interactive session: linear regression & a population growth model

To get a feel for working in R we'll use linear regression to fit a population growth model to data. Below are some data on the growth of a laboratory population of the green alga *Chlorella vulgaris*. This experiment was run during the system-design phase of the study reported by Fussmann et al. (2000).

```
time(days) = 0 1 2 3 4 5 6 7 8 9 10 12 14 21 25 32 33
10<sup>5</sup> cells/ml: 1.1 1.4 4.1 5.5 5.4 10.7 6.0 22.0 19.8 26.7 31.4 30.9 27.1 40.2 36.1 36.8 31.6
```

To analyze these data in R, first enter them¹ as numerical *vectors*:

```
tvals=c(0,1,2,3,4,5,6,7,8,9,10,12,14,21,25,32,33);
cvals=c(1.1,1.4,4.1,5.5,5.4,10.7,6.0,22.0,19.8,26.7,31.4,30.9,27.1,40.2,36.1,36.8,31.6);
```

The function c() *combines* the individual numbers into a vector; enter tvals in the console to see this.

To see a histogram of the population counts enter > hist(cvals) which opens a graphics window and displays the histogram. There are **many** other built-in statistics functions, for example mean(cvals) gets you the mean, sd(cvals) and median(cvals) return the standard deviation and median, respectively.

To see how the algal population increased over time,

```
> plot(tvals,cvals)
```

creates a plot. By default, only the points are plotted. To connect the points, you can ask for a plot of type "l" for "line",

```
> plot(tvals,cvals,type="1")
```

or of types "b" or "o" (for "both" and "overlay") to plot the points and connecting lines. **Try all of these right now.**

The population seems to exhibit sigmoid growth, up to some maximum density. A simple model often fitted to such growth curves is

$$x(t+1) = Rx(t)(1 - bx(t)) \tag{1}$$

which is sometimes called the "discrete logistic" model. To generate solutions of this model we need three things: values for R and b, and a starting density x(0). How can we get these? Well, when x is small, equation (1) says $x(t+1) \approx Rx(t)$, and the solution to x(t+1) = Rx(t) is exponential growth:

$$x(t) = x(0)R^t \quad \Rightarrow \quad \log x(t) = \log x(0) + t \log R. \tag{2}$$

So if we fit a straight line to the initial part of the data, the slope will be an estimate of $\log R$, and the intercept will be an estimate of $\log x(0)$. So let's plot the log-transformed data:

```
> plot(tvals,log(cvals),xlab="Time (days)",ylab="log Chlorella density");
```

Leave this plot on the screen – we'll be adding to it later.

My eye suggested using the first 5 data points to fit our straight line for the initial exponential growth phase. To fit the line by linear regression, we use the lm() function to create a linear model:

```
> y=log(cvals[1:5]); x=tvals[1:5];
> fit = lm(y~ x)
```

¹To save yourself some time, you can cut-and paste the lines below from the PDF file into the R console.

The effect of [1:5] in the first line is to "pull out" only the first 5 entries of cvals and tvals; we'll learn more about this later. The second line produces no output but it has created fit as an **object**, i.e. a multi-part data structure holding the results of the linear regression analysis. Unlike a typical statistics package, R rarely produces summary output automatically. Data analysis in R is done by creating a model, and then giving additional commands to extract information or display results graphically.

The command summary(fit) prints a summary of the results in the console window. A model object like fit is set up so that the summary function can "see" that fit was created by 1m and produces an appropriate summary for an 1m object:

```
Call: lm(formula = y \sim x)
Residuals:
      1
              2
                      3
                              4
                                      5
-0.04138 -0.25527 0.36420 0.20292 -0.27048
Coefficients:
          Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.1367 0.2505 0.546 0.6233
            0.4550
                      0.1023 4.449 0.0211 *
Χ
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.3234 on 3 degrees of freedom
Multiple R-Squared: 0.8684, Adjusted R-squared: 0.8245
F-statistic: 19.79 on 1 and 3 DF, p-value: 0.02113
```

If you've had a statistics class, all of the above should make sense to you. But for right now, the only things we need are the estimated intercept (0.1367) and slope (0.4550).

The regression line can be added to the data plot by another function that takes fit as its input:

```
> abline(fit)
```

This gives the plot shown in Figure 1.

You can also "interrogate" fit directly. Type > names(fit) to get a list of the components of fit.

Components of an object are extracted using the "\$" symbol. For example >fit\$coefficients yields the regression coefficients

```
(Intercept) x 0.1366926 0.4550453
```

We can use those to get two of the estimates that we need:

```
x0=exp(fit$coef[1]); R=exp(fit$coef[2]);
```

Note that "coefficients" was abbreviated but R figured out what you meant, because no other component of fit started with coef.

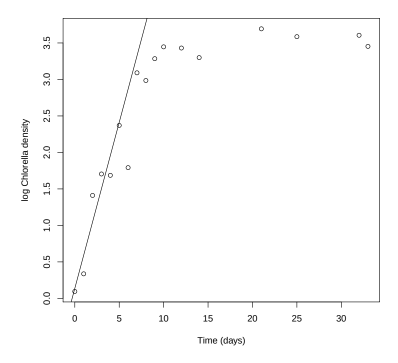


Figure 1: Graphical summary of regression analysis for log Chlorella density as a function of time in days.

We still need to estimate b. Looking at the data, the value of x(t) in the model should stop growing when x(t) is about 35. That is, x(t)=35 then we should get x(t+1)=35 also. Substituting x(t)=35 and x(t+1)=35 into equation (1), we get 35=35R(1-35b). A bit of algebra turns this into $35b=1-\frac{1}{R}$, so we have

$$> b=(1-1/R)/35$$
; b

Exercise 2.1 1-1=0, and (0/R)/35=0. So why does b=(1-1/R)/35 not give b=0?

Exercise 2.2 When we did plot(tvals, log(cvals)) the x and y axis labels were specified using the xlab and ylab arguments. Do ?plot to figure out how to add a title to the plot using the main argument, so that the plot title is **Algal population growth**. \maltese

Exercise 2.3 Now, figure out how to use the pch and col arguments so that the data are plotted as green squares. You can use ?plot and ?par to find out about pch and col.

★

3 Scripts and data files

Modeling and complicated data analyses are more easily done using *scripts*: a series of commands stored in a text file. The Windows and OS X versions of R include a basic script editor (accessed via the **file** menu on the console), but you can also use an external text editor.

Most scripts for working with models or data follow a simple pattern:

1. "Setup" statements.

- 2. Input some data from a file or the keyboard.
- 3. Carry out the calculations that you want.
- 4. Print the results, graph them, or save them to a file.

For example, a script file might

- 1. Load some packages, or run another script file that creates some functions (more on functions later).
- 2. Read in from a text file the parameter values for a predator-prey model, and the numbers of predators and prey at time t=0.
- 3. Calculate the population sizes at times $t = 1, 2, 3, \dots, T$.
- 4. Graph the results and save the graph as a PDF file for including in your term project.

As a first example, the file **Session1.1.R** has the commands from the interactive regression analysis. **Important:** before working with an example file, create a personal copy in some location on your own computer. We will refer to this location as your *temp folder*. At the end of a lab session you **must** save files onto some transportable medium (e.g. a flash drive) or email them to yourself.

Open *your copy of* of **Session1.1.R**, using **File/Open Script** from the console menu. On the **Edit** menu of the script editor, select **Run All**. This equivalent to copying the whole script and pasting it into the console window, so the commands are all executed and a graph is displayed with the results. The source function runs an entire script file without copying it to the console, e.g.

```
> source("c:/temp/Session1.1.R")
```

Source'ing can also be done in point-and-click fashion via File/Source R code on the console window.

Another important time saver is loading data from a text file. Get copies of **Session1.2.R** and **ChlorellaStart.txt** to see an example of how this can be done. In **ChlorellaStart.txt**, time and algal density are entered as two columns of a data matrix. Then instead of typing these in by hand, the command

```
X=read.table("c:/temp/ChlorellaStart.txt")
```

reads the data file and puts the data values into the variable X. Note that the path to the data file has to be specified; you need to edit the script so that it uses the correct path to your data file. In the console window, do

```
> X
```

to see that X has been loaded into R as a data matrix with two columns. V1 and V2 are "default" variable names that R has assigned to the two columns.

The variables are then extracted from X with the commands

```
tvals=X[,1]; cvals=X[,2];
```

These are shorthand for "tvals = everything in column 1 of X", and "cvals = everything in column 2 of X" (you'll learn about working with matrices later). The rest is the same as before, with some additions that specify axis labels and a title.

Exercise 3.1 Make a copy of Session1.2.R under a new name, and modify the copy so that it plots algal density (without log tranformation) versus time, does linear regression of algal density on time, and plots the data appropriately. As in Session1.2.R, use only the first 5 data points for the linear regression. You should end up with a graph sort of like Figure 2.

Data from Fussmann et al. (2000) system

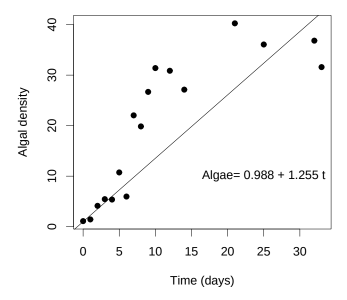


Figure 2: Graphical summary of regression analysis without log transformation.

Exercise 3.2 Run Session1.2.R, then enter the command plot(fit) in the console. R will tell you that it is "Waiting to confirm page change" – help it out by clicking on the plot to do a page change. Next, figure out what just happened by entering ?plot.lm to bring up the Help page for the function plot.lm that carries out a plot command for an object produced by lm. [This is one example of how R uses the fact that data analyses are stored as model objects. fit "knows" what kind of object it is and plot(fit) invokes a function that produces plots suitable for an object produced by lm.] Answer: R produced a series of diagnostic plots exploring whether or not the fitted linear model is a suitable fit to the data. In each of the plots, extreme points (the most likely candidates for "outliers") have been identified according to their sequence in the data set. **

Finally, we want to see how well model (1) fits the data. To see how that can be done, look at **Session1.3.R**. The first sections are familiar: reading in the data and finding the parameter values. Then comes something new: a "for-loop" to solve the model by stepping forward one day at a time, starting from our estimated value at t = 0. We'll learn about this later – for now, just take it for granted:

```
xvals=numeric(34); xvals[1]=x0;
for(j in 2:34) {xvals[j]=R*xvals[j-1]*(1-b*xvals[j-1])}
```

The part to notice now is how to put the data and model solutions onto the same plot:

```
plot(tvals,cvals,type="p",pch=16,cex=1.5, col="green",xlab="Time (days)",
      ylab="Algal density");
points(0:33,xvals,type="o",pch=16,cex=1,col="black",lwd=2);
```

The plot command is familiar with one new wrinkle: the "col" variable sets the plotting color. The points function is just like plot except that it adds new points (or lines) to the most recent plot, rather than making a new plot.

Important NOTE: it is good practice to start each script by clearing R's memory of variables that were created by earlier work, as follows:

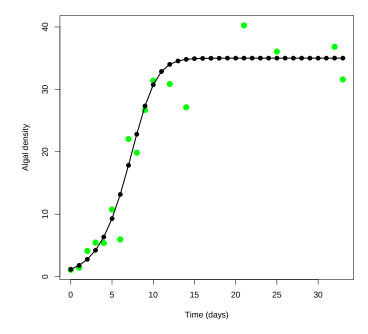


Figure 3: Discrete logistic model (black curve) fitted to experimental data on *Chlorella* growth from the Fussmann et al. (2000) experimental system (green points).

```
rm(list=ls(all=TRUE)); # clear past actions from R's memory
```

This avoids a lot of headaches. A script might run because a variable that it needs was created earlier (not within the current script), and it holds values unrelated to your current work. The script will then run, without any error messages, but the results will be garbage. And, when the TA runs your script, that variable doesn't exist so the script will not run. *You do not want this to happen*. Make sure that *every* script starts with clearing the memory, and that the same is true for *every* exercise if you do several exercises in one script. The best approach is to have a separate file for each exercise; and before you turn it in, close R, restart R, and see if your script runs in a "clean" workspace.

Exercise 3.3 The axes in plots are scaled automatically, but sometimes you want to control them yourself, for example you may want several graphs with exactly the same axes limits. You can control scaling using the xlim and ylim arguments in plot: the general syntax is

$$plot(x,y,xlim=c(x1,x2), ylim=c(y1,y2))$$

For example,

will draw a graph with the x-axis going from 1 to 5, the y-axis from 0.2 to 1.0. The file **ChlorellaGrowth.txt** has data on the maximum algal growth rate r_{max} as a function of light intensity L. Write a script that loads these data into R , and creates a plot of growth rate versus light intensity with the x axis running from 0 to 120, and the y axis running from 1 to 4. Save this script for use in the next exercise.

Exercise 3.4 Several separate graphs can be placed within a single figure by using the par function (short for "parameter") to adjust the layout of the plot. For example the command

divides the plotting area into m rows and n columns. Then as a series of plot commands are used to draw graphs, successive graphs are placed along the top row from left to right, then along the next row, and so on. mfcol=c(m,n) has the same effect except that successive graphs are placed down the first column, then

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down the second column, and so on. par can't be put inside of a plot() command – it has to be separate and it has to come before the plots are drawn.

Save a copy of your script from the last exercise under a new name, and modify it as follows. Use mfcol=c(2,1) to create graphs of (1) growth rate as a function of Light, and (2) log(growth rate) as a function of log(Light), within one figure. Do the same again, using mfcol=c(1,2).

Exercise 3.5 Use ?par to read about other plot control parameters that can be set using par(). Then write a script that draws a 2×2 set of plots, each showing the line y = 5x + 3 with x running from 3 to 8, but with 4 different line styles and 4 different line colors. (Note: setting x=3:8 and y=5*x+3 will create a set of x and y values to draw the line that you need). \maltese

Exercise 3.6 Modify one of your scripts so that at the very end it saves the plot to disk. In Windows, this can be done using savePlot; for other OS's, you can use dev.print. Use the Help system to find out about these functions (e.g., ?savePlot). Note that the argument filename in savePlot, and the argument file in dev.print, can include the path to a folder, for example in Windows you can use

```
filename="c:/temp/MyFigure2".
```

 \mathbb{H}

Note: These are really exercises in using the Help system, with the bonus that you learn about plot. (Let's see, we know plot can graph data points...maybe it can also draw a line to connect the points, or just draw the line and leave out the points. That would be useful. So let's try ?plot and see if it says anything about lines...Hey, it also says that graphical parameters can be given as arguments to plot, so maybe I can set line colors inside the plot command instead of using par all the time....). The Help system is *very* helpful once you get used to it.

4 Vectors

In R, vectors and matrices are just 1- and 2-dimensional rectangular arrays of numbers. Operations with vectors and matrices may seem a bit abstract now, but we need them to do useful things later.

We've already seen two ways to create vectors in R:

1. A command in the console window or a script file listing the values, such as

```
> initialsize=c(1,3,5,7,9,11).
```

2. Using read.table(), for example:

```
initialsize=read.table("c:\\temp\\initialdata.txt")
```

(Note: if the file you're trying to load doesn't exist, this is not going to work!).

Once it has been created, a vector can be used in calculations as if it were a number (more or less)

```
> finalsize=initialsize+1; newsize=sqrt(initialsize); finalsize; newsize;
[1] 2 4 6 8 10 12
[1] 1.000000 1.732051 2.236068 2.645751 3.000000 3.316625
```

Notice that the operations were applied to every entry in the vector. Similarly, commands like initialsize-5, 2*initialsize, initialsize/10 apply subtraction, multiplication, and division to each element of the vector. The same is true for exponentiation,

```
> initialsize^2;
[1] 1 9 25 49 81 121
```

In R the default is to apply functions and operations to vectors in an element by element manner; anything

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else (e.g. matrix multiplication) is done using special notation (discussed below). **NOTE**: this is the *opposite* of MATLAB, where matrix operations are the default and element-by-element requires special notation.

Functions for vector construction

Some of the main functions for creating and working with vectors are listed in Table 3. A set of regularly spaced values can be constructed with the seq function, whose syntax is

```
seq(from, to, by) or seq(from, to, length)
```

The first form generates a vector (from, from+by, from+2*by,...) with the last entry not being larger than to. If a value for by is not specified, its value is assumed to be 1 or -1, depending on whether from or to is larger. The second generates a vector of length evenly-spaced values, running from from to to, for example

```
> seq(1,3,length=6)
[1] 1.0 1.4 1.8 2.2 2.6 3.0
```

There are also two shortcuts for creating vectors with by=1:

```
> 1:8; c(1:8);
[1] 1 2 3 4 5 6 7 8
[1] 1 2 3 4 5 6 7 8
```

A constant vector such as (1,1,1,1) can be created with rep function, whose basic syntax is rep(values,lengths). For example,

```
> rep(3,5)
[1] 3 3 3 3 3
```

created a vector in which the value 3 was repeated 5 times. rep can also be used with a vector of values and their associated lengths, for example

```
> rep( c(3,4),c(2,5) )
[1] 3 3 4 4 4 4 4
```

The value 3 was repeated 2 times, followed by the value 4 repeated 5 times.

R also has many functions for creating vectors of random numbers with various distributions, that are useful in simulating stochastic models. Most of these have a number of **optional arguments**, which means in

seq(from, to, by=1)	Vector of evenly spaced values with specified increment (default = 1)
<pre>seq(from, to, length)</pre>	Vector of evenly spaced values with specified length
c(u,v,)	Combine a set of numbers and/or vectors into a single vector
rep(a,b)	Create vector by repeating elements of a by amounts in b
hist(v)	Histogram plot of value in v
mean(v), var(v), sd(v)	Population mean, variance, standard deviation estimated from values in v
length(v)	Length of vector v
cor(v,w)	Correlation between two vectors

Table 3: Some important R functions for creating and working with vectors. Many of these have other optional arguments; use the help system (e.g. ?cor) for more information. Note that statistical functions such as var regard the values as samples from a population (rather than a list of value for the entire population) and compute an estimate of the population statistic; for example sd(1:3)=1.

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rnorm(n,mean=1,sd=1)	Gaussian distribution(mean=mu, standard deviation=sd)					
runif(n,min=0,max=1)	Uniform distributi	ion on the interva	al (min,max)			
<pre>rbinom(n,size,prob)</pre>	(n, size, prob) Binomial distribution with parameters size=#tri					
	prob=probability	of success p				
rpois(n,lambda)	Poisson distribution	on with mean=la	mbda			
rbeta(n,shape1,shape2)	Beta distribution on the interval $[0,1]$ with shape parameters shape 1,					

Table 4: Some of the main R function for generating vectors of n random numbers. To create random matrices, these vectors can be reshaped using the matrix() function, for example: matrix(rnorm(50*20),50,20) generates a 50×20 matrix of Gaussian(0,1) random numbers.

practice that you can choose to specify their value, or if you don't a default value is assumed. For example, x=rnorm(100) generates 100 random numbers with a Normal (Gaussian) distribution having mean=0, standard deviation=1. But rnorm(100,2,5) yields 100 random numbers from a Gaussian distribution with mean=2, standard deviation=5.

Here, and in the R documentation and help pages, the existence of default values for some arguments of a function is indicated by writing (for example) $\mathbf{rnorm}(\mathbf{n}, \mathbf{mean=0}, \mathbf{sd=1})$. Since no default value is given for n, the user must supply one: $\mathbf{rnorm}()$ gives an error message.

Some of the functions for creating vectors of random numbers are listed in Table 4. Functions to evaluate the corresponding distribution functions are also available. For a listing use the Help system: ?Normal, ?Uniform, ?Lognormal, etc. will list and explain the available functions for each distribution family.

Exercise 4.1 Create a vector $v=(1 \ 5 \ 9 \ 13)$ using seq. Create a vector going from 1 to 5 in increments of 0.2 first by using seq, and then by using a command of the form v=1+b*c(i:j).

Exercise 4.2 Generate a vector of 5000 random numbers from a Gaussian distribution with mean=3, standard deviation=2. Use the functions mean, sd to compute the sample mean and standard deviation of the values in the vector, and hist to visualize the distribution.

Exercise 4.3 The sum of the geometric series $1+r+r^2+r^3+...+r^n$ approaches the limit 1/(1-r) for r<1 as $n\to\infty$. Write a script that computes the sum for r=0.5 and n=10, and write a script that computes the sum. Compare the sum of this vector to the limiting value 1/(1-r). Repeat this for n=50. HINT: look at what v=c(0.5); 2^v ; produces, and figure out why.

Vector addressing

Often it is necessary to extract a specific entry or other part of a vector. This is done using subscripts, for example

```
> q=c(1,3,5,7,9,11); q[3]
[1] 5
```

q[3] extracts the third element in the vector q. You can also access a block of elements using the functions for vector construction, e.g.

```
v=q[2:5]; v
[1] 3 5 7 9
```

This has extracted 2^{nd} through 5^{th} elements in the vector. If you enter v=q[seq(1,5,2)], what will happen?

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Try it and see, and make sure you understand what happened.

Extracted parts of a vector don't have to be regularly spaced. For example

```
> v=q[c(1,2,5)]; v
[1] 1 3 9
```

Addressing is also used to **set specific values within a vector**. For example,

```
> q[1]=12
```

changes the value of the first entry in q while leaving all the rest alone, and

```
> q[c(1,3,5)]=c(22,33,44)
```

changes the 1^{st} , 3^{rd} , and 5^{th} values.

Exercise 4.4 write a one-line command to extract a vector consisting of the second, first, and third elements of q in that order.

Exercise 4.5 Write a script file that computes values of $z=\frac{(x-1)}{(x+1)}$ and $w=\frac{\sin(x^2)}{x^2}$ for $x=1,2,3,\cdots,12$ and plots both of these as a function of x with the points connected by a line.

Vector orientation

You may be wondering if vectors in R are row vectors or column vectors (if you don't know what those are, don't worry: we'll get to it later). The answer is "both and neither". Vectors are printed out as row vectors, but if you use a vector in an operation that succeeds or fails depending on the vector's orientation, R will assume that you want the operation to succeed and will proceed as if the vector has the necessary orientation. For example, R will let you add a vector of length 5 to a 5×1 matrix or to a 1×5 matrix, in either case yielding a matrix of the same dimensions. The fact that R wants you to succeed is both good and bad – good when it saves you needless worry about details, bad when it masks an error that you would rather know about.

5 Matrices

A matrix is a two-dimensional array of numbers. Like vectors, matrices can be created by reading in values from a data file, using the read.table function. Matrices of numbers can also be entered by creating a vector of the matrix entries, and then reshaping them to the desired number of rows and columns using the matrix function. For example

```
> X=matrix(c(1,2,3,4,5,6),2,3)
```

takes the values 1 to 6 and reshapes them into a 2 by 3 matrix.

Note that values in the data vector are put into the matrix column-wise, by default. You can change this by using the optional parameter byrow). For example

<pre>matrix(v,m,n)</pre>	$m \times n$ matrix using the values in v
data.entry(A)	call up a spreadsheet-like interface to edit the values in A
cbind(a,b,c,)	combine compatible objects by binding them along columns
rbind(a,b,c,)	combine compatible objects by binding them along rows
outer(v,w)	"outer product" of vectors v , w: the matrix whose $(i,j)^{th}$ element is
	v[i]*w[j]
diag(n)	$n \times n$ identity matrix
diag(v)	diagonal matrix with vector v on the diagonal
dim(X)	dimensions of matrix X. $dim(X)[1]=\#rows$, $dim(X)[2]=\#columns$
<pre>rowSums(X),colSums(X)</pre>	sum of each row or column of matrix X
<pre>rowMeans(X),colMeans(X)</pre>	mean of each row or column of matrix X
apply(A,MARGIN,FUN)	apply the function FUN to each row of A (if MARGIN=1) or each
	column of A (if MARGIN=2). See ?apply for details and examples.

Table 5: Some important functions for creating and working with matrices. Many of these have additional optional arguments; use the Help system for full details.

```
> A=matrix(1:9,3,3,byrow=T); A
      [,1] [,2] [,3]
[1,] 1 2 3
[2,] 4 5 6
[3,] 7 8 9
```

R will re-cycle through entries in the data vector, if need be, to fill out a matrix of the specified size. So for example matrix(1,50,50) creates a 50×50 matrix of all 1's.

Exercise 5.1 Use a command of the form X=matrix(v,2,4) where v is a data vector, to create the following matrix X

```
[,1] [,2] [,3] [,4]
[1,] 1 1 1 1
[2,] 2 2 2 2
```

Exercise 5.2 Use rnorm and matrix to create a 5×7 matrix of Gaussian random numbers with mean 1 and standard deviation 2.

Another useful function for creating matrices is diag. diag(v,n) creates an $n \times n$ matrix with data vector v on its diagonal. So for example diag(1,5) creates the 5×5 identity matrix, which has 1's on the diagonal and 0 everywhere else.

Finally, one can use the data.entry function. This function can only edit existing matrices, but for example A=matrix(0,3,4); data.entry(A)

will create A as a 3×4 matrix, and then call up a spreadsheet-like interface in which the values can be changed to whatever you need.

5.1 cbind and rbind

If their sizes match, vectors can be combined to form matrices, and matrices can be combined with vectors or matrices to form other matrices. The functions that do this are **cbind** and **rbind**.

cbind binds together columns of two objects. One thing it can do is put vectors together to form a matrix:

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Remember that R interprets vectors as row or column vectors according to what you're doing with them. Here it treats them as column vectors so that columns exist to be bound together. On the other hand,

```
> B=rbind(1:3,4:6); B

[,1] [,2] [,3]

[1,] 1 2 3

[2,] 4 5 6
```

treats them as rows. Now we have two matrices that can be combined.

Exercise 5.3 Verify that rbind(A,B) works, cbind(A,A) works, but cbind(A,B) doesn't. Why not?

5.2 Matrix addressing

Matrix addressing works like vector addressing except that you have to specify both the row and column, or range of rows and columns. For example q=A[2,3] sets q equal to 8, which is the $(2^{nd}$ row, 3^{rd} column) entry of the matrix A, and

```
> A[2,2:3];
[1] 5 8
> B=A[2:3,1:2]; B
      [,1] [,2]
[1,] 2 5
[2,] 3 6
```

There is an easy shortcut to extract entire rows or columns: leave out the limits.

```
> first.row=A[1,]; first.row
[1] 1 4 7
> second.column=A[,2]; second.column;
[1] 4 5 6
```

As with vectors, addressing works in reverse to assign values to matrix entries. For example,

```
A[1,1]=12; A
        [,1] [,2] [,3]
[1,] 12 4 7
[2,] 2 5 8
[3,] 3 6 9
```

The same can be done with blocks, rows, or columns, for example

Exercise 5.4 Use **runif** to construct a 5×5 matrix **B** of random numbers with a uniform distribution between 0 and 1. (a) Extract from it the second row, the second column, and the 3×3 matrix of the values that are not at the margins (i.e. not in the first or last row, or first or last column). (b) Use seq to replace the values in the first row of **B** by 2 5 8 11 14. **

5.3 Matrix operations and matrix-vector multiplication

A numerical function applied to a matrix acts element-by-element.

```
> A=matrix(c(1,4,9,16),2,2); A; sqrt(A);
     [,1] [,2]
[1,] 1 9
[2,] 4 16
     [,1] [,2]
[1,] 1 3
[2,] 2 4
```

The same is true for scalar multiplication and division. Try 2*A, A/3 and see what you get.

If two matrices (or two vectors) are the same size, then you can do element-by-element addition, subtraction, multiplication, division, and exponentiation: (A+B, A-B, A*B, A/B, A B). Matrix × matrix and matrix × vector multiplication (when they are of compatible dimensions) is indicated by the special notation %*%. Remember, **element-by-element is the default in R**. This requires some attention, because R 's eagerness to make things work can sometimes let errors get by without warning. So for example

```
v=1:2; A*v

[,1] [,2]

[1,] 1 9

[2,] 8 32
```

A is a 2×2 matrix, and v is a vector of size 2, so the matrix-vector product Av is legitimate. However, Av should be a vector, not a matrix. Since you (incorrectly) "asked" for element-by-element multiplication, that's what R did, "recycling" through the elements of v when it ran out of entries in v before it ran out of entries in A. What you should have done is

```
> A%*%v
[,1]
[1,] 19
[2,] 36
```

6 Iteration ("Looping")

6.1 For-loops

Loops make it easy to do the same operation over and over again, for example:

- Making population forecasts 1 year ahead, then 2 years ahead, then 3, etc.
- Updating the state of every neuron in a model network based on the inputs it received in the last time interval
- Simulating a biochemical reaction network multiple times with different values for one of the parameters.

There are two kinds of loops in R : for loops, and while loops. A for loop runs for a specified number of steps. These are written as

```
for (var in seq) {
    commands
}
```

Here's an example (in **Loop1.R**):

```
# initial population size
initsize=4;

# create vector to hold results and store initial size
popsize=rep(0,10); popsize[1]=initsize;

# calculate population size at times 2 through 10, write to Command Window
for (j in 2:10) {
    popsize[j]=2*popsize[j-1];
    x=log(popsize[j]);
    cat(j,x,"\n");
}
plot(1:10,popsize,type="l");
```

The first time through the loop, j=2. The line popsize[j]=2*popsize[j-1]; then does popsize[2]=2*popsize[1] so that the solution at time 2 is calculated from the solution at time 1. The second time through the loop, second time through, j=3, so the solution at time 3 is calculated from the solution at time 2. This continues for $j=4,5,6,\cdots,10$, and then the loop is finished R moves on to executing any commands that occur after the end of the loop. The result that the vector popsize holds the values of population size in generations 2 through 10.

Note also the cat function (short for "concatenate") for printing results to the console window. cat converts its arguments to character strings, concatenates them, and then prints them. The " \n " argument is a line-feed character so that each (j, x) pair is put on a separate line.

Several for loops can be nested within each other, which is needed for working with matrices as in the example below. It is important to notice that the second loop is **completely** within the first. Loops must be either **nested** (one completely inside the other) or **sequential** (one starts after the previous one ends).

Type this into a script file and run it; the result should be

```
[,1] [,2] [,3]
[1,] 1 2 3
[2,] 2 4 6
[3,] 3 6 9
```

Line 1 creates A as a matrix of all zeros - this is an easy way to create a matrix of whatever size you need, which can then be filled in with meaningful values as your program runs. Then two nested loops are used to fill in the entries. Line 2 starts a loop over the rows of A, and immediately in line 3 a loop over the columns is started. To fill in the matrix we need to consider all possible values for the pair (row, col). So for row=1, we need to consider col=1,2,3. Then for row=2 we also need to consider col=1,2,3, and the same for row=3. That's what the nested for-loops accomplish. For row=1 (as requested in line 2), the loop in lines 3-5 is executed until it ends. Then we get to the end in line 6, at which point the loop in line 2 moves on to row=2, and so on.

Nested loops also let us automate the process of running a simulation many times, for example with different parameters or to look at the average over many runs of a stochastic model. For example (**Loop2.R**),

```
p=rep(0,5);
                                   (1)
for (init in c(1,5,9)){
                                   (2)
      p[1]=init;
                                   (3)
      for (j in 2:5) {
                                   (4)
         p[j]=2*p[j-1]
                                   (5)
         cat(init,j,p[j],"\n");
                                   (6)
      }
                                   (7)
}
                                   (8)
```

Line 1 creates the vector p. Line 2 starts a loop over initial population sizes Lines 4-7 does one "population growth" simulation Line 8 then closes the loop over initial sizes

The result when you run **Loop2.R** is that the "population growth" calculation is done repeatedly, for a series of values of the initial population size. To make the output a bit nicer we can add some headings as the program runs - run **Loop3.R** and then look at the file to see how the formatting was done.

If this discussion of looping doesn't make sense to you, **stop now and get help**. Loops are essential from here on out.

Exercise 6.1: Imagine that while doing fieldwork in some distant land you and your assistant have picked up a parasite that grows exponentially until treated. Your case is more severe than your assistant's: on return to Ithaca there are 400 of them in you, and only 120 in your assistant. However, your field-hardened immune system is more effective. In your body the number of parasites grows by 10 percent each day, while in your assistant's it increases by 20 percent each day. That is, for you n(j+1) = 1.1n(j) starting from n=400 on day 1, while for your assistant m(j+1) = 1.2m(j) starting from m=120 on day 1.

Write a script file that uses a for-loop to compute the number of parasites in your body and your assistant's over the next 30 days, and draws a single plot of both on log-scale (i.e. log(n(j)) and log(m(j)) versus time for 30 days). The function matplot is useful for putting both n and m on one graph. Save a copy of the script, with the name **Parasite1.R**, for use in future exercises.

NOTE: this can be done without loops, but for the sake of learning, use a for-loop to sequentially compute n(j) and m(j) for j=2 to 31. \maltese

Exercise 6.2: Write a script file that uses for-loops to create the following 5×5 matrix A. Think first: do

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x < y	less than
x > y	greater than
$x \le y$	less than or equal to
x >= y	greater than or equal to
x == y	equal to

Table 6: Some comparison operators in R . Use ?Comparison to learn more.

you want to use nested loops, or sequential?

 \mathbf{H}

Exercise 6.3 Write a script file that uses a for-loop to calculate solutions of the difference equation model

$$N(j+1) = 2e^{-0.2j}N(j)/(1+N(j)), \quad N(1) = 1$$

up to j=13. Remember that in R, e^x is computed using the exp function. Then, have your script plot N(j) versus time j using type="b" in plot, so that N(t) is plotted as points connected by lines. \maltese

7 Branching

7.1 if-else blocks

Sometimes the rules for "what happens next" in a model need to depend on the current values of state variables. The **if** statement lets us do this; the basic format is

```
if(condition) {
    some commands
}else{
    some other commands
}
```

An if block can be set up in other ways, but the layout above, with the }else{ line to separate the two sets of commands, can always be used.

If the "else" is to do nothing, you can leave it out:

```
if(condition) {
    commands
}
```

The conditions controlling a **while** loop are built up from operators that compare two variables (Table 6). These operators return a logical value of TRUE or FALSE. For example, try:

```
> a=1; b=3; c=a<b; d=(a>b); c; d;
```

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The parentheses around (a>b) are optional but can be used to improve readability in script files.

When we compare two vectors or matrices of the same size, or compare a number with a vector or matrix, comparisons are done element-by-element. For example,

```
> x=1:5; b=(x<=3); b
[1] TRUE TRUE TRUE FALSE FALSE</pre>
```

R also does arithmetic on logical values, treating TRUE as 1 and FALSE as 0. So sum(b) returns the value 3, telling us that 3 entries of x satisfied the condition (x <= 3). This is useful for running multiple simulations and seeing how often one outcome occurred rather than another.

Exercise 7.1 Look at and source a copy of Branch1.R to see an if statement which makes the population growth rate depend on the current population size.

More complicated decisions can be built up by nesting one if block within another, i.e. the "other commands" under else can include a second if block. **Branch2.R** uses this method to have population growth tail off in several steps as the population size increases:

```
for (i in 1:50) {
                                      (1)
    if(popnow<250){</pre>
                                      (2)
        popnow=popnow*2;
                                      (3)
    }else{
                                      (4)
        if(popnow<500){</pre>
                                      (5)
            popnow=popnow*1.5
                                      (6)
        }else{
                                      (7)
            popnow=popnow*0.95
                                      (8)
        }
                                      (9)
    }
                                      (10)
    popsize=c(popsize,popnow);
                                      (11)
}
                                      (12)
```

What does this accomplish?

- If popnow is still < 250, then line 3 is executed growth by a factor of 2 occurs. Since the if condition was satisfied, the entire else block (line numbers 5-10 above) isn't looked at; R jumps line (11) and continues from there.
- If popnow is not < 250, R moves on to the else on line 4, and immediately encounters the if on line 5.
- If popnow is < 500 the growth factor of 1.5 applies, and R then jumps to the **end** and continues from there
- If neither of the two if conditions is satisfied, the final else block is executed and population declines by 5% instead of growing.

Exercise 7.2 Starting from a copy of your script Parasite1.R, modify the script so that there is random variation in parasite success, depending on whether or not conditions on a given day are stressful. Specifically, on "bad days" the parasites increase by 10% while on "good days" they are beaten down by your immune system and they go down by 10%, and similarly for your assistant. Specifically,

```
If day j is bad: n(j+1) = 1.1 \times n(j), and m(j+1) = 1.2 \times m(j)
If day j is good: n(j+1) = 0.9 \times n(j), and m(j+1) = 0.8 \times m(j)
```

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Do this by using runif(1) and an if statement to one "toss a coin" for each day: if the random value produced by runif for that day is < 0.5 it's a good day, and otherwise it's bad. Make sure that your script does a new "coin toss" for each day, but that the same toss applies to both you and your assistant. \maltese

7.2 While-loops [advanced]

A while loop lets an iteration continue until some condition is satisfied. For example, we can solve a model until some variable reaches a threshold. The format is

```
while(condition){
     commands
}
```

The loop repeats as long as the condition remains true. **Loop4.R** contains an example similar to the for-loop example; source it to get a graph of population sizes over time. A few things to notice about the program:

- 1. Although the condition in the while loop said while(popsize<1000) the last population value was > 1000. That's because the loop condition is checked **before** the commands in the loop are executed. When the population size was 640 in generation 6, the condition was satisfied so the commands were executed again. After that the population size is 1280, so the loop is finished and the program moves on to statements following the loop.
- 2. Since we don't know in advance how many iterations are needed, we couldn't create in advance a vector to hold all the results. Instead, a vector of results was constructed by starting with the initial population size and appending each new value as it is calculated.
- 3. When the loop ends and we want to plot the results, the "y-values" are popsize, and the x values need to be 0:something. To find "something", the length function is used to find the length of popsize.

Within a while-loop it is often helpful to have a **counter** variable that keeps track of how many times the loop has been executed. In the following code, the counter variable is n:

```
n=1;
while(condition) {
    commands
    n=n+1;
}
```

The result is that n=1 is true while the commands (whatever they are) are being executed for the first time. Afterward n is set to 2, and this remains true during the second time that the commands are executed, and so on. One use of counters is to store a series of results in a vector or matrix: on the n^{th} time through the commands, put the results in the n^{th} entry of the vector, n^{th} row of the matrix, etc.

Exercise 7.3 Write a script file that uses a while-loop to compute the number of parasites in your body and your assistant's so long as you are sicker than your assistant (i.e. so long as n > m) and stops when your assistant is sicker than you. Use a copy of the original, unmodified **Parasite1.R** as your starting point. \maltese

7.3 Logical operators [advanced]

More complicated conditions in if-else structures and while loops can be built by using **logical operators** to combine comparisons:

```
! Negation
```

eigen(A)	eigenvalues and eigenvectors
t(A)	transpose of A
solve(A)	inverse of matrix A
solve(A,B)	solution x of the linear system $Ax = b$ for each column b of the
	matrix B
det(A)	determinant of the matrix A
norm(A)	matrix norm of A (several options)
exp(A)	matrix exponential $\exp(A) = \sum_{n \geqslant 0} \frac{A^n}{n!}$ (in package fMultiVar)
<pre>mtx.exp(A,n)</pre>	efficiently compute powers of a matrix A^n (not element-by-element)
	(in package BioDem)
apply(A,margin,fun)	apply a function fun to the rows (margin=1) or columns (margin=2)
	of matrix A, and return all resulting values
<pre>sapply(v,fun)</pre>	apply a function fun to all elements in vector v, returning a vector of
	values

Table 7: Some important functions for matrix algebra calculations. Many of these have additional optional arguments; use the Help system for full details.

&	&&	AND
Ι		OR

OR is **non-exclusive**, meaning that $x \mid y$ is true if x is true, if y is true, or if both x and y are true. For example: $\Rightarrow a=c(1,2,3,4)$; b=c(1,1,5,5); (a<b)&(a>3); (a<b)&(a>3);

An alternative to (x==y) is the identical function. identical(x,y) returns TRUE if x and y are exactly the same, else FALSE. The difference between these is that if (for example) x and y are vectors (x==y) will return a vector of values for element-by-element comparisons, while identical(x,y) returns a single value: TRUE if each entry in x equals the corresponding entry in y, otherwise FALSE. You can use ?Logical to read more about logical operators.

Exercise 7.4 Use the identical function to construct a one-line command that returns TRUE if each entry in a vector rnorm(5) is positive, and otherwise returns FALSE. **Hint:** rep works on logical variables, so rep(TRUE,5) returns the vector (TRUE,TRUE,TRUE,TRUE,TRUE).

8 Numerical Matrix Algebra

R has functions for matrix-algebra calculations that use "industry standard" numerical libraries (LINPACK, EISPACK, BLAS, LAPACK) libraries. Some of these functions are listed in Table 7.

Some of R 's matrix functions only work on square matrices and will return an error if A is not square, in particular functions for computing eigenvalues and eivenvectors. For the remainder of this section we only consider square matrices.

8.1 Eigenvalues and eigenvectors

Because eigenvalues are so important for studying dynamic models, we will now study eigen in some detail. Recall that if $Aw = \lambda w$ (for A a square matrix, w a nonzero column vector, and λ a real or complex number) then λ is called an *eigenvalue* and w is the corresponding *eigenvector* of A. If v is a row-vector such that $vA = \lambda v$, then v is called a *left eigenvector* of A. The left eigenvalues for a matrix are the same as the (right)

eigenvalues.

We are often most interested in the dominant eigenvalue, which depending on context (discrete versus continuous time models) means either the one with the largest absolute value, or the one with the largest real part.² In R, the abs function computes the absolute value of a real or complex number (and acts element-by-element on vectors and matrices).

eigen returns eigenvalues sorted by absolute value, with the largest first. For example,

As with the output from 1m in our first interactive session (you remember 1m...) vA is a compound *object* composed of two *components*, whose names are values and vectors. The "\$" is used to extract components of a compound object. For example vA\$values is the values part of vA, a vector consisting of the sorted eigenvalues:

```
vA$values
[1] 1.611684e+01 -1.116844e+00 -4.054215e-16
```

As noted above, va\$values[1] is the eigenvalue with the largest absolute value.

The vectors component is a matrix whose columns are the corresponding eigenvectors, sorted in the same order as the eigenvalues. That is,

This calculation verifies the definition of eigenvalues and eigenvectors: $Av_1 = \lambda_1 v_1$, so $A\vec{v}_1 - \lambda_1 \vec{v}_1 = \vec{0}$. The [2,] and [3,] values above are really zeros, but because eigenvalues and eigenvectors are computed numerically, there is a small amount of numerical error.

Exercise 8.1 Verify that the output is also (0,0,0), apart from numerical error, for j=2 and j=3. Explain in words what these calculations show.

Exercise 8.2 Enter the command names (vA) and see what results. From this, infer what the names function does. Enter the command names (A) and infer the meaning of the object NULL in R. Check your guess by using the Help menu on the console window: select R language (standard) and type NULL into the popup window that appears. \maltese

²The general definition of absolute value, which covers both real and complex numbers, is that |a+ib| is the positive square root of (a^2+b^2) , where $i=\sqrt{-1}$.

To get the left eigenvectors of a matrix, we use the fact that the left eigenvectors of a matrix A are the eigenvectors of its transpose, t(A). So

vLA=eigen(t(A))\$vectors

gets you the left eigenvectors of A.

Exercise 8.3 Compute vLA[,j]%*%A-vA\$values[j]*vLA[,j] to verify the last claim about left eigenvectors, for j=1 to 3. \maltese

Eigenvector scalings: For a transition matrix model, the dominant right eigenvector w (i.e. the eigenvector corresponding to the eigenvalue with largest absolute value) is the *stable stage distribution*, so we are most interested in relative proportions. To get those, w=w/sum(w). The dominant left eigenvector \mathbf{v} is the reproductive value, and it is conventional to scale those relative to the reproductive value of a newborn. If newborns are class 1: v=v/v[1].

Exercise 8.4: Write a script file which applies the above to the matrices

$$A = \begin{pmatrix} 1 & -5 & 0 \\ 6 & 4 & 0 \\ 0 & 0 & 2 \end{pmatrix} \quad B = \begin{pmatrix} 0 & 1 & 5 \\ 0.6 & 0 & 0 \\ 0 & 0.4 & 0.9 \end{pmatrix}$$

finding **all** the eigenvalues and then extracting the dominant one and the corresponding left and right eigenvectors. For B, use the scalings defined above. \maltese

8.2 Eigenvalue sensitivities and elasticities

For an $n \times n$ matrix **A** with entries a_{ij} , the sensitivities s_{ij} and elasticities e_{ij} can be computed as

$$s_{ij} = \frac{\partial \lambda}{\partial a_{ij}} = \frac{v_i w_j}{\langle v, w \rangle} \qquad e_{ij} = \frac{a_{ij}}{\lambda} s_{ij}$$
 (3)

where λ is the dominant eigenvalue, \mathbf{v} and \mathbf{w} are dominant left and right eigenvectors, and $\langle v, w \rangle$ is the inner product of \mathbf{v} and \mathbf{w} , computed in R as sum(v*w). So once λ , \mathbf{v} , and \mathbf{w} have been found and stored as variables, it just takes some for-loops to compute the sensitivies and elasticities.

```
vA=eigen(A); lambda=vA$values[1];
w=vA$vectors[,1]; w=w/sum(w);
v=eigen(t(A))$vectors[,1]; v=v/v[1];
vdotw=sum(v*w);
s=A; n=dim(A)[1];
for(i in 1:n) {
    for(j in 1:n) {
        s[i,j]=v[i]*w[j]/vdotw;
    }
}
e=(s*A)/lambda;
```

Note how all the elasticities are computed at once in the last line. In R that kind of "vectorized" calculation is *much* quicker than computing entries one-by-one in a loop. Even better is to use a built-in function that operates at the vector or matrix level. In this case we can use outer to completely eliminate the nested do-loops:

```
vA=eigen(A); lambda=vA$values[1];
w=vA$vectors[,1]; w=w/sum(w);
v=eigen(t(A))$vectors[,1]; v=v/v[1];
s=outer(v,w)/sum(v*w);
e=(s*A)/lambda;
```

Vectorizing code to avoid or minimize loops is an important aspect of efficient R programming.

Exercise 8.5 Construct the transition matrix A, and then find λ , \mathbf{v} , \mathbf{w} for an age-structured model with the following survival and fecundity parameters. Then use those to construct the elasticity matrix for \mathbf{A} .

```
Ages 0-5 are genuine ages with survival probabilities (p_0, p_1, \dots, p_5) = (0.3, 0.4, 0.5, 0.6, 0.6, 0.7)
```

Note that $p_j = a_{j+1,j}$, the chance of surviving from age j to age j+1, for these ages. You can create a vector p with the values above and then use a for-loop to put those values into the right places in A.

Age 6 (or older) are adults with survival 0.9 and fecundity 12.

$$A = \left(\begin{array}{ccccccc} 0 & 0 & 0 & 0 & 0 & 0 & 12 \\ .3 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & .4 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & .5 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & .6 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & .6 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & .7 & .9 \end{array}\right)$$

 \mathbb{H}

Results (with some rounding):

```
\lambda = 1.042
w = (0.6303, 0.1815, 0.0697, 0.0334, 0.0193, 0.0111, 0.055)
v = (1, 3.47, 9.05, 18.85, 32.73, 56.83, 84.59)
```

Exercise 8.6 For the projection matrix

```
A = matrix(c(
0.44, 6.0, 67.4, 183.1,
0.0048, 0.44, 3.04, 8.3,
0, 0.21, 0.042, 0.083,
0, 0.0089, 0.031, 0.014),4,4,byrow=TRUE);
```

use the **which** function with arr.ind=TRUE to find all entries of the elasticity matrix for A that are above the average of the positive entries of the elasticity matrix. HINT: see what A[A>0] returns, and use that construction in computing the average of the positive entries.

8.3 Finding the eigenvalue with largest real part [advanced]

For the Jacobian matrix of a differential equation model, the dominant eigenvalue is the one with the largest real part. To find this, and the associated eigenvector, we need to extract the real parts of the eigenvalues and locate the largest one.

Use the help system – (?complex) – to see the R functions for working with complex numbers. The one we need now is Re, which extracts the real parts of complex numbers.

Use data. entry to create the matrix

$$A = \left(\begin{array}{ccc} 3 & 0 & 0 \\ 2 & 2 & -3 \\ 0 & 3 & 1 \end{array}\right)$$

and you should find that eigen(A)\$values are

```
[1] 1.5+2.95804i 1.5-2.95804i 3.0+0.00000i.
```

The first two are a complex conjugate pair with absolute value 3.316625 (mod(eigen(A)\$values) gets you the absolute values of the eigenvalues), but the third one has the largest real part.

To have R do this for you, we use the which function to find which eigenvalue is dominant (i.e., which has the largest real part). Run the code below, and follow what it's doing; see ?which and ?max if you need to.

```
vA=eigen(A)$values;  # find the eigenvalues
rmax=max(Re(vA));  # find the largest Real part of any eigenvalue
j=which(Re(vA)==rmax);  # which eigenvalue is dominant?
lmax=vA[j];  # dominant eigenvalue
vmax=eigen(A)$vectors[,j];  # dominant eigenvector
```

Exercise 8.7 Try the above on A=diag(c(1,2,3)) and see what you get for j,r,lmax and vmax. Try it again for A=diag(c(1,2,sqrt(as.complex(-9)))) and A=diag(c(3,1,3)), and make sure that you understand what happens in each case.

9 Writing your own functions

A lot of R 's power comes from all its built-in functions, such as plot and lm. A function gets some information from you, and then uses it to do something (like draw a figure) or to compute something (like a set of eigenvalues).

R also lets you write your own functions, which then act like built-in functions (except that they go away when you close an R session). Sometimes this is just a convenience. Using functions makes it easier to see the logical flow of a large program. But sometimes functions are essential, such as when you want to solve a set of differential equations.

R code for defining a function has a very specific structure, but it can go anywhere in a script file. It goes like this example:

```
plotLine=function(x,y) {
    plot(x,y,type="p",cex=1.5)
    fit = lm(y~ x)
    abline(fit,col="red")
}
```

The first line of code specifies the name of the function, plotLine, and its *arguments*: the information it is going to work with (in this case, numeric vectors x and y of the same length). In between the { and the } are the R commands that the function will carry out. To see what this function accomplishes, do the following Exercise right now. It should create a plot like Figure 4.

Exercise 9.1 Open up a new empty script file, copy-paste the code above into your script, and run the script. Then go into the R console window and enter the following code:

```
y = x+0.5*rnorm(10)
plotLine(x,y)
```

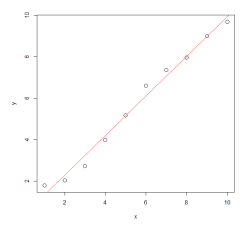


Figure 4: Figure produced by the plotLine function

 \mathbf{H}

Functions can also be written so that they return things that they calculated, like the eigen function does. Here is an example:

```
plotLine2=function(x,y) {
    plot(x,y,type="p",cex=1.5)
    fit = lm(y ~ x)
    abline(fit,col="red")
    return(fit$coef)
}
```

Again, copy-paste this code into a script, run the script, and then in the R console window, enter

```
pars = plotLine2(x,y)
pars
```

You should see that the return statement in plotLine2 resulted in the regression coefficients (namely fit\$coef) getting returned from the function, so that the command pars=plotLine2(x,y) put the coefficients vector into the variable pars.

Exercise 9.2 Write a function whatzit=function(x,y,z) that takes three numbers (or vectors) x, y, z as its arguments, and returns x + y + z. Make sure that it works by running the function like you did with plotLine, and then in the R console window the command whatzit(1,2,3) should give you 6 as the result:

```
> whatzit(1,2,3)
[1] 6
¥
```

Functions can return several different values, by combining them into a multi-part object with named parts. For example,

```
mysquare=function(v,w) {
    q=v^2; r=w^2
    return(list(vSquared=q,wSquared=r))
}
```

You can then extract the components in the R 's usual way.

```
> x=mysquare2(1:4,2:5); names(x);
[1] "vSquared" "wSquared"
> x$vSquared
[1] 1 4 9 16
```

Exercise 9.3 Write a function stats(v) that takes as input a single vector, and returns a list with named components average (mean of the values in the vector), and variance (population variance estimated from the values in the vector, using var). Verify that once you've run the script with the function definition, you get

```
> stats(1:21);

$average

[1] 11

$variance

[1] 38.5
```

10 A simulation project

This section is an optional "capstone" project putting into use the programming skills that have been covered so far. Nothing new about R *per se* is covered in this section.

The exercise is to first write, and then use, a script file that simulates a simple model for density-independent population growth with spatial variation. The model is as follows. The *state variables* are the numbers of individuals in a series of L=20 patches along a line (L stands for "length of the habitat").

1 2	. 3	4							L-1	L	
-----	-----	---	--	--	--	--	--	--	-----	---	--

Let $N_j(t)$ denote the number of individuals in patch j (j = 1, 2, ..., L) at time t (t = 1, 2, 3, ...), and let λ_j be the geometric growth rate in patch j. The *dynamic equations* for this model consist of two steps:

1. Geometric population growth within patches:

$$M_i(t) = \lambda_i N_i(t)$$
 for all j . (4)

2. Dispersal of some individuals between neighboring patches:

$$N_j(t+1) = (1-2d)M_j(t) + dM_{j-1}(t) + dM_{j+1}(t) \quad \text{for } 2 \le j \le L-1$$
 (5)

where 2d is the "dispersal rate". We need special rules for the end patches. For this exercise we assume *reflecting boundaries*: those who exit into the void have the sense to come back. That is, there is no leftward dispersal out of patch 1 and no rightward dispersal out of patch L:

$$N_1(t+1) = (1-d)M_1(t) + dM_2(t)$$

$$N_L(t+1) = (1-d)M_L(t) + dM_{L-1}(t)$$
(6)

Exercise 10.1 As a first step, write a function reflecting that implements the dispersal phase, equations 5 and . That is, the arguments to the function are a pre-dispersal population vector $\mathbf{M}\mathbf{t}$ and the dispersal parameter d, and the function returns the population vector after dispersal has taken place.

```
reflecting <- function(Mt,d) {
    ...
    ... your code to compute Nt1, the population vector N at time t+1
    ...
return(Nt1)
}</pre>
```

This function takes a vector Mt of length L as input, and returns a vector of equal length as output. Equation 5 says that Nt1[2:(L-1)] is the sum of 3 terms: individuals that stay in the same patch, individuals that come in from the left (patches 1 to L-2), and individuals that come in from the right (patches 3 to L). Think about how you can compute each of these 3 terms without using any loops, and then add them up to compute most values in Nt1. After that, you need two more lines to compute the values for patches 1 and L. Note that the code in reflecting has to "figure out" what the value of L is – so what is the R function that tells you the length of a vector?

Exercise 10.2 Using your reflecting function write a script to simulate the model.

- Write your script to <u>start</u> with 5 individuals in each patch at time t=1, <u>iterate</u> the model up to t=100, and <u>graph</u> the log of the total population size (the sum over all patches) over time. Use the following growth rates: $\lambda_j = 0.9$ in the left half of the patches, and $\lambda_j = 1.2$ in the right.
- Write your program so that d and L are parameters, in the sense that the first line of your script file reads d=0.1; L=20;

and the program will still work if these are changed to other values.

Note that this model is not totally different from iterating a matrix population model, in that you start with a founding population at time 1 (described by a vector) and use a loop to compute successive populations at times 2,3,4, and so on. So the script can be structured like the scripts for matrix models. After you specify the values of d and L, and create the vector of λ values, you should create a matrix N with L rows and 100 columns, so that N[j,t] will be $N_j(t)$. The initial population goes into the first column of N, and the model is simulated by doing a loop over time (denoted by k in the code below)

```
for(k in 2:200){ # k is the time index
    ... compute Mt from column k-1 of N, and the lambda values
    ... compute Nt1 using the reflecting function
    ... put Nt1 into N, in the right place
}
```

As in the reflecting function, try to vectorize! Vector/matrix operations are much faster than loops. For example, set things up so that in the for-loop, computing the entire Mt vector is a one-line statement of the form a=b*c.

Exercise 10.3 Use the model (modified as necessary) to ask how the spatial arrangement of good versus bad habitat patches affects the population growth rate. For example, does it matter if all the good sites ($\lambda > 1$) are at one end or in the middle? What if they aren't all in one clump, but are spread out evenly (in some sense) across the entire habitat? Be a theoretician: (a) Patterns will be easiest to see if good sites and bad sites are very different from each other. (b) Patterns will be easiest to see if you come up with a nice way to compare growth rates across different spatial arrangements of patches. (c) Don't confound the experiment by also changing the proportion of good versus bad patches, at the same time you're changing the spatial arrangement.

11 Coin tossing and Markov Chains

The exercises on coin tossing and Markov chains in Chapter 3 of the textbook can be used as the basis for a computer-lab session. For convenience we also include them here. All of the R functions and programming methods required for these exercises have been covered in previous sections, but it is useful to "remember"

- how to generate sets of random uniform and Gaussian random numbers using runif and rnorm.
- how logical operators can be used to convert a vector of numbers into a vector of 1's and 0's according to whether or not a condition holds.
- how to find the places in a vector where the value changes, using logicals and which.

```
>> x = rnorm(20);
>> y = as.numeric(x<0.3);
>> z = which(y[2:20]!=y[1:19])
```

Take a look at x, y, z and make sure you understand why each one is what it is.

11.1 Coin tossing

Exercise 11.1 Experiment with sequences of coin flips produced by a random number generator:

- Generate a vector x of 1000 random numbers uniformly distributed in the unit interval [0, 1].
- Compute and plot a histogram for the values with 10 equal bins of length 0.1. How much variation is there in values of the histogram? Does the histogram make you suspicious that the numbers are not independent and uniformly distributed random numbers?
- Now compute sequences of 10000 and 100000 random numbers uniformly distributed in the unit interval [0, 1], and a histogram for each with ten equal bins. Are your results consistent with the prediction that the range of variation between bins in the histogram is proportional to the square root of the sequence length?

Note: q=hist(runif(1000),10) will plot the first histogram you need, and q\$counts will be a vector of the number in each bin of the histogram, so instead of eyeballing the range of variation you can use range and sd. ★

The theoretical "benchmark" for this experiment, and its relation to coin tossing, goes as follows. Let X_i be 1 if the i^{th} random number falls into the first bin of the histogram, otherwise $X_i = 0$, for $i = 1, 2, 3, \dots n$. So each X_i is a coin toss with P(H) = 0.1. The fraction of tosses that fall in the first bin is then $f = (X_1 + X_2 + \dots + X_n)/n$. Using the basic properties of means and variances, we can derive that E(f) = E(X) and Var(f) = Var(X)/n. The 10 bins of the histogram that you constructed can be regarded as (approximately) 10 repetitions of this experiment, because there's nothing special about bin 1: they all have the same "probability of Heads". So if you compute

```
n=10000; q=hist(runif(n),10); f1=q$counts/n; sqrt(var(f1)) and then compute f2 with n=100,000, you should see (what?)
```

Exercise 11.2 (a) Recall that coin tossing is modeled by the binomial distribution: the probability of k heads in a sequence of n tosses, with p = 0.6 being the probability of heads, is given by

$$c_k(0.6)^k(0.4)^{1000-k}$$
 where $c_k = {1000 \choose k} = \frac{1000!}{k!(1000-k)!}$.

In R , the binomial coefficient
$$\binom{n}{k}$$
 is computed using choose(n,k), e.g. choose(1000,5) gives $\binom{1000}{5}$.

Calculate the probability of k heads for values of k between 500 and 700 in a sequence of 1000 independent tosses. Plot your results with k on the x-axis and the probability of k heads on the y-axis. Comment on the shape of the plot.

- (b) Generate a sequence of 1000 uniformly distributed random numbers r, as in the last exercise, convert them into a sequence of coin tosses in which the probability of heads is 0.6 and the probability of tails is 0.4, and compute the total number of heads in the 1000 coin tosses. Write a script that does this 1000 times (i.e., 1000 repetitions of tossing 1000 coins) and for each repetition stores (in a vector) the total number of heads. Now test the binomial distribution by plotting a histogram of the number of heads obtained in each repetition, and compare the results with the predictions of the binomial distribution.
- (c) Modify the last experiment by doing 10000 repetitions of 100 coin tosses. Comment on the differences you observe between this histogram and the histogram for 1000 repetitions of tossing 1000 coins.

 ★

Tossing multi-sided coins Uniform random numbers can also be used to simulate "coin-toss" experiments where the coin has 3 or more sides. Conceptually this is easy. If there are n sides with probabilities p_1, p_2, \dots, p_n , you generate r=runif(1) and declare that

```
Side 1 occurs if r \le p_1.

Side 2 occurs if p_1 < r \le p_1 + p_2

Side 3 occurs if p_1 + p_2 < r \le p_1 + p_2 + p_3

...
Side n occurs if p_1 + p_2 + \cdots + p_{n-1} < r.
```

To code this compactly in R, note that the outcome depends on the cumulative sums

```
c_1 = p_1, \quad c_2 = p_1 + p_2, \quad c_3 = p_1 + p_2 + p_3, \quad \cdots, c_n = 1.
```

If r is larger than exactly k of these, then the outcome is Side k+1. Cumulative sums of a vector are computed using the function cumsum.

As an example, the following code tosses a 5-sided coin with probabilities given by the vector p defined in the first line:

```
p=c(.3,.1,.3,.1,.2); b=cumsum(p);
side=sum(runif(1)>b)+1
```

To toss the coin repeatedly you can use the sapply function, which applies an arbitrary function to all elements in a vector.

```
side=sapply(runif(1000),FUN=function(x) sum(x>b)+1)
```

In the code above the function to be applied is defined within the call to sapply. More complicated functions can be defined separately, prior to the call, as in the following example:

```
side.choose=function(x) {sum(x>b) +1}
side=sapply(runif(1000), FUN=side.choose)
```

Exercise 11.3 Generate 10000 tosses of a 3-sided coin (with your choice of probabilities), and plot a histogram of the results to verify that your coin is behaving the way it should.

11.2 Markov chains and residence times

The purpose of the following exercises is to generate synthetic data for single channel recordings from finite state Markov chains, and to explore patterns in the "data". Single channel recordings give the times that a Markov chain makes a transition from a closed to an open state or vice versa. The histogram of expected residence times for each state in a Markov chain is exponential, with different mean residence time for different states. To observe this in the simplest case, we again consider coin tossing. The two outcomes, heads or tails, are the different states in this case. Therefore the histogram of residence times for heads and tails should each be exponential. The following steps are taken to compute the residence times:

- Generate sequences of independent coin tosses based on given probabilities.
- Look at the number of *transitions* that occur in each of the sequences (a *transition* is when two successive tosses give different outcomes).
- Calculate the residence times by counting the number of tosses between each transition.

Exercise 11.4 Find the script cointoss.R. This program calculates the residence times of coin tosses by the above methodology, and plots the frequency distribution of residence times in two different ways: a histogram of the number of residence times of each length, and a semi-log plot of the frequency of residence times of each length. Are these results consistent with the prediction that the histograms decreases exponentially? To answer this question, write an R script to make a plot that compares the predicted results with the simulated residence times stored in the vectors hhist and thist.

The predicted result about the histogram for H residence times comes from the following argument: a run of exactly k H's in a row occurs whenever there is a T followed by exactly k H's in a row, followed by another T. The probability of this chain of events is $(1-p)^2p^k$, where p is the probability of getting H on a single toss. So how many runs of exactly k H's in a row should occur in a sequence of nt coin tosses?

Your script should overlay predicted values on the histograms plotted by cointoss.R, and on the semi-log plots of frequences. \maltese

Models for stochastic switching among conformational states of membrane channels are more complicated than a series of independent coin tosses. There are usually more than 2 states, and the transition probabilities are state dependent. Moreover, in measurements some states cannot be distinguished from others. We can observe transitions from an open state to a closed state and vice versa, but transitions between open states (or between closed states) are "invisible".

Here we will simulate data from a Markov chain with 3 states, and then collapse that data to remove the distinction between 2 of the states. We will then analyze the data, as in the previous exercise, to see that the residence time distributions are not consistent with a Markov chain with just two states. We can then use the distributions of residence times in the observations, to reach conclusions about how many states we actually have.

We will consider a membrane that has three states: two closed states C_1 and C_2 , and one open state O. We assume that direct transitions between C_1 and O are impossible, but that the intermediate state C_2 has a shorter residence time than C_1 and O. Here is the transition matrix of a Markov chain we will use to simulate these conditions:

$$\begin{array}{cccc}
C_1 & C_2 & O \\
.98 & .1 & 0 \\ .02 & .7 & .05 \\
0 & .2 & .95
\end{array}$$

$$\begin{array}{cccc}
C_1 \\
C_2 \\
O$$

You can see from the matrix that the probability 0.7 of staying in state C_2 is much smaller than the probability 0.98 of staying in state C_1 or the probability 0.95 of remaining in state C_2 .

Exercise 11.5 Generate a set of 100000 samples from the Markov chain with these transition probabilities. We will label the state C_1 by 1, the state C_2 by 2 and the state O by 3. This can be done by a modification of the method that we used to toss coins with 3 or more sides. The modification is that the probabilities of each side depend on the current state of the membrane:

```
nt = 100000;
A = matrix(c(0.98, 0.10, 0, 0.02, 0.7, 0.05, 0, 0.2, 0.95),3,3,byrow=T);
B = apply(A,2,cumsum); #cumulative sums of each column
A; B;
states=numeric(nt+1); rd=runif(nt);
states[1] = 3; # Start in open state
for(i in 1:nt) {
   b=B[,states[i]]; #cumulative probabilities for current state
   states[i+1]=sum(rd[i]>b)+1 # do the ''coin toss'' based on current state
}
plot(states[1:1000],type="s");
```

Notice the use of apply to compute the cumulative sum of each column of the transition matrix, and type="s" to get a "stairstep" plot of the state transitions (see ?plot).

Exercise 11.6 Compute the eigenvalues and eigenvectors of the matrix A. Compute the total time that your "data" in the vector states spends in each state (use vector operations to do this!) and compare the results with predictions coming from the dominant right eigenvector of A. \maltese

Exercise 11.7 Produce a new vector rstates by taking the vector states that you previously generated (from a 3-state Markov chain) and "reducing" the data so that states 1 and 2 are indistinguishable. You can do that by defining rstates=states, changing every 2 in rstates to 1, then changing every 3 to 2. The states of rstates will be called "closed" (when rstates=1) and "open" (when rstates=2).

Exercise 11.8 Plot the frequency distribution of residence times of the open and closed states in rstates by applying the methods used in cointoss. R. Comment on the shapes of the distributions in each case. Using your knowledge of the transition matrix A, make a prediction about what the residence time distributions of the open states should be, and compare this prediction with the data. Then, show that the residence time distribution for the closed states is not fit well by an exponential distribution.

12 The Hodgkin-Huxley model

The purpose of this section is to develop an understanding of the components of the Hodgkin-Huxley model for the membrane potential of a space-clamped squid giant axon. It goes with the latter part of Chapter 3 in the textbook, and with the **Recommended reading:** Hille, Ion Channels of Excitable Membranes, Chapter 2.

The Hodgkin-Huxley model is the system of differential equations

$$C\frac{dv}{dt} = i - \left[m^{3}hg_{Na}\left(v - v_{Na}\right) + n^{4}g_{K}\left(v - v_{K}\right) + g_{L}\left(v - v_{L}\right)\right]$$

$$\frac{dm}{dt} = 3^{\frac{T-6.3}{10}} \left[(1-m)\Psi\left(\frac{-v-35}{10}\right) - 4m\exp\left(\frac{-v-60}{18}\right)\right]$$

$$\frac{dn}{dt} = 3^{\frac{T-6.3}{10}} \left[0.1\left(1-n\right)\Psi\left(\frac{-v-50}{10}\right) - 0.125n\exp\left(\frac{-v-60}{80}\right)\right]$$

$$\frac{dh}{dt} = 3^{\frac{T-6.3}{10}} \left[0.07(1-h)\exp\left(\frac{-v-60}{20}\right) - \frac{h}{1+\exp(-0.1(v+30))}\right]$$
where
$$\Psi(x) = \frac{x}{\exp(x) - 1}.$$
(7)

The state variables of the model are the membrane potential v and the ion channel gating variables m, n, and h, with time t measured in msec. Parameters are the membrane capacitance C, temperature T, conductances g_{Na}, g_{K}, g_{L} , and reversal potentials v_{Na}, v_{K}, v_{L} . The gating variables represent channel opening probabilities and depend upon the membrane potential. The parameter values used by Hodgkin and Huxley are

g_{Na}	g_K	g_L	v_{Na}	v_K	V_L	T	C
120	36	0.3	55	-72	-49.4011	6.3	1

Most of the data used to derive the equations and estimate the parameter values comes from voltage clamp experiments of the membrane, e.g Figure 2.7 of Hille.

In this set of exercises, we want to see that the model reproduces the voltage clamp data well, and examine some of the approximations and limitations of the parameter estimation. Note that because T=6.3 in the parameter set we are using, the prefactor $3^{\frac{T-6.3}{10}}$ in the equations for m,n and h equals 1, and it can be omitted in all of the exercises. Also, the exercises all consider voltage clamp experiments in which the membrane potential v(t) is externally imposed, and is constant except for instantaneous jumps from one value to another. So for the situation we are considering here, the model (7) reduces to:

$$\frac{dm}{dt} = (1 - m)\Psi\left(\frac{-v - 35}{10}\right) - 4m\exp\left(\frac{-v - 60}{18}\right)$$

$$\frac{dn}{dt} = 0.1 (1 - n)\Psi\left(\frac{-v - 50}{10}\right) - 0.125n\exp\left(\frac{-v - 60}{80}\right)$$

$$\frac{dh}{dt} = 0.07(1 - h)\exp\left(\frac{-v - 60}{20}\right) - \frac{h}{1 + \exp(-0.1(v + 30))}$$
(8)

In voltage clamp, the membrane potential v is held constant, so $\frac{dv}{dt} = 0$ and the first line of (7) gives the following formula for the current:

$$i(t) = m^{3}(t)h(t)g_{Na}(v - v_{Na}) + n^{4}(t)g_{K}(v - v_{K}) + g_{L}(v - v_{L})$$
(9)

When the membrane potential v is constant, the equations for the gating variables m, n, h are first order linear differential equations that can be rewritten in the form

$$\tau_x \frac{dx}{dt} = -(x - x_\infty) \tag{10}$$

where x is m, n or h. The solution to equation (10) is

$$x(t) = x_{\infty} + (x(0) - x_{\infty}) \exp\left(\frac{-t}{\tau_x}\right). \tag{11}$$

Before you start doing these exercises, please read the suggestions for getting started in the following subsection.

Exercise 12.1 Re-write the differential equations for m, n, and h in the form above, thereby obtaining expressions for τ_m , τ_n , τ_h and m_{∞} , n_{∞} , h_{∞} as functions of v.

Exercise 12.2 Write an R script that computes and plots τ_m , τ_n , τ_h and m_∞ , n_∞ , h_∞ as functions of v, for v varying from -100mV to 75mV. You should obtain graphs that look like Figure 2.17 of Hille.

Exercise 12.3 Write an R script to compute and plot as a function of time the current i(t) obtained from voltage clamp experiments in which the membrane is held at a potential of $v_0 = -60 \text{mV}$ and then stepped to a higher potential v_s for 6msec. (While the membrane is at its holding potential -60 mV, the values of m, n, h approach $m_{\infty}(-60), n_{\infty}(-60), h_{\infty}(-60)$. Use these as the initial values m(0), n(0), h(0)). As in Figure 2.7 of Hille, compute the current as a function of time for $v_s = -30, -10, 10, 30, 50, 70, 90$, and plot each of the curves of current on the same graph.

Exercise 12.4 Separate the currents obtained from the voltage clamp experiments by plotting on separate graphs each of the sodium, potassium and leak currents.

Exercise 12.5 Hodgkin and Huxley's 1952 papers explain their choice of the complicated functions in their model, but they had no computers available to analyze their data. In this exercise and the next, we examine procedures for estimating from experimental data the sodium current parameters m_{∞} , h_{∞} , τ_m , τ_h . However, the data that we will use will be generated by the model itself.

As in the previous Exercise, compute and plot the Hodgkin-Huxley sodium current generated by a voltage clamp experiment with a holding potential of v0 = -90 mV and steps to $v_s = -80, -70, -60, -50, -40, -30, -20, -10, 0$. This is your "data". Then using the expression $m^3 h g_{Na}(v-v_{Na})$ for the sodium current, estimate m_{∞} , τ_m , h_{∞} and τ_h as functions of voltage from this simulated data. The most commonly used methods assume that τ_m is much smaller than τ_h , so that the activation variable m reaches its steady state before h changes much. So for times just after the step occurs, you can assume that m is changing but h is holding constant at its initial value. For times long after the step (many multiples of τ_m), you can assume that h is changing but m is holding constant at its steady state value.

Explain the procedures that you used. Some of the parameters are difficult to determine, especially over certain ranges of membrane potential. Why? How do your estimates compare with the values computed in Exercise 12.1?

Exercise 12.6 Challenge: For the parameters that you had difficulty estimating in the previous exercise, design and simulate voltage clamp protocols that help you estimate these parameters better. (See Hille, pp. 44-45.) Describe your protocols and how you estimate the parameters. Plot the currents produced by the model for your new experiments, and give the parameter estimates that you obtain using the additional "data" from your experiments. Further investigation of these procedures is a good topic for a course project.

³Analyzing data that come from a model in order to estimate parameters that you already know may sounds a bit crazy at first, but statisticians do this all the time and many of them are not crazy. Before working with real data, it's a good idea to generate artificial "data" from the model with known parameters, and make sure that your approach to fitting the model allows you to correctly recover the parameters that generated the "data".

12.1 Getting started

We offer here some suggestions for completing the exercises in this section.

Exercise 12.1 Complicated expressions are often built by composing simpler expressions. In any programming language, it helps to introduce intermediate variables. Here, let's look at the gating variable h first. We have

$$\frac{dh}{dt} = 0.07 \exp\left(\frac{-v - 60}{20}\right) (1 - h) - \frac{h}{1 + \exp(-0.1(v + 30))}$$

Introduce the intermediate expressions

$$a_h = 0.07 \exp\left(\frac{-v - 60}{20}\right)$$

and

$$b_h = \frac{1}{1 + \exp(-0.1(v + 30))}$$

Then

$$\frac{dh}{dt} = a_h(1 - h) - b_h h = a_h - (a_h + b_h)h$$

We can then divide this equation by $(a_h + b_h)$ to obtain the desired form

$$\tau_h \frac{dh}{dt} = -(h - h_\infty)$$

as

$$\frac{1}{a_h + b_h} \frac{dh}{dt} = \frac{a_h}{a_h + b_h} - h.$$

Comparing these two expressions we have

$$\tau_h = \frac{1}{a_h + b_h}, \qquad h_\infty = \frac{a_h}{a_h + b_h}.$$

We can implement this in R to compute the values of $h_{\infty}(-45)$ and $\tau_h(-45)$ as follows:

```
v = -45;

ah = 0.07*exp((-v-60)/20);

bh = 1/(1+exp(-0.1*(v+30)));

tauh = 1/(ah+bh);

hinf = ah/(ah+bh);
```

Evaluation of this script gives tauh = 4.6406 and hinf = 0.1534.

Exercise 12.2 To do the second exercise, you could use for-loops to compute τ_m , τ_h , etc. at a set of v values running from -100 to 75. But a better approach is to write a set of R functions tau.m, tau.h, etc., that can take as input a vector of v values, and return a vector of function values, e.g.

```
tau.m=function(v) {
     <stuff>
     return(something)
}
```

Having done that, you can then make a plot of $\tau_m(v)$ with

```
v=seq(-100,75,by=1)
plot(v,tau.m(v),type="1")
```

To put several curves on the same plot, you can chind them into a matrix and use matplot:

```
v=seq(-100,75,by=1)
M=cbind(tau.m(v),tau.h(v))
matplot(v,M,type="1")
```

For $m_{\infty}, \tau_m, n_{\infty}, \tau_n$ there is an added twist: the formula for the function Ψ gives the *indeterminate* value 0/0 when x=0, so R cannot evaluate it there. Nonetheless, using *l'Hopital's rule* from calculus, we can define $\Psi(0)=1$ to make Ψ a smooth function. So the *very first* thing you should do is write a function to evaluate $\Psi(x)$, using ifelse to set $\Psi=1$ when x=0 and $x/(e^x-1)$ when $x\neq 0$. The *very next* thing you should do, is use your function to plot $\Psi(x)$ over a range of values including 0, to make sure that it's working properly. Whenever you write a function, take a minute to test it and make sure that it's doing what it's supposed to.

Exercises 12.3,12.4. To compute currents for the rest of the exercises, it is again helpful to begin by writing functions mvt, nvt, hvt that evaluate the gating variables m, n, h as functions of time, v_0 , and v_s (the holding and step-up potentials). Equation (11) says that, for example,

$$m(t) = m_{\infty}(v_s) + (m_{\infty}(v_0) - m_{\infty}(v_s)) \exp(-t/\tau_m(v_s)), \qquad t \ge 0.$$
 (12)

and your mvt=function(times,v0,vs) needs to evaluate this formula. Once these functions have been defined, you can use them to write functions i.Na,i.K,i.leak for the individual currents, and then use those functions to write a function for the total current. Breaking up the computation of total current into a series of small pieces increases your odds of success, and leaves you with code that's easier to debug, understand and use.

Exercise 12.5 The last exercises require much more ingenuity than the previous ones. To get started, repeat computations like those of Exercise 12.4 to generate the sodium current "data" used in the exercise. For this exercise we assume that v_{Na} is known, so the current can be divided by $(v - v_{Na})$ to obtain the conductance $g_{Na}m^3h$. As your first step, create a matrix in which each column holds the conductance values for one value of v_s at a series of closely-spaced times from t=0 to t=6 msec, and plot those conductance curves so you can see what your "data" look like.

After this is done, strategies must be developed to estimate $m_{\infty}, h_{\infty}, \tau_m, \tau_h$. Frequently used procedures assume that

- 1. the experiment starts at a potential sufficiently hyperpolarized that there is no inactivation (i.e. h=1) and
- 2. activation is so fast relative to inactivation that m reaches its steady state before h has changed significantly.

This separation of time scales – m changing much faster than h because $\tau_m \ll \tau_h$ – means that early, rapid changes in conductance are caused by changes in m, while later, slower changes in conductance are caused by changes in h. One can then estimate m_∞ and τ_m from the increasing portion of the conductance traces, assuming that h=1 during this entire period of time, so that the conductance is $g_{Na}m^3$. Under this assumption, what does the maximum value of the conductance (at a given v_s) tell you about the parameters affecting m(t)?

Then to estimate τ_h , we assume that the decreasing "tail" of the conductance curve is given by $g_{Na}m_{\infty}^3h(t)$, because m(t) has converged to its steady state. The rate at which this curve converges to its limiting value $g_{Na}m_{\infty}^3h_{\infty}$ provides an estimate of τ_h .

It is easier to estimate h_{∞} from a different set of voltage traces, having different values of the holding potential v_0 , followed by a step to a step-up potential v_s that is the same for each trace. In this protocol we start with h partially inactivated (i.e. h < 1), so the peak conductance during the subsequent trace is proportional to the initial value of h, which is $h_{\infty}(v_0)$. A plot of peak conductance as a function of v_0 is then *proportional to* a plot of h_{∞} as a function of v_0 . The constant of proportionality can be inferred from the fact that the maximum value of $h_{\infty}(v)$ is 1. Consult Hille for further descriptions of these protocols.

13 Solving systems of differential equations

Built-in R functions make it relatively easy to do some complicated things. One important example is finding numerical solutions for a system of differential equations

$$\frac{dx}{dt} = f(t, x).$$

Here x is a vector assembled from quantities that change with time, and the *vector field* f gives their rates of change. The Hodgkin-Huxley model from the last section is one example. Here we start with the simple model of a gene regulation network from Gardner et al. (2000) that is described in the textbook. The model is

$$\frac{du}{dt} = -u + \frac{\alpha_u}{1 + v^{\beta}}$$

$$\frac{dv}{dt} = -v + \frac{\alpha_v}{1 + u^{\gamma}}$$
(13)

The variables u, v in this system are functions of time. They represent the concentrations of two repressor proteins P_u, P_v in bacteria that have been infected with a plasmid containing genes that code for P_u and P_v . The plasmid also contains promoters, with P_u a repressor of the promoter of the gene coding for P_v and vice-versa.

The equations are a simple compartment model describing the rates at which u and v change with time. P_u degrades at rate 1 (so the loss rate is $-1 \times u$) and it is produced at a rate $\frac{\alpha_u}{1+v^\beta}$, which is a decreasing function of v. The exponent β models the *cooperativity* in the repression of P_u synthesis by P_v . The processes of degradation and synthesis combine to give the equation for $\frac{du}{dt}$, and the equation for $\frac{dv}{dt}$ is similar.

There are no explicit formulas to solve this pair of equations, but we can interpret what the equations mean geometrically. At each point of the (u,v) plane, we regard $(\frac{du}{dt},\frac{dv}{dt})$ as a **vector** that gives the direction and magnitude for how fast (u,v) jointly change as a function of t. Solutions to the equations give rise to parametric curves (u(t),v(t)) whose tangent vectors $(\frac{du}{dt},\frac{dv}{dt})$ are those specified by the equations.

To plot the vector field, first run the script DMBpplane.R (this script is based on pplane.R by Daniel Kaplan, Department of Mathematics, Macalester College, and has been modified and used here with his permission). This script includes a function that computes the vector field for model (13) with the assumption that $\alpha_u = \alpha_v$:

```
toggle=function(u,v,parms) {
  du= -u + parms[1]/(1+v^parms[2]);
```

```
dv= -v + parms[1]/(1+u^parms[3]);
    return(c(du,dv));
}
```

Note how this function is set up:

- Its input arguments are the two state variables u, v and a parameter vector parms. Even if parms is not used by the function, it must be included in the list of arguments (you can set parms=0).
- The function returns the vector field as a vector.
- The computations are set up so they will go through if u and v are matrices of equal size. This eliminates the needs for for-loops in many of the computations that we will use to study vector fields.

Use this setup for any vector field that you want to study using the functions in DMBpplane.R.

The function phasearrows (also in DMBpplane.R) can then be used to plot the vector field, as in this example:

```
phasearrows(fun=toggle,xlim=c(0,3),ylim=c(0,3),resol=25,parms=c(3,2,2))
```

In this statement, fun is the name of the function that calculates the vector field, xlim and ylim define the range of values for the plot, and setting resol=25 means that arrows showing the vector field will be drawn at a 25×25 grid of points within the plotting region. The parameter vector parms is passed to the vector field function – this argument can be omitted if the vector field function does not use parms.

Exercise 13.1 Run DMBpplane.R to create the toggle function, and then use the command above to plot the vector field.

We can think of solutions to (13) as curves in the plane that "follow the arrows". Given a starting point (u_0, v_0) , the mathematical theory discussed in the textbook proves that there is a unique solution (u(t), v(t)) with $(u(0), v(0)) = (u_0, v_0)$. The process of finding the solution numerically is called *numerical integration*. Methods for numerical integration build up an approximate solution by adding short segments of an approximate solution curve, one after another.

R's numerical ODE solvers are in the **deSolve** package. Download this (if necessary) from CRAN, and then use the command library(deSolve) to load it into your R session.

Exercise 13.2 After you load the deSolve package, use ?rk4 to look at the syntax for this function.

To use R 's ODE solvers, the first step is to write a function to evaluate the vector field *in the specific format required by the solvers*. Here's what that looks like for the toggle switch model:

```
Toggle=function(t,y,parms) {
    u=y[1]; v=y[2];
    du= -u + parms[1]/(1+v^parms[2]);
    dv= -v + parms[1]/(1+u^parms[3]);
    dY=c(du,dv);
    return(list(dY));
}
```

There's a bit to digest in that. Here are the things to note:

1. The function <u>must</u> have input arguments (t,y,parms), even if only the state vector y is used to compute the vector field. Here t is time, y is the state vector, and

- 2. parms is again a vector of parameter values for the function. This allows you to see how solutions change as parameters are varied, without having to re-write and re-run the function. It also simplifies the process of fitting differential equations to data.
- 3. The vector field value has to be returned <u>as a list</u>, which is why we need both *toggle* and *Toggle*. There's a good reason for this; if you're curious and have *way* too much free time on your hands, ?1soda gives the explanation. You've been warned.

The "basic" ODE solver in R is **rk4**, which implements the 4th-order Runge Kutta method with a fixed time step. The format is

```
out=rk4(x0,times,func,parms)
```

Here times is a vector of the times at which you want solution values, x0 is the value of the state vector at the initial time (i.e. times[1]), func func is the function specifying the model (such as Toggle), and parms is the vector of parameter values that will be passed to func.

Here is an example using our Toggle function:

```
x0=c(.2,.1); times=seq(0,50,by=0.2); parms=c(3,2,2);
out=rk4(x0,times,Toggle,parms)
matplot(out[,1],out[,2:3],type="l",ylim=c(0,3),xlab="time t",ylab="u,v");
```

Exercise 13.3 Write a script toggle.R with the commands above, and run them to see some solution trajectories as a function of time. Which plotted curve is u, and which is v? How do you know? Look at the matrix out to see how it is set up: the first column is a list of times, and the other columns are the computed (approximate) values of the state variables at each time. \maltese

The state trajectories both seem to be approaching constants. A second way of looking at the trajectories (**Exercise:** do this yourself right now) is:

```
plot(out[,2],out[,3],type="l",xlab="u",ylab="v")
```

This kind of plot is called a *phase portrait*. It shows the path in the (u, v) *phase plane* taken by the trajectory, but we lose track of the times at which the trajectory passes through each point on this path.

Exercise 13.4 Use rk4 again with initial conditions (0.2, 0.3) to produce a new output matrix out2 and plot phase portraits of both solutions. Do this first for time interval [0, 50] and again for [0, 200].

The trajectories appear to end at the same places, indicating that they didn't go anywhere after t=50. We can explain this by observing that the differential equations vanish at these endpoints. The curves where $\frac{du}{dt}=0$ and $\frac{dv}{dt}=0$ are called *nullclines* for the vector field. They intersect at *equilibrium points*, where both $\frac{du}{dt}=0$ and $\frac{dv}{dt}=0$. The solution with initial point an equilibrium is constant. Here, the equilibrium points are (asymptotically) stable, meaning that trajectories close to the equilibria approach them as t increases.

Nullclines for a general vector field can be plotted using the function nullclines in DMBpplane.R. The syntax is the same as phasearrows, for example:

```
nullclines(fun=toggle,xlim=c(0,3),ylim=c(0,3),resol=250,parms=c(3,2,2))
```

It is a good idea to use a large value of resol so that the nullclines are found accurately.

Exercise 13.5 Without erasing the nullclines, add to that plot (using points) the two solution trajectories with different initial conditions that you have computed (out and out2 from previous exercises). You should see that each trajectory converges to an equilibrium point where the nullclines intersect.

Exercise 13.6 There is a third equilibrium point where the two nullclines intersect, in addition to the two at the ends of the trajectories that you have computed. Experiment with different initial conditions, to see if you can find any trajectories that converge onto this third equilibrium (Hint: what happens in this model if u(0) = v(0) for these parameter values?)

Exercise 13.7 Change the value of α from 3 to 1.5. How does the phase portrait change? Plot the nullclines to help answer this question.

13.1 Always use Isoda! (well, almost always)

The "industrial strength" solver in R is 1soda. This is a front end to a general-purpose differential equation solver (called, oddly enough, Isoda) that was developed at Lawrence Livermore National Laboratory. The full calling format for Isoda in the deSolve package is

```
lsoda(y, times, func, parms, rtol = 1e-6, atol = 1e-6,
  jacfunc = NULL, jactype = "fullint", verbose = FALSE,
  tcrit = NULL, hmin = 0, hmax = NULL, hini = 0, ynames = TRUE,
  maxordn = 12, maxords = 5, bandup = NULL, banddown = NULL,
  maxsteps = 5000, dllname = NULL, initfunc = dllname,
  initpar = parms, rpar = NULL, ipar = NULL, nout = 0,
  outnames = NULL, forcings = NULL, initforc = NULL,
  fcontrol = NULL, events = NULL, lags = NULL,...)
```

Don't panic. Sensible defaults are provided for everything but the arguments required by rk4, so 1soda can be called just like rk4:

```
out=lsoda(x0,times,Toggle,parms=c(3,2,2))
matplot(out[,1],out[,2:3],type="1",ylim=c(0,3));
```

Usually you can get away with doing this, and here we always will. The options rtol and atol can be used to control the accuracy that the numerical integration tries to achieve, by using smaller time steps. 1soda automatically adjusts step sizes to achieve the desired error tolerance (based on error estimates that it calculates), whereas rk4 always goes directly from one value in times to the next.

One key reason for using 1soda rather than rk4 is *stiffness*. Differential equations are called stiff if they have some variables or combinations of variables changing much faster than others. Stiff systems are harder to solve than non-stiff systems and require special techniques. The 1soda solver monitors for signs of stiffness, and automatically switches to a stiff-system solver when necessary. Many biological models are at least mildly stiff, so for real work you should (almost) *always* use 1soda rather than rk4. The only time to try rk4 is when 1soda fails on your problem, returning an error message rather than solution values. You may get a clue as to why that happened by trying rk4 with a very small time step and seeing how the solutions behave, e.g., does a state variable blow up to infinity in finite time?

The only reason for not using 1soda is if you know your equations are stiff. In that case it is more efficient to use a solver that assumes stifness rather than checking for it, such as vode with default settings. See ?ode for a list of available solvers, if you're curious.

Exercise 13.8 Write a script to solve the Lotka-Volterra competition model

$$dx_1/dt = x_1(r_1 - x_1 - ax_2)$$
$$dx_2/dt = x_2(r_2 - x_2 - bx_1)$$

in which the parameters $r_1, r_2, a, b > 0$ are all passed as parameters via the argument parms. Have your script generate solutions for the same parameter values using first rk4 and then 1soda, and make a plot to compare the results. \maltese

Exercise 13.9 Write a script that uses 1soda to solve the constant population size SIR model with births,

$$dS/dt = \mu(S + I + R) - \beta SI - \mu S$$

$$dI/dt = \beta SI - (\gamma + \mu)I$$

$$dR/dt = \gamma I - \mu R$$

For parameter values $\mu=1/60, \gamma=25$ (corresponding to a mean lifetime of 60 years, and disease duration of 1/25 of a year) and population size $S(0)+I(0)+R(0)=10^6$, explore how the dynamics of the disease prevalence I(t) (starting from a low initial disease prevalence) changes as you increase the value of β slowly from 0. NOTE: slowly here means *really slowly*. For example, the rate of new cases/week when $I=10, S=10^6-10$ is roughly $10^7\beta/52\approx 200,000\beta$.

13.2 The logs trick

In many biological models the state variables always must be non-negative, but a numerical ODE solver doesn't know this. If a variable decreases rapidly to near-zero values in the exact solution, a numerical approximate solution might overshoot to a negative value, leading to nonsense. For example, suppose that the number of infectives becomes negative in a standard SIR-type infectious disease model, the transmission rate βSI becomes negative. So contacts between susceptibles and infectives still occur, but their effect in the model is to make the sick one become healthy, pushing I even more negative. Once a model goes down the rabbit hole, it may never come back.

This problem can often be fixed by a simple trick that is often used but rarely written down. The trick is to transform the model onto natural-log scale, solve the transformed model, and back-transform the output. This is much easier than it sounds, because of the fact that for any state variable x(t),

$$\frac{d(\log x(t))}{dt} = \frac{1}{x(t)} \frac{dx}{dt}.$$
(14)

This means that you can compute the untransformed vector field dx/dt, and then get the tranformed vector field with a single element-by-element division. For example, here is the function that computes the toggle-switch model vector field on log scale. The input argument logy is the log-transformed state vector (log u, log v).

```
Toggle.log=function(t,logy,parms) {
     y=exp(logy);
     u=y[1]; v=y[2];
     du= -u + parms[1]/(1+v^parms[2]);
     dv= -v + parms[1]/(1+u^parms[3]);
     dY=c(du,dv);
     return(list(dY/y));
}
```

There are only two differences from the original Toggle: the first line that back-transforms from logy to y, and the last line that uses equation (14) to compute the vector field of the log-transformed state vector. And there are just two minor differences in the call to lsoda: log-transforming the initial conditions, and back-transforming the output.

```
x0=log(c(.2,.1)); times=seq(0,50,by=1); parms=c(3,2,2);
out=lsoda(x0,times,Toggle.log,parms);
out[,-1]=exp(out[,-1]) #column 1 is time, which was not transformed
matplot(out[,1],out[,2:3],type="l",ylim=c(0,3),xlab="time t",ylab="u,v");
```

The logs trick is not a panacea. It can even *create* problems that weren't there originally. If a positive state variable converges to 0 in non-transformed numerical solutions, then on log scale it is diverging to $-\infty$, potentially leading to numerical overflow errors.

Exercise 13.10 Write a script to solve the Lotka-Volterra model (in the previous subsection) using the log trick. Note that for this model you can do (14) algebraically and avoid the dY/y calculation in the R code. Have your model generate solutions using 1soda, and plot the species abundances as a function of time.

14 Equilibrium points and linearization

This section continues our study of differential equations with R. We will investigate the computation of equilibrium points and their linearization. Recall that an equilibrium point of the system $\frac{dx}{dt} = f(x)$ is a vector \bar{x} where $f(\bar{x}) = 0$. If the model's state vector has dimension n, the equilibrium condition $f(\bar{x}) = 0$ is a system of n equations in n unknowns that may have multiple solutions, or no solutions at all.

Solving nonlinear equations is a difficult task for which there are no sure-fire algorithms. Newton's method is a simple iterative algorithm that is usually very fast when it works, but it doesn't always work. Newton's method takes as its input a starting guess at the equilibrium x_0 , ideally a good guess that is close to the equilibrium we see where $f(\bar{x}) = 0$. The method begins by evaluating $y_0 = f(x_0)$, and terminates if the magnitude of y_0 is smaller than a specified tolerance. If y_0 is larger than the tolerance, then a new value x_1 is computed from the solution of the linear approximation to f at x_0 :

$$f(x) \approx L(x) = f(x_0) + Df(x_0)(x - x_0).$$

Writing out the model as

$$\frac{dx_i}{dt} = f_i(x_1, x_2, \cdots, x_n),\tag{15}$$

 $Df(x_0)$ is the $n \times n$ Jacobian matrix whose (i, j) entry is the partial derivative of f_i with respect to x_j , evaluated at the equilibrium. If $Df(x_0)$ is an invertible matrix, then we can solve the linear system L(x) = 0 for x, yielding the new value of x:

$$x_1 = x_0 - [Df(x_0)]^{-1} f(x_0). (16)$$

We then replace x_0 by x_1 , and start over again by evaluating $f(x_1)$ and checking if its value is small enough. If so we stop, otherwise we take x_1 as our new "starting value" and repeat the process to find a point x_2 that is (hopefully) closer to being a solution of f(x) = 0.

Success or failure of Newton's method depends on whether or not x_0 is close enough to an equilibrium \bar{x} at which the Jacobian $Df(\bar{x})$ is an invertible matrix. When that is true, Newton's method converges "quadratically", meaning that the error after one additional iteration is proportional to the current error squared, which is a very good thing ocne the error has become small. When that is not true, Newton's method can continue "forever", never converging onto an equilibrium.

The function newton in DMBpplane.R implements Newton's method on R functions that are set up in the format used for solving differential equations with rk4 or lsoda. So we can apply Newton's method to our toggle switch model (13) using the commands

```
newton(Toggle, x0=c(2.5, 0.4), parms=c(3, 2, 2))
```

The function newton also has some optional arguments for which default values are provided; these include the maximum number of iterations niter, the convergence tolerance tol, and the increment inc used to compute Jacobian matrices Df by finite-difference approximation to the derivatives.

Exercise 14.1 Download and run DMBpplane.R, and then run the command above. Note that the values of the state variables are displayed at each iteration. Recall that for these parameter values there are three equilibrium points. Write a script that finds the two other equilibrium points of the toggle switch model for those parameter values, by choosing different values of x_0 .

The file repress.R implements the six dimensional repressilator model of Elowitz and Leibler:

```
repress=function(t,y,p){
    dy = rep(0,6);
    dy[1] = -y[1] + p[1]/(1.+y[6]^p[4])+ p[2];
    dy[2] = -y[2] + p[1]/(1.+y[4]^p[4])+ p[2];
    dy[3] = -y[3] + p[1]/(1.+y[5]^p[4])+ p[2];
    dy[4] = -p[3]*(y[4]-y[1]);
    dy[5] = -p[3]*(y[5]-y[2]);
    dy[6] = -p[3]*(y[6]-y[3]);
    return(list(dy))
}
```

Note that to save typing parms is replaced here by p – the name doesn't matter so long as it's in the right place in the list of arguments.

Exercise 14.2 Write a script that uses 1 soda and repress to reproduce the textbook figure that shows oscillations in this model by computing and graphing a trajectory for this model with parameters parms = c(50,0,0.2,2). Almost any initial conditions should work – try x0 = 2*runif(6).

Exercise 14.3 Write a script that uses Newton's method to compute an equilibrium point of the repressilator for the same values of the parameters. \(\mathbb{H} \).

We can use eigenvalues and eigenvectors as tools to study solutions of a vector field near an equilibrium point \bar{x} , as discussed in the textbook. The basic idea is that we approximate the vector field by the *linear* system

$$\frac{dw}{dt} = Aw$$

where $w = x - \bar{x}$ and A is the $n \times n$ Jacobian matrix $Df(x_0)$ that newton computes for us. In many circumstances the phase portrait of this linear system will look similar to the phase portrait of $\frac{dx}{dt} = f(x)$.

If v is an eigenvector of A with eigenvalue λ , the curve

$$w(t) = \exp(t\lambda)v$$

is a solution of $\frac{dw}{dt} = Aw$ because $Av = \lambda v$. If the eigenvalue λ is negative, then $\exp(t\lambda) \to 0$ as $t \to \infty$. Complex eigenvalues give solutions that have trigonometric terms: $\exp(it) = \cos(t) + i\sin(t)$. Whenever the real parts of all the eigenvalues are negative, the equilibrium point is *linearly stable*. Otherwise it is unstable.

Exercise 14.4 Write a script to compute the eigenvalues of the Jacobian at the equilibrium point that you found for the repressilator model (recall that the Jacobian at the equilibrium is one of the components of the

Parameter	Set 1	Set 2
g_{Ca}	4.4	5.5
g_K	8	8
g_L	2	2
v_{Ca}	120	120
v_K	-84	-84
v_L	-60	-60
C	20	20
φ	0.04	0.22
i	90	90
v_1	-1.2	-1.2
v_2	18	18
v_3	2	2
v_4	30	30

Table 8: Parameter sets for the Morris-Lecar model.

object returned by newton). Then, change the parameters to parms = c(50,1,0.2,2) and recompute the equilibrium point and its eigenvalues.

Exercise 14.5 Write a script to compute and print the eigenvalues of the three equilibrium points for the toggle switch model with parms = c(3,2,2). Plotting the nullclines will let you pick starting conditions for newton near each of the equilibria. So have your script plot the nullclines, then find all three equilibria and compute the eigenvalues of the Jacobian at each equilibrium. You should find that the two asymmetric equilibria are stable. The symmetric equilibrium has one positive and one negative eigenvalue, making it a saddle.

Exercise 14.6 Continuing with the toggle switch model: choose points $x=\bar{x}\pm\epsilon w_1$ where $0<\epsilon\ll 1$ and w_1 is the eigenvector of the Jacobian corresponding to the positive eigenvalue. Write a script that computes and plots (on the same graph as a plot of the nullclines) solution trajectories starting at these points. These trajectories are an approximation to the *unstable manifold* of the saddle, the solution curve that converges to the equilibrium as $t\to -\infty$.

Exercise 14.7 Next do the same for the eigenvector w_2 corresponding to the negative eigenvalue, but integrate *backward* in time; i.e., choose a negative final time for your integration. These trajectories approximate the *stable manifold* of the saddle, the solution curve that converges to the equilibrium as $t \to \infty$.

15 Phase-plane analysis and the Morris-Lecar model

Recommended reading: Rinzel and Ermentrout, Analysis of Neural Excitability and Oscillations. In: Koch and Segev, Methods in Neuronal Modeling: From Synapses to Networks (2nd edition). MIT Press, Cambridge, MA (1998).

In this section we continue to study phase portraits of two-dimensional vector fields, using the Morris-Lecar model for the membrane potential of barnacle muscle fiber. The differential equations for the Morris-Lecar model are

$$C\frac{dv}{dt} = i - g_{Ca}m_{\infty}(v)(v - v_{Ca}) - g_K w(v - v_K) - g_L(v - v_L)$$

$$\tau_w(v)\frac{dw}{dt} = \phi(w_{\infty}(v) - w)$$

$$m_{\infty}(v) = 0.5\left(1 + \tanh(\frac{v - v_1}{v_2})\right)$$

$$w_{\infty}(v) = 0.5\left(1 + \tanh(\frac{v - v_3}{v_4})\right)$$

$$\tau_w(v) = \frac{1}{\cosh\left(\frac{v - v_3}{2v_4}\right)}$$
(17)

The parameters used in the textbook are listed in table 8.

For phase-plane analysis we recommend stepping outside R and using a specialized tool. One option is John Polking's pplane, which is currently distributed as a Java application. Another fine option is Bard Ermentrout's open-source program XPP (Ermentrout 2002). Finally, we provide an R utility called Rpplane that includes the essential tools for the phase-plane exercises here and in the textbook, but not much else. Our past experience suggests that Rpplane is more reliable than pplane when the nullclines are highly nonlinear.

Using pplane At this writing, pplane is available at http://math.rice.edu/~dfield/dfpp.html. Download pplane.jar, find the directory where you saved the file and double-click on the downloaded file. Alternatively, run pplane from the command line with the command: java -jar pplane.jar (this seems to be the better option on a Mac). Running pplane requires the Java Runtime Environment. Older versions of pplane for Matlab are available from the same web page, but these are not fully compatible with the current version of Matlab.

To load the Morris-Lecar model into a Matlab version of pplane, download the file ML.pps from the text-book webpage. In the Pplane setup window, use File/load a system... and select ML.pps. The "default" system in the setup window will then be replaced by the Morris-Lecar model, with two parameters free to be varied, g_{Ca} and ϕ . Click on the Proceed bar at the bottom right of the setup window; this will open a graph window in which you can work with the phase portrait. Everything you need is in the menus, but it will take a bit of looking around or a guided tour by your instructor.

Using XPP The home page for XPP is www.math.pitt.edu/~bard/xpp/xpp.html; you can download the source code and binaries for various operating platforms, including (at this writing) iPad and iPhone. Yes, you can now solve differential equations on your phone. For Windows PCs, the best option at this writing appears to be the one described at www.math.pitt.edu/~bard/xpp/ezwin.html, with links to all required files. Also download the file MLdmb.ode from the textbook web page, and save it into any convenient folder. If you follow Bard's directions for installing XPP, you'll be able to launch the program by first starting the X server that you installed, and then drag-and-drop a model-definition file, such as MLdmb.ode, onto the xpp.bat icon on your desktop. There's some online documentation and a tutorial for XPP at its home page, and full details in Ermentrout (2002).

Using Rpplane The function Rpplane is included in DMBpplane.R. The syntax is Rpplane(fun,xlim,ylim,parms=NULL,add=F,ngrid=5,maxtime=100)

• fun is the function defining the vector field, which needs to be in the format of the toggle function of the last section.

- xlim, ylim define the plotting region.
- parms is the vector of any parameter values that are used by fun.
- add indicates whether the results should be displayed on the currently active graphics device.
- ngrid is the number of grid points used for plotting a grid of trajectories
- maxtime is the time interval over which solution trajectories are computed

So Rpplane(ml,c(-60,40),c(-0,1),parms=c(5.5,0.22),maxtime=250); will launch Rpplane with parameters $g_{Ca}=5.5, \phi=0.22$ using the ml function that defines the vector field in DMBpplane.R, drawing trajectories out to time t=250, and default values for the other arguments. A high value of maxtime is needed in this system for drawing the stable/unstable manifolds of saddle points.

Always start by drawing the nullclines or phase arrows. You can then add to those plots by selecting from the menu. To start a trajectory, find a fixed point or get the local stable/unstable manifold for a saddle, you need to click on the current graph to select the starting point.

Plotting symbols distinguish different kinds of fixed points.

- **Spirals** (complex conjugate eigenvalues) are plotted as circles (closed = stable, open = unstable).
- **Nodes** (two real eigenvalues with the same sign) are plotted as triangles (closed = stable, open=unstable).
- Saddles are plotted as open diamonds.

The coordinates of the fixed point and the eigenvalues are printed to the console window. The unstable manifold of a saddle is plotted in red, and the stable manifold in black.

Trajectories going backwards in time are the way to hunt for unstable periodic orbits, because they are stable in the time-reversed system. So for example if a stable fixed point is surrounded by an unstable periodic orbit, a backwards trajectory starting near the fixed point and inside the periodic orbit will converge onto the period orbit from the inside. A backward trajectory starting just outside the periodic orbit will converge onto it from the outside.

Exercise 15.1 Compute phase portraits for the Morris-Lecar model at the two different tabulated sets of parameter values. Label

- Each of the equilibrium points by type,
- The stable and unstable manifolds of any saddle points
- The stability of the periodic orbits.

It's OK to print the graph out and do the labeling by hand, or (in Rpplane) you can use the text function in the Console window to add text to an existing plot.

Bifurcations of a system occur at parameters where the number of equilibria or periodic orbits change. The typical bifurcations encountered while varying a single parameter at a time in a system with at most a single saddle point are

- 1. **Saddle-node bifurcation:** The Jacobian at an equilibrium point has a zero eigenvalue. At the bifurcation point two equilibria appear, or two equilibria collide and then disappear.
- 2. **Hopf bifurcation:** The Jacobian at an equilibrium point has a pair of pure imaginary eigenvalues. The equilibrium goes from being a stable spiral to an unstable spiral, or vice-versa.
- 3. **Homoclinic bifurcation:** There is a trajectory in both the stable and unstable manifold of a saddle.

4. **Saddle-node of limit cycle bifurcation:** A periodic orbit has double eigenvalue 1.

The changes in dynamics that occur at each kind of bifurcation are discussed in Chapter 5 of the textbook.

Exercise 15.2 At saddle-node bifurcations, two equilibria appear or disappear. Figure 5.14 of the textbook shows that as g_{Ca} is varied with $\phi = 0.22$, saddle-node bifurcations occur near $g_{Ca} = 5.32$ and $g_{Ca} = 5.64$. Compute phase portraits for values of g_{Ca} near these bifurcations, describing in words how the phase portraits change.

Exercise 15.3 Now set $g_{Ca} = 5.5$ and vary ϕ in the range from (0.04, 0.22). Show that both Hopf and homoclinic bifurcations occur in this range. What are the approximate parameter values at which bifurcations occur? Draw labeled phase portraits on both sides of the bifurcations, indicating the changes that occur. It will be useful to systematically draw phase portraits at a series of ϕ values, and look for changes between two successive phase portraits.

Exercise 15.4 Hopf bifurcations are *supercritical* if stable periodic orbits emerge from the equilibrium (which is then unstable) and *subcritical* if unstable periodic orbits emerge from the equilibrium (which is then stable). Is the Hopf bifurcation you located in the previous exercise subcritical or supercritical? Explain how you know.

Exercise 15.5 With g_{Ca} set to 4.4, show that the two periodic orbits you computed in the first exercise approach each other and coalesce as ϕ is increased. This is a saddle-node of limit cycles (snlc) bifurcation. Draw phase portraits on the two sides of the bifurcation.

Exercise 15.6 For parameter values $\phi = 0.33$ and g_{Ca} varying near the saddle-node value (≈ 5.64), the saddle-node is a *snip* bifurcation, where *ip* stands for "infinite period". Explain what this bifurcation is, using computed phase portraits as an illustration, and explain why the name is appropriate.

16 Simulating Discrete-Event Models

This section is an introduction to simulating models that track discrete agents (organisms, molecules, neurons) as they change in state, as an alternative to compartment models that assume large numbers of agents. The material here can can be regarded as a warmup for simulating finite-population disease models (Chapter 6 in the textbook), or as some simple examples of agent-based models (Chapter 8 in the textbook).

Figure 5 shows a compartment model for biased movement of particles between two compartments. The corresponding system of differential equations is

$$\frac{dx_1}{dt} = Lx_2 - Rx_1$$

$$\frac{dx_2}{dt} = Rx_1 - Lx_2$$
(18)

$$\begin{array}{c|c}
 & \xrightarrow{\rho_{21}=Rx_1} \\
\hline
 & & \downarrow \\
 & \rho_{12}=Lx_2
\end{array}$$

Figure 5: Compartment diagram for biased movements between 2 compartments.

Even for molecules – but much more so for individuals catching a disease – changes in state are discrete events in which individuals move one by one from one compartment to another. In some cases, such as molecular diffusion, transitions can really occur at any instant. But for modeling purposes we can have

transitions occurring at closely-spaced times $dt, 2dt, 3dt, \cdots$ for some *short* time step dt, and allow each individual to follow a Markov chain with transition matrix

$$A = \begin{bmatrix} 1 - (R \times dt) & L \times dt \\ R \times dt & 1 - (L \times dt) \end{bmatrix}$$

In TwoState.R the movement decisions for many particles are made by using runif to toss many coins at once. If there are N particles in compartment 1, each with probability Rdt of moving to compartment 2, then sum(runif(N)<R*dt) simulates the combined outcome (number moving) from all of the "coin tosses". Note in TwoState.R that at each time step, first all coins are tossed – for all particles in all compartments – and only then are particles moved to update the state variables. Note also the use of **ifelse** to avoid trying to toss zero coins (and see ?ifelse).

Each simulation of the model will have a different outcome, but some properties will be more or less constant. In particular

- 1. Once dt is small enough to approximate a continuous-time process, further decreases in dt have essentially no effect on the behavior of simulations. Roughly, dt is small enough to model continuous time if there would be practically no chance of an individual in continuous time doing 2 or more things in a time interval of length dt. For this model, that means that we must have $(Rdt) \times (Ldt) \ll 1$, i.e. $dt \ll 1/\sqrt{RL}$.
- 2. A compartment's typical range of departures from solutions of the differential equation are of order $1/\sqrt{n}$ where n is the expected number of particles in the compartment.

If there are many particles, coin-tossing with **runif** becomes slow. Instead, we can recall that the binomial random variable $\mathbf{B}(N,p)$ is the number of heads in N coin-tosses with probability p of heads on each toss. In R , binomial-distributed random numbers are generated by the rbinom function. The syntax is:

This generates a vector of n random numbers from a $\mathbf{B}(N,p)$ distribution. The script file TwoState2.R uses **rbinom** instead of **runif** to do the coin tosses.

Exercise 16.1 The pure death process in discrete time tracks a population of initial size N in which the only events are the occasional death of individuals. Between times t and t+1, each individual alive at time t has probability p of death, and probability 1-p of surviving to time t+1, independent of what happens to any other individual in the population. Eventually everybody is dead, so the main summary measure of a simulation run is the time it takes until the population is extinct.

Write an R script to simulate the pure death process by using coin-tossing as in the two-compartment example above. That is, if it is now time t and there are N(t) individuals still alive, we do N(t) coin-tosses with probability p of Heads(=death) and subtract those from the current population. This continues until everyone is dead. The first line of your m-file should specify the values of N and p, e.g. N=250; p=0.05;

Exercise 16.2 One run of a stochastic model is not the whole story, because the next run will be different. One needs to do multiple runs, and look at the distribution of possible outcomes. Extend your script file from the last exercise so that it does 100 runs of the pure death process (all having the same values of N and p), stores the time at which the population goes extinct in each run, computes the mean and standard deviation of extinction times across runs, and plots a histogram of the extinction times. \maltese

Exercise 16.3 Let's give the two-compartment example a totally different interpretation. We have n poten-

	a	-0.5	-0.5	-0.5	-0.5	-0.5	-0.5	-0.5	-0.5
Ĩ	b	1.25	1.25	1.25	1.25	1.25	1.25	1.25	1.25
								1.948	
ĺ	\overline{j}	-0.5	-0.49	-0.48	-0.45	-0.45	-0.45	-0.45	-0.45

Table 9: Eight sets of parameter values for FHN system

tial sites for local populations in a landscape, which can either be empty (state=1) or occupied (state=2). Then R is the chance that an empty site becomes occupied, and L is the chance that an occupied site becomes extinct. It's plausible that L is constant over time, but the rate at which empty sites are occupied should depend on the how many sites are occupied. So modify TwoState2.R so that in each time step, R is proportional to the number of occupied (state=2) sites. Note that you should choose parameters so that R is never bigger than 1 – that is, even if only one site is empty, that site has some probability of remaining empty. \maltese

Exercise 16.4 Modify your script from the last exercise so that it runs until all sites are in state 1 or until t=100, whichever comes first, and prints out the extinction time (the population as a whole goes extinct when all sites are in state 1). Find parameters such that extinction at or before t=100 is likely but not 100% certain.

17 Phase-plane analysis and the FitzHugh-Nagumo model

We continue our study of phase portraits and bifurcations of two-dimensional vector fields, using the FitzHugh-Nagumo system. These equations give a simple model of an excitable system:

$$\frac{dx}{dt} = c\left(x - \frac{1}{3}x^3 - y + j\right), \qquad \frac{dy}{dt} = \frac{1}{c}\left(x + a - by\right) \tag{19}$$

Here x(t) represents the level of excitation, and y(t) represents recovery. Notice that if $c \gg 0$, dx/dt is much larger than dy/dt, so x is the fast variable and y is the slow variable. The parameter j is a stimulus – for example, current applied to a neuron.

When the parameters a and b satisfy 0 < b < 1 and $1 - \frac{2b}{3} < a < 1$, there is exactly one equilibrium point. In this exercise, we investigate a different range of parameters.

Exercise 17.1 Implement the system in pplane. Figure 6 shows a phase portrait with parameters (a, b, c, j) = (-0.5, 0.5, 1, -0.5) that you can use to check that you've entered the equations correctly.

Exercise 17.2 Compute (and turn in) a complete phase portrait for the system with (a, b, c, j) = (-0.5, 1.25, 2, -0.45), showing

- The equilibrium points and their stability.
- Any limit cycles and their stability.
- The limit sets of the stable and unstable manifolds of any saddles.

Exercise 17.3 Compute phase portraits for the parameter values given in Table 9. Turn in a write-up that identifies exactly what type of bifurcation (or bifurcations) occur between the parameter values in each successive pair of columns, and what components (equilibria, periodic orbits, limit sets of stable and unstable manifolds of a saddle point) differ for each pair of phase portraits. For example, you might contrast two successive phase portraits by saying "There is a saddle-node bifurcation, which results in the disappearance

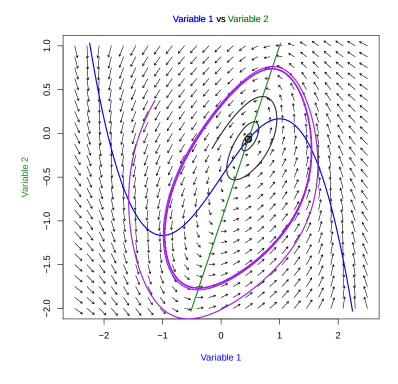


Figure 6: Phase portrait of FHN system with (a, b, c, j) = (-0.5, 0.5, 1, -0.5)

of two equilibria". Note that only one parameter changes between each successive pair of columns, so the bifurcations that happen in this exercise will be of the types discussed in lecture. \maltese

Exercise 17.4 For which sets of parameter values in the table is the system bistable? (Bistability means there are two distinct stable limit sets, not necessarily equilibrium points. A saddle point is not stable.)

18 Simulating the dynamics of spatial pattern formation

In this section we will study how the reaction-diffusion mechanism can create dynamic patterns in space. As an example, we will use a system of equations where the reaction mechanism is the Fitzhugh-Nagumo model, which is an approximation to the Hodgkin-Huxley equations for the action potential in nerve cells. So that x and y can be used for spatial coordinates, we use y and y for the state variables (note that y here corresponds to y in the previous section, and the applied current y is absent):

$$\frac{\partial u}{\partial t} = \frac{1}{e} (u - \frac{u^3}{3} - v) + D \left(\frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} \right)$$

$$\frac{\partial v}{\partial t} = e(u + b - 0.5v)$$
(20)

In the electrophysiology interpretation of the model, v represents the gating variable of an ion channel (fraction of open channels), which does not move; u represents the membrane potential, which changes due to the combined effects of transmembrane currents (ion flow) and diffusion of ions in the tissue (which could be the surface of the heart, or (in one space dimension) a nerve axon). The model therefore has diffusion in u but not in v, which is the extreme limit of the differing diffusion constants that are required for pattern formation by the Turing mechanism.

18.1 Finite difference approximation

The simplest way to get approximate numerical solutions of equation (20) is to discretize both space and time, replacing the derivatives in the equations by finite difference approximations. To explain this approach in the simplest setting, consider *one-dimensional* diffusion,

$$\frac{du}{dt} = D\frac{\partial^2 u}{\partial x^2}, \quad 0 \le x \le 1, 0 \le t \le T \tag{21}$$

with initial conditions $u(x,0) = u_0(x)$. For the spatial derivatives, we break up the interval [0,1] into m cells of width k = 1/m, centered at the *mesh points*

$$x_i = (i - 0.5)k, \quad i = 1, 2, \dots, m.$$

To estimate the second spatial derivative, note that

$$\frac{u(x_{i+1},t) - u(x_i,t)}{k} \tag{22}$$

is a finite-difference approximation to $\partial u/\partial x$ at the point halfway between x_i and x_{i+1} . Similarly,

$$\frac{u(x_i, t) - u(x_{i-1}, t)}{k} \tag{23}$$

is an approximation to $\partial u/\partial x$ at the point halfway between x_{i-1} and x_i . The difference between (22) and (23), divided by k, is therefore an approximation to $\partial^2 u/\partial x^2$ at x_i . That is,

$$\frac{\partial^{2} u}{\partial x^{2}}(x_{i},t) \approx \frac{[u(x_{i+1},t) - u(x_{i},t)]/k - [u(x_{i},t) - u(x_{i-1},t)]/k}{k} \\
= \frac{u(x_{i+1},t) + u(x_{i-1},t) - 2u(x_{i},t)}{k^{2}} \tag{24}$$

Note that both the first and second derivatives are estimated by a centered difference approach,

$$f'(x) \approx \frac{f(x+\varepsilon) - f(x-\varepsilon)}{2\varepsilon}$$
 (25)

The Taylor series $f(x\pm\varepsilon)=f(x)\pm\varepsilon f'(x)+\frac{1}{2}\varepsilon^2f''(x)+O(\varepsilon^3)$ shows that the centered difference (25) equals $f'(x)+O(\varepsilon^3)$. In contrast, the one-sided difference $\frac{f(x+\varepsilon)-f(x)}{\varepsilon}$ equals $f'(x)+O(\varepsilon^2)$, so centered differences are more accurate for small ε .

You may be wondering: what do we do at x_1 and x_m , where (24) involves the nonexistent x_0 and x_{m+1} ? Recall that the solution to a system of partial differential equations depends on the boundary conditions – what we assume about conditions at and beyond the boundary of the spatial domain. The boundary conditions that are most often encountered in practice determine the values at the mesh points x_0 and x_{m+1} that are outside the spatial domain.

For our FHN example, we will consider *no flux* boundary conditions: no material should flow out of the domain due to diffusion. The terms $[u(x_{i+1},t)-u(x_i,t)]$ and $[u(x_i,t)-u(x_{i-1},t)]$ in the first line of (24) represent net material flowing between two spatial cells. To make this flow zero at the domain boundaries x=0, x=1, we apply the finite-difference approximation (24) with $u(x_0,t)=u(x_1,t)$ and $u(x_{m+1},t)=u(x_m,t)$ for all t. Using those values, we can compute (24) at all of the mesh points inside the spatial domain (i.e., x_1 to x_m).

Exercise 18.1 What should we do if the boundary conditions are that $u(x,t) \equiv 0$ for x outside [0,1]? What should we do for *periodic* boundary conditions $u(0,t) \equiv u(1,t)$?

For the time derivatives, let h be the time step for solutions of (21) – that is, we want values of $u(x_i, t)$ at times $t = 0, h, 2h, \cdots$. To accomplish this we can use *Euler's method* which is based on the estimate

$$\frac{\partial u}{\partial t}(x,t) \approx \frac{u(x,t+h) - u(x,t)}{h} \implies u(x_i,t+h) \approx u(x_i,t) + h\frac{\partial u}{\partial t}.(x_i,t) \tag{26}$$

Applying (26) and (24) to (21) at the mesh points x_i , we get

$$u(x_{i}, t + h) \approx u(x_{i}, t) + hD \frac{\partial^{2} u}{\partial x^{2}}$$

$$\approx u(x_{i}, t) + hD \frac{u(x_{i+1}, t) + u(x_{i-1}, t) - 2u(x_{i}, t)}{k^{2}}.$$
(27)

Starting from the initial conditions $u(x_i, 0) = u_0(x_i)$, applying (27) at all mesh points gives an approximate solution at time h. Starting from there, we do the same again to get an approximate solutionat time 2h, and so on.

18.2 General method of lines

The solution method (27) consists of three steps:

- 1. Space is discretized so that the model only "lives" at a set of mesh points x_i .
- 2. Values of state variables at the grid points are used to approximate the spatial derivatives in the dynamic equations (20).
- 3. We numerically solve the system of ODEs that result from steps 1 and 2.

This general approach is called the *Method of Lines*. This name comes from imagining a set of lines running forward in time, one based at each grid point, along which we solve the ODEs. In (27) we used evenly-spaced mesh points for step 1, finite-difference approximation for step 2, and Euler's method for step 3. All of those choices favor simplicity over efficiency, and more accurate results can be gotten with a bit more work.

Step 3 can be done with a more accurate method for solving systems of ODEs. Substituting (24) into the model (equation (21)) we have the (approximate) system of ODEs

$$\frac{du(x_i,t)}{dt} = D\frac{u(x_{i+1},t) + u(x_{i-1},t) - 2u(x_i,t)}{k^2}.$$
 (28)

We can then get numerical solutions with any method for solving systems of ODEs – in principle. In practice, the systems resulting from finite-difference approximations to PDEs are often hard to solve numerically, as we discuss below.

Step 2 can be done more accurately using *spectral methods* (Trefethen 2000). Spectral methods do a spatial interpolation of each state variable at each time point, fitting a smooth function that matches the current values at the mesh points. Spatial derivates of the fitted function are used as the approximate spatial derivatives of the state variable. Spectral methods generally use unevenly spaced mesh points, and the best choice of mesh points and interpolating function depends on the spatial domain and boundary conditions. Spectral methods can be very efficient because the entire (interpolate+differentiate) process can be expressed as a single matrix multiplication, with a matrix that is computed once and then used at all time steps. See Trefethen (2000) for an introduction and pointers to more advanced treatments.

Spectral methods are an efficient way to get high-accuracy solutions on a simple spatial domains, but it is usually necessary to use an implicit or semi-implicit ODE solver (Trefethen 2000). For the complex spatial domains that often arise in engineering applications, other methods are needed that start by breaking up a complex domain into a set of simpler domains. In general, apart from simple PDEs on simple domains (such as the ones in this document), it is better to step outside R and use specialized software for numerical solution of PDEs.

18.3 Spatial Fitzhugh-Nagumo

The script sfn.R uses the finite-difference approach on the spatial Fitzhugh-Nagumo model (20) in a rectangulr domain with no flux boundary conditions. For two spatial dimensions, we break up space into an nx by ny grid of square cells with side k, and put a mesh point at the center of each cell (rectangular cells could be used instead). To apply the no-flux boundary conditions, we surround the domain by a "ring" of sites where the state variables take the same values as those at the adjacent site (in the x or y direction) in the interior of the domain. Using this trick, the following R function calculates the right hand side of the finite-difference approximation for the right-hand side of (20):

```
sfn=function(u,v) {
    ny = dim(u)[1];    nx = dim(u)[2];
    uec = cbind(u[,1],u,u[,nx]);  # u with extra columns
    uer = rbind(u[1,],u,u[ny,]);  # u with extra rows
    ul = uer[3:(ny+2),]+uer[1:ny,]+uec[,1:nx]+uec[,3:(nx+2)]-4*u;
    uf = (u-0.3333*u*u*u-v)/e + dx2*ul;
    vf = e*(u + b - 0.5*v);
    return(list(uf=uf,vf=vf));
}
```

ul is the finite difference approximation to $\frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2}$ that results from applying (24) in both the x and y directions, and dx2 is value of D/k^2 .

An important practical feature is that there are *no loops* in sfn. Everything is written as matrix operations, making the calculations much faster. Simulating the system is still quite slow; loops would make it intolerable.

As in the example of 1-dimensional diffusion, we use Euler's method for the time derivatives:

```
sfn.run=function(nsteps,init) {
    u=init$u; v=init$v;
    for (i in 1:nsteps){
        out = sfn(u,v);
        u = u + h*out$uf; v = v + h*out$vf;
        if (i%%5 == 1) image(1:dim(u)[2],1:dim(u)[1],t(u),col=rainbow(n=1000,start=0,end=0.7))
        cat(i,"\n");
    }
    return(list(u=u,v=v));
}
```

Because each time step depends on the results from the previous step, we *do* need a for-loop to iterate the model over time. The value returned by sfn.run is the final state of the model, suitable for use as a new "initial condition" to continue the run.

The script sfn.R includes everything needed to produce and plot simulations of the model. First, select and run the block of code with the functions sfn and sfn.run. Then select and run the code for the first parameter set/initial condition at the bottom of the script. The commands

```
pdf(file="sfn.pdf",onefile=TRUE);
nsteps=1000; # or however many time steps you want
out=sfn.run(nsteps,init1);
dev.off();
```

run the model with the first parameters and initial conditions for 1000 iterations, and save an image of the system state to a PDF file every 5 time steps. To continue the model run, you can use sfn.run again, starting from the model state at the last time step:

```
out=sfn.run(nsteps,out)
```

Exercise 18.2 Here are some exercises you can do with the sfn.R script:

- (a) Restart the model using the second set of parameters and initial conditions, and see what happens when spiral waves collide with with one another.
- (b) Experiment with changing the spatial discretization parameter k. What effect do you expect to see on the spatial pattern?
- (c) Experiment with gradually increasing the time step h. You should discover that solutions quickly go from reasonable to nonsensical, or that the computation crashes. It is characteristic of explicit solution methods that very small values of h are needed to keep numerical solutions of PDEs from developing errors that grow exponentially fast ("explicit" means that the set of values at the next time step is a computable function of the values at the current time; in "implicit" methods the values at the next time step are the solution to a system of nonlinear equations, which must be found numerically).

The script sfnLines.R uses Method of Lines to solve the spatial Fitzhugh Nagumo model by Method of Lines. The order-4 Runga-Kutta method is used to solve the ODE system, by re-writing the sfn function so it can be used with the rk4 function. Because rk4 is an explicit method like Euler's method, it still requires short time steps, but it gives more accurate solution of the ODE system. Only implicit solution methods, such as the widely used Crank-Nicholson method, remain numerically stable for long time steps.

Exercise 18.3 Modify sfnLines.R to generate numerical solutions of the Klausmeier (1999) model for pattern formation in vegetation on hillsides in semiarid regions. A nondimensionalized version of the model is:

$$\frac{\partial w}{\partial t} = a - w - wn^2 + v \frac{dw}{dx}
\frac{\partial n}{\partial t} = wn^2 - mn + \frac{\partial^2 n}{\partial x^2} + \frac{\partial^2 n}{\partial y^2}$$
(29)

Here w is water and n is vegetation. In the w equation a>0 represents supply rate (rainfall), wn^2 is uptake by plants and -w is other loss processes; vdw/dx is downhill (leftward) flow of water at velocity v. In the n equation wn^2 is vegetation growth, mn is vegetation mortality, and the diffusion term represents local expansion of vegetated areas through growth of established plants and new recruits that establish in the neighborhood of their parents.

Have your script simulate the model on a 60×60 grid of square cells of side dx=dy=1 (physically, these are 1 m^2 cells) with parameter values a=2, v=80, m=0.5. Use periodic boundary conditions in both x and y, that is

$$u(0, y, t) = u(60, y, t), \quad u(x, 0, t) = u(x, 60, t).$$

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In other respects, use the same methods as in sfnLines.R for generating the approximate numerical solution of this model.

For initial conditions: if nx=60 is the number of grid cells in each direction, use

```
n0=seq(0,1,length=nx); n0=(4*n0*(1-n0))^2;
n0 = 0.5+0.25*outer(n0,n0);
w0 = matrix(parms["a"],nx,nx);
image(1:nx,1:nx,t(n0)/max(n0),main="Vegetation",col=rainbow(n=1000,start=0,end=0.7));
```

The initial water concentration w0 is the steady state in the absence of vegetation; the last line above shows you what the initial vegetation pattern is. You will need to use a short time step (h = 0.01, for example) to get valid solutions. \maltese

Exercise 18.4 What pattern does the model produce, if you let it run long enough? (If you have not run it long enough, you can continue a simulation by taking the final state as your new initial conditions). Turn in a script that runs "long enough" and a plot illustrating the pattern that forms. \(\mathbf{H}\)

Exercise 18.5 Write a script that explores the question: is the final pattern sensitive to initial conditions, or do you find the same behavior in the long run regardless of what initial conditions you choose? Remember that the initial conditions have to satisfy the periodic boundary conditions, and only consider initial conditions where vegetation is not completely absent.

Exercise 18.6 Write a script to explore the question: how does the final pattern change as parameters a, v, m are changed? Experiment by making small changes in these, one at a time, and determining what the effects are. For example, do things happen faster or slower? Do biologically interesting features become larger, smaller, or disappear?

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