Practical R for Epidemiologists

Second Edition

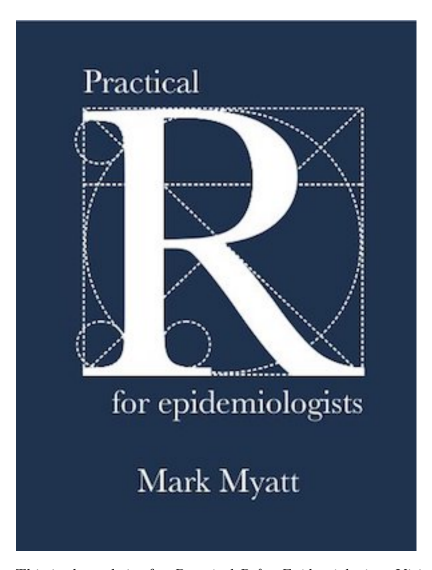
Mark Myatt and Ernest Guevarra 2018-04-29

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Welcome



This is the website for $Practical\ R$ for Epidemiologists. Visit the GitHub repository for this site or buy it as a Kindle ebook on Amazon.

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Introduction

These notes are intended as a practical introduction to using the R environment for data analysis and graphics to work with epidemiological data. Topics covered include univariate statistics, simple statistical inference, charting data, two-by-two tables, stratified analysis, chi-square test for trend, logistic regression, survival analysis, computer-intensive methods, and extending R using user-provided functions. You should be able to follow the material if you are reasonably familiar with the mechanics of statistical estimation (e.g. calculation of odds ratios and confidence intervals) and require a system that can perform simple or complex analyses to your exact specifications.

These notes are split into ten sections:

Introduction: You are reading this section now!

Introducing R: Some information about the R system, the way the R system works, how to get a copy of R, and how to start R.

Exercise 1: Read a dataset, producing descriptive statistics, charts, and perform simple statistical inference. The aim of the exercise is for you to become familiar with R and some basic R functions and objects.

Exercise 2: In this exercise we explore how to manipulate R objects and how to write functions that can manipulate and extract data and information from R objects and produce useful analyses.

Exercise 3: In this exercise we explore how R handles generalised linear models using the example of logistic regression as well as seeing how R can perform stratified (i.e. Mantel-Haenszel) analysis as well as analysing data arising from matched case-control studies.

Exercise 4: In this exercise we use R to analyse a small dataset using the methods introduced in the previous exercises.

Exercise 5: In this exercise we explore how R can be extended using add-in packages. Specifically, we will use an add-in package to perform a survival analysis.

Exercise 6: In this exercise we explore how to make your own R functions behave like R objects so that they return a data-structure that can be manipulated or interrogated by other R functions.

Exercise 7: In this exercise we explore how you can use R to produce custom graphical

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functions.

Exercise 8: In this exercise we explore some more graphical functions and create custom graphical functions that produce two variable plots, pyramid charts, Pareto charts, charts with error bars, and simple mesh-maps.

Exercise 9: In this exercise we explore ways of implementing computer-intensive methods, such as the bootstrap and computer based simulation, using standard R functions.

If you are interested in a system that is flexible, can be tailored to produce exactly the analysis you want, provides modern analytical facilities, and have a basic understanding of the mechanics of hypothesis testing and estimation then you should consider following this material.

Chapter 1

Introducing R

R is a system for data manipulation, calculation, and graphics. It provides:

- Facilities for data handling and storage
- A large collection of tools for data analysis
- Graphical facilities for data analysis and display
- A simple but powerful programming language

R is often described as an environment for working with data. This is in contrast to a package which is a collection of very specific tools. R is not strictly a statistics system but a system that provides many classical and modern statistical procedures as part of a broader data-analysis tool. This is an important difference between R and other statistical systems. In R a statistical analysis is usually performed as a series of steps with intermediate results being stored in objects. Systems such as SPSS and SAS provide copious output from (e.g.) a regression analysis whereas R will give minimal output and store the results of a fit for subsequent interrogation or use with other R functions. This means that R can be tailored to produce exactly the analysis and results that you want rather than produce an analysis designed to fit all situations.

R is a language based product. This means that you interact with R by typing commands such as:

table(SEX, LIFE)

rather than by using menus, dialog boxes, selection lists, and buttons. This may seem to be a drawback but it means that the system is considerably more flexible than one that relies on menus, buttons, and boxes. It also means that every stage of your data management and analysis can be recorded and edited and re-run at a later date. It also provides an audit trail for quality control purposes.

R is available under UNIX (including Linux), the Macintosh operating system OS X, and Microsoft Windows. The method used for starting R will vary from system to system. On UNIX systems you may need to issue the R command in a terminal session or click on an

icon or menu option if your system has a windowing system. On Macintosh systems R will be available as an application but can also be run in a terminal session. On Microsoft Windows systems there will usually be an icon on the Start menu or the desktop.

R is an open source system and is available under the *GNU general public license* (GPL) which means that it is available for free but that there are some restrictions on how you are allowed to distribute the system and how you may charge for bespoke data analysis solutions written using the R system. Details of the general public license are available from http://www.gnu.org/copyleft/gpl.html.

R is available for download from http://www.r-project.org/.

This is also the best place to get extension packages and documentation. You may also subscribe to the R mailing lists from this site. R is supported through mailing lists. The level of support is at least as good as for commercial packages. It is typical to have queries answered in a matter of a few hours.

Even though R is a free package it is more powerful than most commercial packages. Many of the modern procedures found in commercial packages were first developed and tested using R or S-Plus (the commercial equivalent of R).

When you start R it will issue a prompt when it expects user input. The default prompt is:

```
>
```

This is where you type commands that call functions that instruct R to (e.g.) read a data file, recode data, produce a table, or fit a regression. For example:

```
> table(SEX, LIFE)
```

If a command you type is not complete then the prompt will change to:

+

on subsequent lines until the command is complete:

```
> table(
+ SEX, LIFE +)
```

The > and + prompts are not shown in the example commands in the rest of this material.

The example commands in this material are often broken into shorter lines and indented for ease of understanding. The code still works as lines are split in places where R knows that a line is not complete. For example:

```
table(SEX,
     LIFE)
```

could be entered on a single line as:

```
table(SEX, LIFE)
```

In this example R knows that the command is not complete until the brackets are closed. The following example could also be written on one line:

```
salex.lreg.coeffs <-
coef(summary(salex.lreg))</pre>
```

In this case R knows that the <- operator at the end of the first line needs further input.

R maintains a history of previous commands. These can be recalled and edited using the up and down arrow keys.

Output that has scrolled off the top of the output / command window can be recalled using the window or terminal scroll bars.

Output can be saved using the sink() function with a file name: sink("results.out") to start recording output. Use the sink() function without a file name to stop recording output: sink()

You can also use clipboard functions such as copy and paste to (e.g.) copy and then paste selected chunks of output into an editor or word processor running alongside R.

All the sample data files used in the exercises in this manual are space delimited text files using the general format:

```
ID AGE IQ
1 39 94
2 41 89
3 42 83
4 30 99
5 35 94
6 44 90
7 31 94
8 39 87
```

R has facilities for working with files in different formats including (through the use of extension packages) ODBC (open database connectivity) and SQL data sources, EpiInfo, EpiData, Minitab, SPSS, SAS, S-Plus, and Stata format files.

1.1 Retrieving data

All of the exercises in this manual assume that the necessary data files are located in the current working directory. All of the data files that you require to follow this material are in a ZIP archive that can be downloaded from:

http://www.brixtonhealth.com/prfe/prfe.zip

A command such as:

```
read.table("data/fem.dat", header = TRUE)
```

retrieves the data stored in the file named fem.dat which is stored in the current working directory.

To retrieve data that is stored in files outside a different directory you need to specify the full path to the file. For example:

```
read.table("~/prfe/fem.dat", header = TRUE)
```

will retrieve the data stored in the file named fem.dat stored in the prfe directory under the user's home directory on UNIX, Linux, and OS X systems.

R follows many UNIX operating and naming conventions including the use of the backslash (\) character to specify special characters in strings (e.g. using \n to specify a new line in printed output). Windows uses the backslash (\) character to separate directory and file names in paths. This means that Windows users need to escape any backslashes in file paths using an additional backslash character. For example:

```
read.table("c:\\prfe\\fem.dat", header = TRUE)
```

will retrieve the data that is stored in the file named fem.dat which is stored in the prfe directory off the root directory of the C: drive. The Windows version of R also allows you to specify UNIX-style path names (i.e. using the forward slash (/) character as a separator in file paths). For example:

```
read.table("c:/prfe/fem.dat", header = TRUE)
```

Path names may include shortcut characters such as:

- . The current working directory
- .. Up one level in the directory tree
- ~ The user's home directory (on UNIX-based systems)

R also allows you to retrieve files from any location that may be represented by a standard uniform resource locator (URL) string. For example:

```
read.table("file://~/prfe/fem.dat", header = TRUE)
```

will retrieve the data stored in the file named fem.dat stored in the prfe directory under the users home directory on UNIX-based systems.

All of the data files used in this section are stored in the /prfe directory Brixton Health's website. This means, for example, that you can use the read.table() function specifying

```
"http://www.brixtonhealth.com/prfe/fem.dat"
```

as the URL to retrieve the data that is stored in the file named fem.dat which is stored in the /prfe directory of this guide's GitLab repository.

Chapter 2

Getting acquainted with R

In this exercise we will use R to read a dataset and produce some descriptive statistics, produce some charts, and perform some simple statistical inference. The aim of the exercise is for you to become familiar with R and some basic R functions and objects.

The first thing we will do, after starting R, is issue a command to retrieve an example dataset:

```
fem <- read.table("fem.dat", header = TRUE)</pre>
```

This command illustrates some key things about the way R works.

We are instructing R to assign (using the <- operator) the output of the read.table() function to an object called fem.

The fem object will contain the data held in the file fem.dat as an R data.frame object:

```
class(fem)
```

```
## [1] "data.frame"
```

You can inspect the contents of the fem data.frame (or any other R object) just by typing its name:

fem

```
ID AGE IQ ANX DEP SLP SEX LIFE
                                             WT
##
          39 94
## 1
       1
                   2
                        2
                             2
                                  1
                                        1
                                           2.23
       2
          41 89
                    2
                        2
                             2
                                  1
                                           1.00
          42 83
                   3
                        3
                             2
                                           1.82
## 3
       3
                                  1
                                        1
      4
          30 99
                   2
                             2
                                        1 -1.18
## 5
      5
          35 94
                   2
                        1
                             1
                                  1
                                        2 - 0.14
                             2
       6
          44 90
                                  2
                                          0.41
## 6
                  NA
```

Note that the fem object is built from other objects. These are the named vectors (columns) in the dataset:

```
names(fem)
```

```
## [1] "ID" "AGE" "IQ" "ANX" "DEP" "SLP" "SEX" "LIFE" "WT"
```

The [1] displayed before the column names refers to the numbered position of the first name in the output. These positions are known as indexes and can be used to refer to individual items. For example:

```
names(fem)[1]
```

[1] "ID"

names(fem)[8]

[1] "LIFE"

names(fem)[2:4]

[1] "AGE" "IQ" "ANX"

The data consist of 118 records:

nrow(fem)

[1] 118

each with nine variables:

ncol(fem)

[1] 9

for female psychiatric patients.

The columns in the dataset are:

ID	Patient ID
AGE	Age in years
IQ	IQ score
ANX	Anxiety (1=none, 2=mild, 3=moderate, 4=severe)
\mathbf{DEP}	Depression (1=none, 2=mild, 3=moderate or severe)
SLP	Sleeping normally (1=yes, 2=no)
\mathbf{SEX}	Lost interest in sex (1=yes, 2=no)
${f LIFE}$	Considered suicide (1=yes, 2=no)
\mathbf{WT}	Weight change (kg) in previous 6 months

The first ten records of the fem data.frame are:

```
IQ ANX DEP SLP SEX LIFE
##
       ID AGE
                                               WT
                94
                      2
                          2
                               2
                                         1
                                             2.23
## 1
           39
                                    1
## 2
        2
           41
                89
                      2
                          2
                               2
                                    1
                                         1
                                            1.00
```

```
## 3
         3
             42
                  83
                        3
                              3
                                   2
                                        1
                                                   1.82
                                               1
                        2
                              2
                                   2
         4
             30
                  99
                                        1
                                               1 - 1.18
##
##
         5
             35
                  94
                         2
                                   1
                                        1
                                               2 - 0.14
   5
                              1
                                        2
##
         6
             44
                  90
                       NA
                              1
                                   2
                                                   0.41
                              2
##
   7
         7
             31
                  94
                        2
                                 NA
                                        1
                                               1 - 0.68
                        3
                              2
                                   2
                                        1
                                               2
##
   8
         8
             39
                  87
                                                   1.59
## 9
         9
             35
                -99
                         3
                              2
                                   2
                                                 -0.55
                                        1
                                               1
                              2
             33
                         2
                                   2
                                        1
                                               1
## 10 10
                  92
                                                   0.36
```

You may check this by asking R to display all columns of the first ten records in the fem data.frame:

```
fem[1:10, ]
```

```
WT
##
       ID AGE
                 IQ ANX DEP SLP
                                    SEX LIFE
## 1
        1
            39
                 94
                        2
                             2
                                  2
                                       1
                                              1
                                                 2.23
   2
        2
                        2
                             2
                                  2
##
            41
                 89
                                       1
                                              1
                                                  1.00
##
   3
        3
            42
                 83
                        3
                             3
                                  2
                                       1
                                              1
                                                  1.82
##
   4
        4
            30
                 99
                        2
                             2
                                  2
                                       1
                                               -1.18
                                              1
        5
                        2
                             1
                                  1
                                       1
##
            35
                 94
                                              2 - 0.14
                                  2
                                       2
                                              2
##
   6
        6
            44
                 90
                       NA
                             1
                                                 0.41
        7
                        2
                             2
##
   7
            31
                 94
                                 NA
                                       1
                                              1 - 0.68
## 8
        8
            39
                 87
                        3
                             2
                                  2
                                       1
                                              2
                                                 1.59
                             2
                                  2
## 9
        9
            35
                -99
                        3
                                       1
                                              1 - 0.55
## 10 10
            33
                 92
                        2
                             2
                                  2
                                       1
                                              1
                                                 0.36
```

The space after the comma is optional. You can think of it as a *placeholder* for where you would specify the indexes for columns you wanted to display. For example:

```
fem[1:10,2:4]
```

displays the first ten rows and the second, third and fourth columns of the fem data.frame:

```
##
       AGE
             IQ ANX
## 1
        39
             94
                   2
## 2
        41
             89
                   2
## 3
        42
             83
                   3
                   2
## 4
        30
             99
                   2
## 5
        35
             94
## 6
        44
             90
                  NA
## 7
                   2
        31
             94
## 8
        39
             87
                   3
## 9
        35
            -99
                   3
                   2
## 10
        33
             92
```

NA is a special value meaning not available or missing.

You can access the contents of a single column by name:

```
fem$IQ
##
      [1]
            94
                 89
                      83
                           99
                                94
                                      90
                                           94
                                                87 -99
                                                          92
                                                               92
                                                                    94
                                                                         91
                                                                              86
                                                                                   90
                                                                                       -99
                                                                                             91
##
     [18]
            82
                 86
                      88
                           97
                                96
                                      95
                                           87
                                              103 -99
                                                          91
                                                               87
                                                                    91
                                                                         89
                                                                              92
                                                                                   84
                                                                                        94
                                                                                             92
##
     [35]
            96
                 96
                      86
                           92
                               102
                                      82
                                           92
                                                90
                                                     92
                                                          88
                                                               98
                                                                    93
                                                                         90
                                                                              91
                                                                                  -99
                                                                                        92
                                                                                             92
##
     [52]
            91
                 91
                      86
                           95
                                91
                                      96
                                         100
                                                99
                                                     89
                                                          89
                                                               98
                                                                    98 103
                                                                              91
                                                                                   91
                                                                                        94
                                                                                             91
     [69]
                 92
                      96
                                87
                                           95
                                                     95
                                                                   -99
                                                                       -99
                                                                                   92
                                                                                             93
##
            85
                           90
                                      95
                                                87
                                                          88
                                                               94
                                                                              87
                                                                                        86
     [86]
##
            92 106
                      93
                           95
                                95
                                     92
                                           98
                                               92
                                                     88
                                                          85
                                                               92
                                                                    84
                                                                         92
                                                                              91
                                                                                   86
                                                                                        92
                                                                                             89
    [103] -99
                 96
                      97
                            92
                                92
                                      98
                                           91
                                                91
                                                     89
                                                          94
                                                               90
                                                                    96
                                                                         87
                                                                                   89 -99
                                                                              86
fem$IQ[1:10]
```

```
## [1] 94 89 83 99 94 90 94 87 -99 92
```

The \$ sign is used to separate the name of the data.frame and the name of the column of interest. Note that R is case-sensitive so that IQ and iq are **not** the same.

You can also access rows, columns, and individual cells by specifying row and column positions. For example, the IQ column is the third column in the fem data.frame:

```
fem[ ,3]
##
      [1]
            94
                 89
                      83
                           99
                                94
                                     90
                                          94
                                               87 -99
                                                         92
                                                              92
                                                                   94
                                                                        91
                                                                             86
                                                                                  90
                                                                                     -99
                                                                                            91
##
     [18]
            82
                 86
                      88
                           97
                                96
                                     95
                                          87
                                              103 -99
                                                         91
                                                              87
                                                                   91
                                                                        89
                                                                             92
                                                                                  84
                                                                                       94
                                                                                            92
                           92 102
                                                                                -99
##
     [35]
            96
                 96
                      86
                                     82
                                          92
                                               90
                                                    92
                                                         88
                                                              98
                                                                   93
                                                                        90
                                                                             91
                                                                                       92
                                                                                            92
##
     [52]
            91
                 91
                      86
                           95
                                91
                                     96 100
                                               99
                                                    89
                                                         89
                                                              98
                                                                   98 103
                                                                             91
                                                                                  91
                                                                                       94
                                                                                            91
##
     [69]
            85
                 92
                      96
                           90
                                87
                                     95
                                          95
                                               87
                                                    95
                                                         88
                                                              94
                                                                  -99
                                                                      -99
                                                                             87
                                                                                  92
                                                                                       86
                                                                                            93
                                95
                                          98
##
     [86]
            92 106
                      93
                           95
                                     92
                                               92
                                                    88
                                                         85
                                                              92
                                                                   84
                                                                        92
                                                                             91
                                                                                  86
                                                                                       92
                                                                                            89
## [103]
           -99
                 96
                      97
                           92
                                92
                                     98
                                          91
                                               91
                                                    89
                                                         94
                                                              90
                                                                   96
                                                                        87
                                                                             86
                                                                                  89 -99
fem[9,]
                IQ ANX DEP SLP SEX LIFE
                                                 WT
##
      ID AGE
## 9
       9
           35 -99
                      3
                           2
                                2
                                     1
                                           1 - 0.55
fem[9,3]
```

```
## [1] -99
```

There are missing values in the IQ column which are all coded as -99. Before proceeding we must set these to the special NA value:

```
fem$IQ[fem$IQ == -99] <- NA
```

The term inside the square brackets is also an index. This type of index is used to refer to subsets of data held in an object that meet a particular condition. In this case we are instructing R to set the contents of the IQ variable to NA if the contents of the IQ variable is -99.

Check that this has worked:

fem\$IQ

```
##
      [1]
            94
                 89
                      83
                           99
                                 94
                                      90
                                           94
                                                87
                                                          92
                                                               92
                                                                    94
                                                                         91
                                                                              86
                                                                                   90
                                                                                        NA
                                                                                             91
                                                     NA
##
     [18]
            82
                 86
                      88
                           97
                                 96
                                      95
                                           87
                                               103
                                                     NA
                                                          91
                                                               87
                                                                    91
                                                                         89
                                                                              92
                                                                                   84
                                                                                        94
                                                                                             92
                                                                                             92
##
     [35]
            96
                 96
                      86
                           92 102
                                      82
                                           92
                                                90
                                                     92
                                                          88
                                                               98
                                                                    93
                                                                         90
                                                                              91
                                                                                   NA
                                                                                        92
##
     [52]
            91
                 91
                      86
                           95
                                91
                                      96
                                         100
                                                99
                                                     89
                                                          89
                                                               98
                                                                    98
                                                                        103
                                                                              91
                                                                                   91
                                                                                        94
                                                                                             91
     [69]
##
            85
                 92
                      96
                                 87
                                      95
                                           95
                                                87
                                                     95
                                                               94
                                                                    NA
                                                                         NA
                                                                              87
                                                                                   92
                                                                                        86
                                                                                             93
                           90
                                                          88
     [86]
##
            92 106
                      93
                           95
                                 95
                                      92
                                           98
                                                92
                                                     88
                                                          85
                                                               92
                                                                    84
                                                                         92
                                                                              91
                                                                                   86
                                                                                        92
                                                                                             89
   [103]
                 96
                      97
                            92
                                92
                                      98
                                           91
                                                91
                                                     89
                                                          94
                                                               90
                                                                    96
                                                                         87
##
            NA
                                                                              86
                                                                                   89
                                                                                        NA
```

We can now compare the groups who have and have not considered suicide. For example:

```
by(fem$IQ, fem$LIFE, summary)
```

Look at the help for the by() function:

```
help(by)
```

Note that you may use ?by as a shortcut for help(by).

The by() function applies another function (in this case the summary() function) to a column in a data frame (in this case fem\$IQ) split by the value of another variable (in this case fem\$LIFE).

It can be tedious to always have to specify a data.frame each time we want to use a particular variable. We can fix this problem by 'attaching' the data.frame:

attach(fem)

```
## The following objects are masked from fem (pos = 3):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
## The following objects are masked from fem (pos = 5):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
  The following objects are masked from fem (pos = 6):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
##
  The following objects are masked from fem (pos = 11):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
```

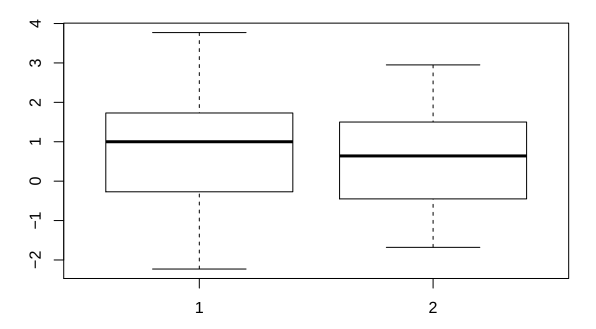
We can now refer to the columns in the fem data.frame without having to specify the name of the data.frame. This time we will produce summary statistics for WT by LIFE:

```
by(WT, LIFE, summary)
```

```
## LIFE: 1
```

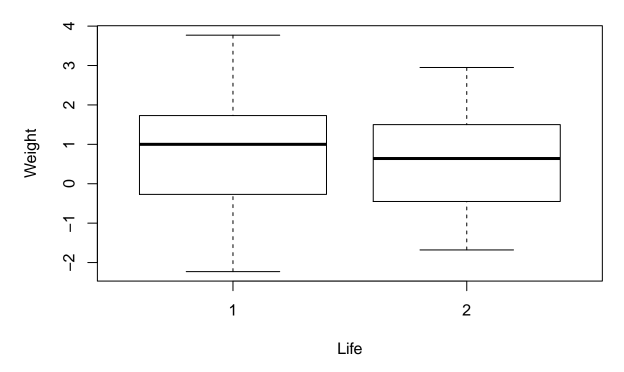
We can view the same data as a box and whisker plot:

```
boxplot(WT ~ LIFE)
```



We can add axis labels and a title to the graph:

Weight BY Life



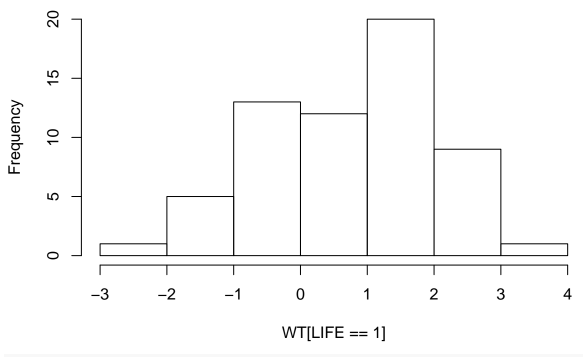
A more descriptive title might be "Weight Change BY Considered Suicide".

The groups do not seem to differ much in their medians and the distributions appear to be reasonably symmetrical about their medians with a similar spread of values.

We can look at the distribution as histograms:

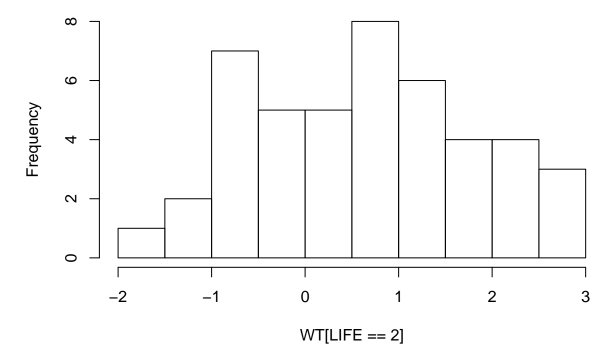
```
hist(WT[LIFE == 1])
```

Histogram of WT[LIFE == 1]



hist(WT[LIFE == 2])

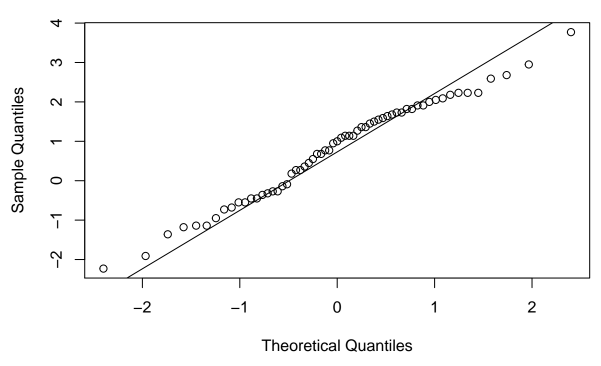
Histogram of WT[LIFE == 2]



and check the assumption of normality using quantile-quantile plots:

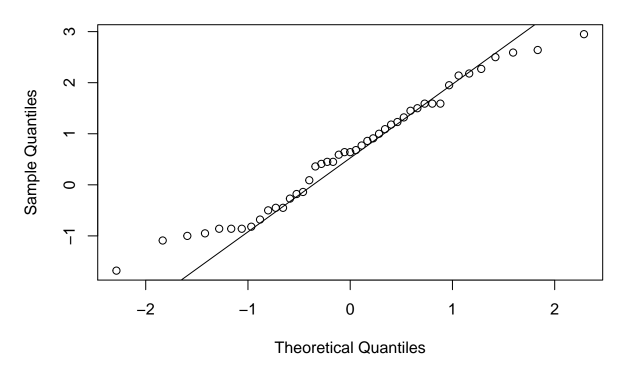
```
qqnorm(WT[LIFE == 1])
qqline(WT[LIFE == 1])
```

Normal Q-Q Plot



```
qqnorm(WT[LIFE == 2])
qqline(WT[LIFE == 2])
```

Normal Q-Q Plot



or by using a formal test:

```
shapiro.test(WT[LIFE == 1])
##
##
    Shapiro-Wilk normality test
##
          WT[LIFE == 1]
## data:
## W = 0.98038, p-value = 0.4336
shapiro.test(WT[LIFE == 2])
##
##
    Shapiro-Wilk normality test
##
          WT[LIFE == 2]
## data:
## W = 0.97155, p-value = 0.3292
```

Remember that we can use the by() function to apply a function to a data.frame, including statistical functions such as shapiro.test():

```
by(WT, LIFE, shapiro.test)

## LIFE: 1
##
## Shapiro-Wilk normality test
##
```

```
## data: dd[x,]
## W = 0.98038, p-value = 0.4336
##
## ------
## LIFE: 2
##
## Shapiro-Wilk normality test
##
## data: dd[x,]
## W = 0.97155, p-value = 0.3292
```

We can also test whether the variances differ significantly using *Bartlett's test* for the homogeneity of variances:

```
bartlett.test(WT, LIFE)
```

```
##
## Bartlett test of homogeneity of variances
##
## data: WT and LIFE
## Bartlett's K-squared = 0.32408, df = 1, p-value = 0.5692
```

There is no significant difference between the two variances.

Many functions in R have a *formula interface* that may be used to specify multiple variables and the relations between multiple variables. We could have used the formula interface with the bartlett.test() function:

```
bartlett.test(WT ~ LIFE)
```

```
##
## Bartlett test of homogeneity of variances
##
## data: WT by LIFE
## Bartlett's K-squared = 0.32408, df = 1, p-value = 0.5692
```

Having checked the normality and homogeneity of variance assumptions we can proceed to carry out a t-test:

```
t.test(WT ~ LIFE, var.equal = TRUE)
```

```
##
## Two Sample t-test
##
## data: WT by LIFE
## t = 0.59869, df = 104, p-value = 0.5507
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.3382365 0.6307902
```

groups:

LIFE

##

```
## sample estimates:
## mean in group 1 mean in group 2
##
         0.7867213
                         0.6404444
```

There is no evidence that the two groups differ in weight change in the previous six months.

We could still have performed a t-test if the variances were not homogenous by setting the var.equal parameter of the t.test() function to FALSE:

```
t.test(WT ~ LIFE, var.equal = FALSE)
##
##
   Welch Two Sample t-test
##
## data: WT by LIFE
## t = 0.60608, df = 98.866, p-value = 0.5459
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.3326225 0.6251763
## sample estimates:
## mean in group 1 mean in group 2
         0.7867213
                          0.6404444
##
or performed a non-parametric test:
wilcox.test(WT ~ LIFE)
##
   Wilcoxon rank sum test with continuity correction
##
##
## data: WT by LIFE
## W = 1488, p-value = 0.4622
## alternative hypothesis: true location shift is not equal to 0
An alternative, and more general, non-parametric test is:
kruskal.test(WT ~ LIFE)
##
##
   Kruskal-Wallis rank sum test
##
## data: WT by LIFE
## Kruskal-Wallis chi-squared = 0.54521, df = 1, p-value = 0.4603
We can use the table() function to examine the differences in depression between the two
```

table(DEP, LIFE)

```
## DEP 1 2
## 1 0 26
## 2 42 24
## 3 16 1
```

##

##

data: tab

Fisher's Exact Test for Count Data

The two distributions look very different from each other. We can test this using a chi-square test on the table:

```
chisq.test(table(DEP, LIFE))
##
##
    Pearson's Chi-squared test
##
## data: table(DEP, LIFE)
## X-squared = 43.876, df = 2, p-value = 2.968e-10
Note that we passed the output of the table() function directly to the chisq.test()
function. We could have saved the table as an object first and then passed the object to the
chisq.test() function:
tab <- table(DEP, LIFE)
chisq.test(tab)
##
##
    Pearson's Chi-squared test
##
## data: tab
## X-squared = 43.876, df = 2, p-value = 2.968e-10
The tab object contains the output of the table() function:
class(tab)
## [1] "table"
tab
##
      LIFE
           2
## DEP
        1
##
     1
        0 26
     2 42 24
##
     3 16 1
##
We can pass this table object to another function. For example:
fisher.test(tab)
```

```
## p-value = 1.316e-12
## alternative hypothesis: two.sided
```

When we are finished with the tab object we can delete it using the rm() function:

```
rm(tab)
```

You can see a list of available objects using the ls() function:

```
ls()
```

```
## [1] "fem"
```

This should just show the fem object.

Fisher's Exact Test for Count Data

We can examine the association between loss of interest in sex and considering suicide in the same way:

```
tab <- table(SEX, LIFE)
tab
      LIFE
##
## SEX 1
           2
     1 58 38
##
     2 5 12
##
fisher.test(tab)
##
   Fisher's Exact Test for Count Data
##
##
## data: tab
## p-value = 0.03175
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
     1.080298 14.214482
##
## sample estimates:
## odds ratio
     3.620646
##
```

Note that with a two-by-two table the fisher.test() function produces an estimate of, and confidence intervals for, the odds ratio. Again, we will delete the tab object:

```
rm(tab)
```

We could have performed the Fisher exact test without creating the tab object by passing the output of the table() function directly to the fisher.test() function:

```
fisher.test(table(SEX, LIFE))
##
```

```
##
## data: table(SEX, LIFE)
## p-value = 0.03175
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 1.080298 14.214482
## sample estimates:
## odds ratio
## 3.620646
```

Choose whichever method you find easiest but remember that it is easy to save the results of any function for later use.

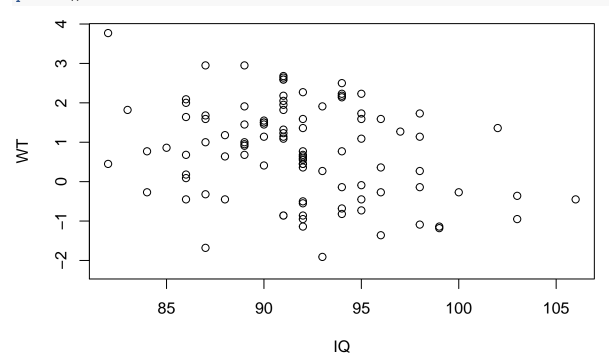
We can explore the correlation between two variables using the cor() function:

```
cor(IQ, WT, use = "pairwise.complete.obs")
```

```
## [1] -0.2917158
```

or by using a scatter plot:

plot(IQ, WT)



and by a formal test:

```
cor.test(IQ, WT)
```

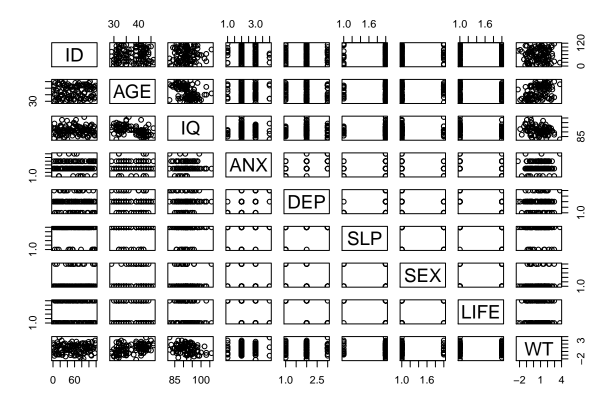
```
##
## Pearson's product-moment correlation
##
```

```
## data: IQ and WT
## t = -3.0192, df = 98, p-value = 0.003231
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## -0.4616804 -0.1010899
## sample estimates:
## cor
## -0.2917158
```

With some functions you can pass an entire data frame rather than a list of variables:

```
cor(fem, use = "pairwise.complete.obs")
```

```
##
              ID
                       AGE
                                   ΙQ
                                             ANX
## ID
       1.00000000 0.03069077 0.0370598672 -0.02941825 -0.0554147209
## AGE
       0.03069077
                 1.00000000 -0.4345435680 0.06734300 -0.0387049246
       0.03705987 -0.43454357
                          1.0000000000 -0.02323787 -0.0001307404
## IQ
      1.00000000 0.5437946347
## ANX
## DEP
      -0.05541472 -0.03870492 -0.0001307404 0.54379463 1.0000000000
## SLP
      -0.07268743  0.02606547  0.0812993104  0.22317875
                                                 0.5248724551
## SEX
       ## LIFE -0.05604349 -0.10300193 -0.0915396469 -0.34211268 -0.6139017253
## WT
       0.02640131 0.41574411 -0.2917157832 0.11817532
                                                 0.0233742465
              SLP
##
                        SEX
                                 LIFE
## ID
      ## AGE
       ## IQ
       0.081299310 - 0.05365587 - 0.09153965 - 0.291715783
## ANX
       0.223178752 -0.21062493 -0.34211268 0.118175321
## DEP
       0.524872455 -0.30584223 -0.61390173
                                     0.023374247
## SLP
       1.000000000 -0.29053971 -0.35186578 -0.009259774
## SEX
      -0.290539709 1.00000000 0.22316967 -0.027826514
## LIFE -0.351865775 0.22316967 1.00000000 -0.058605326
## WT
      -0.009259774 -0.02782651 -0.05860533 1.000000000
pairs(fem)
```



The output can be a little confusing particularly if it includes categorical or record identifying variables. To avoid this we can create a new object that contains only the columns we are interested in using the column binding cbind() function:

```
newfem <- cbind(AGE, IQ, WT)
cor(newfem, use = "pairwise.complete.obs")</pre>
```

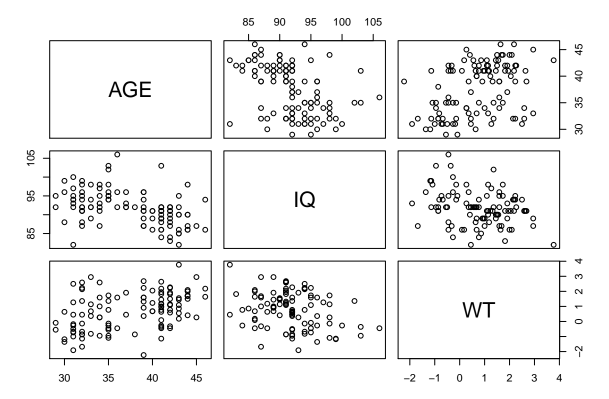
```
## AGE IQ WT

## AGE 1.0000000 -0.4345436 0.4157441

## IQ -0.4345436 1.0000000 -0.2917158

## WT 0.4157441 -0.2917158 1.0000000

pairs(newfem)
```



When we have finished with the newfem object we can delete it:

```
rm(newfem)
```

There was no real need to create the newfem object as we could have fed the output of the cbind() function directly to the cor() or pairs() function:

```
cor(cbind(AGE, IQ, WT), use = "pairwise.complete.obs")
```

```
## AGE IQ WT

## AGE 1.0000000 -0.4345436 0.4157441

## IQ -0.4345436 1.0000000 -0.2917158

## WT 0.4157441 -0.2917158 1.0000000

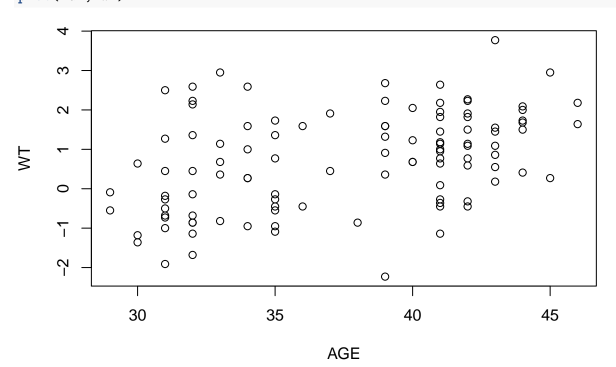
pairs(cbind(AGE, IQ, WT))
```



It is, however, easier to work with the newfem object rather than having to retype the cbind() function. This is particularly true if you wanted to continue with an analysis of just the three variables.

The relationship between AGE and WT can be plotted using the $\operatorname{plot}()$ function:

plot(AGE, WT)



And tested using the cor() and cor.test() functions:

```
cor(AGE, WT, use = "pairwise.complete.obs")
## [1] 0.4157441
cor.test(AGE, WT)
##
##
   Pearson's product-moment correlation
##
## data: AGE and WT
## t = 4.6841, df = 105, p-value = 8.457e-06
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.2452434 0.5612979
## sample estimates:
##
         cor
## 0.4157441
Or by using the linear modelling lm() function:
summary(lm(WT ~ AGE))
##
## Call:
## lm(formula = WT ~ AGE)
##
## Residuals:
                  1Q
                       Median
                                    3Q
                                            Max
## -3.10678 -0.85922 -0.05453 0.71434 2.70874
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -3.25405
                           0.85547 -3.804 0.00024 ***
## AGE
                           0.02261
                                   4.684 8.46e-06 ***
                0.10592
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.128 on 105 degrees of freedom
     (11 observations deleted due to missingness)
## Multiple R-squared: 0.1728, Adjusted R-squared:
## F-statistic: 21.94 on 1 and 105 DF, p-value: 8.457e-06
```

We use the **summary()** function here to extract summary information from the output of the **lm()** function.

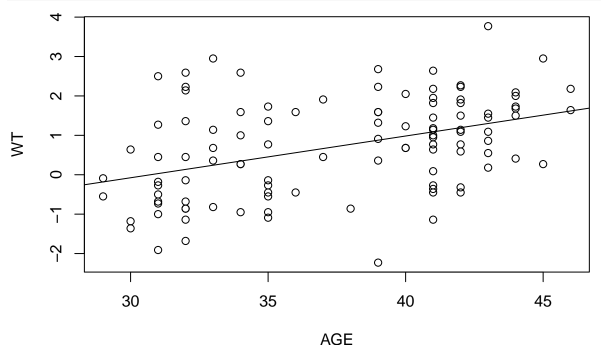
It is often more useful to use lm() to create an object:

```
fem.lm \leftarrow lm(WT \sim AGE)
```

And use the output in other functions:

```
summary(fem.lm)
```

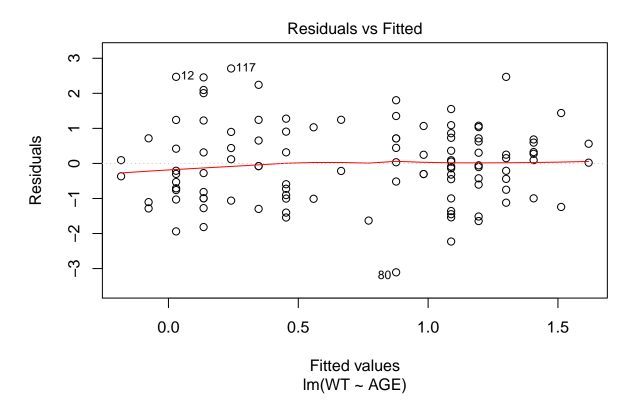
```
##
## Call:
## lm(formula = WT ~ AGE)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                     3Q
                                             Max
## -3.10678 -0.85922 -0.05453
                               0.71434
                                        2.70874
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -3.25405
                           0.85547
                                    -3.804 0.00024 ***
## AGE
                0.10592
                           0.02261
                                      4.684 8.46e-06 ***
## ---
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 1.128 on 105 degrees of freedom
     (11 observations deleted due to missingness)
## Multiple R-squared: 0.1728, Adjusted R-squared: 0.165
## F-statistic: 21.94 on 1 and 105 DF, p-value: 8.457e-06
plot(AGE, WT)
abline(fem.lm)
```

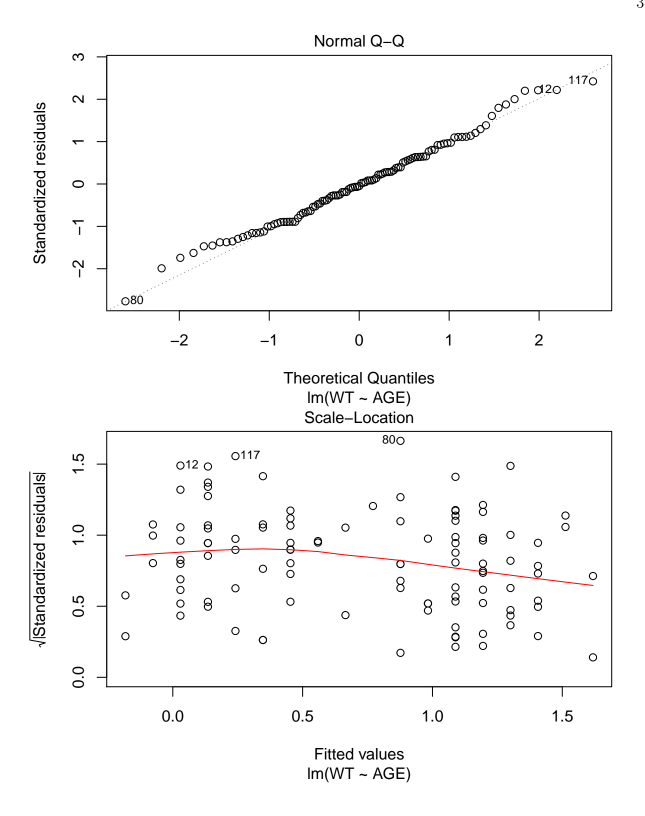


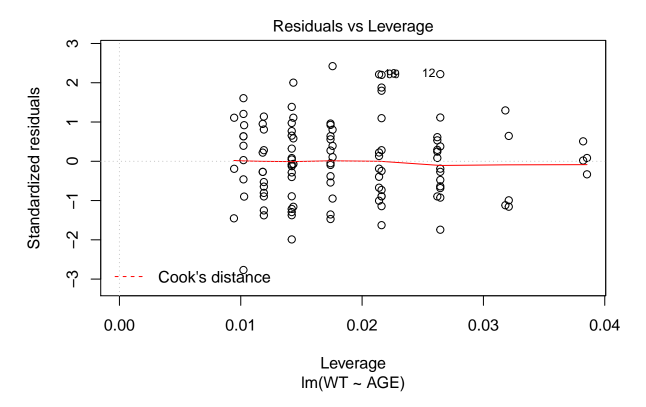
In this case we are passing the intercept and slope information held in the fem.lm object to the abline() function which draws a regression line. The abline() function adds to an existing plot. This means that you need to keep the scatter plot of AGE and WT open before issuing the abline() function call.

A useful function to apply to the fem.lm object is plot() which produces diagnostic plots of the linear model:

plot(fem.lm)







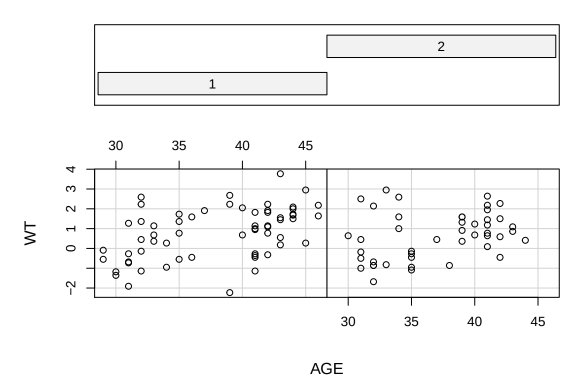
Objects created by the lm() function (or any of the modelling functions) can use up a lot of memory so we should remove them when we no longer need them:

```
rm(fem.lm)
```

It might be interesting to see whether a similar relationship exists between AGE and WT for those who have and have not considered suicide. This can be done using the coplot() function:

```
coplot(WT ~ AGE | as.factor(LIFE))
```

Given: as.factor(LIFE)

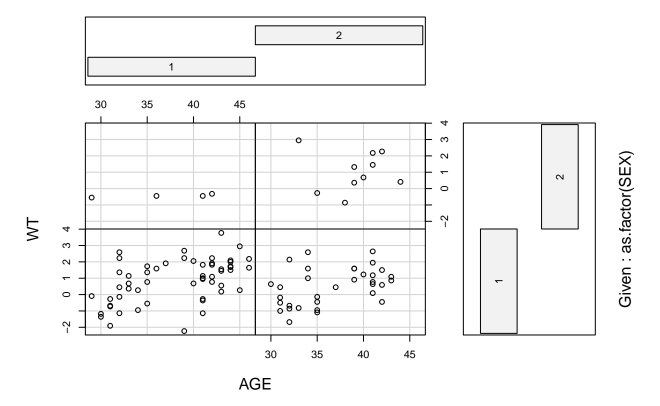


##
Missing rows: 21, 22, 31, 43, 44, 45, 69, 81, 101, 104, 114, 115

The two plots looks similar. We could also use coplot() to investigate the relationship between AGE and WT for categories of both LIFE and SEX:

```
coplot(WT ~ AGE | as.factor(LIFE) * as.factor(SEX))
```





##
Missing rows: 12, 17, 21, 22, 31, 43, 44, 45, 66, 69, 81, 101, 104, 105, 114, 115
although the numbers are too small for this to be useful here.

We used the as.factor() function with the coplot() function to ensure that R was aware that the LIFE and SEX columns hold categorical data.

We can check the way variables are stored using the data.class() function:

data.class(fem\$SEX)

[1] "numeric"

We can 'apply' this function to all columns in a data frame using the sapply() function:

sapply(fem, data.class)

```
## ID AGE IQ ANX DEP SLP SEX
## "numeric" "numeric" "numeric" "numeric" "numeric" "numeric"
## LIFE WT
## "numeric" "numeric"
```

The **sapply()** function is part of a group of functions that apply a specified function to data objects:

Function(s)	Applies a function to
apply()	rows and columns of matrices, arrays, and tables
lapply()	components of lists and data.frames
<pre>sapply()</pre>	components of lists and data.frames
<pre>mapply()</pre>	components of lists and data.frames
tapply()	subsets of data

Related functions are aggregate() which compute summary statistics for subsets of data, by() which applies a function to a data.frame split by factors, and sweep() which applies a function to an array.

The parameters of most R functions have default values. These are usually the most used and most useful parameter values for each function. The cor.test() function, for example, calculates Pearson's product moment correlation coefficient by default. This is an appropriate measure for data from a bivariate normal distribution. The DEP and ANX variables contain ordered data. An appropriate measure of correlation between DEP and ANX is Kendall's tau. This can be obtained using:

```
cor.test(DEP, ANX, method = "kendall")
```

```
##
## Kendall's rank correlation tau
##
## data: DEP and ANX
## z = 5.5606, p-value = 2.689e-08
## alternative hypothesis: true tau is not equal to 0
## sample estimates:
## tau
## 0.4950723
```

Before we finish we should save the fem data.frame so that next time we want to use it we will not have to bother with recoding the missing values to the special NA value. This is done with the write.table() function:

```
write.table(fem, file = "newfem.dat", row.names = FALSE)
```

Everything in R is either a function or an object. Even the command to quit R is a function:

```
q()
```

When you call the q() function you will be asked if you want to save the workspace image. If you save the workspace image then all of the objects and functions currently available to you will be saved. These will then be automatically restored the next time you start R in the current working directory.

For this exercise there is no need to save the workspace image so click the **No** or **Don't Save** button (GUI) or enter **n** when prompted to save the workspace image (terminal).

2.1 Summary

- R is a functional system. Everything is done by calling functions.
- R provides a large set of functions for descriptive statistics, charting, and statistical inference.
- Functions can be chained together so that the output of one function is the input of another function.
- R is an object oriented system. We can use functions to create objects that can then be manipulated or passed to other functions for subsequent analysis.

Chapter 3

Manipulating objects and creating new functions

In this exercise we will explore how to manipulate R objects and how to write functions that can manipulate and extract data and information from R objects and produce useful analyses.

Before we go any further we should start R and retrieve a dataset:

```
salex <- read.table("salex.dat", header = TRUE, na.strings = "9")</pre>
```

Missing values are coded as 9 throughout this dataset so we can use the na.strings parameter of the read.table() function to replace all 9's with the special NA code when we retrieve the dataset. Check that this works by examining the salex data.frame:

salex

##		ILL	HAM	BEEF	EGGS	MUSHROOM	PEPPER	PORKPIE	PASTA	RICE	LETTUCE	OTAMOT
##	1	1	1	1	1	1	1	2	2	2	2	2
##	2	1	1	1	1	2	2	1	2	2	2	1
##	3	1	1	1	1	1	1	1	1	1	1	2
##	4	1	1	1	1	2	2	2	2	2	1	1
##	5	1	1	1	1	1	1	1	1	1	1	1
##	6	1	1	1	1	2	2	2	2	2	2	1
##	7	1	1	1	1	1	1	1	2	2	2	2
##	8	1	1	2	1	1	1	2	1	1	1	2
##	9	1	1	1	1	2	1	1	2	1	2	2
##	10	1	1	1	1	2	1	1	1	1	1	1
##	11	1	2	2	1	1	1	2	2	2	1	1
##	12	1	1	1	1	2	2	2	2	2	2	2
##	13	2	2	1	2	2	2	1	2	2	2	1
##	14	1	1	1	1	2	2	2	1	1	2	1
##	15	1	1	1	1	1	1	2	1	1	2	2

									_	_		_
##	16	1	1	1	1	1	1	1	2	2	2	2
##	17	1	1	1	1	1	1	1	1	1	1	1
##	18	2	1	1	2	2	2	2	2	2	2	2
##	19	2	1	1	1	1	2	2	1	1	2	1
##	20	2	1	1	2	2	2	2	2	2	2	2
##	21	2	2	2	2	2	2	2	2	2	2	2
##	22	1	1	1	1	2	2	2	2	2	1	1
	23	1	2	1	2	2	2	2	1	1	2	1
##		1	1	1	1	2	1	2	1	1	2	2
##		1	1	1	2	1	1	1	1	1	1	1
##	26	1	1	2	1	1	1	2	2	2	1	1
##	27	1	1	1	1	2	2	1	2	1	1	1
	28	1	1	1	1	1	1	2	1	1	2	2
##	29	1	2	1	1	1	NA	2	1	1	1	1
##	30	1	1	1	2	2	2	1	2	2	2	2
##	31	1	1	1	1	1	2	2	1	1	2	2
##	32	1	1	1	1	1	2	NA	2	1	1	1
##	33	1	1	1	1	2	2	2	1	2	2	2
##	34	1	1	1	1	1	2	2	2	2	1	1
##	35	1	1	1	1	1	1	1	1	2	2	1
##	36	2	2	1	2	2	2	2	2	2	2	2
	37	1	1	1	1	1	1	2	1	1	1	1
##		1	1	1	2	2	2	1	1	1	1	2
##		1	1	1	1	1	1	1	2	2	1	2
##		1	1	1	1	1	1	1	2	2	1	1
	41	1	1	1	2	2	1	2	1	1	1	1
##		1	1	1	2	2	2	2	2	2	2	2
##		1	1	1	1	1	1	2	1	1	1	1
##		1	2	1	2	2	2	1	2	2	1	2
##		1	1	1	1	1	2	2	2	1	1	1
##		1	1	1	2	2	2	2	1	1	1	1
##		1	1	1	1	2	2	2	2	1	1	2
##		1	1	1	1	1	NA	1	1	1	2	2
##		1	1	1	1	2	1	2	2	1	1	1
##	50	1	2	1	1	2	2	2	1	2	2	1
##	51	2	2	1	2	2	2	2	2	2	2	2
##	52	2	1	1	2	2	2	2	1	2	2	1
##	53	2	1	1	2	2	2	1	2	2	2	1
##	54	2	1	1	2	1	2	1	2	2	2	1
##		2	1	1	1	1	1	2	2	1	2	2
##		2	1	1	2	2	2	2	2	2	2	1
##		2	1	1	1	1	1	1	2	2	2	2
##		2	1	1	1	2	2	1	2	1	2	2
##		2	1	1	2	2	2	2	2	2	2	2
##		2	2	2	2	2	2	1	2	2	2	2
##	00	2	_	۷	_	۷	۷	1	_	۷	2	

##	61	2	1	1	2 2	2	1	2	2	2	2
##	62	2	1	2	2 2	2	2	2	2	1	1
##	63	1	1	1	1 1	. 1	2	2	2	2	1
##	64	2	1	1	2 2	2	2	2	2	2	2
##	65	2	1	1	1 1	. 2	1	2	1	2	2
##	66	2	2	1	2 2	2	2	2	2	2	2
##	67	2	2	1	2 2	2	2	2	2	2	2
##	68	2	1	1	2 1	. 1	1	1	2	2	1
##	69	2	2	1	2 2	2	2	2	2	2	2
##	70	2	2	1	2 2	2	2	2	2	2	2
##	71	1	1	2	2 2	2	1	2	1	2	2
##	72	2	1	2	1 NA	NA.	2	2	2	2	1
##	73	1	1	1	1 2		1	2	2	2	2
##		1	1	2	1 NA		2	1	1	1	1
##		1	1	2	2 2		2	1	2	1	1
##		1	1	1	1 2		1	1	2	2	2
##		1	1	1 1	NA NA		1	2	1	1	1
##		COLES			PEACHCAKE				ALMONDS		
##	1		2	2	2	2	2	2	2		
##			2	2	2	2		2	2		
##			2	1	2	1	2	2	2		
##			2	2	2	1	2	2	2		
##			1	2	2	1	2	1	2		
##			1	1	2	1	2	2	2		
##			1	1	1	2		2	2		
##			1	1	2	2		1	2		
##			2	2	2	2		1	2		
	10		1	1	2	2		1	1		
##	11		2	2	2	2		2	N A		
##	12		2	1	2	1	2	2	2		
##			2	1	2	2		2	NA		
##			1	1	2	2		1	2		
##			1		2	2		1			
##			1	2		2		2			
##			1			2		2			
##			2			2		2			
##			1			2		2			
##			2			1		2			
##			2			2		2			
##			2			1		2			
##			1			2		2			
##			1			2		1			
##			1	2		2		1			
##			1	2		2		1	2		
##			1	1		1			2		
			_	-	-	_	_	-	_		

##	28	2	1	2	2	2	2	NA
##	29	1	1	2	2	2	2	NA
##	30	2	2	2	2	2	2	2
##	31	2	2	2	2	2	2	2
##	32	2	2	2	2	2	2	2
##	33	1	2	2	2	2	2	2
##	34	1	2	2	2	2	1	2
##	35	1	2	2	2	2	1	2
##	36	2	2	2	2	2	2	NA
##	37	1	1	2	1	2	1	2
##	38	2	2	2	2	2	2	2
##	39	2	2	2	1	2	2	2
##	40	1	2	2	2	2	2	2
##	41	1	1	2	2	NA	1	NA
##	42	2	2	2	2	2	2	NA
##	43	1	1	2	2	2	2	NA
##	44	2	2	2	2	2	2	2
##	45	1	2	2	2	2	1	2
##	46	1	2	2	2	2	1	2
##	47	2	2	2	NA	2	1	2
##	48	2	1	2	2	2	2	2
	49	1	1	2	2	2	1	2
##		NA	2	2	1	2	1	1
##		2	2	2	2	2	2	NA
	52	2	2	2	1	2	2	1
##		2	2	2	2	1	2	2
	54	2	2	2	2	2	2	2
##		2	1	2	2	2	2	2
	56	2	2	2	2	1	2	2
##	57	1	2	2	2	2	2	1
	58	2	1	1	2	2	2	2
##		2	1	1	2	2	1	2
##		2	2	2	2	2	2	2
##		2	1	2	2	2	1	1
##		2	1	2	2	2	2	2
##		1	2	2	1	1	2	2
##		2	2	2	2	2	2	2
##		1	1	2	2	2	1	2
##		2	2	2	2	2	1	NA
##		2	1	2	2	2	2	2
##		2	2	2	2	2	2	2
##		2	1	2	2	2	2	2
##		2	2	2	2	2	2	2
##	71	2	2	2	2	2	2	2
	72	2	2	2	2	2	1	2
			_	2	2	_	_	

##	73	2	2	2	2	2	2	2
##	74	1	1	2	1	2	2	2
##	75	1	1	2	2	2	2	NA
##	76	2	2	2	2	2	2	NA
##	77	1	1	2	2	2	2	2

names(salex)

##	[1]	"ILL"	"HAM"	"BEEF"	"EGGS"	"MUSHROOM"
##	[6]	"PEPPER"	"PORKPIE"	"PASTA"	"RICE"	"LETTUCE"
##	[11]	"OTAMOT"	"COLESLAW"	"CRISPS"	"PEACHCAKE"	"CHOCOLATE"
##	Г16Т	"FRUTT"	"TRIFIE"	"ALMONDS"		

This data comes from a food-borne outbreak. On Saturday 17th October 1992, eighty-two people attended a buffet meal at a sports club. Within fourteen to twenty-four hours, fifty-one of the participants developed diarrhoea, with nausea, vomiting, abdominal pain and fever.

The columns in the dataset are as follows:

ILL	Ill or not-ill
HAM	Baked ham
BEEF	Roast beef
EGGS	Eggs
MUSHROOM	Mushroom flan
PEPPER	Pepper flan
PORKPIE	Pork pie
PASTA	Pasta salad
RICE	Rice salad
LETTUCE	Lettuce
TOMATO	Tomato salad
COLESLAW	Coleslaw
CRISPS	Crisps
PEACHCAKE	Peach cake
CHOCOLATE	Chocolate cake
FRUIT	Tropical fruit salad
TRIFLE	Trifle
ALMONDS	Almonds

Data is available for seventy-seven of the eighty-two people who attended the sports club buffet. All of the variables are coded 1=yes, 2=no.

We can use the attach() function to make it easier to access our data:

attach(salex)

The following objects are masked from salex (pos = 11):

```
##
```

```
## ALMONDS, BEEF, CHOCOLATE, COLESLAW, CRISPS, EGGS, FRUIT, HAM,

ILL, LETTUCE, MUSHROOM, PASTA, PEACHCAKE, PEPPER, PORKPIE,

RICE, TOMATO, TRIFLE
```

The two-by-two table is a basic epidemiological tool. In analysing data from a food-borne outbreak collected as a retrospective cohort study, for example, we would tabulate each exposure (suspect foodstuffs) against the outcome (illness) and calculate risk ratios and confidence intervals. R has no explicit function to calculate risk ratios from two-by-two tables but we can easily write one ourselves.

The first step in writing such a function would be to create the two-by-two table. This can be done with the table() function. We will use a table of HAM by ILL as an illustration:

```
table(HAM, ILL)
```

This command produces the following output:

```
## ILL
## HAM 1 2
## 1 46 17
## 2 5 9
```

We can manipulate the output directly but it is easier if we instruct R to save the output of the table() function in an object:

```
tab <- table(HAM, ILL)
```

The tab object contains the output of the table() function:

tab

```
## ILL
## HAM 1 2
## 1 46 17
## 2 5 9
```

As it is stored in an object we can examine its contents on an item by item basis.

The tab object is an object of class table:

```
class(tab)
```

```
## [1] "table"
```

We can extract data from a table object by using indices or row and column co-ordinates:

```
tab[1,1]
```

```
## [1] 46
```

```
tab[1,2]
```

```
## [1] 17
```

```
tab[2,1]
```

[1] 5

The numbers in the square brackets refer to the **position** (as row and column co-ordinates) of the data item in the table **not** the **values** of the variables. We can extract data using the values of the row and column variables by enclosing the index values in double quotes ("). For example:

```
tab["1","1"]
```

[1] 46

The two methods of extracting data may be combined. For example:

```
tab[1,"1"]
```

```
## [1] 46
```

We can calculate a risk ratio using the extracted data:

```
(tab[1,1]/(tab[1,1]+tab[1,2]))/(tab[2,1]/(tab[2,1]+tab[2,2]))
```

Which returns a risk ratio of

```
## [1] 2.044444
```

This is a tedious calculation to have to type in every time you need to calculate a risk ratio from a two-by-two table. It would be better to have a function that calculates and displays the risk ratio automatically. Fortunately, R allows us to do just that.

The function() function allows us to create new functions in R:

```
tab2by2 <- function(exposure, outcome) {}</pre>
```

This creates an empty function called tab2by2 that expects two parameters called exposure and outcome. We could type the whole function in at the R command prompt but it is easier to use a text editor:

```
fix(tab2by2)
```

This will start an editor with the empty tab2by2() function already loaded. We can now edit this function to make it do something useful:

```
function(exposure, outcome)
  {
  tab <- table(exposure, outcome)
  a <- tab[1,1]
  b <- tab[1,2]
  c <- tab[2,1]
  d <- tab[2,2]</pre>
```

```
rr <- (a / (a + b)) / (c / (c + d))
print(tab)
print(rr)
}</pre>
```

Once you have made the changes shown above, check your work, save the file, and quit the editor. Before proceeding we should examine the tab2by2() function to make sure we understand what the function will do:

- The first line defines tab2by2 as a function that expects to be given two parameters which are called exposure and outcome.
- The body of the function (i.e. the work of the function) is enclosed within curly brackets ({}).
- The first line of the body of the function creates a table object (tab) using the variables specified when the tab2by2() function is called (these are the parameters exposure and outcome).
- The next line creates four new objects (called a, b, c, and d) which contain the values of the four cells in the two-by-two table.
- The following line calculates the risk ratio using the objects a, b, c, and d and stores the result of the calculation in an object called rr.
- The final two lines print the contents of the tab and rr objects.

Let's try the tab2by2() function with our test data:

```
tab2by2(HAM, ILL)
```

```
## outcome

## exposure 1 2

## 1 46 17

## 2 5 9

## [1] 2.044444
```

The tab2by2() function displays a table of HAM by ILL followed by the risk ratio calculated from the data in the table.

Try producing another table:

```
tab2by2(PASTA, ILL)
```

```
## outcome
## exposure 1 2
## 1 25 3
## 2 26 23
## [1] 1.682692
```

Have a look at the R objects available to you:

ls()

```
## [1] "fem" "salex" "tab" "tab2by2"
```

Note that there are no a, b, c, d, or rr objects.

Examine the tab object:

tab

```
## ILL
## HAM 1 2
## 1 46 17
## 2 5 9
```

This is the table of HAM by ILL that you created earlier **not** the table of PASTA by ILL that was created by the tab2by2() function.

The tab, a, b, c, d, and rr objects in the tab2by2() function are local to that function and do not change anything outside of that function. This means that the tab object inside the function is independent of any object of the same name outside of the function.

When a function completes its work, all of the objects that are local to that function are automatically removed. This is useful as it means that you can use object names inside functions that will not interfere with objects of the same name that are stored elsewhere. It also means that you do not clutter up the R workspace with temporary objects.

Just to prove that tab in the tab2by2() function exists only in the tab2by2() function we can delete the tab object from the R workspace:

```
rm(tab)
```

Now try another call to the tab2by2() function:

tab2by2(FRUIT, ILL)

```
## outcome

## exposure 1 2

## 1 1 4

## 2 49 22

## [1] 0.2897959
```

Now list the R objects available to you:

ls()

```
## [1] "fem" "salex" "tab2by2"
```

Note that there are no tab, a, b, c, d, or rr objects.

The tab2by2() function is very limited. It only displays a table and calculates and displays a simple ratio. A more useful function would also calculate and display a confidence interval

for the risk ratio. This is what we will do now. Use the fix() function to edit the tab2by2() function:

```
fix(tab2by2)
```

We can now edit this function to calculate and display a 95% confidence interval for the risk ratio.

```
function(exposure, outcome) {
   tab <- table(exposure, outcome)
   a <- tab[1,1]
   b <- tab[1,2]
   c <- tab[2,1]
   d <- tab[2,2]
   rr <- (a / (a + b)) / (c / (c + d))
   se.log.rr <- sqrt((b / a) / (a + b) + (d / c) / (c + d))
   lci.rr <- exp(log(rr) - 1.96 * se.log.rr)
   uci.rr <- exp(log(rr) + 1.96 * se.log.rr)
   print(tab)
   print(rr)
   print(lci.rr)
   print(uci.rr)
}</pre>
```

Once you have made the changes shown above, check your work, save the file, and quit the editor. We should test our revised function:

```
tab2by2(EGGS, ILL)
```

which produces the following output:

```
## outcome
## exposure 1 2
## 1 40 6
## 2 10 20
## [1] 2.608696
## [1] 1.553564
## [1] 4.38044
```

The function works but the output could be improved. Use the fix() function to edit the tab2by2() function:

```
function(exposure, outcome) {
  tab <- table(exposure, outcome)
  a <- tab[1,1]
  b <- tab[1,2]
  c <- tab[2,1]
  d <- tab[2,2]</pre>
```

Once you have made the changes shown above, save the file and quit the editor.

Now we can test our function again:

```
tab2by2(EGGS, ILL)
```

Which produces the following output:

```
## outcome

## exposure 1 2

## 1 40 6

## 2 10 20

##

## RR : 2.608696

## 95% CI : 1.553564 4.38044
```

The tab2by2() function displays output but does not behave like a standard R function in the sense that you cannot save the results of the tab2by2() function into an object:

```
test2by2 <- tab2by2(EGGS, ILL)
```

```
## outcome
## exposure 1 2
## 1 40 6
## 2 10 20
##
## RR: 2.608696
## 95% CI: 1.553564 4.38044
```

displays output but does not save anything in the test2by2 object:

```
test2by2
```

```
## NULL
```

The returned value (NULL) means that test2by2 is an empty object. We will not worry about this at the moment as the tab2by2() function is good-enough for our current purposes. In Exercise 6 we will explore how to make our own functions behave like standard R functions.

We will now add the calculation of the odds ratio and its 95% confidence interval to the tab2by2() function using the fix() function.

There are two ways of doing this. We could either calculate the odds ratio from the table and use (e.g.) the method of Woolf to calculate the confidence interval:

or use the output of the fisher.test() function:

Note that we can refer to components of a function's output using the same syntax as when we refer to columns in a data.frame (e.g. ft\$estimate to examine the estimate of the odds ratio from the fisher.test() function stored in the object ft).

The names of elements in the output of a standard function such as fisher.test() can be found in the documentation or the help system. For example:

```
help(fisher.test)
```

Output elements are listed under the Value heading.

Revise the tab2by2() function to include the calculation of the odds ratio and the 95% confidence interval. The revised function will look something like this:

```
function(exposure, outcome) {
  tab <- table(exposure, outcome)</pre>
  a \leftarrow tab[1,1]
  b < -tab[1,2]
  c < - tab[2,1]
  d < -tab[2,2]
  rr \leftarrow (a / (a + b)) / (c / (c + d))
  se.log.rr \leftarrow sqrt((b / a) / (a + b) + (d / c) / (c + d))
  lci.rr \leftarrow exp(log(rr) - 1.96 * se.log.rr)
  uci.rr \leftarrow \exp(\log(rr) + 1.96 * \text{se.log.rr})
  or <- (a / b) / (c / d)
  se.log.or \leftarrow sqrt(1 / a + 1 / b + 1 / c + 1 / d)
  lci.or \leftarrow exp(log(or) - 1.96 * se.log.or)
  uci.or \leftarrow \exp(\log(\text{or}) + 1.96 * \text{se.log.or})
  ft <- fisher.test(tab)</pre>
  cat("\n")
  print(tab)
```

Once you have made the changes shown above, check your work, save the file, and quit the editor.

Test the tab2by2() function when you have added the calculation of the odds ratio and its 95% confidence interval.

Now that we have a function that will calculate risk ratios and odds ratios with confidence intervals from a two- by-two table we can use it to analyse the salex data:

```
tab2by2(HAM, ILL)
```

##

```
##
##
           outcome
## exposure 1 2
          1 46 17
##
##
            5 9
##
## Relative Risk
                   : 2.044444
## 95% CI
                     : 0.9964841 4.194501
##
## Sample Odds Ratio : 4.870588
## 95% CI
                     : 1.428423 16.60756
##
## MLE Odds Ratio
                  : 4.75649
## 95% CI
                      : 1.22777 20.82921
tab2by2(BEEF, ILL)
##
##
           outcome
## exposure
            1 2
##
          1 45 22
##
          2 6 4
##
## Relative Risk
                   : 1.119403
## 95% CI
                     : 0.6568821 1.907592
```

```
## Sample Odds Ratio : 1.363636
## 95% CI : 0.3485746 5.334594
##
## MLE Odds Ratio : 1.357903
## 95% CI : 0.2547114 6.428414
tab2by2(EGGS, ILL)
##
## outcome
## exposure 1 2
## 1 40 6
##
       2 10 20
##
## Relative Risk : 2.608696
## 95% CI : 1.553564 4.38044
##
## Sample Odds Ratio : 13.33333
## 95% CI
           : 4.240168 41.92706
##
## MLE Odds Ratio : 12.74512
## 95% CI
          : 3.762787 50.05419
tab2by2(MUSHROOM, ILL)
##
##
   outcome
## exposure 1 2
## 1 24 6
## 2 25 19
##
## Relative Risk : 1.408
## 95% CI : 1.028944 1.926697
##
## Sample Odds Ratio : 3.04
## 95% CI
          : 1.037274 8.909506
## MLE Odds Ratio : 2.995207
## 95% CI
                 : 0.9421008 10.7953
tab2by2(PEPPER, ILL)
##
        outcome
## exposure 1 2
##
      1 24 3
```

##

2 23 22

```
##
## Relative Risk : 1.73913
## 95% CI
             : 1.26876 2.383882
##
## Sample Odds Ratio : 7.652174
## 95% CI
            : 2.013718 29.07844
## MLE Odds Ratio : 7.448216
## 95% CI
                   : 1.861728 44.12015
tab2by2(PORKPIE, ILL)
##
##
          outcome
## exposure 1 2
##
        1 21 9
##
         2 29 17
##
## Relative Risk : 1.110345
## 95% CI
                  : 0.8044752 1.532509
## Sample Odds Ratio : 1.367816
## 95% CI
            : 0.5113158 3.659032
##
## MLE Odds Ratio : 1.362228
## 95% CI
                   : 0.4636016 4.190667
tab2by2(PASTA, ILL)
##
##
         outcome
## exposure 1 2
##
         1 25 3
         2 26 23
##
##
## Relative Risk : 1.682692
                  : 1.255392 2.255433
## 95% CI
##
## Sample Odds Ratio : 7.371795
## 95% CI
                 : 1.964371 27.66451
##
## MLE Odds Ratio : 7.195422
## 95% CI
                   : 1.829867 42.07488
tab2by2(RICE, ILL)
```

```
##
       outcome
## exposure 1 2
## 1 28 4
## 2 23 22
##
## Relative Risk : 1.711957
## 95% CI : 1.250197 2.344268
##
## Sample Odds Ratio : 6.695652
## 95% CI : 2.017327 22.22335
##
## MLE Odds Ratio : 6.532868
## 95% CI : 1.852297 29.84928
tab2by2(LETTUCE, ILL)
##
## outcome
## exposure 1 2
## 1 28 1
      2 23 25
##
##
## Relative Risk : 2.014993
## 95% CI : 1.488481 2.727744
## Sample Odds Ratio : 30.43478
## 95% CI : 3.826938 242.041
##
## MLE Odds Ratio : 29.32825
## 95% CI : 4.161299 1284.306
tab2by2(TOMATO, ILL)
##
## outcome
## exposure 1 2
## 1 29 9
##
      2 22 17
## Relative Risk : 1.352871
         : 0.974698 1.877771
## 95% CI
##
## Sample Odds Ratio : 2.489899
## 95% CI : 0.9347213 6.632562
##
## MLE Odds Ratio : 2.459981
```

```
: 0.8467562 7.558026
## 95% CI
tab2by2(COLESLAW, ILL)
##
##
        outcome
## exposure 1 2
      1 29 3
##
        2 21 23
##
## Relative Risk : 1.89881
## 95% CI
                 : 1.366876 2.63775
##
## Sample Odds Ratio : 10.5873
## 95% CI
          : 2.806364 39.9417
##
## MLE Odds Ratio : 10.26269
         : 2.600771 60.35431
## 95% CI
tab2by2(CRISPS, ILL)
##
##
        outcome
## exposure 1 2
## 1 21 10
        2 30 16
##
##
## Relative Risk : 1.03871
## 95% CI
          : 0.7529065 1.433004
##
## Sample Odds Ratio : 1.12
        : 0.4258139 2.945888
## 95% CI
##
## MLE Odds Ratio : 1.118358
## 95% CI
         : 0.3858206 3.340535
tab2by2(PEACHCAKE, ILL)
##
        outcome
## exposure 1 2
        1 2 2
##
        2 49 24
##
##
## Relative Risk : 0.744898
## 95% CI : 0.27594 2.010846
##
```

```
## Sample Odds Ratio : 0.4897959
## 95% CI : 0.06497947 3.691936
##
## MLE Odds Ratio : 0.4947099
## 95% CI : 0.03393887 7.209143
tab2by2(CHOCOLATE, ILL)
##
## outcome
## exposure 1 2
## 1 12 2
##
        2 38 24
##
## Relative Risk : 1.398496
## 95% CI : 1.045064 1.871456
##
## Sample Odds Ratio : 3.789474
## 95% CI
           : 0.7791326 18.43089
##
## MLE Odds Ratio : 3.733535
## 95% CI
           : 0.7318646 37.28268
tab2by2(FRUIT, ILL)
##
##
   outcome
## exposure 1 2
## 1 1 4
## 2 49 22
##
## Relative Risk : 0.2897959
## 95% CI : 0.04985828 1.684408
##
## Sample Odds Ratio : 0.1122449
## 95% CI
          : 0.01185022 1.06318
## MLE Odds Ratio : 0.1157141
## 95% CI
                 : 0.002240848 1.256134
tab2by2(TRIFLE, ILL)
##
        outcome
## exposure 1 2
##
      1 19 5
##
        2 32 21
```

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```
##
## Relative Risk : 1.311198
## 95% CI : 0.9718621 1.769016
##
## Sample Odds Ratio : 2.49375
## 95% CI : 0.8067804 7.708156
##
## MLE Odds Ratio : 2.465794
## 95% CI : 0.7363311 9.778463
```

tab2by2(ALMONDS, ILL)

```
##
##
           outcome
                 2
## exposure
             1
             3
                3
##
          1
          2 38 19
##
##
## Relative Risk
                      : 0.75
## 95% CI
                      : 0.3300089 1.7045
## Sample Odds Ratio : 0.5
## 95% CI
                      : 0.09203498 2.716358
##
## MLE Odds Ratio
                      : 0.505905
## 95% CI
                       : 0.06170211 4.141891
```

Make a note of any positive associations (i.e. with a risk ratio > 1 with a 95% confidence intervals that does not include one). We will use these for the next exercise when we will use logistic regression to analyse this data.

Save the tab2by2() function:

```
save(tab2by2, file = "tab2by2.r")
```

We can now quit R:

```
q()
```

For this exercise there is no need to save the workspace image so click the **No** or **Don't Save** button (GUI) or enter **n** when prompted to save the workspace image (terminal).

3.1 Summary

- R objects contain information that can be examined and manipulated.
- R can be extended by writing new functions.

- New functions can perform simple or complex data analysis.
- New functions can be composed of parts of existing function.
- New functions can be saved and used in subsequent R sessions.
- Objects defined within functions are local to that function and only exist while that function is being used. This means that you can re-use meaningful names within functions without them interfering with each other.

Chapter 4

Logistic regression and stratified analysis

In this exercise we will explore how R handles generalised linear models using the example of logistic regression. We will continue using the salex dataset. Start R and retrieve the salex dataset:

```
salex <- read.table("salex.dat", header = TRUE, na.strings = "9")</pre>
```

When we analysed this data using two-by-two tables and examining the risk ratio and 95% confidence interval associated with each exposure we found many significant positive associations:

Variable	RR	95% CI	
EGGS	2.61	1.55, 4.38	
MUSHROOM	1.41	1.03, 1.93	
PEPPER	1.74	1.27, 2.38	
PASTA	1.68	1.26, 2.26	
RICE	1.72	1.25, 2.34	
LETTUCE	2.01	1.49, 2.73	
COLESLAW	1.89	1.37, 2.64	
CHOCOLATE	1.39	1.05, 1.87	

Some of these associations may be due to *confounding* in the data. We can use logistic regression to help us identify independent associations.

Logistic regression requires the dependent variable to be either 0 or 1. In order to perform a logistic regression we must first recode the ILL variable so that 0=no and 1=yes:

```
table(salex$ILL)
```

##

1 2

```
## 51 26
```

```
salex$ILL[salex$ILL == 2] <- 0
table(salex$ILL)

##
## 0 1
## 26 51</pre>
```

We could work with our data as it is but if we wanted to calculate odds ratios and confidence intervals we would calculate their reciprocals (i.e. odds ratios for non-exposure rather than for exposure). This is because of the way the data has been coded (1=yes, 2=no).

In order to calculate meaningful odds ratios the exposure variables should also be coded 0=no, 1=yes. The actual codes used are not important as long as the value used for 'exposed' is one greater than the value used for 'not exposed'.

We could issue a series of commands similar to the one we have just used to recode the ILL variable. This is both tedious and unnecessary as the structure of the dataset (i.e. all variables are coded identically) allows us to recode all variables with a single command:

```
salex <- read.table("salex.dat", header = TRUE, na.strings = "9")</pre>
salex[1:5, ]
##
                     EGGS MUSHROOM PEPPER PORKPIE PASTA RICE LETTUCE TOMATO
      ILL HAM
               BEEF
## 1
                         1
                                                      2
                                                              2
                                                                    2
                                                                             2
        1
             1
                   1
                                    1
                                             1
                                                                                      2
                                             2
                                                                             2
##
   2
             1
                   1
                         1
                                    2
                                                      1
                                                              2
                                                                    2
                                                                                      1
        1
## 3
             1
                   1
                         1
                                    1
                                             1
                                                      1
                                                              1
                                                                    1
                                                                             1
                                                                                      2
        1
                                    2
                                             2
                                                      2
                                                              2
                                                                    2
        1
             1
                   1
                         1
                                                                             1
## 4
                                                                                      1
##
   5
        1
             1
                   1
                         1
                                    1
                                             1
                                                      1
                                                              1
                                                                    1
                                                                                      1
##
      COLESLAW
                CRISPS PEACHCAKE CHOCOLATE FRUIT TRIFLE ALMONDS
## 1
              2
                       2
                                   2
                                               2
                                                      2
                                                               2
                                                                        2
## 2
              2
                       2
                                   2
                                               2
                                                      2
                                                               2
                                                                        2
              2
                                   2
                                                      2
                                                                         2
                                                               2
## 3
                       1
                                               1
## 4
              2
                       2
                                   2
                                               1
                                                      2
                                                               2
                                                                        2
                                                      2
## 5
              1
                       2
                                   2
                                               1
                                                                         2
                                                               1
salex <- 2 - salex
salex[1:5, ]
```

```
ILL HAM BEEF EGGS MUSHROOM PEPPER PORKPIE PASTA RICE LETTUCE TOMATO
##
##
   1
        1
             1
                    1
                          1
                                     1
                                              1
                                                        0
                                                               0
                                                                      0
                                                                               0
                                                                                        0
## 2
        1
             1
                    1
                          1
                                     0
                                              0
                                                        1
                                                               0
                                                                      0
                                                                               0
                                                                                        1
                          1
                                     1
                                              1
                                                        1
                                                                      1
                                                                                1
##
   3
        1
             1
                    1
                                                               1
                                                                                        0
## 4
        1
             1
                    1
                          1
                                     0
                                              0
                                                        0
                                                               0
                                                                      0
                                                                                1
                                                                                        1
   5
        1
             1
                    1
                          1
                                     1
                                              1
                                                        1
                                                               1
                                                                      1
                                                                                1
                                                                                        1
##
      COLESLAW CRISPS PEACHCAKE CHOCOLATE FRUIT TRIFLE ALMONDS
##
              0
                       0
                                    0
                                                0
                                                        0
                                                                0
## 1
## 2
              0
                       0
                                    0
                                                0
                                                        0
                                                                0
                                                                          0
```

##	3	0	1	0	1	0	0	0
##	4	0	0	0	1	0	0	0
##	5	1	0	0	1	0	1	0

WARNING: The attach() function works with a copy of the data.frame rather than the original data.frame. Commands that manipulate variables in a data.frame may not work as expected if the data.frame has been attached using the attach() function.

It is better to manipulate data *before* attaching a data.frame. The detach() function may be used to remove an attachment prior to any data manipulation.

Many R users avoid using the attach() function altogether.

We can now use the generalised linear model glm() function to specify the logistic regression model:

The method used by the glm() function is defined by the family parameter. Here we specify binomial errors and a logit (logistic) linking function.

We have saved the output of the glm() function in the salex.lreg object. We can examine some basic information about the specified model using the summary() function:

```
summary(salex.lreg)
```

```
##
## Call:
## glm(formula = ILL ~ EGGS + MUSHROOM + PEPPER + PASTA + RICE +
       LETTUCE + COLESLAW + CHOCOLATE, family = binomial(logit),
##
##
       data = salex)
##
## Deviance Residuals:
##
        Min
                   1Q
                         Median
                                        3Q
                                                 Max
## -1.92036 -0.49869
                        0.06877
                                   0.40906
                                             2.07182
##
## Coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) -2.021864
                                     -2.988
                           0.676606
                                              0.00281 **
## EGGS
                3.579366
                           1.267870
                                       2.823
                                             0.00476 **
## MUSHROOM
               -3.584345
                           1.728999
                                     -2.073
                                              0.03817 *
## PEPPER
                2.348074
                           1.428177
                                       1.644
                                              0.10015
## PASTA
                1.774818
                           1.162762
                                       1.526 0.12692
## RICE
                           1.193840
                                       0.096
                                             0.92381
                0.114180
                3.401828
## LETTUCE
                           1.234060
                                       2.757
                                              0.00584 **
## COLESLAW
                0.763857
                           1.024373
                                       0.746
                                              0.45586
## CHOCOLATE
                0.009782
                           1.314683
                                       0.007
                                              0.99406
```

```
## ---
                 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 91.246
                              on 69
                                     degrees of freedom
## Residual deviance: 41.260
                              on 61
                                     degrees of freedom
     (7 observations deleted due to missingness)
## AIC: 59.26
##
## Number of Fisher Scoring iterations: 7
```

We will use *backwards elimination* to remove non-significant variables from the logistic regression model. Remember that previous commands can be recalled and edited using the up and down arrow keys – they do not need to be typed out in full each time.

CHOCOLATE is the least significant variable in the model so we will remove this variable from the model. Storing the output of the glm() function is useful as it allows us to use the update() function to add, remove, or modify variables without having to describe the model in full:

```
salex.lreg <- update(salex.lreg, . ~ . - CHOCOLATE)
summary(salex.lreg)</pre>
```

```
##
## Call:
## glm(formula = ILL ~ EGGS + MUSHROOM + PEPPER + PASTA + RICE +
       LETTUCE + COLESLAW, family = binomial(logit), data = salex)
##
## Deviance Residuals:
##
        Min
                          Median
                                        3Q
                                                  Max
                    1Q
## -1.92561
            -0.49859
                         0.07555
                                              2.07200
                                   0.38723
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
                                     -3.053
## (Intercept)
                -2.0223
                             0.6623
                                             0.00226 **
## EGGS
                 3.5890
                             1.2188
                                      2.945
                                              0.00323 **
## MUSHROOM
                -3.5992
                             1.6885
                                     -2.132
                                             0.03305 *
## PEPPER
                 2.3544
                             1.4275
                                      1.649
                                              0.09910 .
## PASTA
                 1.7770
                             1.1215
                                      1.585
                                              0.11308
## RICE
                 0.1170
                             1.1388
                                      0.103
                                              0.91819
## LETTUCE
                 3.4109
                             1.2316
                                      2.770
                                              0.00561 **
## COLESLAW
                 0.7630
                             1.0224
                                      0.746
                                             0.45547
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
```

```
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 92.122 on 70
                                      degrees of freedom
## Residual deviance: 41.273 on 63
                                      degrees of freedom
     (6 observations deleted due to missingness)
## AIC: 57.273
##
## Number of Fisher Scoring iterations: 7
RICE is now the least significant variable in the model so we will remove this variable from
the model:
salex.lreg <- update(salex.lreg, . ~ . - RICE)</pre>
summary(salex.lreg)
##
## Call:
## glm(formula = ILL ~ EGGS + MUSHROOM + PEPPER + PASTA + LETTUCE +
       COLESLAW, family = binomial(logit), data = salex)
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
                                         2.0697
## -1.8877 -0.4999
                      0.0786
                                0.3897
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                -2.0169
                            0.6600
                                    -3.056 0.00224 **
                                      3.026
## EGGS
                 3.6142
                             1.1944
                                             0.00248 **
## MUSHROOM
                             1.6134 -2.201
                -3.5508
                                             0.02774 *
                 2.3002
                            1.3200
## PEPPER
                                      1.743
                                             0.08141 .
                 1.8230
                                      1.773
## PASTA
                             1.0280
                                             0.07617 .
## LETTUCE
                 3.4199
                             1.2273
                                      2.787
                                             0.00533 **
## COLESLAW
                 0.7611
                             1.0203
                                      0.746
                                             0.45571
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 92.122
                              on 70
                                      degrees of freedom
## Residual deviance: 41.283
                              on 64
                                     degrees of freedom
     (6 observations deleted due to missingness)
## AIC: 55.283
##
## Number of Fisher Scoring iterations: 6
```

COLESLAW is now the least significant variable in the model so we will remove this variable

from the model:

```
salex.lreg <- update(salex.lreg, . ~ . - COLESLAW)</pre>
summary(salex.lreg)
##
## Call:
## glm(formula = ILL ~ EGGS + MUSHROOM + PEPPER + PASTA + LETTUCE,
       family = binomial(logit), data = salex)
##
##
## Deviance Residuals:
        Min
##
                   1Q
                         Median
                                       3Q
                                                 Max
## -1.98481 -0.50486
                        0.08871
                                  0.36910
                                             2.06065
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
                            0.6545 -3.049 0.00230 **
## (Intercept) -1.9957
## EGGS
                 3.8152
                            1.1640 3.278 0.00105 **
## MUSHROOM
                -3.4008
                            1.5922 -2.136 0.03269 *
## PEPPER
                 2.3520
                            1.3269 1.773 0.07631 .
## PASTA
                            0.9922 1.986 0.04701 *
                 1.9706
## LETTUCE
                 3.4786
                            1.2246 2.841 0.00450 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 92.982 on 71 degrees of freedom
## Residual deviance: 41.895 on 66
                                     degrees of freedom
     (5 observations deleted due to missingness)
## AIC: 53.895
## Number of Fisher Scoring iterations: 6
PEPPER is now the least significant variable in the model so we will remove this variable from
the model:
salex.lreg <- update(salex.lreg, . ~ . - PEPPER)</pre>
summary(salex.lreg)
##
## Call:
## glm(formula = ILL ~ EGGS + MUSHROOM + PASTA + LETTUCE, family = binomial(logit),
       data = salex)
##
##
## Deviance Residuals:
##
      Min
                                   3Q
                 1Q
                      Median
                                           Max
```

```
## -2.0920 -0.5360
                    0.1109
                               0.4876
                                        2.0056
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
                            0.6128 -3.048 0.002306 **
## (Intercept)
                -1.8676
                 3.7094
## EGGS
                            1.0682
                                     3.473 0.000515 ***
## MUSHROOM
                -1.6165
                            1.0829 -1.493 0.135524
## PASTA
                 1.8440
                            0.9193
                                     2.006 0.044864 *
## LETTUCE
                 3.2458
                            1.1698
                                     2.775 0.005527 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 94.659
                              on 73
                                     degrees of freedom
## Residual deviance: 45.578
                              on 69
                                     degrees of freedom
     (3 observations deleted due to missingness)
## AIC: 55.578
##
## Number of Fisher Scoring iterations: 6
MUSHROOM is now the least significant variable in the model so we will remove this variable
from the model:
salex.lreg <- update(salex.lreg, . ~ . - MUSHROOM)</pre>
summary(salex.lreg)
##
## Call:
## glm(formula = ILL ~ EGGS + PASTA + LETTUCE, family = binomial(logit),
       data = salex)
##
##
## Deviance Residuals:
##
       Min
                 10
                      Median
                                    3Q
                                            Max
## -2.2024 -0.5108
                      0.2038
                               0.4304
                                         2.0501
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                -1.9710
                            0.6146 -3.207 0.00134 **
## EGGS
                 2.6391
                            0.7334
                                     3.599
                                            0.00032 ***
## PASTA
                 1.6646
                            0.8376
                                     1.987
                                             0.04689 *
## LETTUCE
                 3.1956
                            1.1516
                                     2.775 0.00552 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
```

```
##
## Null deviance: 97.648 on 75 degrees of freedom
## Residual deviance: 50.529 on 72 degrees of freedom
## (1 observation deleted due to missingness)
## AIC: 58.529
##
## Number of Fisher Scoring iterations: 6
```

There are now no non-significant variables in the model.

Unfortunately R does not present information on the model coefficients in terms of odds ratios and confidence intervals but we can write a function to calculate them for us.

The first step in doing this is to realise that the **salex.lreg** object contains essential information about the fitted model. To calculate odds ratios and confidence intervals we need the regression coefficients and their standard errors. Both:

summary(salex.lreg)\$coefficients

and:

coef(summary(salex.lreg))

extract the data that we require. The preferred method is to use the coef() function. This is because some fitted models may return coefficients in a more complicated manner than (e.g.) those created by the glm() function. The coef() function provides a standard way of extracting this data from all classes of fitted objects.

We can store the coefficients data in a separate object to make it easier to work with:

```
salex.lreg.coeffs <- coef(summary(salex.lreg))
salex.lreg.coeffs</pre>
```

```
## Estimate Std. Error z value Pr(>|z|)
## (Intercept) -1.970967 0.6145691 -3.207071 0.0013409398
## EGGS 2.639115 0.7333899 3.598515 0.0003200388
## PASTA 1.664581 0.8375970 1.987330 0.0468858898
## LETTUCE 3.195594 1.1516159 2.774879 0.0055222320
```

We can extract information from this object by addressing each piece of information by its row and column position in the object. For example:

```
salex.lreg.coeffs[2,1]
```

```
## [1] 2.639115
```

Is the regression coefficient for EGGS, and:

```
salex.lreg.coeffs[3,2]
```

```
## [1] 0.837597
```

is the standard error of the regression coefficient for PASTA. Similarly:

```
salex.lreg.coeffs[ ,1]
```

```
## (Intercept) EGGS PASTA LETTUCE
## -1.970967 2.639115 1.664581 3.195594
```

Returns the regression coefficients for all of the variables in the model, and:

```
salex.lreg.coeffs[ ,2]
```

```
## (Intercept) EGGS PASTA LETTUCE
## 0.6145691 0.7333899 0.8375970 1.1516159
```

Returns the standard errors of the regression coefficients.

The table below shows the indices that address each cell in the table of regression coefficients:

```
matrix(salex.lreg.coeffs, nrow = 4, ncol = 4)
```

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

## [1,] -1.970967 0.6145691 -3.207071 0.0013409398

## [2,] 2.639115 0.7333899 3.598515 0.0003200388

## [3,] 1.664581 0.8375970 1.987330 0.0468858898

## [4,] 3.195594 1.1516159 2.774879 0.0055222320
```

We can use this information to calculate odds ratio sand 95% confidence intervals:

```
or <- exp(salex.lreg.coeffs[ ,1])
lci <- exp(salex.lreg.coeffs[ ,1] - 1.96 * salex.lreg.coeffs[ ,2])
uci <- exp(salex.lreg.coeffs[ ,1] + 1.96 * salex.lreg.coeffs[ ,2])</pre>
```

and make a single object that contains all of the required information:

```
lreg.or <- cbind(or, lci, uci)
lreg.or</pre>
```

```
## or lci uci
## (Intercept) 0.1393221 0.0417723 0.4646777
## EGGS 14.0008053 3.3256684 58.9423019
```

```
## PASTA 5.2834608 1.0231552 27.2832114
## LETTUCE 24.4246856 2.5559581 233.4018193
```

We seldom need to report estimates and confidence intervals to more than two decimal places. We can use the round() function to remove the excess digits:

```
round(lreg.or, digits = 2)
```

```
## or lci uci

## (Intercept) 0.14 0.04 0.46

## EGGS 14.00 3.33 58.94

## PASTA 5.28 1.02 27.28

## LETTUCE 24.42 2.56 233.40
```

We have now gone through all the necessary calculations step-by-step but it would be nice to have a function that did it all for us that we could use whenever we needed to.

First we will create a template for the function:

```
lreg.or <- function(model, digits = 2) {}</pre>
```

and then use the fix() function to edit the lreg.or() function:

```
fix(lreg.or)
```

We can now edit this function to add a calculation of odds ratios and 95% confidence intervals:

```
function(model, digits = 2) {
   lreg.coeffs <- coef(summary(model))
   OR <- exp(lreg.coeffs[ ,1])
   LCI <- exp(lreg.coeffs[ ,1] - 1.96 * lreg.coeffs[ ,2])
   UCI <- exp(lreg.coeffs[ ,1] + 1.96 * lreg.coeffs[ ,2])
   lreg.or <- round(cbind(OR, LCI, UCI), digits = digits)
   lreg.or
}</pre>
```

```
lreg.or <- function(model, digits = 2) {
   lreg.coeffs <- coef(summary(model))
   OR <- exp(lreg.coeffs[ ,1])
   LCI <- exp(lreg.coeffs[ ,1] - 1.96 * lreg.coeffs[ ,2])
   UCI <- exp(lreg.coeffs[ ,1] + 1.96 * lreg.coeffs[ ,2])
   lreg.or <- round(cbind(OR, LCI, UCI), digits = digits)
   lreg.or
}</pre>
```

Once you have made the changes shown above, check your work, save the file, and quit the editor.

We can test our function:

```
lreg.or(salex.lreg)
```

Which produces the following output:

```
lreg.or(salex.lreg)
```

```
## OR LCI UCI
## (Intercept) 0.14 0.04 0.46
## EGGS 14.00 3.33 58.94
## PASTA 5.28 1.02 27.28
## LETTUCE 24.42 2.56 233.40
```

The digits parameter of the lreg.or() function, which has digits = 2 as its default value, allows us to specify the precision with which the estimates and their confidence intervals are reported:

```
lreg.or(salex.lreg, digits = 4)
```

```
## OR LCI UCI
## (Intercept) 0.1393 0.0418 0.4647
## EGGS 14.0008 3.3257 58.9423
## PASTA 5.2835 1.0232 27.2832
## LETTUCE 24.4247 2.5560 233.4018
```

Before we continue, it is probably a good idea to save this function for later use:

```
save(lreg.or, file = "lregor.r")
```

Which can be reloaded whenever it is needed:

```
load("lregor.r")
```

An alternative to using logistic regression with data that contains associations that may be due to confounding is to use stratified analysis (i.e. *Mantel-Haenszel* techniques). With several potential confounders, a stratified analysis results in the analysis of many tables which can be difficult to interpret. For example, four potential confounders, each with two levels would produce sixteen tables. In such situations, logistic regression might be a better approach. In order to illustrate Mantel-Haenszel techniques in R we will work with a simpler dataset.

On Saturday, 21st April 1990, a luncheon was held in the home of Jean Bateman. There was a total of forty-five guests which included thirty-five members of the Department of Epidemiology and Population Sciences at the London School of Hygiene and Tropical Medicine. On Sunday morning, 22nd April 1990, Jean awoke with symptoms of gastrointestinal illness; her husband awoke with similar symptoms. The possibility of an outbreak related to the luncheon was strengthened when several of the guests telephoned Jean on Sunday and reported illness. On Monday, 23rd April 1990, there was an unusually large number of department members absent from work and reporting illness. Data from this outbreak is stored in the file bateman.dat.

The variables in the file bateman.dat are:

ILL	Ill?
CHEESE	Cheddar cheese
CRABDIP	Crab dip
CRISPS	Crisps
BREAD	French bread
CHICKEN	Chicken (roasted, served warm)
RICE	Rice (boiled, served warm)
CAESAR	Caesar salad
TOMATO	Tomato salad
ICECREAM	Vanilla ice-cream
CAKE	Chocolate cake
JUICE	Orange juice
WINE	White wine
COFFEE	Coffee

Data is available for all forty-five guests at the luncheon. All of the variables are coded 1=yes, 2=no. Retrieve and attach the bateman dataset in R:

```
bateman <- read.table("bateman.dat", header = TRUE)
bateman</pre>
```

##		ILL	CHEESE	CRABDIP	CRISPS	BREAD	CHICKEN	RICE	CAESAR	TOMATO	ICECREAM
##	1	1	1	1	1	2	1	1	1	1	1
##	2	2	1	1	1	2	1	2	2	2	1
##	3	1	2	2	1	2	1	2	1	2	1
##	4	1	1	2	1	1	1	2	1	2	1
##	5	1	1	1	1	2	1	1	1	1	2
##	6	1	1	1	1	1	1	2	1	1	2
##	7	1	2	1	1	2	1	1	1	1	1
##	8	2	1	1	1	2	1	1	2	1	1
##	9	2	1	1	1	2	1	1	2	1	1
##	10	2	2	1	1	2	1	2	2	2	1
##	11	1	1	2	1	1	1	1	1	1	1
##	12	1	1	1	1	1	1	1	1	1	1
##	13	2	2	1	1	2	1	1	2	2	1
##	14	1	2	1	1	1	1	1	1	1	1
##	15	1	1	1	1	2	2	1	1	1	2
##	16	1	2	2	2	2	1	1	1	1	1
##	17	2	1	2	1	1	1	1	2	2	1
##	18	1	2	1	1	2	1	1	1	1	1
##	19	1	1	2	2	1	1	1	2	1	1
##	20	2	2	2	2	2	2	2	2	2	2
##	21	2	1	2	2	1	2	1	1	2	2

		_	_		_	_	_	_	_	_	_	
##	22	2	2		2	2	2	2	2	2	2	1
	23	2	2		2	2	2	2	2	2	2	1
	24	1	2		1	1	2	1	1	1	2	1
	25	1	1		2	2	1	1	1	1	1	1
	26	2	2		1	1	1	1	1	2	2	2
##	27	2	2		1	1	1	1	1	2	2	2
##	28	1	2		1	2	2	1	1	2	2	1
##	29	1	1		2	2	1	1	1	2	2	1
##	30	1	2		1	1	2	1	1	1	1	1
##	31	1	2		1	1	2	1	1	1	1	1
##	32	1	1		2	2	2	1	1	1	1	2
##	33	2	1		2	1	1	1	1	1	1	1
##	34	1	2		1	1	2	1	1	1	1	2
##	35	1	1		2	1	2	1	1	1	1	2
##	36	2	1		2	1	1	2	1	1	1	2
##	37	1	2		1	1	2	1	1	1	1	1
##	38	1	1		2	2	2	1	2	1	1	1
##	39	2	2		1	1	1	1	1	1	1	2
	40	1	1		1	1	2	1	2	1	1	2
	41	2	2		1	1	1	1	2	1	1	2
	42	1	1		2	2	1	2	2	1	1	1
	43	1	2		1	1	2	1	2	1	1	2
	44	1	2		1	1	2	2	1	1	1	2
	45	1	2		1	1	2	2	1	1	1	1
##	10	_	JUICE	WINE	COFFEE	-	2	2	_	_	-	_
##	1	1	1	1	1							
##		1	1	1	2							
##		1	2	1	2							
##		1	2	1	2							
##		1	1	1	1							
##		1	1	2	2							
##		1	2	1	1							
##		1	2	1	1							
##		1	2	1	1							
	10		1	2								
##		2 1	1	1	1 2							
	12	1	2	1								
	13		2	1	1 1							
		1										
	14	2	2	1	1							
	15	1	1	1	1							
	16	1	2	1	2							
	17	2	2	1	2							
	18	1	2	1	1							
	19	1	2	1	1							
##	20	1	2	2	1							

```
2
                  2
                        1
## 21
                                 1
## 22
                  2
                        1
                                 2
           1
## 23
           1
                  2
                        1
                                 2
## 24
          2
                  2
                        1
                                 1
## 25
          2
                  2
                        1
                                 1
                        1
## 26
           1
                  1
                                 1
## 27
                        1
                                 1
          1
                  1
                        2
## 28
          2
                  1
                                 1
                        2
## 29
          1
                  1
                                 1
## 30
           1
                  2
                        2
                                 1
## 31
          2
                  2
                        2
                                 1
## 32
           1
                  2
                        2
                                 1
                  2
                        2
## 33
                                 1
          1
## 34
                  2
                        1
                                 2
           1
                  2
## 35
                        1
          1
                                 1
## 36
          2
                  2
                        1
                                 1
## 37
          2
                  2
                        1
                                 2
                  2
                                2
## 38
          1
                        1
                        2
## 39
          1
                  1
                                 2
                  2
## 40
          1
                        1
                                 1
                  2
          2
                        1
                                 1
## 41
## 42
          2
                  2
                        1
                                2
## 43
           1
                  2
                        1
                                2
## 44
           1
                  2
                        1
                                2
## 45
           1
                  1
                        2
                                 2
```

attach(bateman)

```
## The following objects are masked from salex (pos = 3):
##
## CRISPS, ILL, RICE, TOMATO
## The following objects are masked from salex (pos = 12):
##
## CRISPS, ILL, RICE, TOMATO
```

We will use our tab2by2() function to analyse this data. Retrieve this function:

```
load("tab2by2.r")
```

Use the tab2by2() function to analyse the data:

```
tab2by2(CHEESE, ILL)
```

```
## outcome
## exposure 1 2
## 1 15 7
```

2 14 9 ## ## Relative Risk : 1.12013 : 0.7253229 1.729838 ## 95% CI ## ## Sample Odds Ratio : 1.377551 ## 95% CI : 0.4037553 4.699992 ## ## MLE Odds Ratio : 1.367743 ## 95% CI : 0.3427732 5.649399 tab2by2(CRABDIP, ILL) ## ## outcome ## exposure 1 2 ## 1 18 9 ## 2 11 7 ## ## Relative Risk : 1.090909 ## 95% CI : 0.6921784 1.719329 ## ## Sample Odds Ratio : 1.272727 ## 95% CI : 0.3682028 4.3993 ## ## MLE Odds Ratio : 1.265848 ## 95% CI : 0.3042941 5.188297 tab2by2(CRISPS, ILL) ## ## outcome ## exposure 1 2 1 21 12 ## ## 2 8 4 ## **##** Relative Risk : 0.9545455 ## 95% CI : 0.5930168 1.536478 ## ## Sample Odds Ratio : 0.875 ## 95% CI : 0.2170373 3.527619 ## ## MLE Odds Ratio : 0.8775841

: 0.1587568 4.184763

95% CI

##

Sample Odds Ratio : 1.885714

tab2by2(BREAD, ILL) ## ## outcome ## exposure 1 2 ## 1 9 8 2 20 8 ## ## **##** Relative Risk : 0.7411765 : 0.4469843 1.228997 ## 95% CI ## Sample Odds Ratio : 0.45 : 0.1280647 1.581232 ## 95% CI ## ## MLE Odds Ratio : 0.4584416 : 0.1072622 1.897017 ## 95% CI tab2by2(CHICKEN, ILL) ## ## outcome ## exposure 1 2 1 25 11 2 4 5 ## ## Relative Risk : 1.5625 ## 95% CI : 0.7293 : 0.7293337 3.347448 ## ## Sample Odds Ratio : 2.840909 ## 95% CI : 0.637796 12.65415 ## ## MLE Odds Ratio : 2.76979 ## 95% CI : 0.4912167 16.93409 tab2by2(RICE, ILL) ## ## outcome ## exposure 1 2 ## 1 22 10 2 7 6 ## ## ## Relative Risk : 1.276786 : 0.7330759 2.223756 ## 95% CI

```
## 95% CI : 0.5027038 7.073586
##
## MLE Odds Ratio : 1.85813
## 95% CI
                 : 0.4026256 8.531602
tab2by2(CAESAR, ILL)
##
##
       outcome
## exposure 1 2
##
       1 26 5
##
        2 3 11
##
## Relative Risk : 3.913978
          : 1.418617 10.7987
## 95% CI
##
## Sample Odds Ratio : 19.06667
## 95% CI
          : 3.866585 94.02038
##
## MLE Odds Ratio : 17.33517
## 95% CI : 3.179027 133.7994
tab2by2(TOMATO, ILL)
##
##
        outcome
## exposure 1 2
## 1 24 6
        2 5 10
##
##
## Relative Risk : 2.4
          : 1.14769 5.018775
## 95% CI
##
## Sample Odds Ratio : 8
## 95% CI : 1.97785 32.35836
##
## MLE Odds Ratio : 7.553116
## 95% CI
        : 1.642249 41.02567
tab2by2(ICECREAM, ILL)
##
##
        outcome
## exposure 1 2
##
       1 20 9
##
        2 9 7
##
```

##

outcome

```
## Relative Risk : 1.226054
## 95% CI : 0.7463643 2.01404
##
## Sample Odds Ratio : 1.728395
          : 0.4889138 6.110177
## 95% CI
##
## MLE Odds Ratio : 1.7069
## 95% CI
         : 0.4021245 7.255001
tab2by2(CAKE, ILL)
##
##
        outcome
## exposure 1 2
## 1 22 11
        2 7 5
##
##
## Relative Risk : 1.142857
          : 0.6689315 1.95255
## 95% CI
##
## Sample Odds Ratio : 1.428571
## 95% CI
           : 0.3678242 5.548347
##
## MLE Odds Ratio : 1.416945
## 95% CI
                  : 0.2847257 6.685098
tab2by2(JUICE, ILL)
##
##
   outcome
## exposure 1 2
## 1 8 5
##
       2 21 11
##
## Relative Risk : 0.9377289
## 95% CI : 0.5701453 1.542301
##
## Sample Odds Ratio : 0.8380952
## 95% CI : 0.2206785 3.182927
##
## MLE Odds Ratio : 0.8414367
## 95% CI : 0.185464 4.101313
tab2by2(WINE, ILL)
##
```

```
## exposure 1 2
##
          1 22 12
##
          2 7 4
##
## Relative Risk : 1.016807
## 95% CI
                     : 0.6099343 1.695094
##
## Sample Odds Ratio : 1.047619
## 95% CI
              : 0.2543383 4.315141
##
## MLE Odds Ratio
                  : 1.046515
## 95% CI
                     : 0.1855742 5.186546
tab2by2(COFFEE, ILL)
##
##
           outcome
## exposure 1 2
          1 17 11
##
          2 12 5
##
##
## Relative Risk : 0.860119
## 95% CI
                    : 0.5607997 1.319196
##
## Sample Odds Ratio : 0.6439394
## 95% CI
                    : 0.1772875 2.338901
##
## MLE Odds Ratio : 0.6502015
## 95% CI
                     : 0.1388979 2.729586
Two variables (CAESAR and TOMATO) are associated with ILL.
These two variables are also associated with each other:
tab2by2(CAESAR, TOMATO)
##
##
          outcome
## exposure 1
##
          1 27 4
##
          2 3 11
##
## Relative Risk
                   : 4.064516
## 95% CI
                    : 1.477162 11.1838
##
```

Sample Odds Ratio : 24.75

: 4.738936 129.2616

95% CI

##

##

CAESAR

ILL

1

1 2

3 1

2 2 9

```
##
## MLE Odds Ratio : 22.10962
## 95% CI
                      : 3.850174 183.4671
chisq.test(table(CAESAR, TOMATO))
## Warning in chisq.test(table(CAESAR, TOMATO)): Chi-squared approximation may
## be incorrect
##
   Pearson's Chi-squared test with Yates' continuity correction
##
##
## data: table(CAESAR, TOMATO)
## X-squared = 15.877, df = 1, p-value = 6.759e-05
fisher.test(table(CAESAR, TOMATO))
##
## Fisher's Exact Test for Count Data
##
## data: table(CAESAR, TOMATO)
## p-value = 3.442e-05
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
      3.850174 183.467108
## sample estimates:
## odds ratio
##
     22.10962
This suggests the potential for one of these associations to be due to confounding. We can
perform a simple stratified analysis using the table() function:
table (CAESAR, ILL, TOMATO)
## , , TOMATO = 1
##
##
         ILL
## CAESAR 1
        1 23 4
##
##
        2 1
##
## , TOMATO = 2
##
```

table(TOMATO, ILL, CAESAR)

```
## , CAESAR = 1
##
##
         ILL
## TOMATO
           1
               2
        1 23
               4
##
        2 3 1
##
##
   , , CAESAR = 2
##
##
##
         ILL
## TOMATO
           1
               2
               2
           1
##
        1
        2
##
           2
```

It would be useful to calculate odds ratios for each stratum. We can define a simple function to calculate an odds ratio from a two-by-two table:

```
or <- function(x) \{(x[1,1] / x[1,2]) / (x[2,1] / x[2,2])\}
```

We can use apply() to apply the or() function to the two-by-two table in each stratum:

```
tabC <- table(CAESAR, ILL, TOMATO)
apply(tabC, 3, or)

## 1 2
## 11.5 13.5

tabT <- table(TOMATO, ILL, CAESAR)
apply(tabT, 3, or)</pre>
```

```
## 1 2
## 1.916667 2.250000
```

The 3 instructs the apply() function to apply the or() function to the third dimension of the table objects (i.e. levels of the potential confounder in tabC and tabT).

The mantelhaen.test() function performs the stratified analysis:

```
mantelhaen.test(tabC)
##
```

```
##
## Mantel-Haenszel chi-squared test with continuity correction
##
## data: tabC
## Mantel-Haenszel X-squared = 5.752, df = 1, p-value = 0.01647
## alternative hypothesis: true common odds ratio is not equal to 1
## 95 percent confidence interval:
```

```
1.878994 83.156212
##
## sample estimates:
## common odds ratio
##
                12.5
mantelhaen.test(tabT)
##
   Mantel-Haenszel chi-squared test with continuity correction
##
##
## data:
          tabT
## Mantel-Haenszel X-squared = 0.049144, df = 1, p-value = 0.8246
## alternative hypothesis: true common odds ratio is not equal to 1
## 95 percent confidence interval:
     0.3156862 13.4192331
## sample estimates:
## common odds ratio
            2.058219
##
```

It is likely that CAESAR salad was a vehicle of food-poisoning, and that TOMATO salad was not a vehicle of food-poisoning. Many of those at the luncheon ate both CAESAR salad and TOMATO salad. CAESAR confounded the relationship between TOMATO and ILL. This resulted in a spurious association between TOMATO and ILL.

It only makes sense to calculate a common odds ratio in the absence of interaction. We can check for interaction 'by eye' by examining and comparing the odds ratios for each stratum as we did above.

There does appear to be an interaction between CAESAR, WINE, and ILL:

```
tabW <- table(CAESAR, ILL, WINE)
apply(tabW, 3, or)

## 1 2
## 63.0 2.5</pre>
```

Woolf's test for interaction (also known as Woolf's test for the homogeneity of odds ratios) provides a formal test for interaction.

R does not provide a function to perform Woolf's test for the homogeneity of odds ratios but it is possible to write a function to perform this test.

First we will create a template for the function:

```
woolf.test <- function(x) {}</pre>
```

And then use the fix() function to edit the woolf.test() function:

```
fix(woolf.test)
```

We can now edit this function to make it do something useful:

Once you have made the changes shown above, check your work, save the file, and quit the editor. We can use the woolf.test() function to test for a three-way interaction between CAESAR, WINE, and ILL:

```
woolf.test(tabW)
```

Which returns:

```
woolf.test(tabW)
```

##
Woolf's X2 : 3.319492
p-value : 0.06846297

Which is weak evidence of an interaction.

We should test for interaction between CAESAR, TOMATO, and ILL before accepting the results reported by the mantelhaen.test() function:

```
woolf.test(tabC)
```

Woolf's X2 : 0.0001233783 ## p-value : 0.9911376

We can repeat this analysis using logistic regression.

We need to change the coding of the variables to 0 and 1 before specifying the model:

##	ILL	CHEESE	CRABDIP	CRISPS	BREAD	CHICKEN	RICE	CAESAR	TOMATO	ICECREAM
## 1	1	1	1	1	0	1	1	1	1	1
## 2	0	1	1	1	0	1	0	0	0	1
## 3	1	0	0	1	0	1	0	1	0	1
## 4	1	1	0	1	1	1	0	1	0	1
## 5	1	1	1	1	0	1	1	1	1	0
## 6	1	1	1	1	1	1	0	1	1	0
## 7	1	0	1	1	0	1	1	1	1	1
## 8	0	1	1	1	0	1	1	0	1	1
## 9	0	1	1	1	0	1	1	0	1	1
## 10		0	1	1	0	1	0	0	0	1
## 11		1	0	1	1	1	1	1	1	1
## 12		1	1	1	1	1	1	1	1	1
## 13		0	1	1	0	1	1	0	0	1
## 14		0	1	1	1	1	1	1	1	1
## 15		1	1	1	0	0	1	1	1	0
## 16		0	0	0	0	1	1	1	1	1
## 17		1	0	1	1	1	1	0	0	1
## 18		0	1	1	0	1	1	1	1	1
## 19 ## 20		1	0	0	1	1	1	0	1	1
## 20 ## 21		0	0	0	0	0	0 1	0	0	0
## 22		0	0	0	1	0	0	0	0	0
## 23		0	0	0	0	0	0	0	0	1
## 24		0	1	1	0	1	1	1	0	1
## 25		1	0	0	1	1	1	1	1	1
## 26		0	1	1	1	1	1	0	0	0
## 2		0	1	1	1	1	1	0	0	0
## 28		0	1	0	0	1	1	0	0	1
## 29		1	0	0	1	1	1	0	0	1
## 30		0	1	1	0	1	1	1	1	1
## 3:	l 1	0	1	1	0	1	1	1	1	1
## 32	2 1	1	0	0	0	1	1	1	1	0
## 33	3 0	1	0	1	1	1	1	1	1	1
## 34	1 1	0	1	1	0	1	1	1	1	0
## 35	5 1	1	0	1	0	1	1	1	1	0
## 36	0	1	0	1	1	0	1	1	1	0
## 37	7 1	0	1	1	0	1	1	1	1	1
## 38		1	0	0	0	1	0	1	1	1
## 39		0	1	1	1	1	1	1	1	0
## 40		1	1	1	0	1	0	1	1	0
## 41		0	1	1	1	1	0	1	1	0
## 42		1	0	0	1	0	0	1	1	1
## 43		0	1	1	0	1	0	1	1	0
## 44	1 1	0	1	1	0	0	1	1	1	0

##	45	1	0		1	1	0	0	1	1	1	1
##					COFFEE							
##		1	1	1	1							
##		1	1	1	0							
##		1	0	1	0							
##		1	0	1	0							
##		1	1	1	1							
##		1	1	0	0							
## ##		1	0	1	1							
##		1	0	1 1	1							
	10	1 0	0 1	0	1 1							
##		1	1	1	0							
	12	1	0	1	1							
	13	1	0	1	1							
	14	0	0	1	1							
	15	1	1	1	1							
	16	1	0	1	0							
		0	0	1	0							
	18	1	0	1	1							
##	19	1	0	1	1							
##	20	1	0	0	1							
##	21	0	0	1	1							
##	22	1	0	1	0							
##	23	1	0	1	0							
	24	0	0	1	1							
		0	0	1	1							
					1							
					1							
	39	1	1	0	0							
	40	1	0	1	1							
	41	0	0	1	1							
##	42	0	0	1	0							
##	43	1	0	1	0							
######################################	19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42	1 1 0 1 1 0 0 1 1 1 0 0 1 1 1 0 0 1 1 1 0 0 0 1 1 1 0 0 0 0	0 0 0 0 0 0 0 0 1 1 1 1 0 0 0 0 0 0 0 0	1 1 0 1 1 1 1 1 1 0 0 0 0 0 0 0 1 1 1 1	1 1 1 1 0 0 0 1 1 1 1 1 1 1 1 0 0 0 1 1 1 1 0							

```
## 44
                  1
                         0
              0
        1
                  0
## 45
              1
##
## Call:
## glm(formula = ILL ~ CAESAR + TOMATO, family = binomial(logit),
      data = bateman)
##
## Deviance Residuals:
     Min
              1Q Median 3Q
                                   Max
## -1.960 -0.641
                  0.563 0.563
                                 1.835
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -1.4780
                          0.7101 -2.082 0.03739 *
                                  2.611 0.00904 **
## CAESAR
               2.5202
                          0.9653
## TOMATO
              0.7197
                          0.9552
                                  0.753 0.45116
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 58.574 on 44 degrees of freedom
## Residual deviance: 41.408 on 42 degrees of freedom
## AIC: 47.408
##
## Number of Fisher Scoring iterations: 4
##
## Call:
## glm(formula = ILL ~ CAESAR, family = binomial(logit), data = bateman)
##
## Deviance Residuals:
      Min
               1Q
                    Median
                                3Q
                                        Max
## -1.9103 -0.6945
                    0.5931
                             0.5931
                                     1.7552
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -1.2993 0.6513 -1.995 0.046066 *
## CAESAR
               ## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 58.574 on 44 degrees of freedom
```

```
## Residual deviance: 41.940 on 43 degrees of freedom
## AIC: 45.94
##
## Number of Fisher Scoring iterations: 4
Interactions are specified using the multiply (___*__) symbol in the model formula:
bateman.lreg <- glm(formula = ILL ~ CAESAR + WINE + CAESAR * WINE,
                    family = binomial(logit), data = bateman)
summary(bateman.lreg)
##
## Call:
## glm(formula = ILL ~ CAESAR + WINE + CAESAR * WINE, family = binomial(logit),
##
       data = bateman)
##
## Deviance Residuals:
##
      Min
                 1Q
                     Median
                                   3Q
                                           Max
## -2.0393 -0.4590
                    0.5168
                               0.5168
                                        2.1460
##
## Coefficients:
##
                 Estimate Std. Error z value Pr(>|z|)
## (Intercept) -1.092e-15 1.000e+00
                                       0.000
                                               1.0000
## CAESAR
               9.163e-01 1.304e+00
                                      0.703
                                               0.4822
## WINE
              -2.197e+00 1.453e+00 -1.512
                                               0.1305
## CAESAR:WINE 3.227e+00 1.787e+00
                                     1.806
                                               0.0709 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 58.574 on 44 degrees of freedom
## Residual deviance: 38.508
                                    degrees of freedom
                             on 41
## AIC: 46.508
##
## Number of Fisher Scoring iterations: 4
```

Before we continue, it is probably a good idea to save the woolf.test() function for later use:

```
save(woolf.test, file = "woolf.r")
```

4.1 Matched data

Matching is another way to control for the effects of potential confounding variables. Matching is usually performed during data-collection as part of the design of a study.

In a matched case-control studies, each case is matched with one or more controls which are chosen to have the same values over a set of potential confounding variables. In order to illustrate how matched data may be analysed using tabulation and stratification in R we will start with the simple case of one-to-one matching (i.e. each case has a single matched control):

```
octe <- read.table("octe.dat", header = TRUE)
octe[1:10, ]</pre>
```

```
##
       ID CASE OC
## 1
        1
               1
                  1
               2
## 2
        1
                  1
## 3
        2
               1
                  1
## 4
        2
               2
                  1
## 5
        3
               1
                  1
        3
              2
                  1
## 6
##
        4
               1
                  1
## 8
        4
               2
                  1
        5
               1
                  1
## 9
## 10
        5
               2
                  1
```

This data is from a matched case-control study investigating the association between oral contraceptive use and thromboembolism. The cases are 175 women aged between 15 and 44 years admitted to hospital for thromboembolism and discharged alive. The controls are female patients admitted for conditions believed to be unrelated to oral contraceptive use. Cases and controls were matched on age, ethnic group, marital status, parity, income, place of residence, and date of hospitalisation. The variables in the dataset are:

ID	Identifier for the matched sets of cases and controls
\mathbf{CASE}	Case (1) or control (2)
\mathbf{OC}	Used oral contraceptives in the previous month (1=yes, 2=no)

The dataset consists of 350 records:

```
nrow(octe)
## [1] 350
There are 175 matched sets of cases and controls:
length(unique(octe$ID))
```

```
## [1] 175
```

In each matched set of cases and controls there is one case and one control:

table(octe\$ID, octe\$CASE)

```
##
##
          1 2
##
     1
          1 1
     2
##
          1 1
##
     3
          1 1
     4
##
          1 1
##
     5
          1 1
##
     6
          1 1
     7
##
          1 1
##
     8
          1 1
     9
##
          1 1
##
     10
          1 1
##
     11
          1 1
##
      12
          1 1
     13
##
          1 1
##
      14
          1 1
##
          1 1
     15
##
     16
          1 1
##
     17
          1 1
##
     18
          1 1
##
     19
          1 1
##
     20
          1 1
##
     21
          1 1
##
     22
          1 1
##
     23
          1 1
##
     24
          1 1
##
     25
          1 1
##
     26
          1 1
##
     27
          1 1
##
     28
          1 1
##
     29
          1 1
##
     30
          1 1
##
     31
          1 1
##
     32
          1 1
##
     33
          1 1
##
     34
          1 1
##
     35
          1 1
##
     36
          1 1
##
     37
          1 1
##
     38
          1 1
##
     39
          1 1
```

```
##
    40 1 1
##
    41 1 1
##
    42 1 1
##
    43 1 1
    44 1 1
##
##
    45 1 1
##
    46 1 1
##
    47 1 1
##
    48 1 1
##
    49 1 1
##
    50 1 1
    51 1 1
##
    52 1 1
##
    53 1 1
##
    54 1 1
##
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    55 1 1
##
    56 1 1
##
    57 1 1
    58 1 1
##
    59 1 1
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    60 1 1
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    61 1 1
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    62 1 1
    63 1 1
##
    64 1 1
##
##
    65 1 1
    66 1 1
##
    67 1 1
##
##
    68 1 1
##
    69 1 1
##
    70 1 1
    71 1 1
##
##
    72 1 1
    73 1 1
##
##
    74 1 1
##
    75 1 1
##
    76 1 1
    77 1 1
##
    78 1 1
##
##
    79
        1 1
    80 1 1
##
##
    81 1 1
##
    82 1 1
    83 1 1
##
```

84 1 1

##

```
##
     85
         1 1
##
     86
          1 1
##
     87
          1 1
##
     88
          1 1
          1 1
##
     89
##
     90
          1 1
##
          1 1
     91
##
     92
          1 1
##
     93
          1 1
##
     94
          1 1
##
     95
          1 1
          1 1
##
     96
          1 1
##
     97
     98
          1 1
##
         1 1
##
     99
     100 1 1
##
     101 1 1
##
##
     102 1 1
     103 1 1
##
     104 1 1
##
##
     105 1 1
     106 1 1
##
     107 1 1
##
     108 1 1
##
     109 1 1
##
##
     110 1 1
     111 1 1
##
     112 1 1
##
##
     113 1 1
##
     114 1 1
     115 1 1
##
     116 1 1
##
     117 1 1
##
     118 1 1
##
##
     119 1 1
##
     120 1 1
##
     121 1 1
     122 1 1
##
     123 1 1
##
##
     124 1 1
     125 1 1
##
##
     126 1 1
##
     127 1 1
     128 1 1
##
##
     129 1 1
```

```
##
     130 1 1
##
     131 1 1
##
     132 1 1
     133 1 1
##
     134 1 1
##
##
     135 1 1
     136 1 1
##
     137 1 1
##
     138 1 1
##
##
     139 1 1
##
     140 1 1
##
     141 1 1
     142 1 1
##
##
     143 1 1
     144 1 1
##
##
     145 1 1
##
     146 1 1
##
     147 1 1
##
     148 1 1
##
     149 1 1
##
     150 1 1
##
     151 1 1
##
     152 1 1
     153 1 1
##
##
     154 1 1
##
     155 1 1
##
     156 1 1
##
     157 1 1
##
     158 1 1
##
     159 1 1
     160 1 1
##
     161 1 1
##
     162 1 1
##
     163 1 1
##
##
     164 1 1
     165 1 1
##
##
     166 1 1
##
     167 1 1
##
     168 1 1
     169 1 1
##
##
     170 1 1
##
     171 1 1
##
     172 1 1
```

##

##

173 1 1

174 1 1

```
## 175 1 1
```

This data may be analysed using McNemar's chi-squared test which use the number of discordant (i.e. relative to exposure) pairs of matched cases and controls.

To find the number of discordant pairs we need to split the dataset into cases and controls:

```
octe.cases <- subset(octe, CASE == 1)
octe.controls <- subset(octe, CASE == 2)</pre>
```

Sorting these two datasets (i.e. octe.cases and octe.controls) by the ID variable simplifies the analysis:

```
octe.cases <- octe.cases[order(octe.cases$ID), ]
octe.controls <- octe.controls[order(octe.controls$ID), ]</pre>
```

Since the two datasets (i.e. octe.cases and octe.controls) are now sorted by the ID variable we can use the table() function to retrieve the number if concordant and discordant pairs and store them in a table object:

```
tab <- table(octe.cases$0C, octe.controls$0C)
tab

##
## 1 2
## 1 10 57</pre>
```

This table object (i.e. tab) can then be passed to the mcnemar.test() function:

```
mcnemar.test(tab)
```

```
##
## McNemar's Chi-squared test with continuity correction
##
## data: tab
## McNemar's chi-squared = 26.414, df = 1, p-value = 2.755e-07
```

The mcnemar.test() function does not provide an estimate of the odds ratio. This is the ratio of the discordant pairs:

```
r <- tab[1,2]
s <- tab[2,1]
rdp <- r / s
rdp
```

```
## [1] 4.384615
```

##

2 13 95

A confidence interval can also be calculated:

```
ci.p <- binom.test(r, r + s)$conf.int
ci.rdp <- ci.p / (1 - ci.p)
ci.rdp

## [1] 2.371377 8.731311
## attr(,"conf.level")
## [1] 0.95</pre>
```

This provides a 95% confidence interval. Other (e.g. 99%) confidence intervals can be produced by specifying appropriate values for the conf.level parameter of the binom.test() function:

```
ci.p <- binom.test(r, r + s, conf.level = 0.99)$conf.int
ci.rdp <- ci.p / (1 - ci.p)
ci.rdp

## [1] 2.010478 10.949095
## attr(,"conf.level")
## [1] 0.99</pre>
```

An alternative way of analysing this data is to use the mantelhaen.test() function:

```
tab <- table(octe$CCASE, octe$ID)
mantelhaen.test(tab)</pre>
```

```
##
## Mantel-Haenszel chi-squared test with continuity correction
##
## data: tab
## Mantel-Haenszel X-squared = 26.414, df = 1, p-value = 2.755e-07
## alternative hypothesis: true common odds ratio is not equal to 1
## 95 percent confidence interval:
## 2.400550 8.008521
## sample estimates:
## common odds ratio
## 4.384615
```

The Mantel-Haenszel approach is preferred because it can be used with data from matched case-control studies that match more than one control to each case. Multiple matching is useful when the condition being studied is rare or at the early stages of an outbreak (i.e. when cases are hard to find and controls are easy to find).

We will now work with some data where each case has one or more controls:

```
tsstamp <- read.table("tsstamp.dat", header = TRUE)
tsstamp</pre>
```

```
## ID CASE RBTAMP
## 1 1 1 1
```

##	2	1	2	1
##	3	1	2	1
##	4	2	1	1
##	5	2	2	2
##	6	2	2	1
##	7	2	2	1
##	8	3	1	2
##	9	3	2	2
##	10	3	2	1
##	11	4	1	1
##	12	4	2	2
##	13	5	1	1
##	14	5	2	1
##	15	5	2	2
##	16	6	1	1
##	17	6	2	1
##	18	7	1	1
##	19	7	2	2
##	20	7	2	2
##	21	8	1	1
##	22	8	2	1
##	23	8	2	2
##	24	9	1	2
##	25	9	2	1
##	26	9	2	2
##	27	9	2	2
##	28	10	1	1
##	29	10	2	2
##	30	10	2	2
##	31	11	1	1
##	32	11	2	2
##	33	11	2	2
##	34	12	1	1
##	35	12	2	2
##	36	12	2	2
##	37	12	2	2
##	38	13	1	1
##	39	13	2	2
##	40	13	2	2
##	41	14	1	2
##	42	14	2	2
##	43	14	2	2

This data is from a matched case-control study investigating the association between the use of different brands of tampon and toxic shock syndrome undertaken during an outbreak.

Only a subset of the original dataset is used here. The variables in the dataset are:

ID	Identifier for the matched sets of cases and controls
\mathbf{CASE}	Case (1) or control (2)
RBTAMP	Used Rely brand tampons (1=yes, 2=no)

The dataset consists of forty-three (43) records:

```
nrow(tsstamp)
```

```
## [1] 43
```

There are fourteen (14) matched sets of cases and controls:

```
length(unique(tsstamp$ID))
```

```
## [1] 14
```

Each matched set of cases and controls consists of one case and one or more controls:

```
table(tsstamp$ID, tsstamp$CASE)
```

```
##
##
         1 2
        1 2
##
     2
        1 3
##
     3
       1 2
##
     4 1 1
##
##
     5
       1 2
        1 1
##
     6
        1 2
##
     7
##
     8
        1 2
##
        1 3
     10 1 2
##
     11 1 2
##
     12 1 3
##
     13 1 2
##
##
     14 1 2
```

##

The McNemar's chi-squared test is not useful for this data as it is limited to the special case of one-to-one matching.

Analysing this data using a simple tabulation such as:

```
fisher.test(table(tsstamp$RBTAMP, tsstamp$CASE))
##
## Fisher's Exact Test for Count Data
```

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```
## data: table(tsstamp$RBTAMP, tsstamp$CASE)
## p-value = 0.007805
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 1.542686 53.734756
## sample estimates:
## odds ratio
## 7.709932
```

ignores the matched nature of the data and is, therefore, also not useful for this data.

The matched nature of the data may be accounted by stratifying on the variable that identifies the matched sets of cases and controls (i.e. the ID variable) using the mantelhaen.test() function:

```
mantelhaen.test(table(tsstamp$RBTAMP, tsstamp$CASE, tsstamp$ID))
```

```
##
## Mantel-Haenszel chi-squared test with continuity correction
##
## data: table(tsstamp$RBTAMP, tsstamp$CASE, tsstamp$ID)
## Mantel-Haenszel X-squared = 5.9384, df = 1, p-value = 0.01481
## alternative hypothesis: true common odds ratio is not equal to 1
## 95 percent confidence interval:
## 1.589505 43.191463
## sample estimates:
## common odds ratio
## 8.285714
```

Analysis of several risk factors or adjustment for confounding variables not matched for in the design of a matched case-control study cannot be performed using tabulation-based procedures such as the McNemar's chi-squared test and Mantel-Haenszel procedures. In these situations a special form of logistic regression, called conditional logistic regression, should be used.

We can now quit R:

```
q()
```

For this exercise there is no need to save the workspace image so click the **No** or **Don't Save** button (GUI) or enter **n** when prompted to save the workspace image (terminal).

4.2 Summary

• R provides functions for many kinds of complex statistical analysis. We have looked at using the generalised linear model glm() function to perform logistic regression.

We have looked ar the mantelhaen.test() function to perform stratified analyses and the mantelhaen.test() and mcnemar.test() functions to analyse data from matched case-control studies.

• R can be extended by writing new functions. New functions can perform simple or complex data analysis. New functions can be composed of parts of existing function. New functions can be saved and used in subsequent R sessions. By building your own functions you can use R to build your own statistical analysis system.

Chapter 5

Analysing some data with R

In this exercise we will use the R functions we have already used and the functions we have added to R to analyse a small dataset. First we will start R and retrieve our functions:

```
load("tab2by2.r")
load("lregor.r")
```

And then retrieve and attach the sample dataset:

```
gudhiv <- read.table("gudhiv.dat", header = TRUE, na.strings = "X")
attach(gudhiv)

## The following objects are masked from gudhiv (pos = 11):
##

## CIR, GAMBIAN, GUD, HIV, INJ12M, MARRIED, PARTNERS, SEXPRO,
## TRAVOUT, UTIGC</pre>
```

This data is from a cross-sectional study of 435 male patients who presented with sexually transmitted infections at an outpatient clinic in The Gambia between August 1988 and June 1990. Several studies have documented an association between genital ulcer disease (GUD) and HIV infection. A study of Gambian prostitutes documented an association between seropositivity for HIV-2 and antibodies against Treponema pallidum (a serological test for syphilis). Prostitutes are not the ideal population for such studies as they may have experienced multiple sexually transmitted infections and it is difficult to quantify the number of times they may have had sex with HIV-2 seropositive customers. A sample of males with sexually transmitted infections is easier to study as they have probably had fewer sexual partners than prostitutes and much less contact with sexually transmitted infection pathogens. In such a sample it is also easier to find subjects and collect data. The variables in the dataset are:

MARRIED	Married $(1=yes, 0=no)$
GAMBIAN	Gambian Citizen (1=yes, 0=no)
GUD	History of GUD or syphilis (1=yes, 0=no)
UTIGC	History of urethral discharge (1=yes, 0=no)

CIR	Circumcised (1=yes, 0=no)
TRAVOUT	Travelled outside of Gambia and Senegal (1=yes, 0=no)
SEXPRO	Ever had sex with a prostitute (1=yes, 0=no)
INJ12M	Injection in previous 12 months (1=yes, 0=no)
PARTNERS	Sexual partners in previous 12 months (number)
HIV HIV-2	positive serology (1=yes, 0=no)

Data is available for all 435 patients enrolled in the study.

We will start our analysis by examining pairwise associations between the binary exposure variables and the HIV variable using the tab2by2() function that we wrote earlier:

```
tab2by2(MARRIED, HIV)
tab2by2(GAMBIAN, HIV)
tab2by2(GUD, HIV)
tab2by2(UTIGC, HIV)
tab2by2(CIR, HIV)
tab2by2(TRAVOUT, HIV)
tab2by2(SEXPRO, HIV)
tab2by2(INJ12M, HIV)
```

tab2by2(MARRIED, HIV)

```
##
##
          outcome
             0
                 1
## exposure
         0 321 13
##
##
         1 93
                 8
##
## Relative Risk : 1.043751
## 95% CI
                    : 0.9818512 1.109554
##
## Sample Odds Ratio: 2.124069
## 95% CI
              : 0.8545749 5.279433
##
## MLE Odds Ratio
                 : 2.119801
## 95% CI
                     : 0.7380371 5.714354
```

tab2by2(GAMBIAN, HIV)

```
## outcome
## exposure 0 1
## 0 73 4
## 1 341 17
```

```
##
## Relative Risk : 0.9953155
## 95% CI
            : 0.9400068 1.053879
##
## Sample Odds Ratio : 0.909824
## 95% CI
            : 0.2974059 2.783333
##
## MLE Odds Ratio : 0.9100104
## 95% CI
                  : 0.2853202 3.826485
tab2by2(GUD, HIV)
##
##
          outcome
## exposure 0 1
         0 339 12
         1 72
##
##
## Relative Risk : 1.086538
## 95% CI
                  : 1.003531 1.176412
##
## Sample Odds Ratio : 3.53125
## 95% CI
            : 1.434372 8.693509
##
## MLE Odds Ratio : 3.517408
## 95% CI
                   : 1.258556 9.491924
tab2by2(UTIGC, HIV)
##
##
         outcome
## exposure 0 1
         0 261 12
##
##
         1 151 9
##
## Relative Risk : 1.013027
## 95% CI
                  : 0.9678841 1.060275
##
## Sample Odds Ratio : 1.296358
## 95% CI
               : 0.5338453 3.147997
##
## MLE Odds Ratio : 1.295532
## 95% CI
                   : 0.4703496 3.438842
tab2by2(CIR, HIV)
```

```
## outcome
## exposure 0 1
## 0 10 3
##
      1 392 17
##
## Relative Risk : 0.8025903
## 95% CI
               : 0.5955085 1.081682
##
## Sample Odds Ratio : 0.1445578
## 95% CI : 0.0364195 0.5737851
##
## MLE Odds Ratio : 0.1460183
## 95% CI : 0.03322189 0.899754
tab2by2(TRAVOUT, HIV)
##
## outcome
## exposure 0 1
      0 152 2
##
##
       1 256 19
##
## Relative Risk : 1.060268
## 95% CI
        : 1.02181 1.100173
## Sample Odds Ratio : 5.640625
        : 1.295879 24.55218
## 95% CI
##
## MLE Odds Ratio : 5.624226
        : 1.32716 50.45859
## 95% CI
tab2by2(SEXPRO, HIV)
##
## outcome
## exposure 0 1
## 0 268 13
##
      1 143 8
## Relative Risk : 1.007093
          : 0.9621259 1.054161
## 95% CI
##
## Sample Odds Ratio : 1.153308
## 95% CI : 0.4671083 2.847562
##
## MLE Odds Ratio : 1.152912
```

95% CI : 0.4042323 3.083152

tab2by2(INJ12M, HIV)

```
##
##
           outcome
              0
## exposure
                   7
          0 146
##
          1 268
##
                 14
##
## Relative Risk
                      : 1.004097
## 95% CI
                      : 0.9610996 1.049018
##
## Sample Odds Ratio : 1.089552
## 95% CI
                      : 0.4301305 2.759916
##
## MLE Odds Ratio
                      : 1.089351
## 95% CI
                       : 0.4006202 3.263814
```

Note that our tab2by2() function returns misleading risk ratio estimates and confidence intervals for this dataset. This is because the function expects the exposure and outcome variables to be ordered with exposure-present and outcome-present as the first category (e.g. 1 = present, 2 = absent). This coding is reversed (i.e. 0 = absent, 1 = present) in the gudhiv dataset.

We can produce risk ratio estimates for variables in the gudhiv data using the tab2by2() function and a simple transformation of the exposure and outcome variables. For example:

```
tab2by2(2 - GUD, 2 - HIV)
```

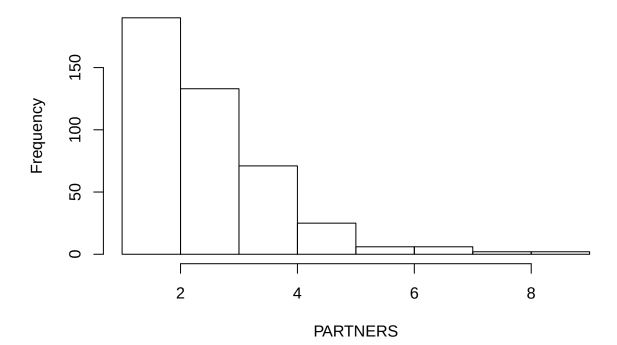
```
##
##
           outcome
              1
## exposure
              9
                 72
##
          1
##
          2
             12 339
##
## Relative Risk
                      : 3.25
## 95% CI
                      : 1.417411 7.451965
##
## Sample Odds Ratio : 3.53125
## 95% CI
                      : 1.434372 8.693509
##
## MLE Odds Ratio
                     : 3.517408
## 95% CI
                       : 1.258556 9.491924
```

The odds ratio estimates returned by the tab2by2() function, with or without this transformation, are correct. The GUD and TRAVOUT variables are associated with HIV.

PARTNERS is a continuous variable and we should examine its distribution before doing anything with it:

```
table (PARTNERS)
hist (PARTNERS)
## PARTNERS
               3
      1
          2
                                           9
##
                    4
                        5
                             6
                                       8
    61 129 133
                                           2
                  71
                       25
                             6
                                  6
                                       2
##
```

Histogram of PARTNERS



The distribution of PARTNERS is severely non-normal. Instead of attempting to transform the variable we will produce summary statistics for each level of the HIV variable and perform a non-parametric test:

```
by (PARTNERS, HIV, summary)
kruskal.test(PARTNERS ~ HIV)
## HIV: O
                                Mean 3rd Qu.
##
      Min. 1st Qu.
                     Median
                                                 Max.
                       3.00
##
               2.00
                                2.72
                                         3.00
                                                 8.00
      1.00
##
## HIV: 1
##
      Min. 1st Qu.
                     Median
                                Mean 3rd Qu.
                                                 Max.
##
     1.000
             4.000
                      5.000
                               5.381
                                       7.000
                                                9.000
##
##
    Kruskal-Wallis rank sum test
```

```
##
## data: PARTNERS by HIV
## Kruskal-Wallis chi-squared = 32.036, df = 1, p-value = 1.514e-08
```

An alternative way of looking at the data is as a tabulation:

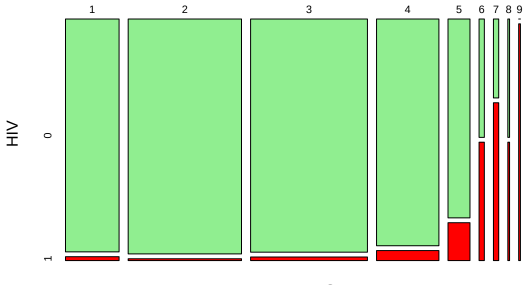
```
table(PARTNERS, HIV)
```

##	H	IIV	
##	PARTNERS	0	1
##	1	60	1
##	2	128	1
##	3	131	2
##	4	68	3
##	5	21	4
##	6	3	3
##	7	2	4
##	8	1	1
##	9	0	2

You can use the plot() function to represent this table graphically:

```
plot(table(PARTNERS, HIV), color = c("lightgreen", "red"))
```

table(PARTNERS, HIV)



PARTNERS

There appears to be an association between the number of sexual PARTNERS in the previous twelve months and positive HIV serology. The proportion with positive HIV serology increases as the number of sexual partners increases:

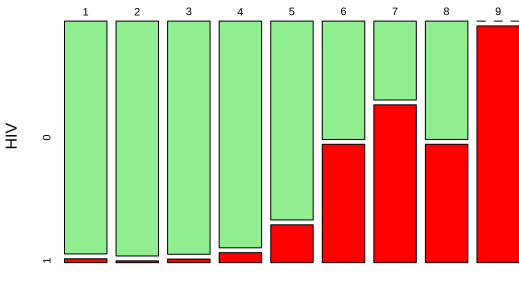
```
prop.table(table(PARTNERS, HIV), 1) * 100
```

```
##
           HIV
## PARTNERS
                                   1
##
             98.3606557
                           1.6393443
          1
##
          2
             99.2248062
                           0.7751938
          3
            98.4962406
                           1.5037594
##
##
          4
            95.7746479
                           4.2253521
##
          5
            84.0000000
                         16.0000000
          6 50.0000000
                         50.0000000
##
##
          7
            33.333333
                         66.666667
             50.0000000
##
          8
                         50.0000000
##
          9
              0.0000000 100.0000000
```

The '1' instructs the prop.table() function to calculate row proportions. You can also use the plot() function to represent this table graphically:

```
plot(prop.table(table(PARTNERS, HIV), 1) * 100, color = c("lightgreen", "red"))
```

prop.table(table(PARTNERS, HIV), 1) * 100



PARTNERS

The *chi-square test for trend* is an appropriate test to perform on this data. The prop.trend.test() function that performs the *chi-square test for trend* requires you to specify the *number of events* and the *number of trials*. In this table:

table (PARTNERS, HIV)

```
## HIV
## PARTNERS 0 1
## 1 60 1
```

```
2 128
##
                      1
            3 131
                      2
##
            4
               68
                      3
##
            5
               21
                      4
##
##
            6
                 3
                      3
                 2
            7
                      4
##
                 1
##
            8
                      1
                 0
                      2
            9
##
```

The *number of events* in each row is in the second column (labelled 1) and the *number of trials* is the total number of cases in each row of the table.

We can extract this data from a table object:

```
tab <- table(PARTNERS, HIV)
events <- tab[ ,2]
trials <- tab[ ,1] + tab[ ,2]

tab <- table(PARTNERS, HIV)
events <- tab[ ,2]
trials <- tab[ ,1] + tab[ ,2]</pre>
```

Another way of creating the trials object would be to use the apply() function to sum the rows of the tab object:

```
trials <- apply(tab, 1, sum)
```

Pass this data to the prop.trend.test() function:

```
prop.trend.test(events, trials)
```

```
##
## Chi-squared Test for Trend in Proportions
##
## data: events out of trials ,
## using scores: 1 2 3 4 5 6 7 8 9
## X-squared = 76.389, df = 1, p-value < 2.2e-16</pre>
```

With a linear trend such as this we can use PARTNERS in a logistic model without recoding or creating indicator variables. We can now specify and fit the logistic regression model:

```
##
## Call:
## glm(formula = HIV ~ GUD + TRAVOUT + PARTNERS, family = binomial(logit))
##
```

```
## Deviance Residuals:
##
        Min
                   1Q
                         Median
                                        3Q
                                                 Max
## -1.70415 -0.19849
                      -0.11148 -0.06247
                                             3.11742
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
                -9.4854
                            1.4663 -6.469 9.86e-11 ***
## (Intercept)
## GUD
                 1.3869
                            0.5937
                                     2.336
                                              0.0195 *
## TRAVOUT
                 2.0867
                            0.9547
                                     2.186
                                              0.0288 *
## PARTNERS
                 1.1605
                            0.2050
                                     5.662 1.50e-08 ***
## ---
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 167.364 on 425
                                       degrees of freedom
## Residual deviance: 99.377
                               on 422 degrees of freedom
     (9 observations deleted due to missingness)
## AIC: 107.38
##
## Number of Fisher Scoring iterations: 8
```

We can use the lreg.or() function that we wrote earlier to calculate and display odds ratios and confidence intervals:

```
lreg.or(gudhiv.lreg)
```

```
## OR LCI UCI
## (Intercept) 0.00 0.00 0.00
## GUD 4.00 1.25 12.81
## TRAVOUT 8.06 1.24 52.35
## PARTNERS 3.19 2.14 4.77
```

PARTNERS is incorporated into the logistic model as a continuous variable.

The odds ratio reported for PARTNERS is the odds ratio associated with a unit increase in the number of sexual PARTNERS. A man reporting five sexual partners, for example, was over three times as likely (odds ratio = 3.19) to have a positive HIV-2 serology than a man reporting four sexual partners.

An alternative approach would be to have created an *indicator* variables:

```
part.gt.5 <- ifelse(PARTNERS > 5, 1, 0)
```

This creates a new variable (part.gt.5) that indicates whether or not an individual subject reported having more than five sexual partners in the previous twelve months:

table(PARTNERS, part.gt.5)

```
##
            part.gt.5
## PARTNERS
               0
                    1
##
              61
                    0
           1
           2 129
##
                    0
##
           3 133
                    0
              71
##
           4
                    0
##
           5
              25
                    0
##
           6
               0
                    6
##
           7
               0
                    6
                    2
               0
##
           8
##
           9
               0
                    2
```

You can also inspect this on a case-by-case basis:

cbind(PARTNERS, part.gt.5)

```
##
           PARTNERS part.gt.5
      [1,]
##
                    2
                               0
##
      [2,]
                    2
                               0
      [3,]
                    1
                                0
##
      [4,]
                    2
                               0
##
      [5,]
                    3
                                0
##
##
      [6,]
                    2
                                0
      [7,]
##
                    4
                               0
                    5
##
      [8,]
                               0
      [9,]
                    2
                               0
##
     [10,]
                    3
##
                               0
     [11,]
                    4
                               0
##
     [12,]
                    3
##
                               0
##
     [13,]
                    1
                                0
                    2
     [14,]
##
                                0
##
     [15,]
                    5
                               0
##
     [16,]
                    8
                                1
##
     [17,]
                    5
                                0
     [18,]
                    3
##
                                0
##
    [19,]
                    2
                               0
##
     [20,]
                    1
                                0
##
     [21,]
                    2
                                0
                    3
     [22,]
                                0
##
    [23,]
                    2
                               0
##
                    3
##
     [24,]
                               0
##
    [25,]
                    4
                               0
##
     [26,]
                    3
                               0
                               0
##
     [27,]
                    4
```

##	[28,]	3	0
##	[29,]	4	0
##	[30,]	5	0
##	[31,]	4	0
##	[32,]	3	0
##	[33,]	4	0
##	[34,]	5	0
##	[35,]	2	0
##	[36,]	4	0
##	[37,]	3	0
##	[38,]	2	0
##	[39,]	1	0
##	[40,]	2	0
##	[41,]	3	0
##	[42,]	4	0
##	[43,]	3	0
##	[44,]	3	0
##	[45,]	2	0
##	[46,]	4	0
##	[47,]	5	0
##	[48,]	4	0
##	[49,]	3	0
##	[50,]	4	0
##	[51,]	1	0
##	[52,]	1	0
##	[53,]	2	0
##	[54,]	3	0
##	[55,]	3	0
##	[56,]	3	0
##	[57,]	3	0
##	[58,]	4	0
##	[59,]	5	0
##	[60,]	4	0
##	[61,]	3	0
##	[62,]	3	0
##	[63,]	5	0
##	[64,]	2	0
##	[65,]	2	0
##	[66,]	3	0
##	[67,]	2	0
##	[68,]	1	0
##	[69,]	2	0
##	[70,]	3	0
##	[71,]	2	0
##	[72,]	3	0

##	[73,]	4	0
##	[74,]	3	0
##	[75,]	3	0
##	[76,]	2	0
##	[77,]	5	0
##	[78,]	4	0
##	[79,]	3	0
##	[80,]	1	0
##	[81,]	2	0
##	[82,]	5	0
##	[83,]	3	0
##	[84,]	7	1
##	[85,]	6	1
##	[86,]	5	0
##	[87,]	5	0
##	[,88]	5	0
##	[89,]	4	0
##	[90,]	3	0
##	[91,]	2	0
##	[92,]	5 1	0
## ##	[93,] [94,]	1	0
##	[95,]	1	0
##	[96,]	1	0
##	[97,]	2	0
##	[98,]	3	0
##	[99,]	4	0
##	[100,]	3	0
##	[101,]	3	0
##	[102,]	2	0
##	[103,]	3	0
##	[104,]	4	0
##	[105,]	3	0
##	[106,]	2	0
##	[107,]	3	0
##	[108,]	4	0
##	[109,]	3	0
##	[110,]	3	0
##	[111,]	4	0
##	[112,]	3	0
##	[113,]	6	1
##	[114,]	3	0
##	[115,]	4	0
##	[116,]	3	0
##	[117,]	3	0

##	[118,]	2	0
##	[119,]	3	0
##	[120,]	4	0
##	[121,]	7	1
##	[122,]	3	0
##	[123,]	2	0
##	[124,]	3	0
##	[125,]	4	0
##	[126,]	3	0
##	[127,]	2	0
##	[128,]	3	0
##	[129,]	4	0
##	[130,]	8	1
##	[131,]	5	0
##	[132,]	6	1
##	[133,]	5	0
##	[134,]	4	0
##	[135,]	4	0
##	[136,]	4	0
##	[137,]	3	0
##	[138,]	4	0
##	[139,]	3	0
##	[140,]	3	0
##	[141,]	2	0
##	[142,]	2	0
##	[143,]	1	0
##	[144,]	2	0
##	[145,]	1	0
##	[146,]	2	0
##	[147,]	3	0
##	[148,]	1	0
##	[149,]	2	0
##	[150,]	3	0
##	[151,]	2	0
##	[152,]	2	0
##	[153,]	2	0
##	[154,]	1	0
##	[155,]	1	0
##	[156,]	2	0
##	[157,]	3	0
##	[158,]	3	0
##	[159,]	2	0
##	[160,]	3	0
##	[161,]	4	0
##	[162,]	2	0

##	[163,]	5	0
	[164,]	4	0
##			
##	[165,]	2	0
##	[166,]	3	0
##	[167,]	2	0
##	[168,]	2	0
##	[169,]	1	0
##	[170,]	4	0
##	[171,]	3	0
##	[172,]	3	0
##	[173,]	2	0
##	[174,]	3	0
##	[175,]	2	0
##	[176,]	4	0
##	[177,]	3	0
##	[178,]	2	0
##	[179,]	3	0
##	[180,]	4	0
##	[181,]	2	0
##	[182,]	3	0
##	[183,]	3	0
##	[184,]	4	0
##	[185,]	2	0
##	[186,]	3	0
##	[187,]	2	0
##	[188,]	2	0
##	[189,]	3	0
##	[190,]	3	0
##	[191,]	2	0
##	[192,]	3	0
##	[193,]	2	0
##	[194,]	4	0
##	[195,]	3	0
##	[196,]	2	0
##	[197,]	2	0
##	[198,]	3	0
##	[199,]	2	0
##	[200,]	3	0
##	[201,]	2	0
##	[202,]	3	0
##	[203,]	3	0
##	[204,]	2	0
##	[205,]	3	0
##	[206,]	2	0
##	[207,]	3	0

##	[208,]	2	0
##	[209,]	1	0
##	[210,]	6	1
##	[211,]	9	1
##	[212,]	1	0
##	[213,]	2	0
##	[214,]	3	0
##	[215,]	4	0
##	[216,]	5	0
##	[217,]	4	0
##	[218,]	5	0
##	[219,]	5	0
##	[220,]	5	0
##	[221,]	4	0
##	[222,]	3	0
##	[223,]	4	0
##	[224,]	3	0
##	[225,]	2	0
##	[226,]	1	0
##	[227,]	2	0
##	[228,]	3	0
##	[229,]	2	0
##	[230,]	1	0
##	[231,]	4	0
##	[232,]	3	0
##	[233,]	4	0
##	[234,]	3	0
##	[235,]	3	0
##	[236,]	2	0
##	[237,]	2	0
##	[238,]	1	0
##	[239,]	2	0
##	[240,]	3	0
##	[241,]	2	0
##	[242,]	1	0
##	[243,]	2	0
##	[244,]	4	0
##	[245,]	3	0
##	[246,]	2	0
##	[247,]	3	0
##	[248,]	2	0
##	[249,]	2	0
##	[250,]	1	0
##	[251,]	2	0
##	[252,]	3	0

##	[253,]	2	0
##	[254,]	3	0
##	[255,]	1	0
##	[256,]	2	0
##	[257,]	3	0
##	[258,]	2	0
##	[259,]	4	0
##	[260,]	3	0
##	[261,]	3	0
##	[262,]	2	0
##	[263,]	2	0
##	[264,]	1	0
##	[265,]	1	0
##	[266,]	1	0
##	[267,]	1	0
##	[268,]	1	0
##	[269,]	1	0
##	[270,]	1	0
##	[271,]	1	0
##	[272,]	1	0
##	[273,]	1	0
##	[274,]	2	0
##	[275,]	3	0
##	[276,]	2	0
##	[277,]	3	0
##	[278,]	2	0
##	[279,]	1	0
##	[280,]	2	0
##	[281,]	3	0
##	[282,]	4	0
##	[283,]	3	0
##	[284,]	2	0
##	[285,]	3	0
##	[286,]	2	0
##	[287,]	1	0
##	[288,]	2	0
##	[289,]	4	0
##	[290,]	7	1
##	[291,]	1	0
##	[292,]	4	0
##	[293,]	1	0
##	[294,]	3	0
##	[295,]	3	0
##	[296,]	4	0
##	[297,]	3	0

##	[298,]	2	0
##	[299,]	2	0
##	[300,]	1	0
##	[301,]	2	0
##	[302,]	3	0
##	[303,]	3	0
##	[304,]	3	0
##	[305,]	2	0
##	[306,]	4	0
##	[307,]	4	0
##	[308,]	5	0
##	[309,]	4	0
##	[310,]	4	0
##	[311,]	9	1
##	[312,]	3	0
##	[313,]	3	0
##	[314,]	2	0
##	[315,]	2	0
##	[316,]	1	0
##	[317,]	1	0
##	[318,]	2	0
##	[319,]	7	1
##	[320,]	3	0
##	[321,]	2	0
##	[322,]	1	0
##	[323,]	2	0
##	[324,]	4	0
##	[325,]	6	1
##	[326,]	5	0
##	[327,]	3	0
##	[328,]	2	0
##	[329,]	3	0
##	[330,]	4	0
##	[331,]	3	0
##	[332,]	2	0
##	[333,]	3	0
##	[334,]	4	0
##	[335,]	3	0
##	[336,]	3	0
##	[337,]	2	0
##	[338,]	3	0
##	[339,]	4	0
##	[340,]	3	0
##	[341,]	2	0
##	[342,]	3	0

##	[343,]	1	0
##	[344,]	1	0
##	[345,]	2	0
##	[346,]	3	0
##		4	
	[347,]	3	0 0
##	[348,]		
##	[349,]	3	0
##	[350,]	2	0
##	[351,]	4	0
##	[352,]	5	0
##	[353,]	4	0
##	[354,]	3	0
##	[355,]	3	0
##	[356,]	2	0
##	[357,]	2	0
##	[358,]	1	0
##	[359,]	4	0
##	[360,]	1	0
##	[361,]	1	0
##	[362,]	4	0
##	[363,]	3	0
##	[364,]	2	0
##	[365,]	1	0
##	[366,]	4	0
##	[367,]	1	0
##	[368,]	2	0
##	[369,]	3	0
##	[370,]	1	0
##	[371,]	5	0
##	[372,]	4	0
##	[373,]	3	0
##	[374,]	2	0
##	[375,]	1	0
##	[376,]	2	0
##	[377,]	3	0
##	[378,]	2	0
##	[379,]	4	0
##	[380,]	2	0
##	[381,]	3	0
##	[382,]	4	0
##	[383,]	7	1
##	[384,]	3	0
##	[385,]	2 4	0
##	[386,]	4	0
##	[387,]	4	0

##	[388,]	3	0
##	[389,]	2	0
##	[390,]	2	0
##	[391,]	1	0
##	[392,]	2	0
##	[393,]	6	1
##	[394,]	7	1
##	[395,]	2	0
##	[396,]	1	0
##	[397,]	2	0
##	[398,]	3	0
##	[399,]	1	0
##	[400,]	2	0
##	[401,]	3	0
##	[402,]	2	0
##	[403,]	1	0
##	[404,]	2	0
##	[405,]	3	0
##	[406,]	2	0
##	[407,]	3	0
##	[408,]	2	0
##	[409,]	3	0
##	[410,]	2	0
##	[411,]	4	0
##	[412,]	2	0
##	[413,]	2	0
##	[414,]	1	0
##	[415,]	2	0
##	[416,]	3	0
##	[417,]	2	0
##	[418,]	3	0
##	[419,]	2	0
##	[420,]	3	0
##	[421,]	2	0
##	[422,]	3	0
##	[423,]	4	0
##	[424,]	2	0
##	[425,]	2	0
##	[426,]	3	0
##	[427,]	4	0
##	[428,]	4	0
##	[429,]	1	0
##	[430,]	2	0
##	[431,]	3	0
##	[432,]	2	0

```
## [433,]
                            0
                 1
## [434,]
                 1
                            0
## [435,]
                 2
                            0
We can now specify and fit the logistic regression model using our indicator variable:
gudhiv.lreg <- glm(formula = HIV ~ GUD + TRAVOUT + part.gt.5,</pre>
                   family = binomial(logit))
summary(gudhiv.lreg)
lreg.or(gudhiv.lreg)
##
## Call:
## glm(formula = HIV ~ GUD + TRAVOUT + part.gt.5, family = binomial(logit))
##
## Deviance Residuals:
                                             Max
##
       Min
                 1Q
                       Median
                                    3Q
## -1.6092 -0.2205
                     -0.2205
                               -0.0719
                                          3.4521
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
                -5.9559
                             0.9850 -6.046 1.48e-09 ***
## (Intercept)
## GUD
                 1.4930
                             0.5805
                                      2.572
                                               0.0101 *
## TRAVOUT
                 2.2514
                             0.9319
                                      2.416
                                               0.0157 *
## part.gt.5
                 4.6791
                             0.7560
                                      6.189 6.05e-10 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 167.36
                               on 425
                                       degrees of freedom
## Residual deviance: 106.43
                               on 422
                                       degrees of freedom
     (9 observations deleted due to missingness)
## AIC: 114.43
##
## Number of Fisher Scoring iterations: 7
                         LCI
                                UCI
##
                   OR
                        0.00
                               0.02
## (Intercept)
                 0.00
## GUD
                 4.45
                        1.43
                              13.89
## TRAVOUT
                 9.50
                       1.53
                              59.02
## part.gt.5
               107.67 24.47 473.84
```

For this exercise there is no need to save the workspace image so click the No or Don't

We can now quit R:

q()

Save button (GUI) or enter n when prompted to save the workspace image (terminal).

5.1 Summary

- $\bullet\,$ Using built-in functions and our own functions we can use R to analyse epidemiological data.
- The power of R is that it can be easily extended. Many user-contributed functions (usually packages of related functions) are available for download over the Internet. We will use one of these packages in the next exercise.

Chapter 6

Extending R with packages

R has no built-in functions for survival analysis but, because it is an extensible system, survival analysis is available as an add-in package. You can find a list of add-in packages at the R website.

```
http://www.r-project.org/
```

Add-in packages are installed from the Internet. There are a series of R functions that enable you to download and install add-in packages.

The survival package adds functions to R that enable it to analyse survival data. This package may be downloaded and installed using install.packages("survival") or from the Packages or Packages & Data menu if you are using a GUI version of R.

Packages are loaded into R as they are needed using the library() function. Start R and load the survival package:

```
library(survival)
```

Before we go any further we should retrieve a dataset:

```
ca <- read.table("ca.dat", header = TRUE)
attach(ca)

## The following objects are masked from ca (pos = 10):
##
## group, status, time</pre>
```

The columns in this dataset on the survival of cancer patients in two different treatment groups are as follows:

$\overline{ ext{time}}$	Survival or censoring time (months)
status	Censoring status (1=dead, 0=censored)
group	Treatment group $(1/2)$

##

##

##

21

23

27

12

11

10

1

1

1

0.5347

0.4861

0.4375

0.1033

0.1047

0.1049

0.36614

0.31866

0.27340

0.781

0.742

0.700

summary(ca.surv)

We next need to create a survival object from the time and status variables using the Surv() function:

```
response <- Surv(time, status)</pre>
```

We can then specify the model for the survival analysis. In this case we state that survival (response) is dependent upon the treatment group:

```
ca.surv <- survfit(response ~ group)</pre>
```

The summary() function applied to a survfit object lists the survival probabilities at each time point with 95% confidence intervals:

Call: survfit(formula = response ~ group) ## ## group=1 ## time n.risk n.event survival std.err lower 95% CI upper 95% CI ## 8 22 1 0.955 0.0444 0.8714 1.000 9 1 ## 21 0.909 0.0613 0.7966 1.000 ## 13 19 0.0744 0.7270 1.000 1 0.861 ## 17 0.0856 0.6591 0.997 14 1 0.811 0.0940 ## 18 16 1 0.760 0.5963 0.968 1 ## 19 15 0.709 0.1005 0.5373 0.936 ## 21 14 1 0.659 0.1053 0.4814 0.901 ## 23 13 1 0.608 0.1087 0.4282 0.863 ## 1 30 10 0.547 0.1136 0.3643 0.822 9 ## 31 1 0.486 0.3046 0.776 0.1161 ## 32 8 0.426 1 0.1164 0.2489 0.727 7 ## 1 34 0.365 0.1146 0.1971 0.675 ## 48 5 1 0.292 0.1125 0.1371 0.621 ## 3 1 56 0.195 0.1092 0.0647 0.585 ## ## group=2 ## time n.risk n.event survival std.err lower 95% CI upper 95% CI 0.9583 ## 4 24 1 0.0408 0.88163 1.000 23 2 ## 5 0.8750 0.0675 0.75221 1.000 ## 6 21 1 0.8333 0.0761 0.69681 0.997 ## 7 20 1 0.7917 0.0829 0.64478 0.972 ## 8 19 2 0.7083 0.0928 0.54795 0.916 ## 9 17 1 0.6667 0.0962 0.50240 0.885 ## 11 16 1 0.6250 0.0988 0.45845 0.852 ## 12 15 1 0.5833 0.1006 0.41598 0.818

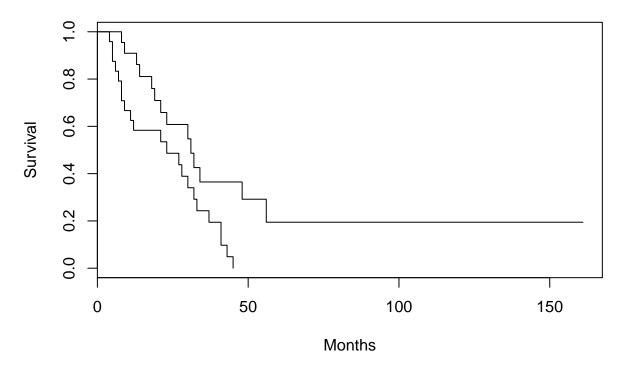
##	28	9	1	0.3889	0.1039	0.23032	0.657
##	30	8	1	0.3403	0.1017	0.18945	0.611
##	32	7	1	0.2917	0.0981	0.15088	0.564
##	33	6	1	0.2431	0.0930	0.11481	0.515
##	37	5	1	0.1944	0.0862	0.08157	0.464
##	41	4	2	0.0972	0.0650	0.02624	0.360
##	43	2	1	0.0486	0.0473	0.00722	0.327
##	45	1	1	0.0000	NaN	NA	NA

Printing the ca.surv object provides another view of the results:

ca.surv

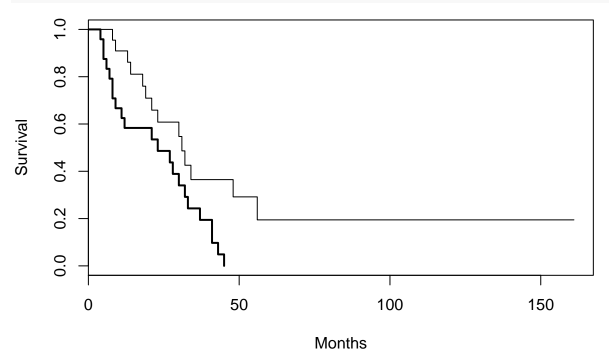
The plot() function with a survfit object displays the survival curves:

```
plot(ca.surv, xlab = "Months", ylab = "Survival")
```



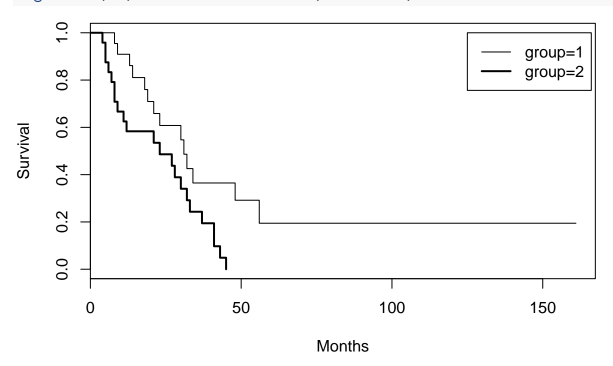
We can make it easier to distinguish between the two lines by specifying a width for each line using the lwd parameter of the plot() function:





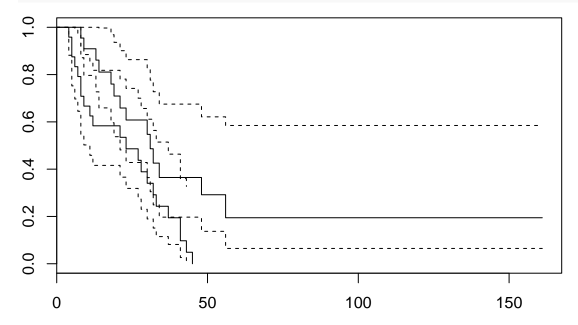
It would also be useful to add a legend:

legend(125, 1, names(ca.surv\$strata), lwd = c(1, 2))



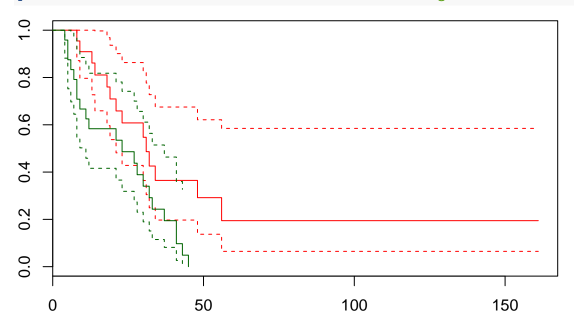
If there is only one survival curve to plot then plotting a survfit object will plot the survival curve with 95% confidence limits. You can specify that confidence limits should be plotted when there is more than one survival curve but the results can be disappointing:

```
plot(ca.surv, conf.int = TRUE)
```



Plots can be improved by specifying different colours for each curve:

```
plot(ca.surv, conf.int = TRUE, col = c("red", "darkgreen"))
```



We can perform a formal test of the two survival times using the **survdiff()** function:

```
survdiff(response ~ group)
```

```
## Call:
## survdiff(formula = response ~ group)
##
## N Observed Expected (O-E)^2/E (O-E)^2/V
```

```
## group=1 22 14 21.1 2.38 6.26
## group=2 24 22 14.9 3.36 6.26
##
## Chisq= 6.3 on 1 degrees of freedom, p= 0.0123
```

We can now quit R:

```
q()
```

For this exercise there is no need to save the workspace image so click the **No** or **Don't Save** button (GUI) or enter **n** when prompted to save the workspace image (terminal).

6.1 Summary

- R can be extended by adding additional packages. Some packages are included with the standard R installation but many others are available and may be downloaded from the Internet.
- You can find a list of add-in packages at the R website: http://www.r-project.org/
- Packages may also be downloaded and installed from this site using the install.packages() function or from the Packages or Packages & Data menu if you are using a GUI version of R.
- Packages are loaded into R as they are needed using the library() function. You can use the search() function to display a list of loaded packages and attached data.frames.

Chapter 7

##

Making your own objects behave like R objects

In the previous exercises we concentrated on writing functions that take some input data, analyse it, and display the results of the analysis. The standard R functions we have used all do this. The fisher.test() function, for example, takes a table object (or the names of two variables) as input and calculates and displays the p- value for Fisher's exact test and the odds ratio and associated confidence interval for two-by-two tables:

```
fem <- read.table("fem.dat", header = TRUE)</pre>
attach(fem)
## The following objects are masked from fem (pos = 6):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
## The following objects are masked from fem (pos = 7):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
   The following objects are masked from fem (pos = 9):
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
##
  The following objects are masked from fem (pos = 10):
##
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
## The following objects are masked from fem (pos = 15):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
fisher.test(SEX, LIFE)
```

```
Fisher's Exact Test for Count Data
##
##
## data: SEX and LIFE
## p-value = 0.03175
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
     1.080298 14.214482
## sample estimates:
## odds ratio
##
     3.620646
The results of the fisher.test() function may also be saved for later use:
ft <- fisher.test(SEX, LIFE)</pre>
ft
##
   Fisher's Exact Test for Count Data
##
##
## data: SEX and LIFE
## p-value = 0.03175
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
     1.080298 14.214482
## sample estimates:
## odds ratio
##
     3.620646
The fisher.test() function returns an object of the class htest:
class(ft)
## [1] "htest"
which is a list containing the output of the fisher.test() function. Each item of output
is stored as a different named item in the list:
names(ft)
str(ft)
## [1] "p.value"
                      "conf.int"
                                     "estimate"
                                                    "null.value" "alternative"
## [6] "method"
                      "data.name"
## List of 7
## $ p.value
                  : num 0.0318
```

\$ conf.int : atomic [1:2] 1.08 14.21 ..- attr(*, "conf.level")= num 0.95

..- attr(*, "names")= chr "odds ratio"

\$ estimate : Named num 3.62

##

##

```
$ null.value : Named num 1
##
    ..- attr(*, "names")= chr "odds ratio"
## $ alternative: chr "two.sided"
## $ method
                 : chr "Fisher's Exact Test for Count Data"
## $ data.name : chr "SEX and LIFE"
## - attr(*, "class")= chr "htest"
Each of these items can be referred to by name:
ft$estimate
ft$conf.int
## odds ratio
##
     3.620646
## [1] 1.080298 14.214482
## attr(,"conf.level")
## [1] 0.95
When you display the output of the fisher.test() function either by calling the function
directly:
fisher.test(SEX, LIFE)
##
   Fisher's Exact Test for Count Data
##
##
## data: SEX and LIFE
## p-value = 0.03175
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
##
     1.080298 14.214482
## sample estimates:
## odds ratio
##
     3.620646
or by typing the name of an object created using the fisher.test() function:
ft
##
##
   Fisher's Exact Test for Count Data
##
## data: SEX and LIFE
## p-value = 0.03175
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
     1.080298 14.214482
##
## sample estimates:
```

```
## odds ratio
## 3.620646
```

The print() function takes over and formatted output is produced. The print() function knows about htest class objects and produces output of the correct format for that class of object. This means that any function that produces an htest object (or any other standard R object) does not need to include R commands to produce formatted output.

All hypothesis testing functions supplied with R produce objects of the htest class and use the print() function to produce formatted output. For example:

```
tt <- t.test(WT ~ LIFE)
class(tt)
tt
  [1] "htest"
##
##
   Welch Two Sample t-test
##
##
## data: WT by LIFE
## t = 0.60608, df = 98.866, p-value = 0.5459
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.3326225 0.6251763
## sample estimates:
## mean in group 1 mean in group 2
##
         0.7867213
                         0.6404444
```

You can use this feature of R in your own functions. We will explore this by writing a function to test the null hypothesis that the *variance to mean ratio* of a vector of numbers is equal to one. Such a test might be used to investigate the spatial distribution (e.g. over natural sampling units such as households) of cases of a disease.

Create a new function using the function() function:

```
v2m.test <- function(data) {}</pre>
```

And start the function editor:

```
fix(v2m.test)
```

Now edit this function to make it do something useful:

```
function(data) {
  nsu <- length(data)
  obs <- sum(data)
  m <- obs / nsu
  v <- var(data)
  vmr <- v / m</pre>
```

```
chi2 \leftarrow sum((data - m)^2) / m
  df <- nsu - 1
  p <- 1 - pchisq(chi2, df)
  names(chi2) <- "Chi-square"</pre>
  names(df) <- "df"</pre>
  names(vmr) <- "Variance : mean ratio"</pre>
  v2m <- list(method = "Variance to mean test",</pre>
                data.name = deparse(substitute(data)),
                statistic = chi2,
                parameter = df,
                p.value = p,
                estimate = vmr)
  class(v2m) <- "htest"</pre>
  return(v2m)
}
v2m.test <- function(data) {</pre>
  nsu <- length(data)</pre>
  obs <- sum(data)
  m <- obs / nsu
  v <- var(data)</pre>
  vmr <- v / m
  chi2 \leftarrow sum((data - m)^2) / m
  df <- nsu - 1
  p <- 1 - pchisq(chi2, df)
```

names(chi2) <- "Chi-square"</pre>

names(vmr) <- "Variance : mean ratio"</pre>

v2m <- list(method = "Variance to mean test",</pre>

statistic = chi2,
parameter = df,
p.value = p,
estimate = vmr)

data.name = deparse(substitute(data)),

names(df) <- "df"</pre>

class(v2m) <- "htest"</pre>

return(v2m)

}

Once you have made the changes shown above, check your work, save the file, and quit the editor.

Before proceeding we should examine the v2m.test() function to make sure we understand what is happening:

1. The first eight lines after the opening curly bracket ({) contain the required calculations.

- 2. The next three lines use the names() function to give our variables names that will make sense in formatted output.
- 3. The next line creates a list of items that the function returns using some of the names used by htest class objects.
- 4. The next line tells R that the list object called v2m is of the class htest.
- 5. The next line causes the function to return the v2m object (i.e. a list of class htest containing the named items method, data.name, statistic, parameter, p.value, and estimate).
- 6. The final line ends the function definition.

Note that objects of class htest may contain items with the following names:

Item	$\mathbf{U}\mathbf{sage}$	
method Text description of the test used to title output		
data.name	Name(s) of data or variables used for the test	
null.value	The null value	
statistic	Value of test statistic	
parameter	A test parameter such as the degrees of freedom of the test	
	statistic	
p.value	The p-value of the test	
estimate An estimate (e.g. the mean)		
conf.int Confidence interval of estimate		
alternative Text describing the alternative hypothesis		
note Text note		

We are now ready to test the v2m.test() function. This table:

Number of cases: 0 1 2 3 4 6 Number of households: 24 29 26 14 5 2

shows the number of cases of chronic (stunting) undernutrition found in a random sample of 100 households.

We can reproduce the data behind this table using a combination of the c() and rep() functions:

```
## stunt
## 0 1 2 3 4 6
## 24 29 26 14 5 2
```

And use it to test our new v2m.test() function:

```
v2m.test(stunt)
```

Which should produce the following output:

```
v2m.test(stunt)
```

```
##
## Variance to mean test
##
## data: stunt
## Chi-square = 110.16, df = 99, p-value = 0.2083
## sample estimates:
## Variance: mean ratio
##
## 1.11274
```

If your vm2.test() function does not produce this output then use the fix() function:

```
fix(v2m.test)
```

to check and edit the vm2.test() function and try again.

The important thing to note from this exercise is that R allows us to specify a class for the output of our functions. This means that we can use standard R classes and functions to (e.g.) produce formatted output without us having to write commands to format the output ourselves.

More importantly, it also means that we can write functions that return values when we need them to return values but can also produce formatted output when we need them to produce formatted output.

Our v2m.test() function can produce values for later use:

```
vm <- v2m.test(stunt)
vm$p.value</pre>
```

```
## [1] 0.2083442
```

or produce formatted output:

```
v2m.test(stunt)
```

```
##
## Variance to mean test
##
## data: stunt
## Chi-square = 110.16, df = 99, p-value = 0.2083
## sample estimates:
## Variance: mean ratio
##

1.11274
```

This way of working is not limited to using standard R classes and functions.

R also allows us to define our own classes. We will explore this by defining functions and a new class to deal with two-by-two tables.

We need to create two functions:

- 1. One function will handle the calculations.
- 2. A second function function will produce formatted output when required.

Create a new function using the function() function:

```
rr22 <- function(exposure, outcome) {}</pre>
```

And start the function editor:

```
fix(rr22)
```

Now edit this function to make it do something useful:

```
function(exposure, outcome) {
  tab <- table(exposure, outcome)
  a <- tab[1,1]
  b <- tab[1,2]
  c <- tab[2,1]
  d <- tab[2,2]
  rr <- (a / (a + b)) / (c / (c + d))
  se.log.rr <- sqrt((b / a) / (a + b) + (d / c) / (c + d))
  lci <- exp(log(rr) - 1.96 * se.log.rr)
  uci <- exp(log(rr) + 1.96 * se.log.rr)
  rr22.output <- list(estimate = rr, ci = c(lci, uci))
  class(rr22.output) <- "rr22"
  return(rr22.output)
}</pre>
```

Once you have made the changes shown above, save the file and quit the editor.

The rr22() function is similar to the tab2by2() function that you created in the second exercise of this tutorial except that the function now returns a list of values instead of formatted output:

```
fem <- read.table("fem.dat", header = TRUE)
attach(fem)
rr22.test <- rr22(SEX, LIFE)
names(rr22.test)
rr22.test$estimate
rr22.test$conf.int
rr22.test$conf.int[1]
rr22.test$conf.int[2]</pre>
```

The following objects are masked from fem (pos = 3):

```
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
##
## The following objects are masked from fem (pos = 7):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
##
   The following objects are masked from fem (pos = 8):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
   The following objects are masked from fem (pos = 10):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
## The following objects are masked from fem (pos = 11):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
## The following objects are masked from fem (pos = 16):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
  [1] "estimate" "ci"
## [1] 2.054167
## NULL
## NULL
## NULL
The function returns a list of class rr22:
class(rr22.test)
## [1] "rr22"
The displayed output from the rr22() function is, however, not pretty:
print(rr22.test)
rr22(SEX, LIFE)
## $estimate
## [1] 2.054167
##
## $ci
## [1] 0.966417 4.366232
## attr(,"class")
## [1] "rr22"
```

```
## $estimate
## [1] 2.054167
##
## $ci
## [1] 0.966417 4.366232
##
## attr(,"class")
## [1] "rr22"
```

This can be fixed by creating a new function:

```
print.rr22 <- function(x) {}</pre>
```

And start the function editor:

```
fix(print.rr22)
```

Now edit this function to make it do something useful:

Once you have made the changes shown above, check your work, save the file, and quit the editor.

The function name print.rr22() indicates that this function contains the print method for objects of class rr22. All objects of class rr22 will use the function print.rr22() instead of the standard R print() function to produce formatted output:

```
rr22(SEX, LIFE)
rr22.test <- rr22(SEX, LIFE)
rr22.test
print(rr22.test)</pre>
```

```
## RR : 2.054167

## 95% CI : 0.966417; 4.366232

## RR : 2.054167

## 95% CI : 0.966417; 4.366232

## RR : 2.054167

## 95% CI : 0.966417; 4.366232
```

Note that we can still extract returned values from an rr22 class object:

```
rr22.test$estimate
```

The print.rr22() function only controls the way an entire rr22 object is displayed.

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You might like to use the save() function to save the v2m.test(), rr22(), and print.rr22() functions before quitting R. We can now quit R:

q()

For this exercise there is no need to save the workspace image so click the **No** or **Don't Save** button (GUI) or enter **n** when prompted to save the workspace image (terminal).

7.1 Summary

- R objects can be assigned a class or type.
- Objects of a specific class or type may share functions that extract and manipulate data common to members of that class. This allows you to write functions that handle data that is common to all members of that class (e.g. to produce formatted output for hypothesis testing functions).
- R provides a set of ready-made classes (e.g. htest) which can be used by standard R functions such as the print() and summary() functions.
- R allows you to create new classes and class-specific functions that can extract and manipulate data common to the new classes.
- Classes allows you to create versatile functions that return values when you need them to return values but can also produce formatted output when you need them to produce formatted output.
- Classes allow you to write functions that can be chained together so that the output of one function is the input of another function.

Chapter 8

Writing your own graphical functions

R provides a pretty full set of graphical functions for plotting data as well as plot() methods for a wide variety of statistical functions. There will be times, however, when you will need to write you own graphical functions to present and analyse data in a specific way. In this exercise we will create a function that produces a plot that may be used for assessing agreement between two methods of clinical measurement as described in:

Bland JM, Altman DG. Statistical Methods for Assessing Agreement Between Two Methods of 310.

Which involves plotting the difference of two measurements against the mean of the two measurements and calculating and displaying limits of agreement.

Start R and retrieve and attach the sample dataset:

```
ba <- read.table("ba.dat", header = TRUE)
attach(ba)

## The following objects are masked from ba (pos = 10):
##
## Mini, Wright</pre>
```

The ba data frame contains measurements (in litres per minute) taken with a Wright peak flow meter and a Mini-Wright peak flow meter. This is the same data that is presented in the referenced Lancet article:

```
##
      Wright Mini
## 1
         494
              512
## 2
         395
             430
## 3
         516
             520
## 4
         434
             428
## 5
         476 500
## 6
         557
              600
## 7
         413
              364
## 8
         442
              380
```

```
## 9
         650 658
## 10
         433
             445
## 11
         417
              432
## 12
              626
         656
## 13
         267
              260
## 14
         478 477
## 15
         178
              259
## 16
         423
              350
## 17
         427
              451
```

You can examine the ba data.frame using the print() and summary() functions:

```
print(ba)
ba
summary(ba)
```

```
##
      Wright Mini
## 1
         494 512
## 2
         395
              430
## 3
         516 520
         434 428
## 4
## 5
         476
              500
         557
              600
## 6
## 7
         413
              364
## 8
         442
              380
## 9
         650
              658
## 10
         433
              445
## 11
         417
              432
              626
## 12
         656
## 13
         267
              260
## 14
         478
              477
              259
## 15
         178
         423
## 16
              350
## 17
         427
              451
##
      Wright Mini
## 1
         494
              512
## 2
         395
              430
## 3
         516
              520
         434
             428
## 4
              500
## 5
         476
## 6
         557
              600
## 7
         413
              364
## 8
         442
              380
## 9
         650
              658
## 10
         433
              445
```

```
## 11
         417
               432
## 12
         656
               626
## 13
         267
               260
## 14
         478
               477
## 15
         178
               259
         423
## 16
               350
## 17
         427
               451
##
        Wright
                           Mini
##
    Min.
            :178.0
                     Min.
                             :259.0
##
    1st Qu.:417.0
                      1st Qu.:380.0
##
    Median :434.0
                     Median :445.0
##
    Mean
            :450.4
                     Mean
                             :452.5
    3rd Qu.:494.0
                      3rd Qu.:512.0
##
            :656.0
##
    Max.
                     Max.
                             :658.0
```

The function() function allows us to create new functions in R:

```
ba.plot <- function(a, b) {}</pre>
```

This creates an empty function called ba.plot() that expects two parameters called a and b. We could type the whole function in at the R command prompt but it is easier to use a text editor:

```
fix(ba.plot)
```

We will start be writing a basic function which we will gradually improve throughout this exercise.

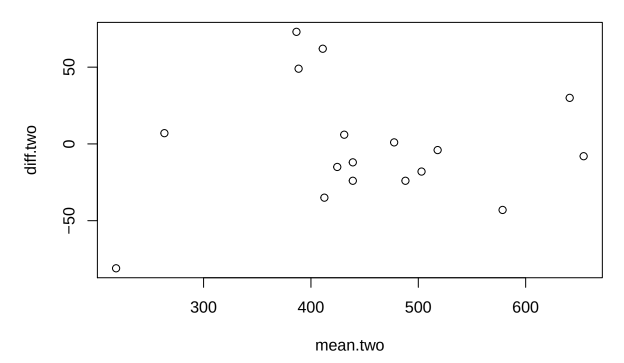
Edit the ba.plot() function to read:

```
function(a, b) {
  mean.two <- (a + b) / 2
  diff.two <- a - b
  plot(mean.two, diff.two)
}</pre>
```

Once you have made the changes shown above, check your work, save the file, and quit the editor.

The function calculates the mean and the difference of the two measures and then plots the results. Lets try the ba.plot() function with the test data:

```
ba.plot(Wright, Mini)
```



The resulting plot is rather plain and lacks meaningful titles and axis labels. Use the fix() function to edit the ba.plot() function:

```
fix(ba.plot)
```

Edit the function to read:

```
function(a, b, title = "Bland and Altman Plot") {
   a.txt <- deparse(substitute(a))
   b.txt <- deparse(substitute(b))
   x.lab <- paste("Mean of", a.txt, "and", b.txt)
   y.lab <- paste(a.txt, "-", b.txt)
   mean.two <- (a + b) / 2
   diff.two <- a - b
   plot(mean.two, diff.two, xlab = x.lab, ylab = y.lab, main = title)
}</pre>
```

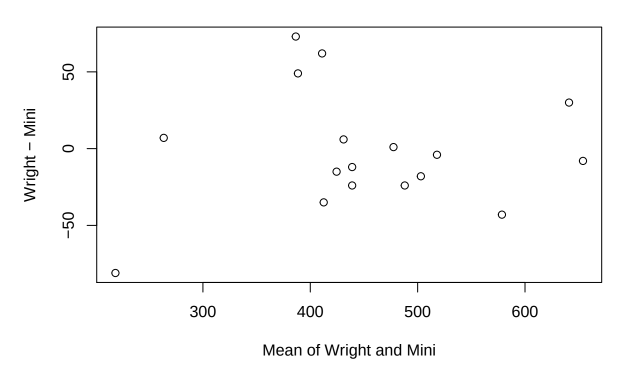
Once you have made the changes shown above, check your work, save the file, and quit the editor.

We have added a new parameter (title) to the function and given this a default value of Bland and Altman Plot. Adding title as a parameter means that we will be able to specify a title for the plot when we call the function. We have also used the function combination deparse(substitute()) to retrieve the names of the vectors passed to parameters a and b. The paste() function pastes pieces of text together. It is used here to create the text for the axis labels used with the plot() function.

Lets try the ba.plot() function with the test data:

ba.plot(Wright, Mini)

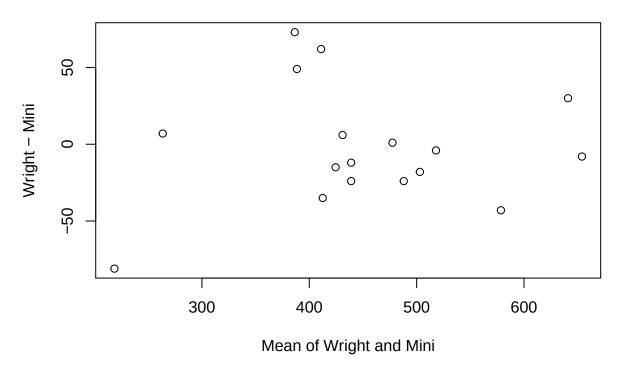
Bland and Altman Plot



We may also specify a title for the plot using the title parameter:

```
ba.plot(Wright, Mini, title = "PEFR data")
```

PEFR data



We can now edit the function to calculate and plot mean, difference, and the limits of agreement. Use the fix() function to edit the ba.plot() function:

```
fix(ba.plot)
```

Edit the function to read:

```
function(a, b, title = "Bland and Altman Plot") {
  a.txt <- deparse(substitute(a))</pre>
  b.txt <- deparse(substitute(b))</pre>
  x.lab <- paste("Mean of", a.txt, "and", b.txt)</pre>
  y.lab <- paste(a.txt, "-", b.txt)</pre>
  mean.two \leftarrow (a + b) / 2
  diff.two <- a - b
  plot(mean.two, diff.two, xlab = x.lab, ylab = y.lab, main = title)
  mean.diff <- mean(diff.two)</pre>
  sd.diff <- sd(diff.two)</pre>
  upper <- mean.diff + 1.96 * sd.diff
  lower <- mean.diff - 1.96 * sd.diff</pre>
  lines(x = range(mean.two), y = c(mean.diff, mean.diff), lty = 3)
  lines(x = range(mean.two), y = c(upper, upper), lty = 3)
  lines(x = range(mean.two), y = c(lower, lower), lty = 3)
}
```

Once you have made the changes shown above, check your work, save the file and quit the editor.

We have used the mean() and sd() functions to calculate the mean and standard deviation of the difference between the two measures and calculated the limits of agreement (upper and lower) assuming that the differences are *Normally* distributed.

The lines() function is then used to plot the mean and the limits of agreement on top of the existing scatter plot.

The parameter lty = 3 used with the lines() function specifies dotted lines.

R provided a great number of graphical parameters that can be used to customise plots. You can see a list of these parameters using:

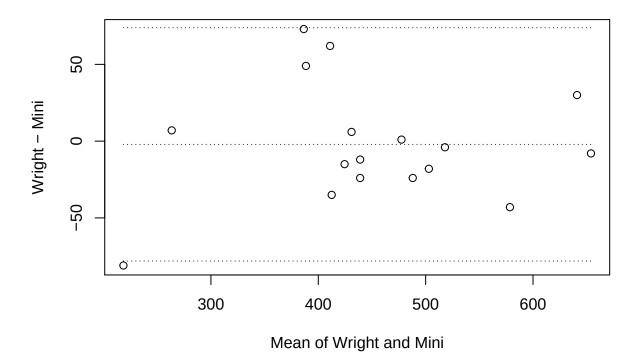
```
help(par)
```

These parameters can be specified for almost all graphical functions.

Lets try the ba.plot() function with the test data:

```
ba.plot(Wright, Mini, title = "Difference vs. mean for PEFR data")
```

Difference vs. mean for PEFR data



The function is almost complete. All that remains to do is to label the lines with the values of the mean difference and the limits of agreement.

Use the fix() function to edit the ba.plot() function:

```
fix(ba.plot)
```

Edit the function to read:

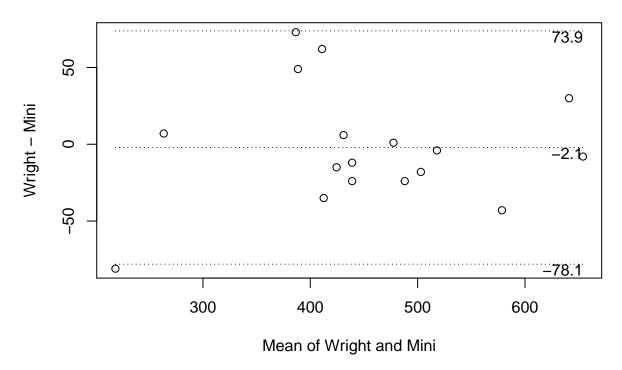
```
function(a, b, title = "Bland and Altman Plot") {
  a.txt <- deparse(substitute(a))</pre>
  b.txt <- deparse(substitute(b))</pre>
  x.lab <- paste("Mean of", a.txt, "and", b.txt)</pre>
  y.lab <- paste(a.txt, "-", b.txt)</pre>
  mean.two <- (a + b) / 2
  diff.two <- a - b
  plot(mean.two, diff.two, xlab = x.lab, ylab = y.lab, main = title)
  mean.diff <- mean(diff.two)</pre>
  sd.diff <- sd(diff.two)</pre>
  upper <- mean.diff + 1.96 * sd.diff
  lower <- mean.diff - 1.96 * sd.diff</pre>
  lines(x = range(mean.two), y = c(mean.diff, mean.diff), lty = 3)
  lines(x = range(mean.two), y = c(upper, upper), lty = 3)
  lines(x = range(mean.two), y = c(lower, lower), lty = 3)
  m.text <- round(mean.diff, digits = 1)</pre>
  u.text <- round(upper , digits = 1)</pre>
  1.text <- round(lower, digits = 1)</pre>
  text(max(mean.two), mean.diff, m.text, adj = c(1,1))
  text(max(mean.two), upper, u.text, adj = c(1,1))
  text(max(mean.two), lower, l.text, adj = c(1,1))
}
```

Once you have made the changes shown above, check your work, save the file, and quit the editor. We have used the round() function to limit the display of the mean difference and the limits of agreement to one decimal place and used the text() function to display these (rounded) values. The adj parameter to the text() function controls the position and justification of text.

Let's try the ba.plot() function with the test data:

```
ba.plot(Wright, Mini, title = "PEFR data")
```

PEFR data



The graphical function is now complete.

One improvement that we could make is for the function to produce a chart and return the values of the mean difference and the limits of agreement.

We would do this in exactly the same way as we would with a non-graphical function. We would return the mean difference and the limits of agreement as members of a list.

We could also specify a class for the returned list and create a class specific **print()** function (or *method*) to produce nicely formatted output.

Use the fix() function to edit the ba.plot() function:

```
fix(ba.plot)
```

Edit the function to read:

```
function(a, b, title = "Bland and Altman Plot") {
   a.txt <- deparse(substitute(a))
   b.txt <- deparse(substitute(b))
   x.lab <- paste("Mean of", a.txt, "and", b.txt)
   y.lab <- paste(a.txt, "-", b.txt)
   mean.two <- (a + b) / 2
   diff.two <- a - b
   plot(mean.two, diff.two, xlab = x.lab, ylab = y.lab, main = title)
   mean.diff <- mean(diff.two)
   sd.diff <- sd(diff.two)</pre>
```

```
upper <- mean.diff + 1.96 * sd.diff
lower <- mean.diff - 1.96 * sd.diff
lines(x = range(mean.two), y = c(mean.diff, mean.diff), lty = 3)
lines(x = range(mean.two), y = c(upper, upper), lty = 3)
lines(x = range(mean.two), y = c(lower, lower), lty = 3)
m.text <- round(mean.diff, digits = 1)
u.text <- round(upper , digits = 1)
l.text <- round(lower, digits = 1)
text(max(mean.two), mean.diff, m.text, adj = c(1,1))
text(max(mean.two), upper, u.text, adj = c(1,1))
text(max(mean.two), lower, l.text, adj = c(1,1))
ba <- list(mean = mean.diff, limits = c(lower, upper))
class(ba) <- "ba"
return(ba)
}</pre>
```

Once you have made the changes shown above, save the file and quit the editor.

Create a print() function for objects of the ba class:

```
print.ba <- function(x) {}</pre>
```

Use the fix() function to edit the new function:

```
fix(print.ba)
```

Edit the function to read:

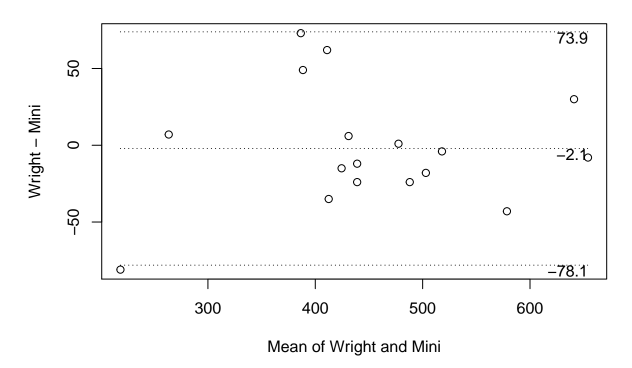
```
function(x) {
  cat("Mean difference : ", x$mean, "\n",
     "Limits of agreement : ", x$limits[1], "; ", x$limits[2], "\n",
     sep = "")
}
```

Once you have made the changes shown above, check your work, save the file, and quit the editor.

Let's try the ba.plot() function with the test data:

```
ba.plot(Wright, Mini, title = "PEFR data")
```

PEFR data



```
## Mean difference : -2.117647
```

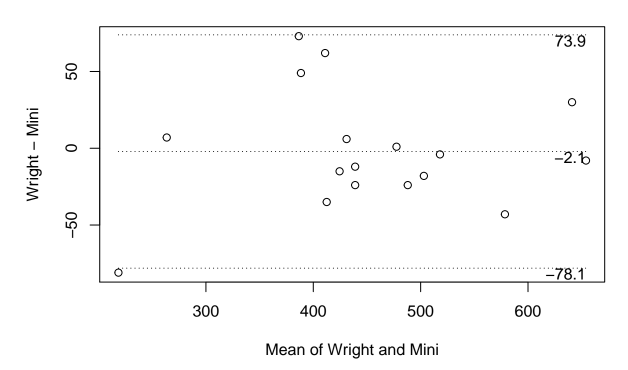
Limits of agreement : -78.0973; 73.86201

The function produces the plot and returns the mean difference and limits of agreement as a list of class ba which is formatted and printed by the print.ba() function.

We can manipulate the returned values just as we would with any other function:

```
ba.test <- ba.plot(Wright, Mini)
print(ba.test)
ba.test
ba.test$mean
ba.test$limits
ba.test$limits[1]
ba.test$limits[2]</pre>
```

Bland and Altman Plot



Mean difference : -2.117647

Limits of agreement : -78.0973; 73.86201

Mean difference : -2.117647

Limits of agreement : -78.0973; 73.86201

[1] -2.117647

[1] -78.09730 73.86201

[1] -78.0973

[1] 73.86201

You might like to use the save() function to save the ba.plot() and print.ba() functions before quitting R.

We can now quit R:

q()

For this exercise there is no need to save the workspace image so click the **No** or **Don't Save** button (GUI) or enter **n** when prompted to save the workspace image (terminal).

8.1 Summary

• R allows you to create functions that produce graphical output.

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• R allows you to create functions that produce graphical output and return values.

- R objects can be assigned a class or type.
- R allows you to create new classes and class-specific functions that can extract and manipulate data common to the new classes.
- Classes allows you to create versatile functions that return values when we need them to return values but can also produce formatted output when we need them to produce formatted output.
- Classes allow you to write functions that can be chained together so that the output of one function is the input of another function.

Chapter 9

More graphical functions

Graphical functions in R are just like any other function in R in the sense that R provides you with a set of functions which can be altered or added to. In this exercise we will experiment with some of the graphical functions provided by R to demonstrate the flexibility of graphical functions in R. We will then use the graphical functions that we experiment with to create some useful graphical functions of our own.

The first function that we will develop will be a function that is capable of plotting two data series on a single graph. We will take this exercise slowly in order to introduce some further graphical functions. Before we go any further we should start R and retrieve a dataset:

```
mal <- read.table("malaria.dat", header = TRUE)
attach(mal)</pre>
```

The file malaria.dat contains data on rainfall (in mm) and the number of cases of malaria reported from health centres in an administrative district of Ethiopia between July 1997 and July 1999. The columns in this dataset are as follows:

Time	Month and year (as text)
Cases	Number of cases of malaria reported
Rain	Rainfall in mm

Examine the dataset:

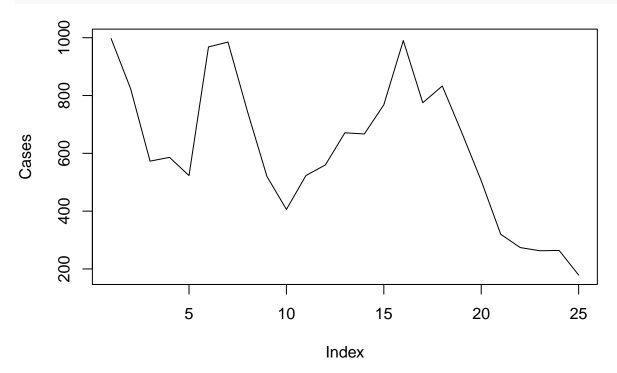
mal

```
##
        Time Cases
                     Rain
## 1
      Jul-97
               997
                     68.5
## 2
      Aug-97
               824 162.1
      Sep-97
               573 138.8
## 4
      Oct-97
               586 222.2
## 5
      Nov-97
               523 115.5
## 6 Dec-97
               968
                    37.2
```

```
## 7
      Jan-98
               985
                     96.6
      Feb-98
               745
                     99.1
## 8
## 9
      Mar-98
               520
                     51.2
## 10 Apr-98
               406
                     80.0
## 11 May-98
               523 112.4
## 12 Jun-98
               560 183.7
## 13 Jul-98
                671 101.0
## 14 Aug-98
                667 252.5
## 15 Sep-98
               768
                     40.8
## 16 Oct-98
                990 193.7
## 17 Nov-98
                775
                     17.5
## 18 Dec-98
               833
                      0.0
## 19 Jan-99
                672
                     33.5
## 20 Feb-99
                505
                      0.0
## 21 Mar-99
                320 106.8
## 22 Apr-99
                274 117.4
## 23 May-99
                263 175.8
## 24 Jun-99
               264 187.5
## 25 Jul-99
               179 283.5
```

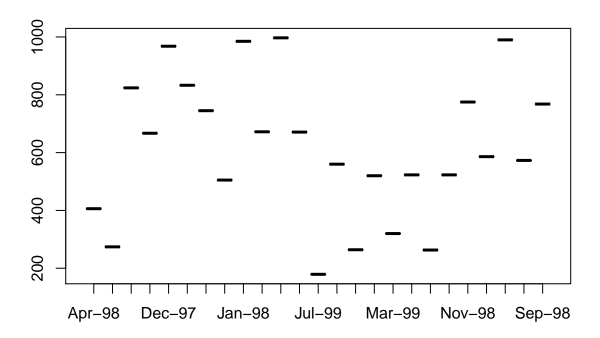
First we will plot the number of cases of malaria seen over time using the plot() function:

plot(Cases, type = "1")



The problem with this plot is that it does not treat the data as a time series. Adding the Time variable to the plot does not solve the problem:

```
plot(Time, Cases, type = "1")
```



Because Time is a factor variable. If you convert Time to a character variable using as.character() or prevent R from converting Time to a factor using the as.is parameter to the read.table() function the plot() function will return an error because it expects a numeric x-axis variable. We should, instead, specify a time series (ts) class object. Rather than change the original data, we will create a new object using the ts() function:

```
cases.ts \leftarrow ts(Cases, start = c(1997, 7), frequency = 12)
```

Examine the cases.ts object:

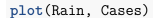
```
cases.ts
```

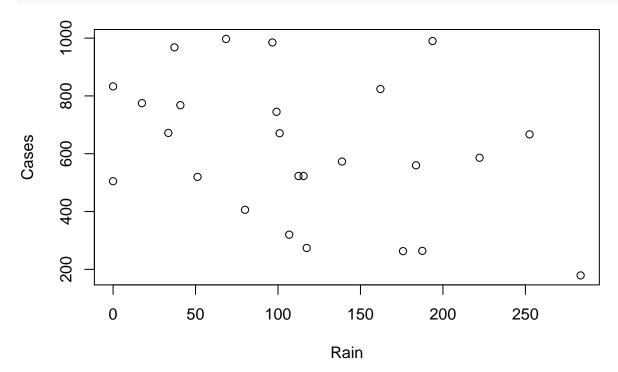
We can now plot cases.ts as a time series:

```
plot(cases.ts)
```



We might want to explore the association between the Rain and Cases variables. A simple scatter plot is not particularly informative:

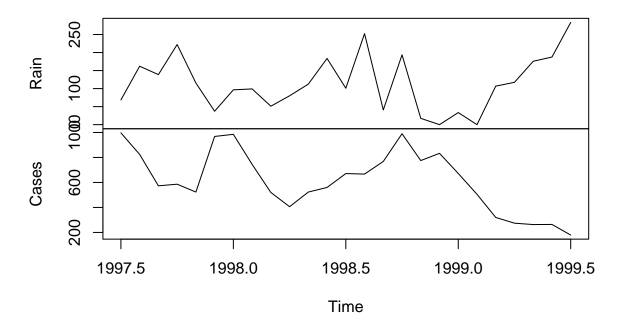




It is better to treat both variables as time series (which they are) and use the built-in plot() methods for objects of class ts:

```
rain.cases.ts <- ts(cbind(Rain, Cases), start = c(1997,7), frequency = 12)
plot(rain.cases.ts)</pre>
```

rain.cases.ts



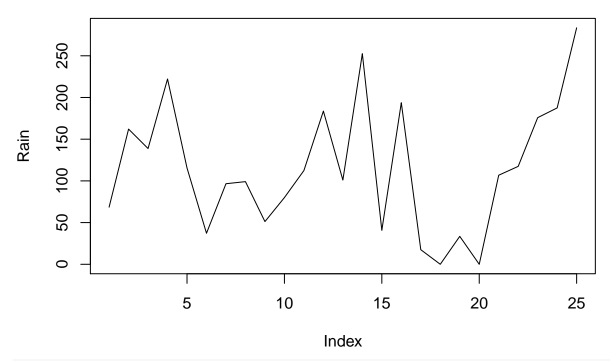
The association between the Rain and Cases variables is now clearer with the number of malaria cases peaking shortly after peaks in rainfall.

The plot() function when used with objects of class ts produces useful output but it is not particularly flexible and the output is, sometimes, not particularly pretty. We can however use basic graphical functions to produce multiple plots. First we will set the mfrow graphical parameter using the par() function:

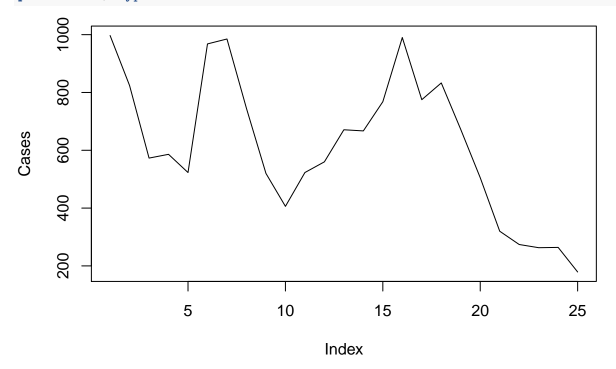
```
par(mfrow = c(2, 1))
```

The par() function sets a graphical parameter. The mfrow parameter is used to set the number of charts that will appear on a page in rows and columns. We have specified two rows with one chart per row. Test this by plotting two charts:

```
plot(Rain, type = "1")
```





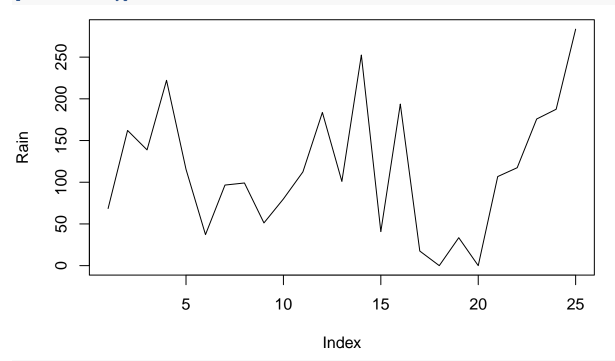


We will want to have tick-marks on the x-axis of each for each record. We can set the number of tick-marks on axes by setting the lab graphical parameter using the par() function:

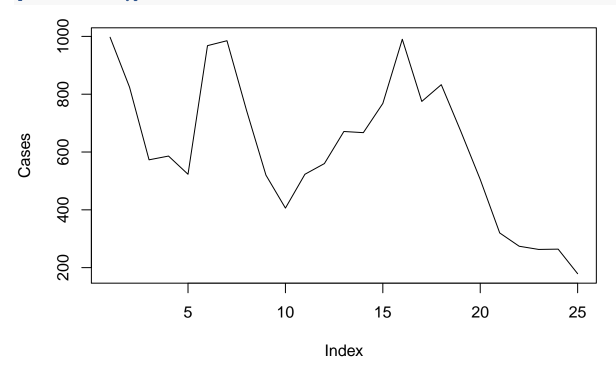
The par() function sets a graphical parameter. The lab parameter is used to set the number tick-marks on the x and y axes and the label size. We have specified a tick-mark on the x-axis for each record (i.e. using length(Time)), ten tick-marks on the y-axis, and a label

length of seven. Test this by plotting two charts:

plot(Rain, type = "1")



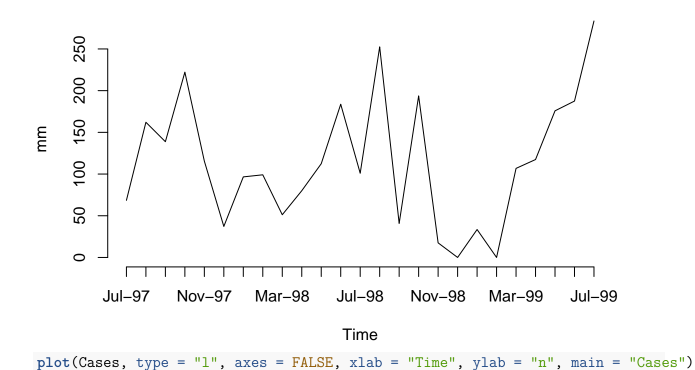
plot(Cases, type = "1")



The problem with these charts is that the month and year are not displayed on the x-axis. We can get round this by plotting a chart without axes and then specifying the axes and labels directly:

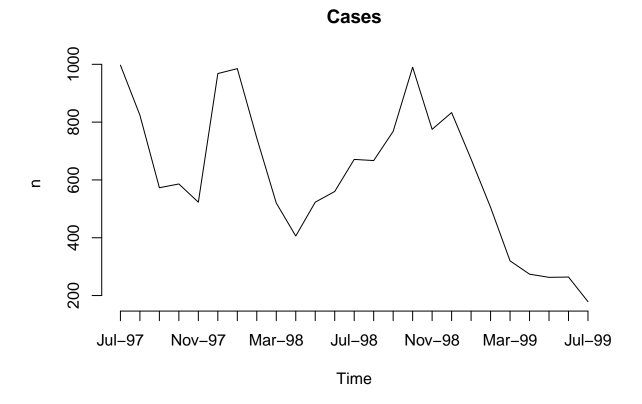
```
plot(Rain, type = "l", axes = FALSE, xlab = "Time", ylab = "mm", main = "Rainfall")
axis(side = 1, labels = as.character(Time), at = 1:length(Time))
axis(side = 2)
```

Rainfall



axis(side = 1, labels = as.character(Time), at = 1:length(Time))

axis(side = 2)



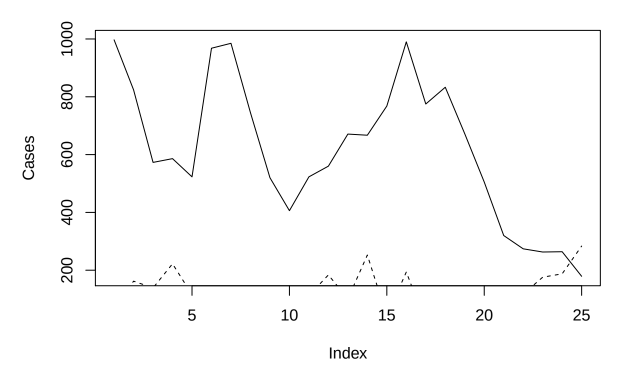
The resulting charts now look much better (you may need to resize the plot to display the x-axis labels correctly) but it would be nice to be able draw the two lines on a single chart.

Before proceeding we will use the par() function to specify one plot per window (using the mfrow parameter) and set the default number of tick-marks on the axes (using the lab parameter):

```
par(mfrow = c(1, 1))
par(lab = c(5, 5, 7))
```

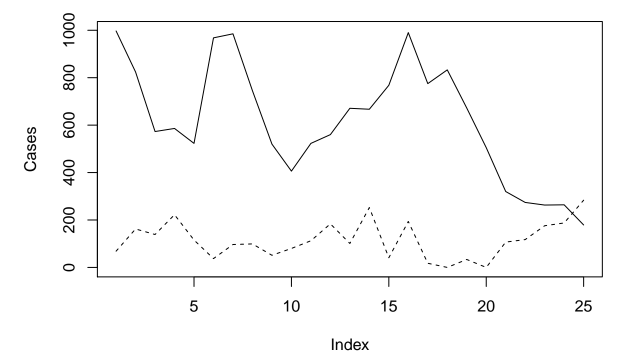
And then use the plot() and lines() function to draw the two lines on the same graph:

```
plot(Cases, type = "1")
lines(Rain, lty = 2)
```



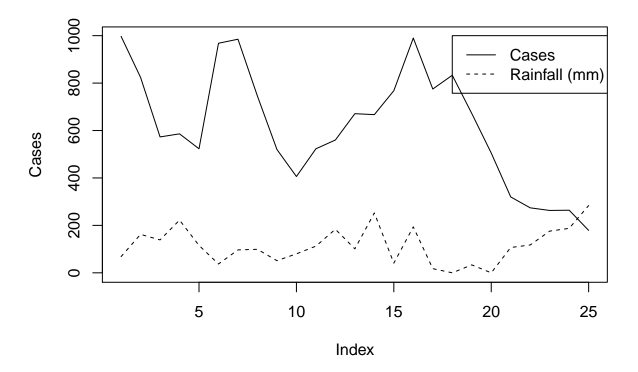
The problem with this is that the ranges of the two variables are different and the plot() function automatically sets the y-axis to the range of the specified variable. To fix this problem we need to set the limits of the y-axis to the minimum and maximum value of both of variables using the ylim parameter of the plot() function:

```
plot(Cases, type = "1", ylim = c(min(Cases, Rain), max(Cases, Rain)))
lines(Rain, lty = 2)
```



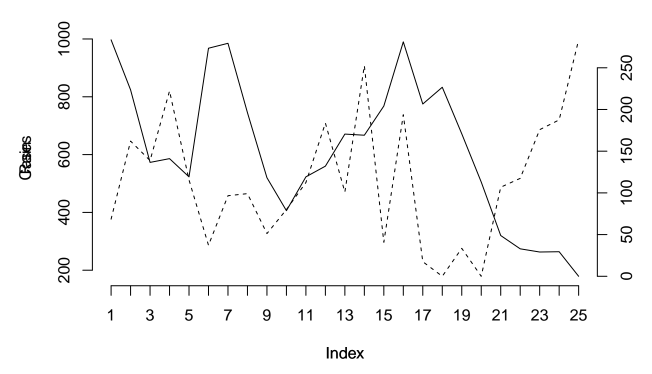
We can improve the chart by adding a legend:

```
legend(18, 1000, legend = c("Cases", "Rainfall (mm)"), lty = c(1,2))
```



We could continue to improve the chart (e.g. by adding labels for the x-axis tick-marks taken from the Time variable, specifying more meaningful axis labels, and specifying a title) but the chart would be more useful if each variable made full use of the plotting area. We can do this by plotting one chart on top of another by using the new graphical parameter:

```
par(lab = c(length(Time), 5, 7))
plot(Cases, type = "l", lty = 1, axes = FALSE)
axis(side = 2)
par(new = TRUE)
plot(Rain, type = "l", lty = 2, axes = FALSE)
axis(side = 4)
axis(side = 1)
```



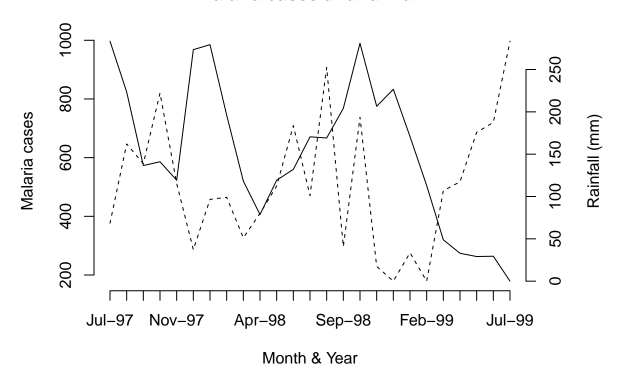
This chart is much clearer but there are still some improvements that could be made:

- The chart should have a title. We can do this using the main parameter of either of the plot() functions.
- The y-axis labels are displayed on top of each other beside the left-hand y-axis. We can solve this problem by preventing the second plot() function from displaying a y-axis label (i.e. by specifying an empty character string for the ylab parameter).
- We will need to make room on the right-hand side of the chart for an axis label (i.e. by setting the mar (margin) graphical parameter) and place the label there ourselves (using the mtext() function).
- The x-axis should display the month and year which are held as character strings in the Time variable. We can do this using the labels parameter of the axis() function after setting the appropriate number of tick-marks using the lab graphical parameter.

The x-axis should be properly labelled. We can do this using the xlab parameters of the plot() functions. An empty string must be specified for one of the plot() functions in order to prevent the default label from being displayed.

Try this now:

Malaria cases and rainfall



Now that we know how to create a two-axis chart, we can write a function that we will be able to use whenever we need to plot two variables on the same chart. Create a new function called plot2var():

```
plot2var <- function() {}</pre>
```

This creates an empty function called plot2var().

Use the fix() function to edit the plot2var() function:

```
fix(plot2var)
```

Edit the function to read:

```
main = paste(y1.lab, "&", y2.lab)) {
  old.par.mar <- par("mar")</pre>
  old.par.lab <- par("lab")</pre>
  par(mar = c(5, 5, 4, 5))
  if(!missing(x.ticks)) {par(lab = c(length(x.ticks), 5, 7))}
  plot(y1, type = "1", lty = 1, axes = FALSE,
       xlab = "", ylab = "", main = main)
  axis(side = 2)
  mtext(text = y1.lab, side = 2, line = 3)
  par(new = TRUE)
  plot(y2, type = "1", lty = 2, axes = FALSE,
       ylab = "", xlab = x.lab)
  axis(side = 4)
  mtext(text = y2.lab, side = 4, line = 3)
  if(!missing(x.ticks)) {
    axis(side = 1, labels = as.character(x.ticks),
         at = 1:length(x.ticks))
  } else {axis(side = 1)}
  par(mar = old.par.mar)
  par(lab = old.par.lab)
}
```

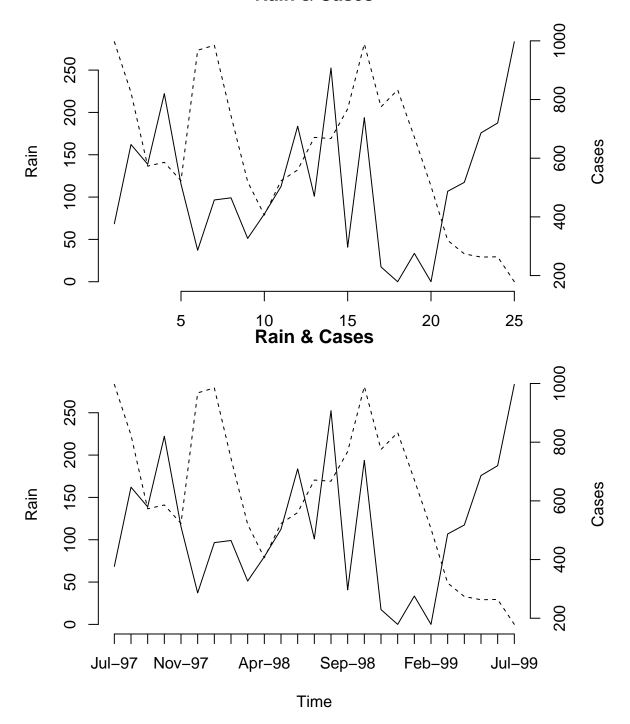
Once you have made the changes shown above, save the file and quit the editor.

Note that with this function we have given some of the parameters default values in the function definition and we have also used the if() function to check whether the user specified a value for the x.ticks parameter. We also save and restore the graphical parameters mar and lab so as to prevent changes to these parameters in the plot2var() function affecting other graphical functions.

Let's try the plot2var() function with the test data:

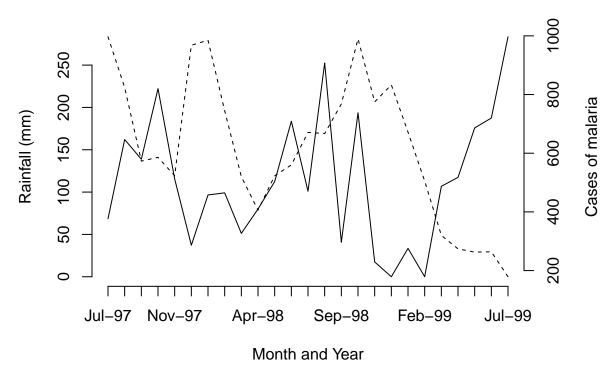
```
plot2var(Rain, Cases)
plot2var(Rain, Cases, Time)
```

Rain & Cases



Note how the function has used default values for the axis labels and chart title. We can override these default values if we want to:





You might like to use the save() function to save the plot2var() function.

As an exercise you might want to edit the plot2var() function to automatically add a legend to the two-axis chart using the legend() function with y1.lab and y2.lab. Before continuing we should detach the mal data.frame:

```
detach(mal)
```

9.1 Population pyramid

A common chart type that is not available in many statistical applications and in R is the population pyramid.

Before we go any further we should retrieve a dataset:

```
pop <- read.table("pop.dat", header = TRUE)
attach(pop)

## The following objects are masked from fem (pos = 4):

##
## AGE, SEX

## The following objects are masked from fem (pos = 5):

##
## AGE, SEX</pre>
```

```
## The following objects are masked from fem (pos = 9):
##
##
       AGE, SEX
## The following objects are masked from fem (pos = 10):
##
##
       AGE, SEX
## The following objects are masked from fem (pos = 12):
##
##
       AGE, SEX
## The following objects are masked from fem (pos = 13):
##
##
       AGE, SEX
  The following objects are masked from fem (pos = 18):
##
##
       AGE, SEX
```

The file pop.dat contains data on the age (in months) and sex of 438 children aged between six and sixty months collected as part of a nutritional anthropometry survey of the Khosh Valley in Northeast Afghanistan.

The columns in this dataset are as follows:

AGE	Age of the child in months
SEX	Sex of the child (M/F)

Examine the first twenty records of the dataset:

```
pop[1:20, ]
```

```
##
      AGE SEX
## 1
         7
             М
        42
## 2
             М
## 3
        60
             М
             F
## 4
        60
## 5
        48
             Μ
## 6
             F
        60
## 7
        18
             М
## 8
        48
             М
## 9
        60
             F
## 10
        36
             Μ
## 11
        24
             F
## 12
        60
             М
## 13
        60
             Μ
## 14
        48
             F
```

##

9

15

0

```
## 15
        18
             Μ
## 16
        60
             М
## 17
         6
             М
         7
## 18
             М
## 19
        12
             М
## 20
        60
             М
```

The first step is to make groups from the AGE variable since many ages are biased towards full years:

```
table(AGE)
barplot(table(AGE), col = "white")
## AGE
##
    6
             9 10 12 13 14 15 17 18 22 23 24 25 26 30 34 36 38 40 42 48 54 60
          8 15
                3 21
                       1
                          3
                             5
                                 1 45
                                       1
                                          1 48
                                                 1
                                                   2 24
                                                          1 80
                                                                2
                                                                    1
                                                                       9 67
                                                                             4 83
80
9
4
20
       6
            8
                10
                     13
                          15
                               18
                                    23
                                         25
                                             30
                                                   36
                                                       40
                                                            48
                                                                 60
```

So we will centre the age-groups around the months representing full years:

0

```
age.group <- cut(AGE, c(0, 17, 29, 41, 53, 99))
```

We can check that the grouping operation has worked as expected by tabulating AGE and age.group:

```
table(AGE, age.group)
##
       age.group
## AGE
         (0,17] (17,29] (29,41] (41,53] (53,99]
     6
              7
                       0
                                 0
                                          0
                                                  0
##
     7
              5
                       0
                                 0
                                          0
                                                  0
##
##
                       0
                                 0
                                          0
                                                  0
     8
              8
```

0

0

##	10	3	0	0	0	0
##	12	21	0	0	0	0
##	13	1	0	0	0	0
##	14	3	0	0	0	0
##	15	5	0	0	0	0
##	17	1	0	0	0	0
##	18	0	45	0	0	0
##	22	0	1	0	0	0
##	23	0	1	0	0	0
##	24	0	48	0	0	0
##	25	0	1	0	0	0
##	26	0	2	0	0	0
##	30	0	0	24	0	0
##	34	0	0	1	0	0
##	36	0	0	80	0	0
##	38	0	0	2	0	0
##	40	0	0	1	0	0
##	42	0	0	0	9	0
##	48	0	0	0	67	0
##	54	0	0	0	0	4
##	60	0	0	0	0	83

We now use the table() function to produce the summary data for the population pyramid: table(age.group, SEX)

```
## SEX

## age.group F M

## (0,17] 34 35

## (17,29] 54 44

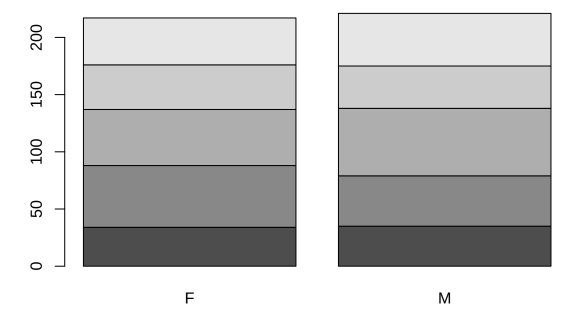
## (29,41] 49 59

## (41,53] 39 37

## (53,99] 41 46
```

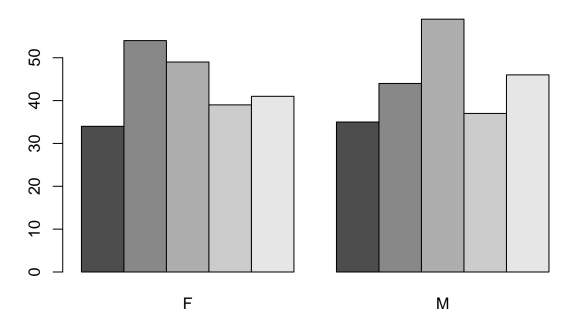
We will construct our population pyramid using the barplot() function:

```
barplot(table(age.group, SEX))
```



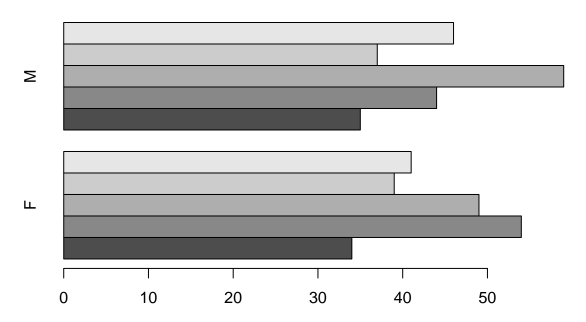
The default behaviour of the barplot() function is to produce stacked bars. We can set the beside parameter to display the bars side-by-side:

```
barplot(table(age.group, SEX), beside = TRUE)
```



We can also use the horiz parameter to present the data as horizontal bars:

```
barplot(table(age.group, SEX), beside = TRUE, horiz = TRUE)
```



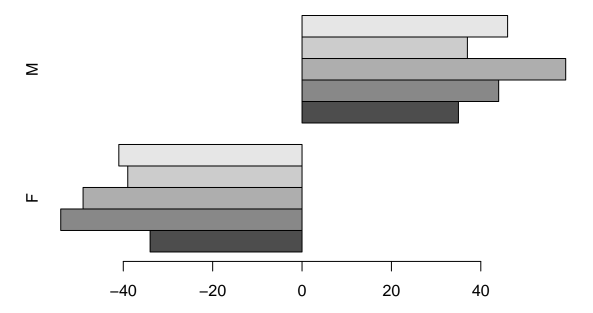
In order to centre the bars around zero we need to make one column of the summary data table contain negative numbers:

```
tab <- table(age.group, SEX)
tab</pre>
```

```
## SEX
## age.group F M
## (0,17] 34 35
## (17,29] 54 44
## (29,41] 49 59
## (41,53] 39 37
## (53,99] 41 46

tab[ ,1] <- -tab[ ,1]
tab</pre>
```

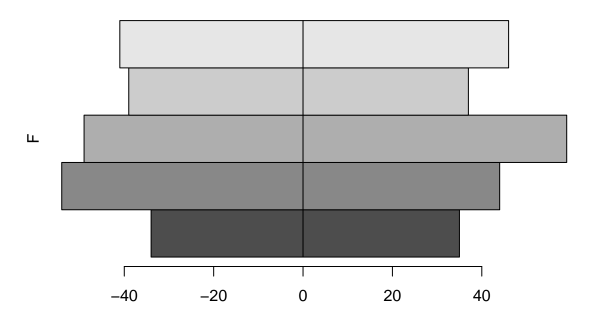
```
##
            SEX
## age.group
                   Μ
     (0,17] -34
##
                  35
     (17,29] -54
##
                  44
     (29,41] -49
##
                  59
     (41,53] -39
##
                  37
##
     (53,99] -41
                  46
barplot(tab, beside = TRUE, horiz = TRUE)
```



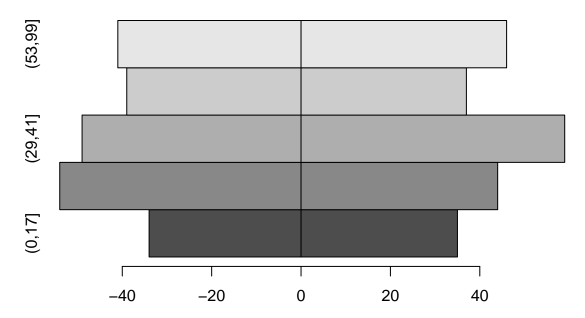
This is looking better.

We still need to shift the second set of bars down beside the first set of bars using the space parameter:

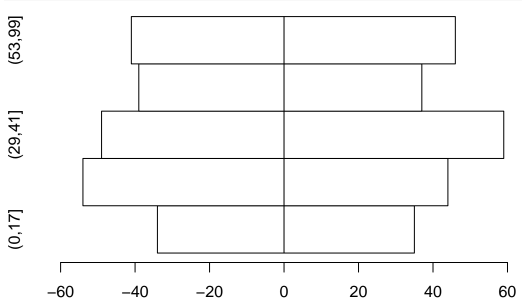
```
barplot(tab, beside = TRUE, horiz = TRUE, space = c(0, -nrow(tab)))
```



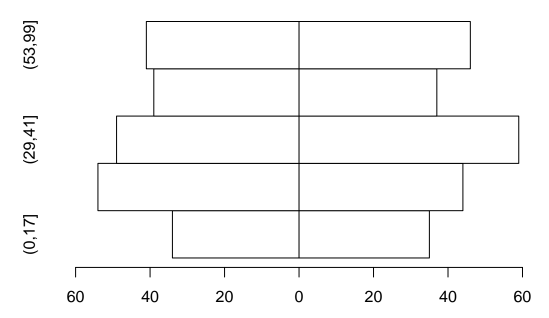
The axis labels are wrong but we can fix that using the names.arg parameter:



The chart can still be improved upon by making the fill-colour of each bar white and by expanding the x-axis slightly:



The chart would be better if the x-axis displayed only positive numbers:



Now we know how to create a population pyramid, we can write a function that we will be able to use whenever we need to plot a population pyramid.

Create a new function called pyramid.plot():

```
pyramid.plot <- function() {}</pre>
```

This creates an empty function called pyramid.plot().

Use the fix() function to edit the pyramid.plot() function:

```
fix(pyramid.plot)
```

Edit the function to read:

```
function(x,
         main = paste("Pyramid plot of", deparse(substitute(x)),
                      "by", deparse(substitute(g))),
         xlab = paste(deparse(substitute(g)),
                      "(", levels(g)[1], "/",levels(g)[2],")"),
         ylab = deparse(substitute(x))) {
  tab <- table(x, g)
  tab[ ,1] <- -tab[ ,1]
  barplot(tab,
          horiz = TRUE,
          beside = TRUE,
          space = c(0, -nrow(tab)),
          names.arg = c(dimnames(tab)$x, dimnames(tab)$x),
          xlim = c(min(tab) * 1.2,
          \max(tab) * 1.2),
          col = "white",
```

```
main = main,
    xlab = xlab,
    ylab = ylab,
    axes = FALSE)

axis(side = 1,
    labels = abs(axTicks(side = 1)),
    at = (axTicks(side = 1)))
}
```

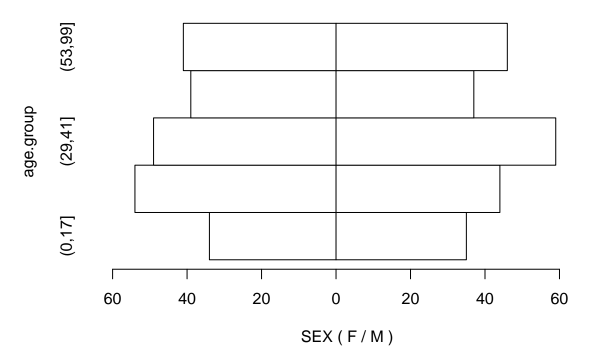
Note that with this function we have given some of the parameters default values in the function definition.

Giving default values to parameters is useful because it means that you do not need to specify parameters such as titles and axis labels unless you want to. Many R functions use default parameters which are usually set to the most frequently used values.

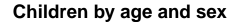
Once you have made the changes shown above, check your work, save the file, and quit the editor. Let's try the pyramid.plot() function with the test data:

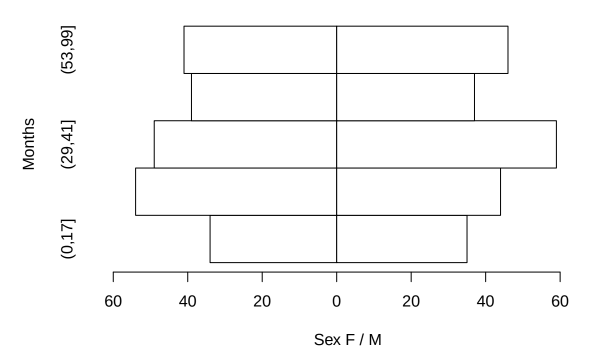
```
pyramid.plot(age.group, SEX)
```

Pyramid plot of age.group by SEX



Note how the function has used default values for the axis labels and chart titles. We can override these default values if we want to:





You might like to use the save() function to save the pyramid.plot() function.

9.2 Pareto chart

Another type of chart that is missing from many statistical applications is the *Pareto* chart which is a bar chart where the bars are sorted by the bar value with the largest bar drawn first. Such a chart is easier to interpret than a pie chart, particularly when there are more than a few categories being plotted.

Before we go any further we should detach the pop data.frame and retrieve a new dataset:

```
detach(pop)
sssw <- read.table("sssw.dat", header = TRUE)
attach(sssw)</pre>
```

The file sssw.dat contains data on the marital status, home circumstances, and ethnic group of 152 persons recruited into a study into the levels of stress experienced by student social workers in the United Kingdom. The columns in this dataset are as follows:

```
children | | | | | 5 = Children | | | | | 6 = Friends or colleagues | + - - + | ethnic | Ethnic group coded as: | | | | | 1 = African | | | | | 2 = West-Indian | | | | | 3 = Indian | | | | | 4 = Pakistani | | | | | 5 = Bangladeshi | | | | | 6 = East African Asian | | | | | 7 = Chinese | | | | | 8 = Cypriot | | | | 9 = Black European | | | | | 10 = White European | | | | 11 = Other | + - - - +
```

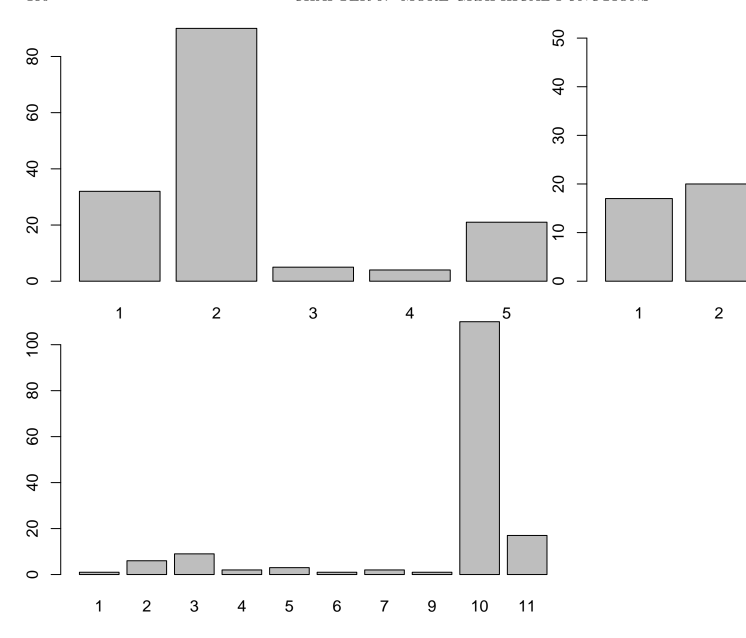
Examine the dataset:

```
sssw[1:20, ]
```

```
marital living ethnic
##
## 1
              2
                      5
                              2
## 2
              1
                      4
                              2
              1
                      1
## 3
                             11
## 4
              1
                      4
                             10
              2
## 5
                      3
                             10
## 6
              2
                      6
                             10
## 7
              1
                      4
                             10
              1
                      4
## 8
                             10
              2
## 9
                      2
                              4
              2
                      6
## 10
                             10
              2
## 11
                      5
                             10
## 12
              1
                      3
                              1
## 13
              5
                      3
                             10
              1
                      4
## 14
                             10
## 15
              2
                      6
                              3
## 16
              1
                      4
                             10
## 17
              1
                      3
                              3
                      3
              5
## 18
                             10
## 19
              3
                      5
                             10
              5
## 20
                      3
                             10
```

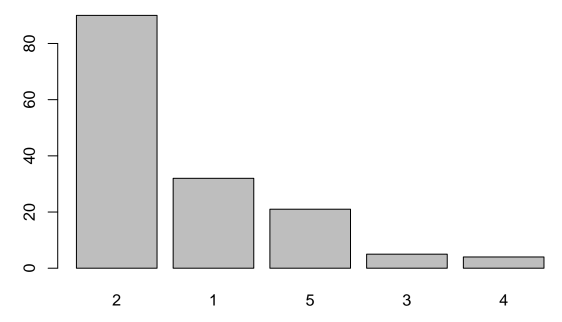
Producing a bar chart from this data is simple as long as we remember to pass summary data (i.e. created using the table() function) to the barplot() function instead of the variable name:

```
barplot(table(marital))
barplot(table(living))
barplot(table(ethnic))
```



Creating a Pareto chart only requires us to sort the summary data. We do this using the rev() and sort() functions:

barplot(rev(sort(table(marital))))



Having to specify rev(sort(table(variable))) each time we want to produce a *Pareto* plot is rather tedious but now that we know how to create a *Pareto* chart, we can write a function that we will be able to use whenever we need to plot a Pareto chart. Create a new function called pareto():

```
pareto <- function() {}</pre>
```

This creates an empty function called pareto(). Use the fix() function to edit the pareto() function:

```
fix(pareto)
```

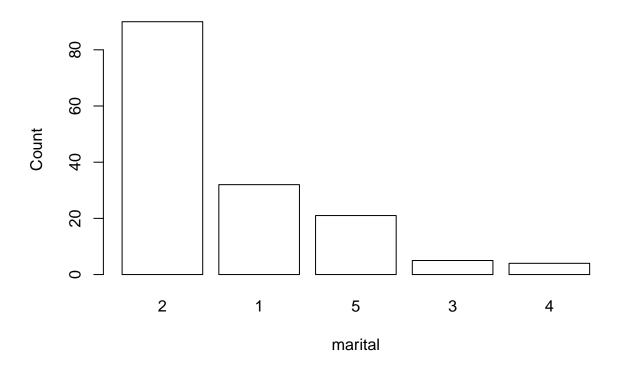
Edit the function to read:

Once you have made the changes shown above, check your work, save the file, and quit the editor.

Let's try the pareto() function with the test data:

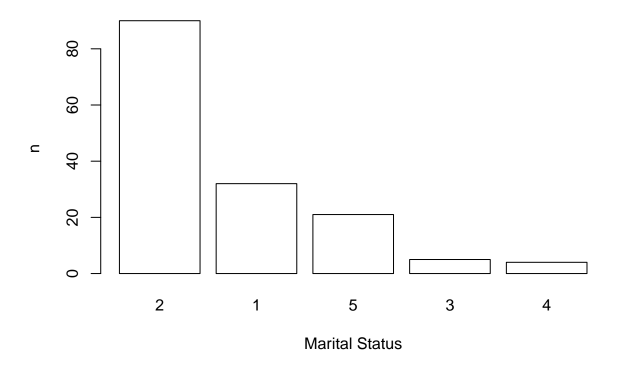
```
pareto(marital)
```

Pareto Chart of marital



Note how the function has used default values for the axis labels and chart titles. We can override these default values if we want to:

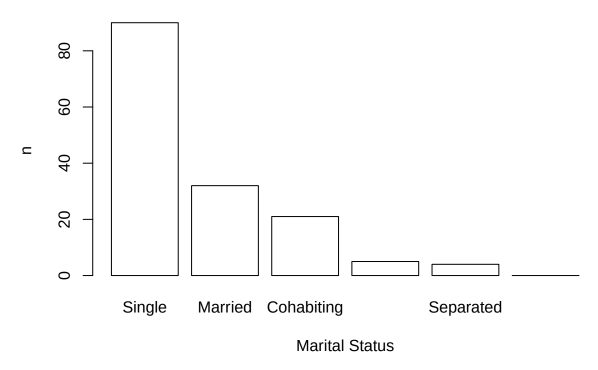
Marital Status



Note that we can use value labels if the variable we plot is a factor with value labels as levels rather than a simple numeric vector:

```
## ms
## Married Single Divorced Separated Cohabiting Widowed
## 32 90 5 4 21 0
```

Marital Status



You may need to resize the plot to display the x-axis labels correctly.

You might like to use the save() function to save the pareto() function.

9.3 Adding confidence intervals or error bars on plots

You may want to plot your data with confidence intervals or error bars. R does not have a function to do this but it is a relatively simple matter to write a function to do so. On the way we will use some of Rs data management functions as well.

Before we go any further we should detach the sssw data.frame and retrieve a new dataset:

```
detach(sssw)
diets <- read.table("diets.dat", header = TRUE)</pre>
```

The file diets.dat contains data from a trial of two different diets undertaken at an adult therapeutic feeding centre in Somalia. The columns in this dataset are as follows:

$\overline{\mathrm{day}}$	The day after start of diet that measurements were taken
oedema	Type of undernutrition coded as: $1 = \text{Oedematous}$; $2 = \text{Marasmic}$
$\operatorname{\mathbf{diet}}$	The trial diets coded as: LP = Low protein; HP = High protein
\mathbf{wt}	Mean weight change weight velocity) in g/kg/day since the previous
	measurement
sd	Standard deviation of weight change in g/kg/day

n Number of subjects at each observation

Examine the dataset:

diets

```
##
      day oedema diet
                           wt
                                 sd
                                      n
## 1
         3
                 1
                     ΗP
                          1.1 11.5
                                     37
## 2
        6
                 1
                     HP
                         1.0 11.0
                                     37
## 3
        9
                 1
                     ΗP
                         1.2
                               8.4
                                     37
## 4
        12
                 1
                     HP -1.0
                               7.4
                                     37
## 5
       15
                     HP -0.9
                               3.2
                                     37
                 1
                     HP -1.7
## 6
        18
                 1
                               5.8
                                     37
## 7
       21
                     HP -1.7
                               4.8
                                     37
                 1
## 8
       24
                     HP -1.7
                                     37
                 1
                               4.8
## 9
       27
                 1
                     HP -1.7
                               5.3
                                     37
## 10
       30
                 1
                     HP -2.5
                               6.8
                                     37
## 11
       33
                 1
                     HP -1.3
                               8.4
                                     37
## 12
                 2
                         5.1 15.1 291
        3
                     ΗP
                         5.1 13.4 291
## 13
                 2
        6
                     HP
## 14
        9
                 2
                     ΗP
                         4.5
                               6.5 291
## 15
                 2
                         5.2
                               9.9 291
        12
                     ΗP
## 16
       15
                 2
                     HP
                         4.6
                               8.2 291
## 17
       18
                 2
                     ΗP
                         5.1 13.4 291
                         5.0 13.4 291
## 18
       21
                 2
                     ΗP
## 19
       24
                 2
                     HP
                          5.2 15.1 291
## 20
                 2
                         5.1 11.6 291
       27
                     ΗP
## 21
                 2
       30
                     ΗP
                         5.0
                               9.2 291
## 22
       33
                 2
                     ΗP
                         5.2
                               8.2 291
## 23
                     LP -3.0 15.7
        3
                 1
                                     65
## 24
        6
                 1
                     LP -2.5 11.9
                                     65
        9
## 25
                 1
                     LP -2.0 13.7
                                     65
## 26
       12
                 1
                     LP
                         2.0 11.0
                                     65
## 27
        15
                 1
                     LP
                         2.1 13.2
                                     65
## 28
                         1.7 16.2
       18
                 1
                     LP
                                     65
## 29
       21
                     LP
                         4.0 11.0
                 1
                                     65
## 30
       24
                 1
                     LP
                         6.5 14.7
                                     65
## 31
       27
                 1
                     LP
                         6.4 10.2
                                     65
## 32
       30
                               8.0
                 1
                     LP
                         6.6
                                     65
## 33
       33
                 1
                     LP
                         6.5 11.0
                                     65
## 34
                 2
        3
                     LP
                          5.1 11.1
                                     86
## 35
        6
                 2
                     LP
                         6.0
                               8.2
                                     86
## 36
        9
                 2
                              8.2
                     LP
                         5.1
                                     86
## 37
       12
                 2
                     LP
                         6.5 11.9
                                     86
```

```
15
                 2
                          6.4
                                      86
## 38
                     LP
                               9.1
## 39
        18
                 2
                     LP
                          5.9 13.8
                                      86
## 40
        21
                 2
                     LP
                          6.1 16.5
                                      86
## 41
        24
                 2
                     LP
                          4.0
                                9.1
                                      86
## 42
        27
                 2
                     LP
                          3.1 12.8
                                      86
## 43
                 2
                     LP
                                7.3
        30
                          4.0
                                      86
## 44
        33
                 2
                     LP
                          5.0
                                9.1
                                      86
```

Note that the dataset contains a summary of the results from the four arms of the trial:

Arm	Oedema	Therapeutic diet
1	Present	High protein
2	Present	Low protein
3	Absent	High protein
4	Absent	Low protein

with observations at 3, 6, 9, 12, 15, 18, 21, 24, 27, 30, and 33 days after admission.

We can calculate a confidence interval for the mean weight velocity (wt) using the data in the sd and n variables. We will use the transform() function to do this:

```
diets <- transform(diets, lci = wt - sd / sqrt(n), uci = wt + sd / sqrt(n))</pre>
```

In this case we are calculating confidence intervals as plus or minus one standard error of the mean.

The transform() function is very useful as it can add columns directly to a data.frame or transform data already stored in a data.frame.

Examine the diets data.frame:

diets

```
##
      day oedema diet
                          wt
                               sd
                                              lci
                                                           uci
                                    n
## 1
        3
                1
                    HP
                         1.1 11.5
                                   37 -0.7905884
                                                    2.99058835
## 2
        6
                    HP
                        1.0 11.0
                                   37 -0.8083889
                                                    2.80838886
                1
## 3
        9
                         1.2
                1
                    ΗP
                              8.4
                                   37 -0.1809515
                                                    2.58095149
## 4
       12
                1
                    HP -1.0
                              7.4
                                   37 -2.2165525
                                                    0.21655251
## 5
       15
                    HP -0.9
                              3.2
                                   37 -1.4260768 -0.37392324
                1
## 6
       18
                1
                    HP -1.7
                              5.8
                                   37 -2.6535141 -0.74648587
## 7
       21
                1
                    HP - 1.7
                              4.8
                                   37 -2.4891151 -0.91088486
## 8
       24
                1
                    HP -1.7
                              4.8
                                   37 -2.4891151 -0.91088486
## 9
       27
                1
                    HP -1.7
                              5.3
                                   37 -2.5713146 -0.82868537
## 10
                    HP - 2.5
                              6.8
                                   37 -3.6179131 -1.38208689
       30
                1
## 11
       33
                    HP -1.3
                              8.4
                                   37 -2.6809515
                1
                                                  0.08095149
                2
## 12
        3
                        5.1 15.1 291
                                        4.2148223
                                                   5.98517768
                        5.1 13.4 291
## 13
        6
                2
                    HP
                                       4.3144781 5.88552191
```

```
## 14
        9
                2
                              6.5 291
                                                     4.88103675
                    ΗP
                         4.5
                                         4.1189633
       12
                2
                         5.2
                              9.9 291
                                                     5.78034828
## 15
                    ΗP
                                         4.6196517
                2
                              8.2 291
                                        4.1193075
                                                     5.08069251
## 16
       15
                    ΗP
                         4.6
## 17
                2
                         5.1 13.4 291
                                        4.3144781
                                                     5.88552191
       18
                    ΗP
## 18
       21
                2
                    HP
                         5.0 13.4 291
                                        4.2144781
                                                     5.78552191
                2
## 19
       24
                    ΗP
                         5.2 15.1 291
                                        4.3148223
                                                     6.08517768
## 20
       27
                2
                    ΗP
                         5.1 11.6 291
                                        4.4199960
                                                     5.78000404
## 21
       30
                2
                    HP
                         5.0
                              9.2 291
                                         4.4606864
                                                     5.53931355
## 22
       33
                2
                              8.2 291
                    ΗP
                         5.2
                                        4.7193075
                                                     5.68069251
## 23
        3
                1
                    LP -3.0 15.7
                                    65 -4.9473453 -1.05265467
## 24
        6
                1
                    LP -2.5 11.9
                                    65 -3.9760133 -1.02398666
## 25
        9
                1
                    LP -2.0 13.7
                                    65 -3.6992759 -0.30072414
## 26
       12
                    LP
                         2.0 11.0
                                    65
                                        0.6356179
                                                     3.36438208
                1
## 27
       15
                1
                    LP
                         2.1 13.2
                                    65
                                        0.4627415
                                                     3.73725850
## 28
       18
                    LP
                         1.7 16.2
                                    65 -0.3093627
                                                     3.70936270
                1
## 29
       21
                1
                    LP
                         4.0 11.0
                                    65
                                        2.6356179
                                                     5.36438208
## 30
       24
                1
                    LP
                         6.5 14.7
                                    65
                                        4.6766894
                                                    8.32331060
## 31
       27
                1
                    LP
                         6.4 10.2
                                    65
                                        5.1348457
                                                     7.66515429
## 32
                         6.6
       30
                1
                    LP
                             8.0
                                    65
                                        5.6077221
                                                     7.59227788
## 33
       33
                    LP
                         6.5 11.0
                                    65
                                        5.1356179
                                                     7.86438208
                1
## 34
                2
                         5.1 11.1
        3
                    LP
                                    86
                                        3.9030562
                                                     6.29694378
## 35
        6
                2
                    LP
                         6.0
                              8.2
                                    86
                                        5.1157713
                                                     6.88422874
## 36
        9
                2
                    LP
                         5.1
                              8.2
                                        4.2157713
                                                     5.98422874
                                    86
## 37
       12
                2
                    LP
                         6.5 11.9
                                    86
                                        5.2167900
                                                     7.78321000
## 38
       15
                2
                    LP
                         6.4
                             9.1
                                    86
                                        5.4187218
                                                     7.38127824
## 39
                2
                         5.9 13.8
       18
                    LP
                                    86
                                        4.4119077
                                                     7.38809227
## 40
       21
                2
                    LP
                         6.1 16.5
                                        4.3207592
                                                    7.87924076
                                    86
## 41
                2
                    LP
                         4.0
                              9.1
       24
                                    86
                                        3.0187218
                                                     4.98127824
## 42
       27
                2
                         3.1 12.8
                    LP
                                        1.7197405
                                                     4.48025950
                                    86
## 43
       30
                2
                    LP
                         4.0
                              7.3
                                    86
                                        3.2128208
                                                     4.78717924
## 44
       33
                2
                    LP
                         5.0
                              9.1
                                    86
                                        4.0187218
                                                     5.98127824
```

Two new columns (lci and uci) have been added.

Now that we have calculated the confidence intervals we should, for convenience, split the diets data.frame into four separate data.frames (one for each arm of the trial):

```
oed.hp <- subset(diets, oedema == 1 & diet == "HP")
oed.lp <- subset(diets, oedema == 1 & diet == "LP")
mar.hp <- subset(diets, oedema == 2 & diet == "HP")
mar.lp <- subset(diets, oedema == 2 & diet == "LP")</pre>
```

Check that each data frame contains the data that you expect it to:

```
oed.hp
oed.lp
mar.hp
```

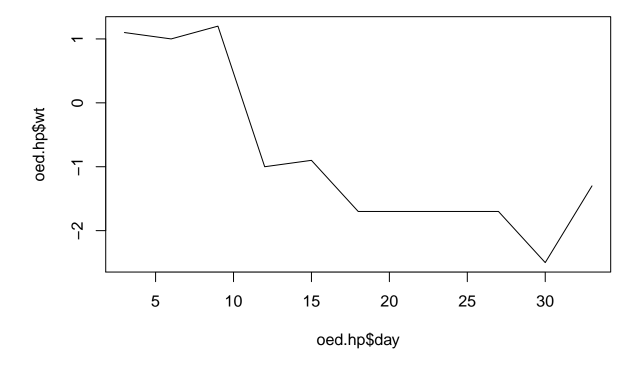
mar.lp

```
##
      day oedema diet
                         wt
                               sd
                                  n
                                            lci
                                                         uci
## 1
        3
                1
                    HP
                        1.1 11.5 37 -0.7905884
                                                  2.99058835
## 2
        6
                1
                    ΗP
                        1.0 11.0 37 -0.8083889
                                                  2.80838886
## 3
        9
                1
                    HP
                        1.2
                             8.4 37 -0.1809515
                                                  2.58095149
## 4
       12
                    HP -1.0
                             7.4 37 -2.2165525
                                                 0.21655251
                1
                    HP -0.9
                             3.2 37 -1.4260768 -0.37392324
## 5
       15
                1
## 6
       18
                    HP - 1.7
                             5.8 37 -2.6535141 -0.74648587
                1
## 7
       21
                             4.8 37 -2.4891151 -0.91088486
                1
                    HP -1.7
                             4.8 37 -2.4891151 -0.91088486
## 8
       24
                1
                    HP -1.7
## 9
       27
                1
                    HP -1.7
                             5.3 37 -2.5713146 -0.82868537
## 10
       30
                    HP -2.5
                             6.8 37 -3.6179131 -1.38208689
                1
## 11
       33
                1
                    HP -1.3
                             8.4 37 -2.6809515 0.08095149
##
      day oedema diet
                         wt
                              sd
                                  n
                                            lci
## 23
        3
                    LP -3.0 15.7 65 -4.9473453 -1.0526547
                1
## 24
        6
                1
                    LP -2.5 11.9 65 -3.9760133 -1.0239867
## 25
        9
                    LP -2.0 13.7 65 -3.6992759 -0.3007241
                1
## 26
       12
                1
                    LP
                        2.0 11.0 65
                                     0.6356179
                                                 3.3643821
## 27
       15
                1
                    LP
                        2.1 13.2 65
                                      0.4627415
                                                 3.7372585
## 28
                        1.7 16.2 65 -0.3093627
       18
                1
                    LP
                                                  3.7093627
                        4.0 11.0 65
## 29
       21
                1
                    LP
                                      2.6356179
                                                 5.3643821
## 30
       24
                    LP
                        6.5 14.7 65
                                      4.6766894
                                                  8.3233106
                1
## 31
       27
                        6.4 10.2 65
                1
                    LP
                                      5.1348457
                                                  7.6651543
## 32
       30
                    LP
                        6.6
                            8.0 65
                                      5.6077221
                                                  7.5922779
                1
## 33
       33
                    LP
                        6.5 11.0 65
                                      5.1356179
                                                 7.8643821
                1
##
      day oedema diet
                        wt
                              sd
                                          lci
                                                    uci
                                   n
## 12
        3
                2
                    HP 5.1 15.1 291 4.214822 5.985178
                2
                    HP 5.1 13.4 291 4.314478 5.885522
## 13
        6
        9
## 14
                2
                    HP 4.5 6.5 291 4.118963 4.881037
       12
                2
                    HP 5.2 9.9 291 4.619652 5.780348
## 15
## 16
       15
                2
                    HP 4.6 8.2 291 4.119307 5.080693
## 17
       18
                2
                   HP 5.1 13.4 291 4.314478 5.885522
                2
                    HP 5.0 13.4 291 4.214478 5.785522
## 18
       21
## 19
       24
                2
                    HP 5.2 15.1 291 4.314822 6.085178
                    HP 5.1 11.6 291 4.419996 5.780004
## 20
       27
                2
## 21
                2
                    HP 5.0 9.2 291 4.460686 5.539314
       30
## 22
                2
                    HP 5.2 8.2 291 4.719307 5.680693
       33
##
      day oedema diet wt
                                                   uci
                              sd n
                                         lci
                    LP 5.1 11.1 86 3.903056 6.296944
## 34
        3
                2
## 35
        6
                2
                    LP 6.0 8.2 86 5.115771 6.884229
                2
                    LP 5.1 8.2 86 4.215771 5.984229
## 36
        9
## 37
       12
                2
                   LP 6.5 11.9 86 5.216790 7.783210
```

```
## 38
      15
                  LP 6.4 9.1 86 5.418722 7.381278
               2
## 39
       18
                  LP 5.9 13.8 86 4.411908 7.388092
## 40
      21
               2
                 LP 6.1 16.5 86 4.320759 7.879241
                 LP 4.0 9.1 86 3.018722 4.981278
## 41
      24
                 LP 3.1 12.8 86 1.719741 4.480259
## 42
      27
              2
                 LP 4.0 7.3 86 3.212821 4.787179
## 43
      30
                  LP 5.0 9.1 86 4.018722 5.981278
## 44
      33
               2
```

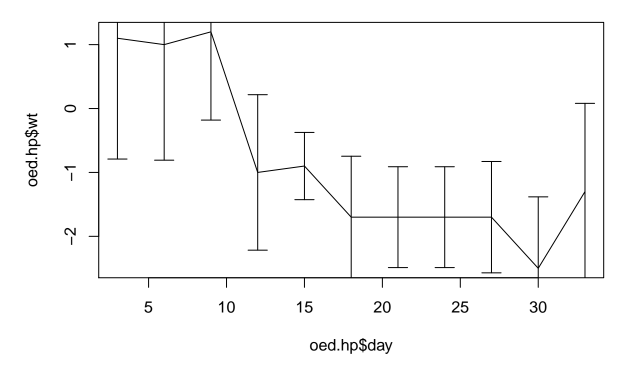
We can now plot the data for one arm of the trial:

```
plot(oed.hp$day, oed.hp$wt, type = "1")
```



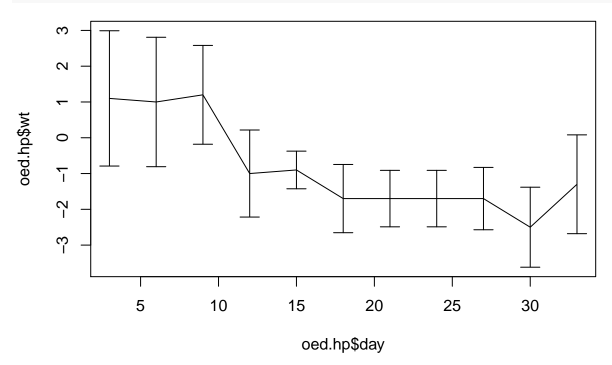
We can add error bars using the arrows() function:

```
arrows(oed.hp$day, oed.hp$lci, oed.hp$day, oed.hp$uci, code = 3, angle = 90, length = 0
```



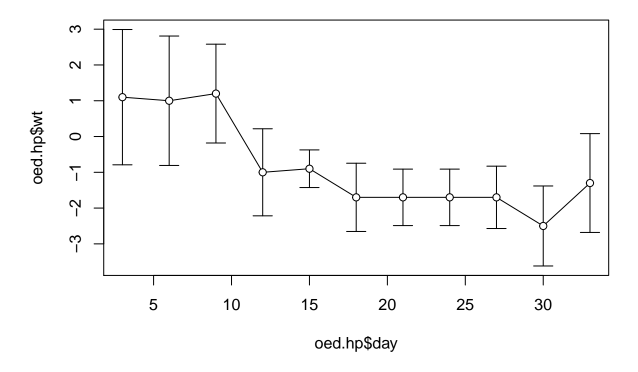
The scale of the y axis is wrong because the plot() function automatically scales axes to the ranges of the x and y data it is given. We can fix this by specifying a different set of limits (from lci and uci) for the y axis using the ylim parameter:

```
plot(oed.hp$day, oed.hp$wt, type = "l", ylim = c(min(oed.hp$lci), max(oed.hp$uci)))
arrows(oed.hp$day, oed.hp$lci, oed.hp$day, oed.hp$uci, code = 3, angle = 90, length = 0.
```



The plot might also be improved by adding plotting symbols:

```
points(oed.hp$day, oed.hp$wt, pch = 21, bg = "white")
```



The axis titles (oed.hp\$day and oed.hp\$wt) are the column (variable) names.

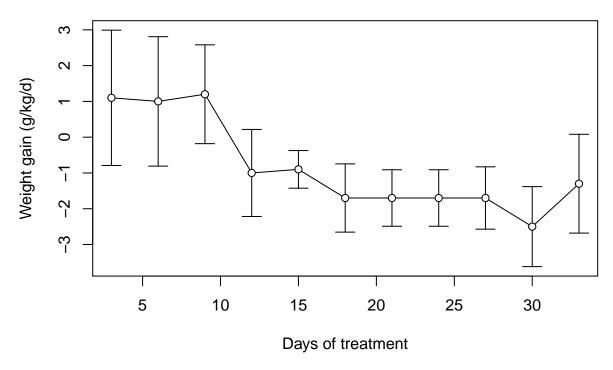
We can specify titles and axes labels in the call to the plot() function:

```
plot(oed.hp$day, oed.hp$wt, type = "l",
    ylim = c(min(oed.hp$lci), max(oed.hp$uci)),
    main = "HP Diet. Oedematous Cases",
    xlab = "Days of treatment", ylab = "Weight gain (g/kg/d)")

arrows(oed.hp$day, oed.hp$lci, oed.hp$day, oed.hp$uci,
    code = 3, angle = 90, length = 0.1)

points(oed.hp$day, oed.hp$wt, pch = 21, bg = "white")
```





Now that we know how to plot error bars, we can write a function that we will be able to use whenever we need to plot data with error bars.

Before continuing we will consider what the new function should be able to do. This will help us when it comes to writing the function. Our new function should:

- 1. Take four numeric vectors (x, y, lower CI for y, and upper CI for y) and plot them.
- 2. Be able to plot the data points as unconnected points or as points joined by lines.
- 3. Calculateappropriatelimitsfortheyaxis.
- 4. Produce a plot without axes so that more than one data series may be plotted on the same chart. 5. Provide sensible default values for axis limits and labels.

From this list we know that we need the function to take several parameters:

Name	Purpose	Default value
X	Data to plot	None
У	Data to plot	None
y.lci	Data to plot	None
y.uci	Data to plot	None
ylim	Limits for y axis	<pre>c(min(y.lci), max(y.uci))</pre>
xlab	Label for x axis	<pre>deparse(substitute(x))</pre>
ylab	Label for y axis	<pre>deparse(substitute(y))</pre>
main	Chart title	paste(ylab, "by", xlab)
type	Type of plot	"1"

Name	Purpose	Default value	
lty	Line type	1	
col	Line and point colour	"black"	
axes	Draw x and y axes	TRUE	
pch	Type of points to plot	1	
bg	Fill colour of point	white	

The parameter names have been chosen to be the same as the parameter names to plot() and points(). This makes the function easier to use. It also makes the function easier to write.

Create a new function called plot.ci():

```
plot.ci <- function() {}</pre>
```

This creates an empty function called plot.ci().

Use the fix() function to edit the plot.ci() function:

```
fix(plot.ci)
```

Edit the function to read:

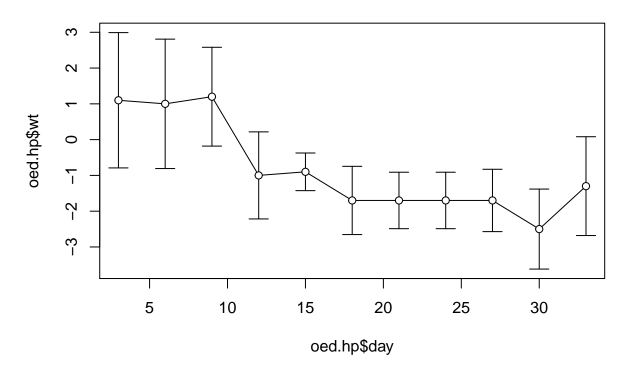
```
function(x,
         y, y.lci, y.uci,
         ylim = c(min(y.lci), max(y.uci)),
         xlab = deparse(substitute(x)),
         ylab = deparse(substitute(y)),
         main = paste(ylab, "by", xlab),
         type = "1",
         lty = 1,
         col = "black",
         axes = TRUE,
         pch = 21,
         bg = "white") {
  plot(x, y, type = type, ylim = ylim, xlab = xlab, ylab = ylab,
       main = main, lty = lty, col = col, axes = axes)
  arrows(x, y.lci, x, y.uci, code = 3, angle = 90, length = 0.1,
         lty = lty, col = col)
  points(x, y, pch = pch, bg = bg, col = col)
}
```

Once you have made the changes shown above, check your work, save the file, and quit the editor.

Let's try the plot.ci() function with the test data:

```
plot.ci(oed.hp$day, oed.hp$wt, oed.hp$lci, oed.hp$uci)
```

oed.hp\$wt by oed.hp\$day

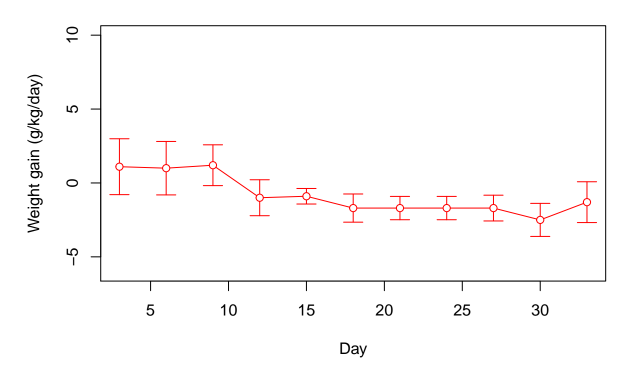


Note how the function has used default values for the axis labels, chart titles, chart limits etc.

We can override these default values if we need to:

```
plot.ci(oed.hp$day, oed.hp$wt, oed.hp$lci, oed.hp$uci,
    ylim = c(-6, 10), xlab = "Day",
    ylab = "Weight gain (g/kg/day)",
    main = "Oedematous", col = "red")
```

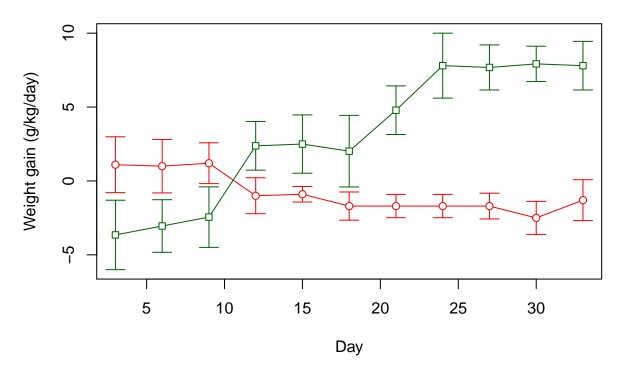
Oedematous



Do not close the plot window.

We should also check that we can plot another data series on this chart:

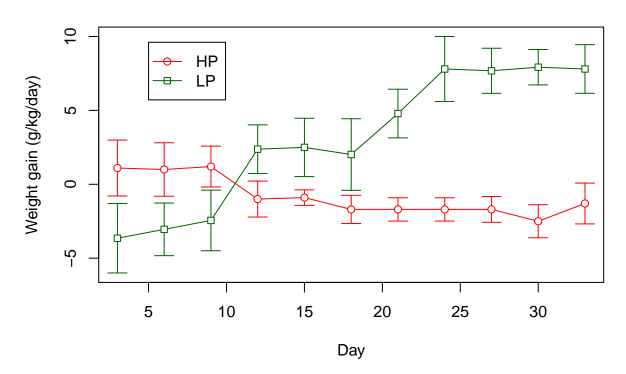
Oedematous



We can also add a legend:

```
legend(5, 8, legend = c("HP", "LP"), lty = c(1, 1),
    pch = c(21, 22), col = c("red", "darkgreen"))
```

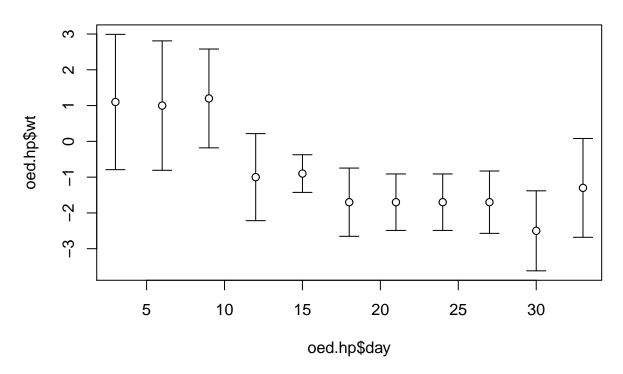
Oedematous



We should also check that we can produce plots of unconnected points:

```
plot.ci(oed.hp$day, oed.hp$wt, oed.hp$lci, oed.hp$uci, type = "p")
```

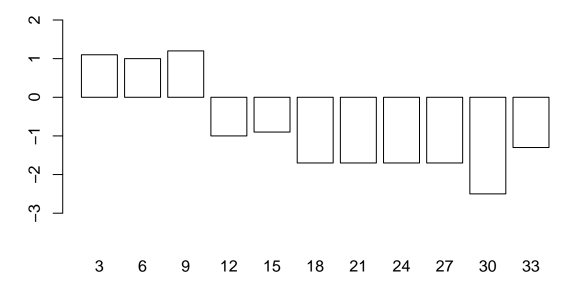
oed.hp\$wt by oed.hp\$day



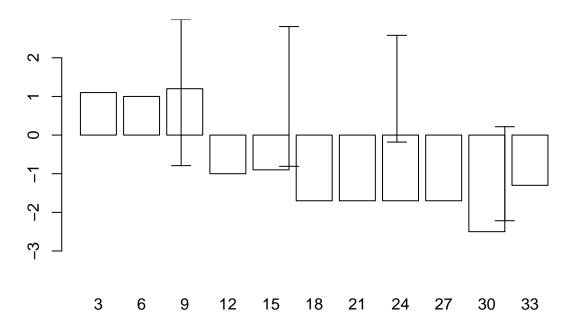
Try plotting the data for the marasmic patients using the plot.ci() function.

You might like to use the save() function to save the plot.ci() function.

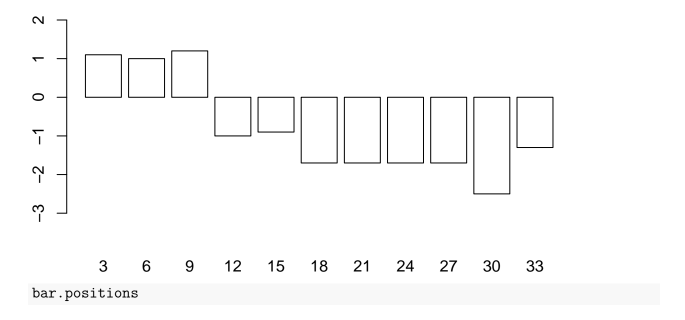
We can use a similar technique to add error bars to different types of plot. If we plot the weight velocities for oedematous patients receiving the high protein diet as a bar chart:



we could add error bars using the arrows() function as we did with a line plot:

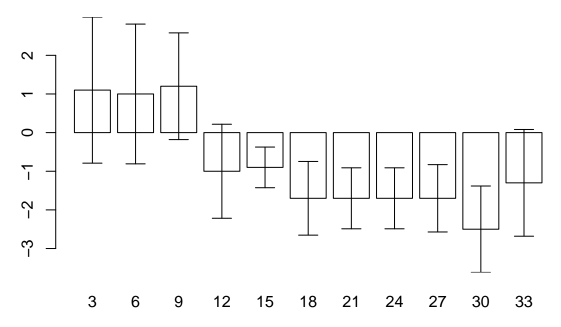


but this does not produce the expected results because the centres of the bars are not placed on the chart at the positions held in oed.hp\$day. This is easily fixed as barplot() returns a numeric vector (or matrix, when beside = TRUE) containing the co-ordinates of the bar midpoints:



```
[,1]
##
##
     [1,]
           0.7
##
     [2,]
           1.9
     [3,]
           3.1
##
##
     [4,]
           4.3
     [5,]
           5.5
##
     [6,]
           6.7
##
##
     [7,]
           7.9
##
     [8,]
           9.1
##
    [9,] 10.3
## [10,] 11.5
## [11,] 12.7
```

We can now use the information stored in bar.positions to specify the positions of the error bars:



Armed with this information, we can write a function that we will be able to use whenever we need to plot a bar chart with error bars.

Create a new function called barplot.ci():

```
barplot.ci <- function() {}</pre>
```

This creates an empty function called barplot.ci().

Use the fix() function to edit the barplot.ci() function:

```
fix(barplot.ci)
```

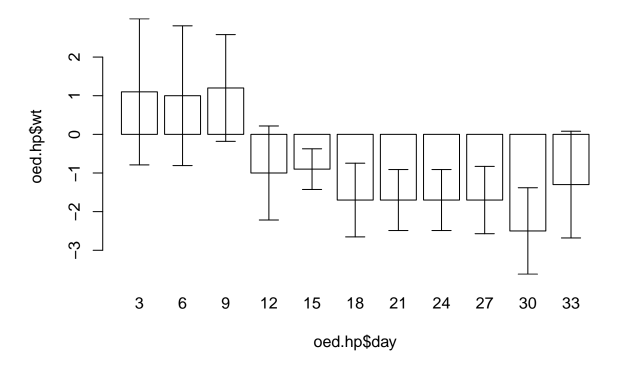
Edit the function to read:

Once you have made the changes shown above, check your work, save the file, and quit the editor.

Let's try the barplot.ci() function with the test data:

barplot.ci(oed.hp\$wt, oed.hp\$day, oed.hp\$lci, oed.hp\$uci)

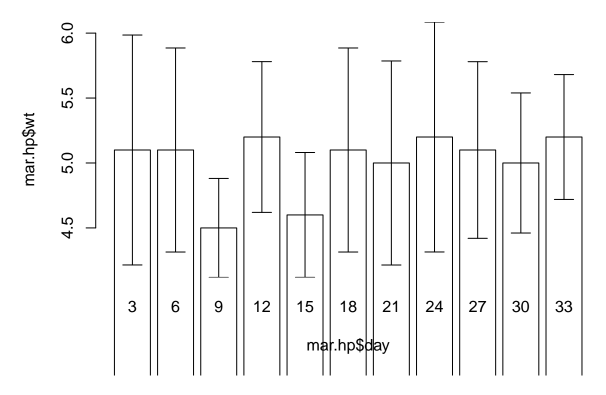
oed.hp\$wt by oed.hp\$day



Try plotting the weight velocities for the marasmic patients receiving the high protein diet using the barplot.ci() function:

barplot.ci(mar.hp\$wt, mar.hp\$day, mar.hp\$lci, mar.hp\$uci)

mar.hp\$wt by mar.hp\$day



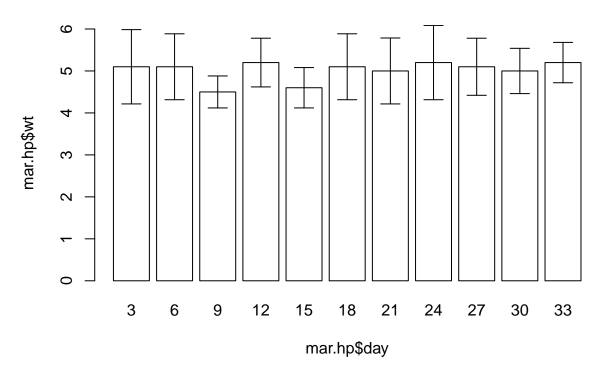
The chart looks wrong. This is because we have set the wrong limits for the y axis. The bars are drawn from zero to the data point but we have specified a limit for the y axis that is not constrained to include zero. This is easy to fix. Edit the barplot.ci() function to read:

Once you have made the changes shown above, check your work, save the file, and quit the editor.

Try plotting the data for the marasmic patients using the barplot.ci() function:

barplot.ci(mar.hp\$wt, mar.hp\$day, mar.hp\$lci, mar.hp\$uci)

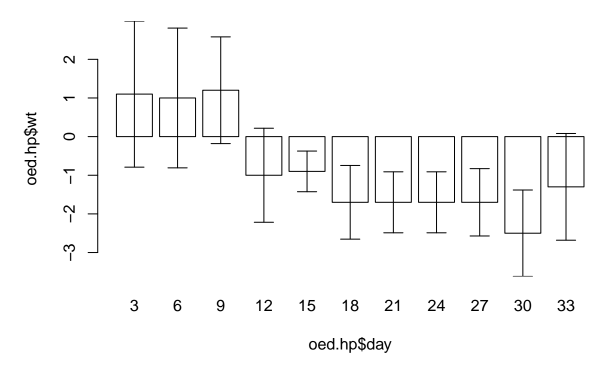
mar.hp\$wt by mar.hp\$day



Check that the function still operates as expected with the data for oedematous patients:

barplot.ci(oed.hp\$wt, oed.hp\$day, oed.hp\$lci, oed.hp\$uci)

oed.hp\$wt by oed.hp\$day



The end of one of the error bars touches the x axis. This can also be fixed by slightly widening the limits for the y axis. It might also be useful to plot the centre position of each error bar. We can use the points() function to do this. Edit the barplot.ci() function to read:

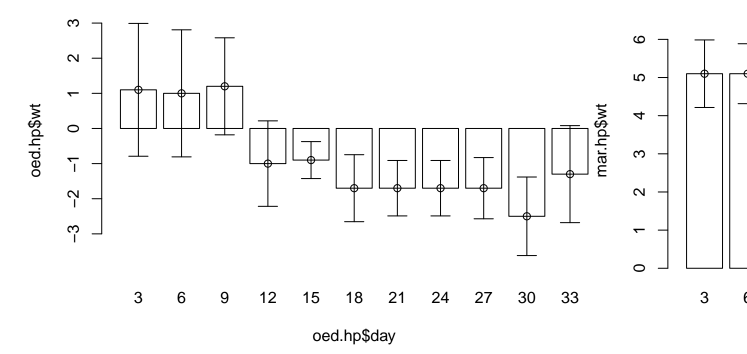
Once you have made the changes shown above, check your work, save the file, and quit the editor.

Check that the function works as expected:

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```
barplot.ci(oed.hp$wt, oed.hp$day, oed.hp$lci, oed.hp$uci)
barplot.ci(mar.hp$wt, mar.hp$day, mar.hp$lci, mar.hp$uci)
```

oed.hp\$wt by oed.hp\$day



The barplot.ci() function now works as expected with both sets of data. It is important when developing your own functions, to test them with different data so as to ensure that they work correctly with a wide range if data.

You might like to use the save() function to save the barplot.ci() function.

9.4 Mesh map

The fact that R provides flexible graphical functions means that, with little extra work, you can use these functions to present your data in appropriate and interesting ways rather than having to rely on a limited set of basic chart types.

In this exercise we will use the plot() function to produce a simple mesh-map.

The file cover.dat contains data from a coverage survey for a therapeutic feeding program (TFP) in central Malawi undertaken in March 2003. Data were collected using the *centric systematic area sampling* method to define sampling locations: A number of communities located closest to the centres of thirty 10 x 10 kilometre grid squares were sampled using active (investigative) case-finding.

The columns in this dataset are as follows:

x	x position of grid square
\mathbf{y}	y position of grid square
cases	Number of cases found in sampled communities in.program Number of cases
	(from above) enrolled in the TFP

Retrieve the dataset:

```
cs <- read.table("cover.dat", header = TRUE)</pre>
```

Examine the dataset:

CS

```
##
      Х
          y cases in.program
## 1
      1
                7
                             2
      2
         5
                             0
## 2
                4
## 3
      2 6
                4
                             1
## 4
      2
         7
                 3
                             1
## 5
      2 8
                 3
                             1
      2
         9
                5
                             1
## 6
## 7
      3
        3
                3
                             0
      3
                 2
## 8
         4
                             0
## 9
      3
          5
                 3
                             0
## 10 3
                             1
          6
                3
## 11 3
                5
                             2
## 12 3
                 2
                             0
## 13 3
                4
                             1
## 14 3 10
                5
                             0
## 15 4
                             2
                 5
## 16 4
          5
                8
                             1
## 17 4
          6
                 6
                             0
## 18 4
         7
                 6
                             1
## 19 4
                 3
                             1
## 20 4
          9
                 5
                             1
                             2
## 21 4 10
                6
## 22 5
                5
                             1
## 23 5
                 3
                             1
## 24 5
          6
                4
                             0
                             3
## 25 5
          7
                 6
## 26 5
         8
                4
                             0
                             2
## 27 6
          3
                8
                             2
## 28 6
          4
                5
## 29 6
          6
                             1
                 6
## 30 6
                 3
```

We should calculate the observed coverage for each grid square:

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```
cs$cvr <- cs$in.program / cs$cases
cs
```

```
##
         y cases in.program
                                     cvr
## 1
      1
         7
                7
                            2 0.2857143
## 2
      2
         5
                4
                            0.0000000
## 3
      2
         6
                4
                            1 0.2500000
      2
         7
## 4
                3
                            1 0.3333333
## 5
      2
         8
                3
                            1 0.3333333
      2
## 6
         9
                5
                            1 0.2000000
## 7
      3
         3
                            0 0.0000000
                3
      3
                2
## 8
         4
                            0 0.000000
## 9
      3
         5
                3
                            0 0.0000000
## 10 3
         6
                3
                            1 0.3333333
## 11 3
                5
                            2 0.4000000
## 12 3
         8
                2
                            0 0.0000000
## 13 3
                4
                            1 0.2500000
## 14 3 10
                5
                            0 0.000000
## 15 4
         4
                5
                            2 0.4000000
## 16 4
         5
                8
                            1 0.1250000
## 17 4
                6
                            0 0.0000000
## 18 4
         7
                            1 0.1666667
                6
## 19 4
         8
                3
                            1 0.3333333
## 20 4
         9
                5
                            1 0.2000000
## 21 4 10
                6
                            2 0.3333333
## 22 5
                5
                            1 0.2000000
## 23 5
         5
                3
                            1 0.3333333
## 24 5
                            0 0.000000
         6
                4
## 25 5
         7
                            3 0.5000000
                6
## 26 5
         8
                4
                            0 0.0000000
## 27 6
         3
                            2 0.2500000
                8
## 28 6
         4
                5
                            2 0.4000000
## 29 6
         6
                6
                            1 0.1666667
## 30 6
                3
                            1 0.3333333
```

Note that some grid squares have zero coverage. It might be useful to use specific plotting characters (e.g. open and filled squares) to indicate zero and non-zero coverage. We can use the ifelse() function to do this:

```
cs$cvr.pch <- ifelse(cs$cvr == 0, 0, 15)
cs
```

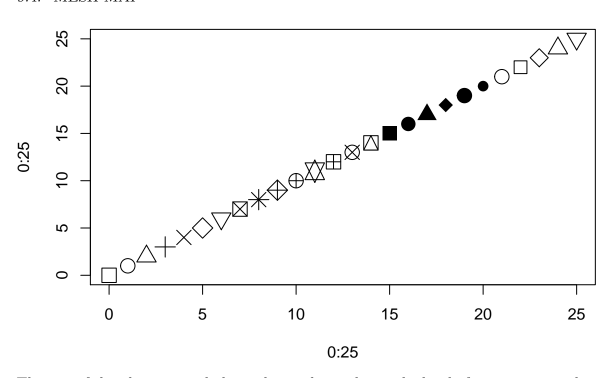
```
##
         y cases in.program
                                    cvr cvr.pch
      Х
## 1
                7
                            2 0.2857143
                                              15
      1
      2
         5
                                               0
## 2
                4
                            0.0000000
## 3
      2
         6
                4
                            1 0.2500000
                                              15
```

##	4	2	7	3	1	0.3333333	15
##	5	2	8	3	1	0.3333333	15
##	6	2	9	5	1	0.2000000	15
##	7	3	3	3	0	0.0000000	0
##	8	3	4	2	0	0.0000000	0
##	9	3	5	3	0	0.0000000	0
##	10	3	6	3	1	0.3333333	15
##	11	3	7	5	2	0.4000000	15
##	12	3	8	2	0	0.0000000	0
##	13	3	9	4	1	0.2500000	15
##	14	3	10	5	0	0.0000000	0
##	15	4	4	5	2	0.4000000	15
##	16	4	5	8	1	0.1250000	15
##	17	4	6	6	0	0.0000000	0
##	18	4	7	6	1	0.1666667	15
##	19	4	8	3	1	0.3333333	15
##	20	4	9	5	1	0.2000000	15
##	21	4	10	6	2	0.3333333	15
##	22	5	4	5	1	0.2000000	15
##	23	5	5	3	1	0.3333333	15
##	24	5	6	4	0	0.0000000	0
##	25	5	7	6	3	0.5000000	15
##	26	5	8	4	0	0.0000000	0
##	27	6	3	8	2	0.2500000	15
##	28	6	4	5	2	0.4000000	15
##	29	6	6	6	1	0.1666667	15
##	30	6	7	3	1	0.3333333	15

A quick way of seeing the code associated with each plotting symbol is:

```
plot(0:25, 0:25, pch = 0:25, cex = 2)
```

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The size of the plotting symbol may be used to indicate the level of coverage in each quadrat but we must ensure that the symbol used for zero-coverage is not invisibly small:

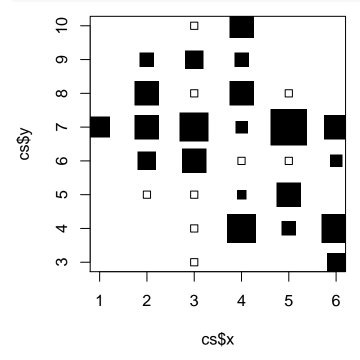
```
cs$cvr.cex <- ifelse(cs$cvr == 0, 1, 10 * cs$cvr)
cs
```

##		х	У	cases	in.program	cvr	cvr.pch	cvr.cex
##	1	1	7	7	2	0.2857143	15	2.857143
##	2	2	5	4	0	0.0000000	0	1.000000
##	3	2	6	4	1	0.2500000	15	2.500000
##	4	2	7	3	1	0.3333333	15	3.333333
##	5	2	8	3	1	0.3333333	15	3.333333
##	6	2	9	5	1	0.2000000	15	2.000000
##	7	3	3	3	0	0.0000000	0	1.000000
##	8	3	4	2	0	0.0000000	0	1.000000
##	9	3	5	3	0	0.000000	0	1.000000
##	10	3	6	3	1	0.3333333	15	3.333333
##	11	3	7	5	2	0.400000	15	4.000000
##	12	3	8	2	0	0.0000000	0	1.000000
##	13	3	9	4	1	0.2500000	15	2.500000
##	14	3	10	5	0	0.0000000	0	1.000000
##	15	4	4	5	2	0.400000	15	4.000000
##	16	4	5	8	1	0.1250000	15	1.250000
##	17	4	6	6	0	0.0000000	0	1.000000
##	18	4	7	6	1	0.1666667	15	1.666667
##	19	4	8	3	1	0.3333333	15	3.333333
##	20	4	9	5	1	0.2000000	15	2.000000
##	21	4	10	6	2	0.3333333	15	3.333333

```
1 0.2000000
## 22 5
         4
                5
                                              15 2.000000
## 23 5
                            1 0.3333333
                                               15 3.333333
         5
                3
## 24 5
         6
                4
                            0 0.0000000
                                               0 1.000000
         7
## 25 5
                6
                            3 0.5000000
                                              15 5.000000
## 26 5
                            0 0.0000000
                                               0 1.000000
         8
                4
## 27 6
         3
                            2 0.2500000
                                               15 2.500000
                8
## 28 6
         4
                5
                            2 0.4000000
                                               15 4.000000
## 29 6
                                               15 1.666667
         6
                6
                            1 0.1666667
## 30 6
                3
                            1 0.3333333
                                               15 3.333333
```

We can now plot the data:

```
par(pty="s")
plot(cs$x, cs$y, cex = cs$cvr.cex, pch = cs$cvr.pch)
```



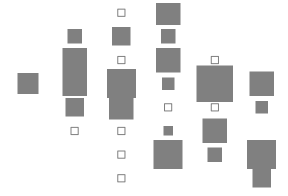
There are some problems with this plot:

- The axes and labels distract from the data.
- The distance between grid-square centres is wider in the x than in the y direction.
- The colour (black) of the plotting symbols is too strong.

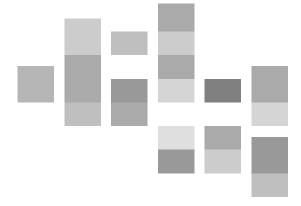
All of these problems can be fixed by specifying values for plot() function parameters:

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```
axes = FALSE,
xlim = c(0,10),
ylim = c(0,10),
col = gray(0.5))
```



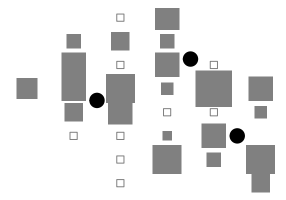
An alternative way of plotting this data is to use shades of grey (rather than the size of the plotting symbol) to represent the level of coverage in each grid-square:



In this context it is useful to show the approximate location of the approximate locat

```
axes = FALSE,
xlim = c(0,10),
ylim = c(0,10),
col = gray(0.5))

points(c(2.5, 4.5, 5.5), c(6.5, 8.25, 5), pch = 19, cex = 2)
```



The techniques introduced in this section allow you to write custom graphical functions but they can also be used to change the default behaviour of standard graphical functions.

9.5 Combining plots

In $Exercise\ 1$ Getting acquainted with R), we saw how the plot() function could be applied to a fitted object:

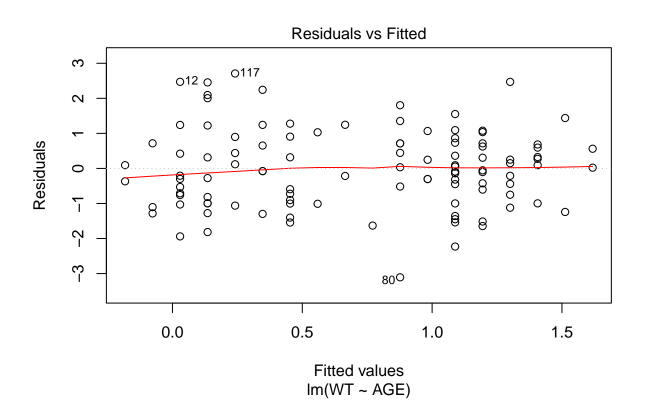
```
fem <- read.table("fem.dat", header = TRUE)</pre>
attach(fem)
## The following objects are masked from fem (pos = 4):
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
##
## The following objects are masked from fem (pos = 5):
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
##
## The following objects are masked from fem (pos = 9):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
## The following objects are masked from fem (pos = 10):
##
##
       AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT
## The following objects are masked from fem (pos = 12):
```

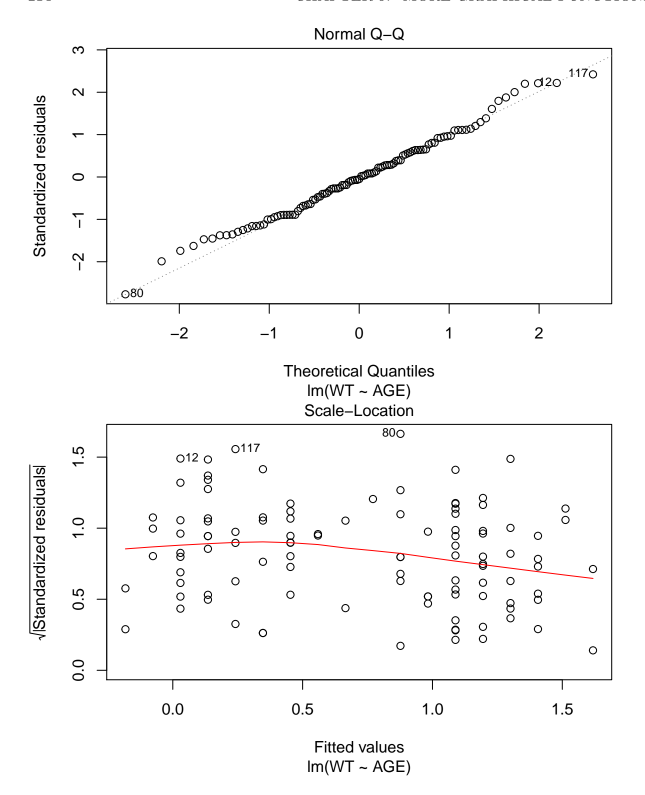
```
##
## AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT

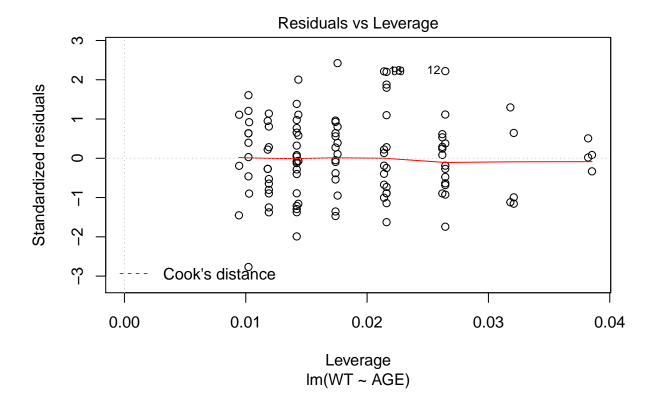
## The following objects are masked from fem (pos = 13):
##
## AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT

## The following objects are masked from fem (pos = 18):
##
## AGE, ANX, DEP, ID, IQ, LIFE, SEX, SLP, WT

fem.lm <- lm(WT ~ AGE)
plot(fem.lm)</pre>
```

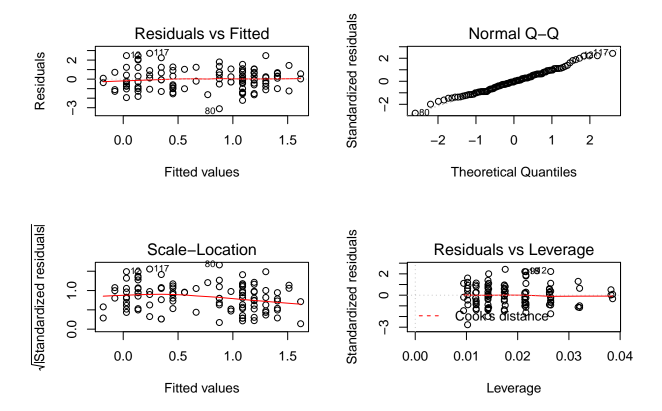






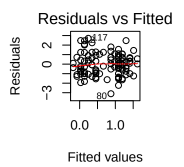
Each of the diagnostic plots are presented as a separate chart. We could use the mfrow parameter of the par() function to present all four diagnostic plots on a single chart:

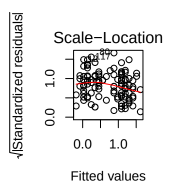
```
par(mfrow = c(2, 2))
plot(fem.lm)
```

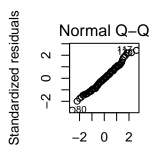


It might improve the appearance of the chart if each of the diagnostic plots were square rather than rectangular:

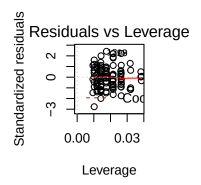
```
par(mfrow = c(2, 2))
par(pty = "s")
plot(fem.lm)
```





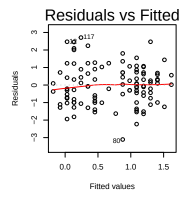


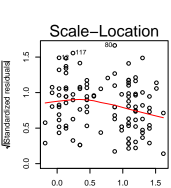
Theoretical Quantiles



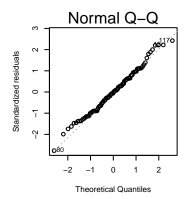
It might improve the appearance of the chart if smaller text and symbols were used:

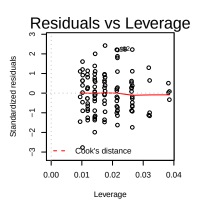
```
par(mfrow = c(2, 2))
par(pty = "s")
par(cex = 0.5)
plot(fem.lm)
```





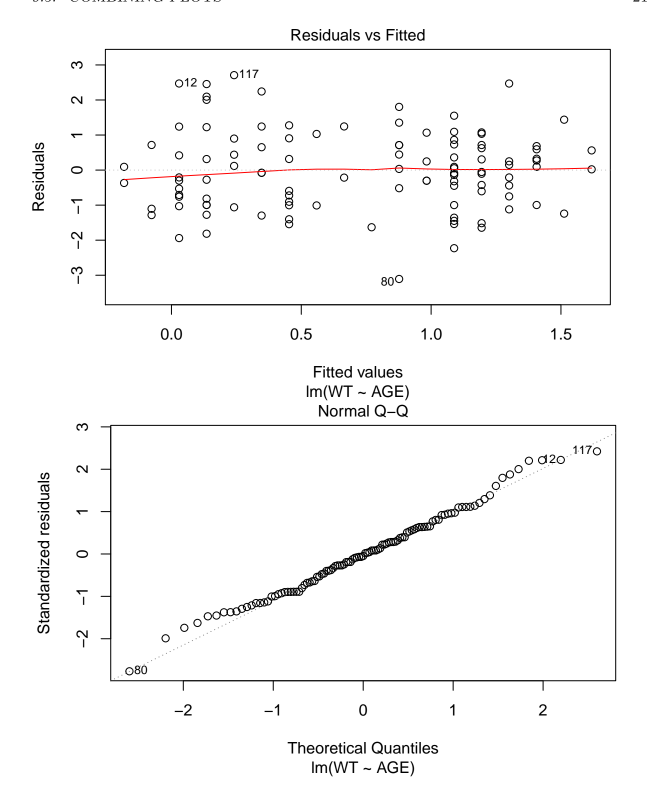
Fitted values

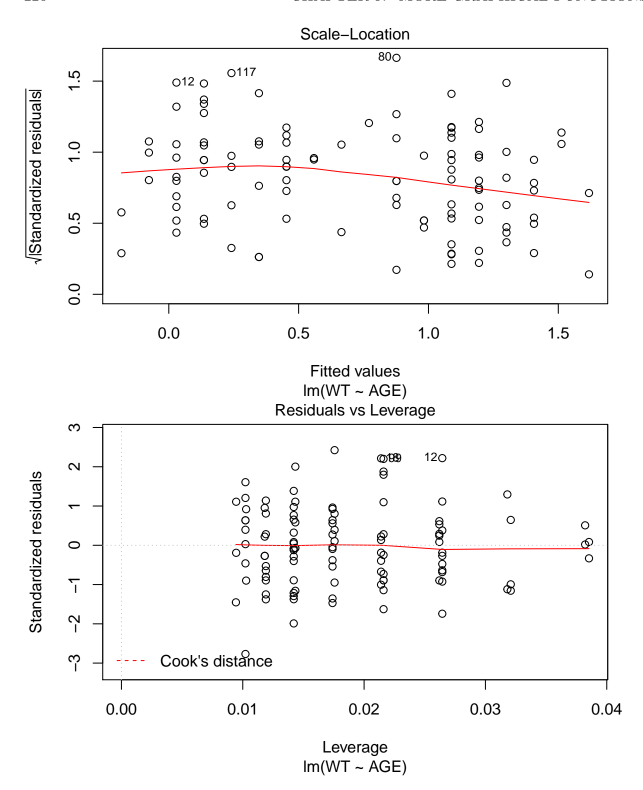




Graphical parameters set using the $\mathtt{par}()$ function affect all subsequent plot commands and must be reset explicitly:

```
par(mfrow = c(1, 1), pty = "m", cex = 1)
plot(fem.lm)
```

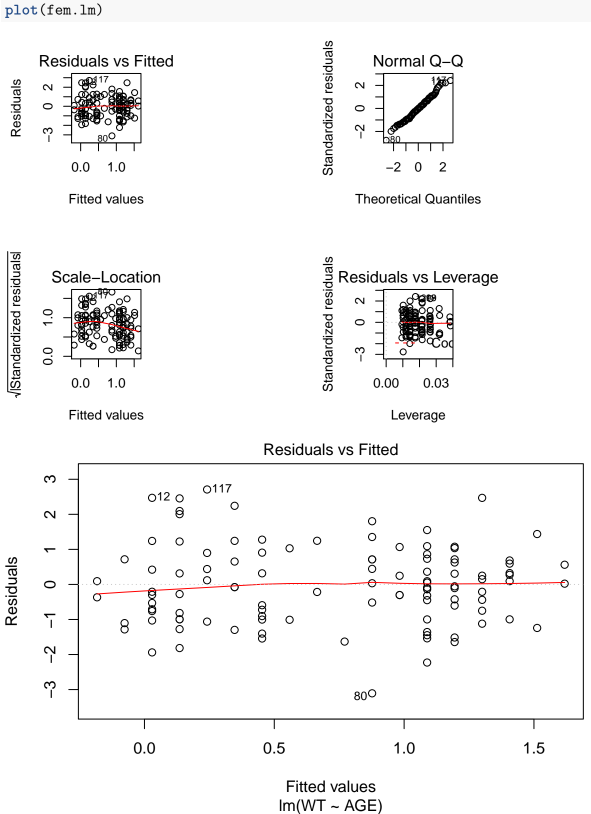


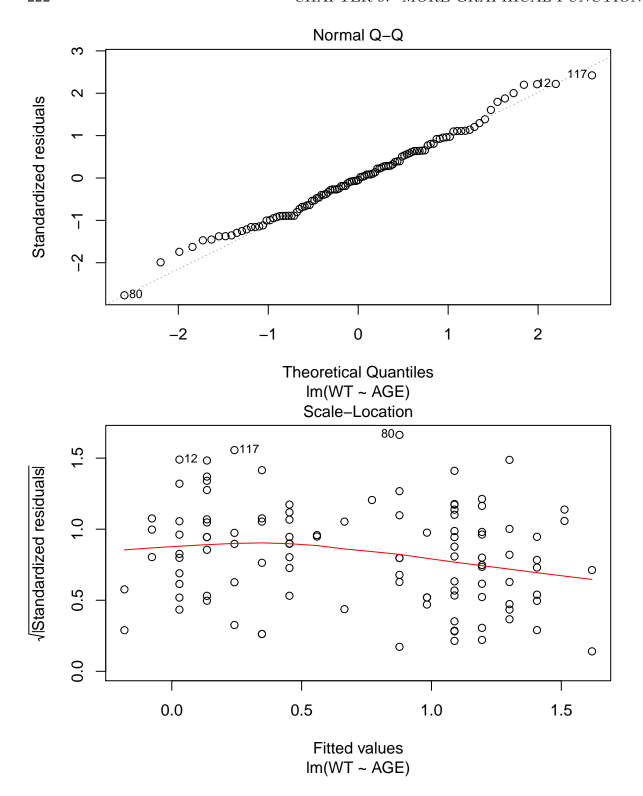


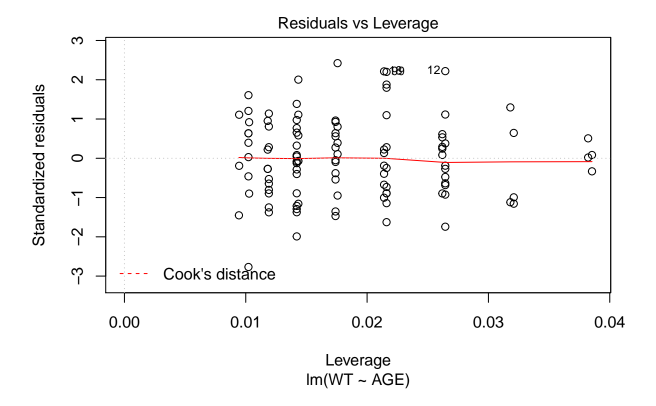
It is possible to save graphical parameters into an R object and use this object to restore original graphical parameters:

```
old.par <- par()
par(mfrow = c(2, 2), pty = "s")
```

plot(fem.lm)
par(old.par)
plot(fem.lm)

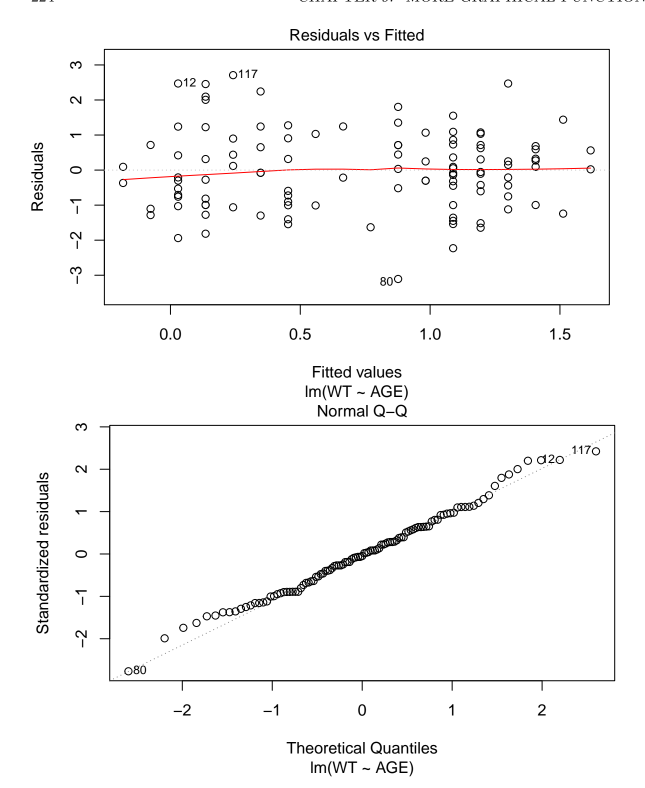


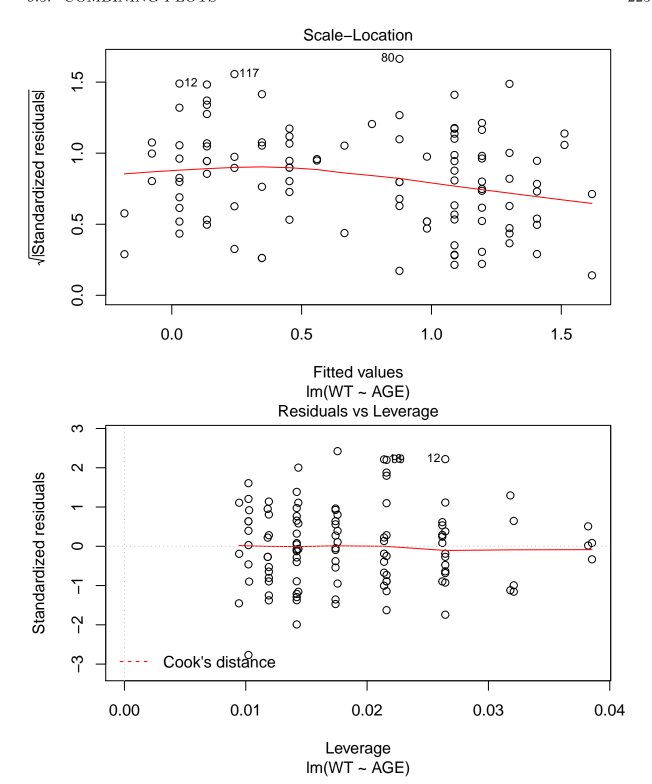


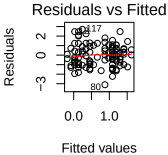


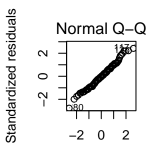
The ability to save and apply graphical parameters means that you can create a library of graphical parameter sets that can be applied with the par() function as required:

```
default.par <- par()
par(mfrow = c(2, 2), pty = "s")
plot.lm.par <- par()
par(default.par)
plot(fem.lm)
par(plot.lm.par)
plot(fem.lm)
par(default.par)
plot(fem.lm)</pre>
```

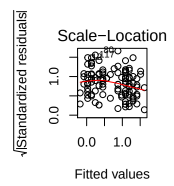


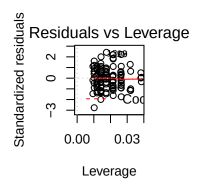


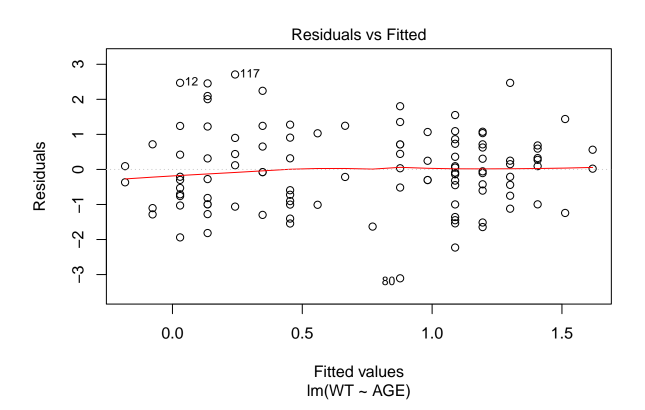


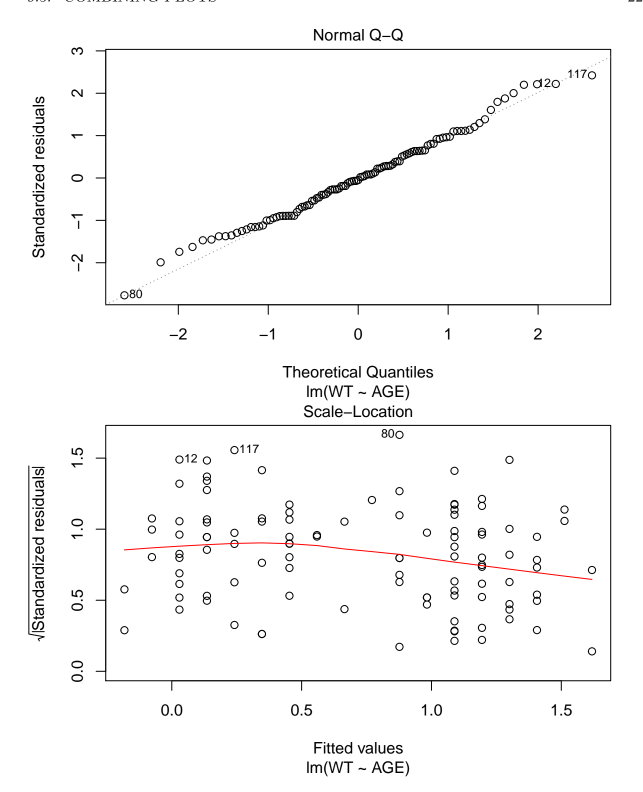


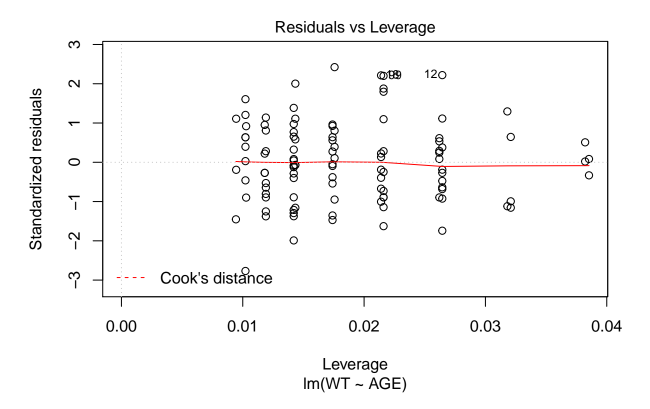








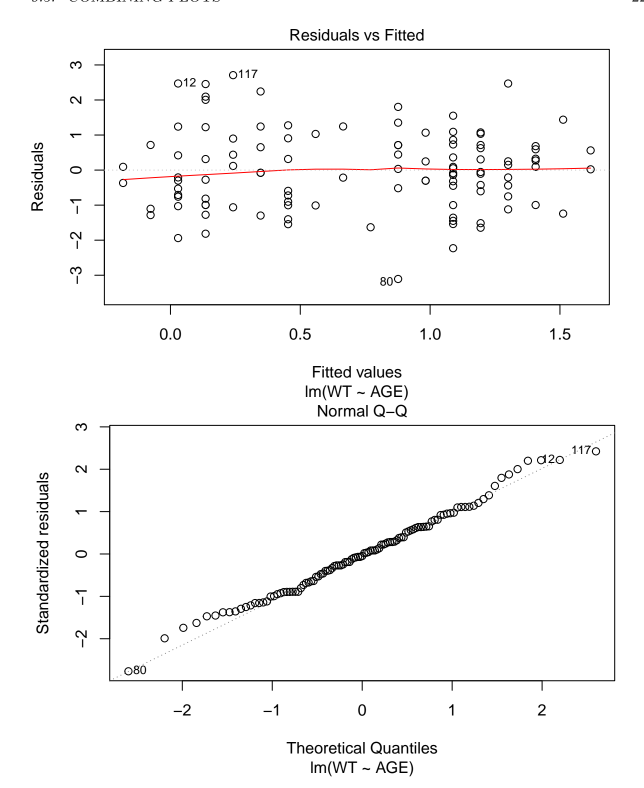


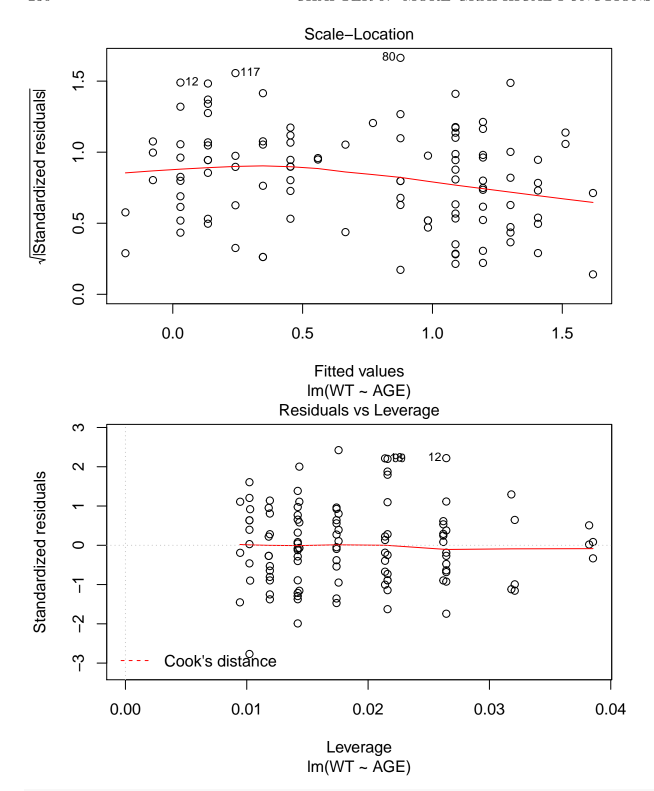


R produces warning messages when you save and restore graphical parameters in this way. This is because some graphical parameters are read only and cannot be changed using the par() function. This has no effect other than to cause R to issue warning messages.

If you do not like the warning messages then you can use the par() function with the no.readonly parameter set to TRUE:

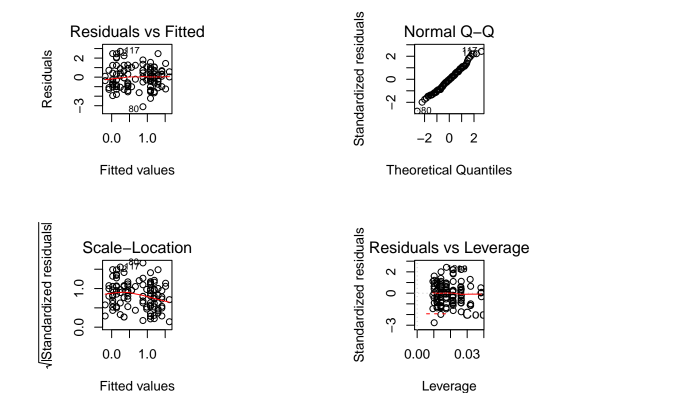
```
default.par <- par(no.readonly = TRUE)
par(mfrow = c(2, 2), pty = "s")
plot.lm.par <- par(no.readonly = TRUE)
par(default.par)
plot(fem.lm)</pre>
```

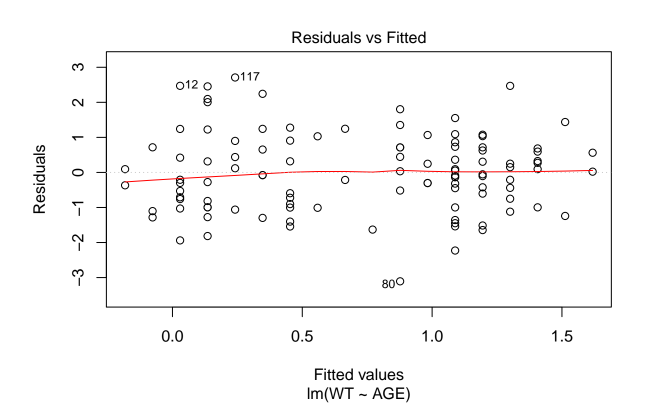


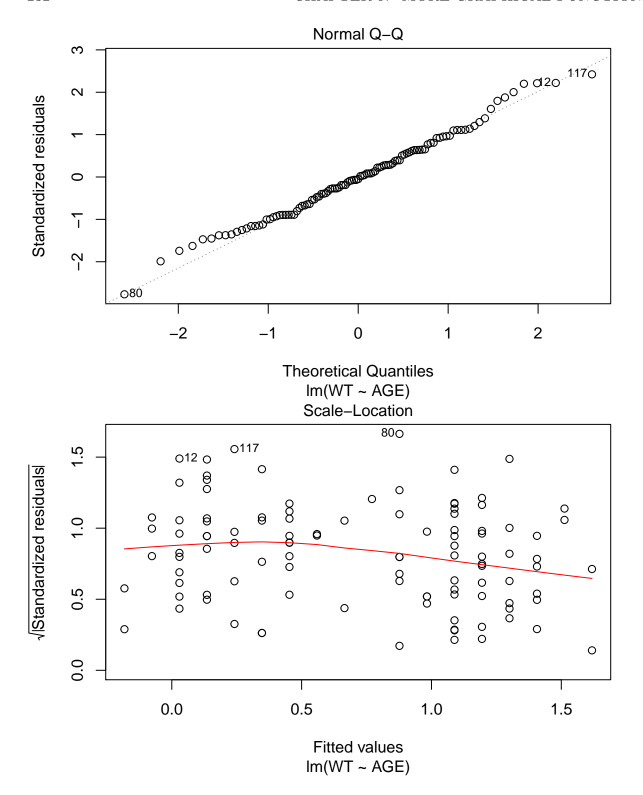


par(plot.lm.par)
plot(fem.lm)

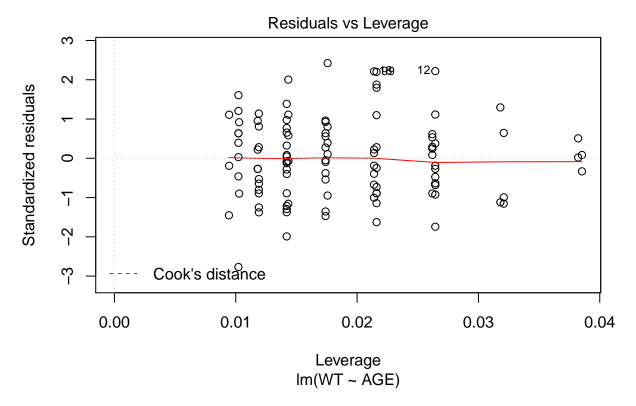
par(default.par)
plot(fem.lm)







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Graphical parameter sets, like any other R object, may be saved and loaded using the save() and load() functions.

9.6 Summary

- R allows you to create functions that produce graphical output.
- R graphical functions are flexible so that you can create functions that can produce chart types that are not available in R or many other statistical applications. Standard plots may also be customised using the par() function.
- R allows you to specify default values for function parameters making functions calls easier by removing the requirement to specify values for every function parameter.

Chapter 10

Computer intensive methods

Chapter 11

What now?