

Statistical Practice in Epidemiology with R, 2006
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Installing R

1.1 Microsoft windows

The current version of R (1 June 2006) is 2.3.1. To install R plus the data and package used in this introduction

- 1. Go to http://cran.r-project.org/mirrors.html and click choose a mirror near you, click on the link to Windows and after that choose base. Download R-2.3.1-win32.exe to your computer
- 2. Run this installation file.
- 3. Fire up R, and at the command prompt type:

```
install.packages("Epi")
```

This will install the Epi package provided you are connected to the net.

4. Go to www.sciviews.org/Tinn-R, download the latest stable setup, and run it. This will install the Tinn-R text editor..

1.2 Gnu/Linux

- 1. Go to www.r-project.org and click on CRAN.
- 2. Choose an rpm or tar file from a mirror, appropriate to your version of GNU/Linux, and use it to install R.
- 3. Fire up R, and at the command prompt type:

```
install.packages("Epi")
```

This will install the Epi package provided you are connected to the net.

1.3 Questions about epidmiology with R

A mailing list has been set up for people who use R for epidemiological analysis. There is a link to it on the Epi homepage, www.pubhealth.ku.dk/~bxc/Epi, or you can join by going to the R-homepage www.r-project.org and then click on "Mailing lists" and find the R-sig-Epi mailing list.

You are welcome to pose questions regarding R and epidemiology there, and we will try to help you as best we can.

Some basic commands in R

2.1 Preliminaries

The purpose of these notes is to describe a small subset of the R language, sufficient to allow someone new to R to get started. The exercises are important because they reinforce basic aspects of R. For further details about R we refer the reader to **An Introduction to R** by W.N.Venables, D.M.Smith, and the R development team. This can be downloaded from the R website at http://www.r-project.org.

To start R click on the R icon. To change your working directory click on the tab with this name and select the directory you want to work in. To get out of R click on the File menu and select Exit. You will be offered the chance to save the work space, but at this stage just exit without saving, then start R again, and change the working directory, as before.

R is case sensitive, so that A is different from a. Commands in R are generally separated by a newline, although a semi-colon can also be used. When using R it makes sense to avoid as much typing as possible by recalling previous commands using the vertical arrow key and editing them.

2.2 Using R as a calculator

Typing 2+2 will return the answer 4, typing 2^3 will return the answer 8 (2 to the power of 3), typing log(10) will return the natural logarithm of 10, which is 2.3026, and typing sqrt(25) will return the square root of 25.

Instead of printing the result you can store it in an object, say

> a <- 2 + 2

which can be used in further calculations. The expression <-, pronounced "gets", is called the assignment operator, and is obtained by typing < and then -. The assignment operator can also be used in the opposite direction, as in

2+2 -> a

The contents of **a** can be printed by typing **a**.

Standard probability functions are readily available. For example, the probability below 1.96 in a standard normal (i.e. Gaussian) distribution is obtained with

> pnorm(1.96)

while

> pchisq(3.84, 1)

will return the probability below 3.84 in a χ^2 distribution on 1 degree of freedom, and

> pchisq(3.84, 1, lower.tail = FALSE)

will return the probability above 3.84.

Exercise 1.

- 1. Calculate $\sqrt{3^2 + 4^2}$.
- 2. Find the probability above 4.3 in a chi-squared distribution on 1 degree of freedom.

2.3 Objects and functions

All commands in R are *functions* which act on *objects*. One important kind of object is a *vector*, which is an ordered collections of numbers, or an ordered collection of character strings. Examples of vectors are 4, 6, 1, 2.2, which is a numeric vector with 4 components, and "Charles Darwin", "Alfred Wallace" which is a vector of character strings with 2 components. The components of a vector must be of the same type (numeric or character). The combine function c(), together with the assignment operator, is used to create vectors. Thus

> v <- c(4, 6, 1, 2.2)

creates a vector \mathbf{v} with components 4, 6, 1, 2.2 by first combining the 4 numbers 4, 6, 1, 2.2 in order and then assigning the result to the vector \mathbf{v} . Collections of components of different types are called *lists*, and are created with the list() function. Thus

> m <- list(4, 6, "name of company")</pre>

creates a list with 3 components. The main differences between the numbers 4, 6, 1, 2.2 and the vector v is that along with v is stored information about what sort of object it is and hence how it is printed and how it is combined with other objects. Try

> v > 3 + v > 3 * v

and you will see that R understands what to do in each case. This may seem trivial, but remember that unlike most statistical packages there are many different kinds of object in R.

You can get a description of the structure of any object using the function str(). For example, str(v) shows that v is numeric with 4 components.

2.4 Sequences

It is not always necessary to type out all the components of a vector. For example, the vector $(15, 20, 25, \dots, 85)$ can be created with

> seq(15, 85, by = 5)

and the vector $(5, 20, 25, \dots, 85)$ can be created with

> c(5, seq(20, 85, by = 5))

You can learn more about functions by typing ? followed by the function name. For example **?seq** gives information about the syntax and usage of the function **seq()**.

Exercise 2.

- 1. Create a vector ${\tt w}$ with components 1, -1, 2, -2
- 2. Print this vector (to the screen)
- 3. Obtain a description of w using str()
- 4. Create the vector w+1, and print it.
- 5. Create the vector (0, 1, 5, 10, 15, ..., 75) using c() and seq().

2.5 The births data

The most important example of a vector in epidemiology is the data on a variable recorded for a group of subjects. To introduce R we use the births data which concern 500 mothers who had singleton births in a large London hospital. These data are available as an R object called **births** in the Epi package.

Some of the variables which make up these data take integer values while others are numeric taking measurements as values. For most variables the integer values are just codes for different categories, such as "male" and "female" which are coded 1 and 2 for the variable sex. You can browse the "births.txt" file by clicking on *Display files* under the *File menu*. It is a tab delimited file, that is the individual items of data are separated by the code for Tab. This is the sort of file you get from a spreadsheet such as Excel. Missing values are left blank, so they appear as TabTab.

2.6 Reading the data

The easiest way to access the births data is first to load the Epi package with

```
> library(Epi)
```

and then to load the data with

```
> data(births)
```

Try

> objects()

to make sure that you have an object called births in your working directory. The function

> str(births)

shows that the object **births** is a data frame with 500 observations of 8 variables. The names and types of the variables are also shown together with the first 10 values of each variable.

Exercise 3.

1. The dataframe "diet" in the Epi package contains data from a follow-up study with coronary heart disease as the end-point. Load these data with

> data(diet)

and print the contents of the data frame to the screen..

- 2. Check that you now have two objects, births, and diet in your work space.
- 3. Obtain a description of the object diet.
- 4. Remove the object diet with the command
 > rm(diet)

Check that you only have the object births left.

| Variable | Units or Coding | Type | Name |
|----------------------------|--------------------------|-------------|---------|
| Subject number | - | categorical | id |
| Birth weight | grams | metric | bweight |
| Birth weight < 2500 g | 1=yes, 0=no | categorical | lowbw |
| Gestational age | weeks | metric | gestwks |
| Gestational age < 37 weeks | 1=yes, 0=no | categorical | preterm |
| Maternal age | years | metric | matage |
| Maternal hypertension | 1=hypertensive, 0=normal | categorical | hyp |
| Sex of baby | 1=male, 2=female | categorical | sex |

Table 2.1: Variables in the births dataset

2.7 Referencing parts of the data frame

Typing births will list the entire data frame - not usually very helpful. Now try

```
> births[1, "bweight"]
```

This will list the value taken by the first subject for the bweight variable. Similarly

```
> births[2, "bweight"]
```

will list the value taken by the second subject for bweight, and so on. To list the data for the first 10 subject for the bweight variable, try

```
> births[1:10, "bweight"]
```

and to list all the data for this variable, try

```
> births[, "bweight"]
```

Exercise 4.

- 1. Print the data on the variable gestwks for subject 7 in the births data frame.
- 2. Print all the data for subject 7.
- 3. Print all the data on the variable gestwks.

2.8 Summaries

A good way to start an analysis is to ask for a summary of the data by typing

```
> summary(births)
```

To see the names of the variables in the data frame try

```
> names(births)
```

Variables in a data frame can be referred to by name, but to do so it is necessary also to specify the name of the data frame. Thus births\$hyp refers to the variable hyp in the births data frame, and typing births\$hyp will print the data on this variable. To summarize the variable hyp try

```
> summary(births$hyp)
```

In most datasets there will be some missing values. These are usually coded using tab delimited blanks to mark the values which are missing. R then codes the missing values using the NA (not available) symbol. The summary shows the number of missing values for each variable.

2.9 Turning a variable into a factor

In R categorical variables are known as *factors*, and the different categories are called the levels of the factor. Variables such as hyp and sex are originally coded using integer codes, and by default R will interpret these codes as numeric values taken by the variables. For R to recognize that the codes refer to categories it is necessary to convert the variables to be factors, and to label the levels. To convert the variable hyp to be a factor, try

```
> hyp <- factor(births$hyp)
> str(births)
```

```
> objects()
```

which shows that hyp is both in your work space (as a factor), and in in the births data frame (as a numeric variable). It is better to use the transform function on the data frame, as in

```
> births <- transform(births, hyp = factor(hyp))
> str(births)
```

which shows that hyp, in the births data frame, is now a factor with two levels, labelled "0" and "1" which are the original values taken by the variable. It is possible to change the labels to (say) "normal" and "hyper" with

Exercise 5.

- 1. Convert the variable sex into a factor
- 2. Label the levels of sex as "male" and "female".

2.10 Frequency tables

When starting to look at any new data frame the first step is to check that the values of the variables make sense and correspond to the codes defined in the coding schedule. For categorical variables (factors) this can be done by looking at one-way frequency tables and checking that only the specified codes (levels) occur. The most useful function for making tables is **stat.table**. This is currently part of the Epi package, so you will need to load this package first with

> library(Epi)

The distribution of the factors hyp and sex can be viewed by typing

```
> stat.table(hyp, data = births)
> stat.table(sex, data = births)
```

Their cross-tabulation is obtained by typing

```
> stat.table(list(hyp, sex), data = births)
```

Cross-tabulations are useful when checking for consistency, but because no distinction is drawn between the response variable and any explanatory variables, they are not useful as a way of presenting data.

2.11 Grouping the values of a metric variable

For a numeric variable like matage it is often useful to group the values and to create a new factor which codes the groups. For example we might cut the values taken by matage into the groups 20–29, 30–34, 35–39, 40–44, and then create a factor called agegrp with 4 levels corresponding to the four groups. The best way of doing this is with the function cut. Try

```
> births <- transform(births, agegrp = cut(matage, breaks = c(20,
+ 30, 35, 40, 45), right = FALSE))
> stat.table(agegrp, data = births)
```

By default the factor levels are labelled [20-25), [25-30), etc., where [20-25) refers to the interval which includes the left hand end (20) but not the right hand end (25). This is the reason for right=FALSE. When right=TRUE (which is the default) the intervals include the right hand end but not the left hand.

It is important to realize that observations which are not inside the range specified in the **breaks()** part of the command result in missing values for the new factor. For example, try

```
> births <- transform(births, agegrp = cut(matage, breaks = c(20,
+ 30, 35), right = FALSE))
> summary(births)
```

Only observations from 20 up to, but not including 35, are included. For the rest, **agegrp** is coded missing. You can specify that you want to cut a variable into a given number of intervals of equal length by specifying the number of intervals. For example

```
> births <- transform(births, agegrp = cut(matage, breaks = 5,
+ right = FALSE))
> stat.table(agegrp, data = births)
```

shows 5 intervals of width 4.

Exercise 6.

- 1. Summarize the numeric variable gestwks, which records the length of gestation for the baby, and make a note of the range of values.
- 2. Create a new factor gest4 which cuts gestwks at 20, 35, 37, 39, and 45 weeks, including the left hand end, but not the right hand. Make a table of the frequencies for the four levels of gest4.
- 3. Create a new factor gest5 which cuts gestwks into 5 equal intervals, and make a table of frequencies.

2.12 Tables of means and other things

To obtain the mean of bweight by sex, try

```
> stat.table(sex, mean(bweight), data = births)
```

The headings of the table can be improved with

> stat.table(sex, list("Mean birth weight" = mean(bweight)), data = births)

To make a two-way table of mean birth weight by sex and hypertension, try

```
> stat.table(list(sex, hyp), mean(bweight), data = births)
```

and to tabulate the count as well as the mean, try

> stat.table(list(sex, hyp), list(count(), mean(bweight)), data = births)

Available functions for the cells of the table are count, mean, weighted.mean, sum, min, max, quantile,median, IQR, and ratio. The last of these is useful for rates and odds. For example, to make a table of the odds of low birth weight by hypertension, try

```
> stat.table(hyp, list(odds = ratio(lowbw, 1 - lowbw, 100)), data = births)
```

The scale factor 100 makes the odds per 100. Margins can be added to the tables, as required. For example,

> stat.table(sex, mean(bweight), data = births, margins = TRUE)

for a one-way table, and

```
> stat.table(list(sex, hyp), mean(bweight), data = births, margins = c(TRUE,
+ FALSE))
> stat.table(list(sex, hyp), mean(bweight), data = births, margins = c(FALSE,
+ TRUE))
> stat.table(list(sex, hyp), mean(bweight), data = births, margins = c(TRUE,
+ TRUE))
```

for a two-way table.

Exercise 7.

- 1. Make a table of median birth weight by sex.
- 2. Do the same for gestation time, but include count as a function to be tabulated along with median. Note that when there are missing values for the variable being summarized the count refers to the number of non-missing observations for the row variable, not the summarized variable.
- 3. Create a table showing the mean gestation time for the baby by hyp and lowbw, together with margins for both.
- 4. Make a table showing the odds of hypertension by sex of the baby.

2.13 Generating new variables

New variables can be produced using assignment together with the usual mathematical operations and functions:

+ - * log exp ^ sqrt

The sign $\hat{}$ means "to the power of", log means "natural logarithm", and sqrt means "square root". The transform() function allows you to transform or generate variables in a data frame. For

example, try

```
> births <- transform(births, num1 = 1, num2 = 2, logbw = log(bweight))</pre>
```

The variable logbw is the natural logarithm of birth weight. Logs base 10 are obtained with log10().

2.14 Logical variables

Logical variables take the values TRUE or FALSE, and behave like factors. New variables can be created which are logical functions of existing variables. For example

```
> births <- transform(births, low = bweight < 2000)
> str(births)
```

creates a logical variable low with levels TRUE and FALSE, according to whether bweight is less than 2000 or not. The logical expressions which R allows are

= < <= > >= !=

The first is logical equals and the last is not equals. One common use of logical variables is to restrict a command to a subset of the data. For example, to list the values taken by bweight for hypertensive women, try

```
> births$bweight[births$hyp == "hyper"]
```

If you want the entire dataframe restricted to hypertensive women try:

```
> births[births$hyp == "hyper", ]
```

The subset() function also allows you to take a subset of a data frame. Try

```
> subset(births, hyp == "hyper")
```

Exercise 8.

- 1. Create a logical variable called **early** according to whether **gestwks** is less than 30 or not.Make a frequency table of **early**.
- 2. Print the id numbers of women with gestwks less than 30 weeks.

2.15 Dates in R

Epidemiological studies often contain date variables which take values such as 2/11/1962. We shall use the diet data to illustrate how to deal with variables whose values are dates.

The important variables in the dataset are chd, which takes the value 1 if the subject develops coronary heart disease during the study the value 0 if the observation is censored, and the three date variables which are date of birth (dob), date of entry (doe) and date of exit (dox). The command

> str(diet)

shows that these three variables are Date variables.

You will also see that the values are just numbers, but if you try

> head(diet)

you will see these variables printed as "real" dates. The variables are internally stored as number of days since 1/1/1970.

To convert a character string (or a character variable) to date format try:

```
> as.Date("14/07/1952", format = "%d/%m/%Y")
> as.numeric(as.Date("14/07/1952", format = "%d/%m/%Y"))
```

The first form shows the date form and the latter the number of days since 1/1/1970, which is a negative number for dates prior to 1/1/1970.

The format parts, "%d" etc., identify elements of the dates, whereas the "/"s are just the separator characters that are in the character string. There are other possibilities for formats, see ?strftime or the section on dates and times in the R command sheet at the end of this document .

Reading dates from an external file is done by reading the fields as character variables and then transforming them to date variables by the function **as.Date**

If you want to enter a fixed date, for example if you want to terminate follow-up at 1st April 1975 you could say:

> newx <- pmin(diet\$dox, as.Date("1975-4-1", format = "%F"))</pre>

The format %F is shorthand for the ISO-standard date representation %Y-m-%d, which is the default, so it can be omitted altogether:

```
> newx <- pmin(diet$dox, as.Date("1975-4-1"))</pre>
```

You can print dates in the format you like by using the function format.Date(), try for example:

```
> bdat <- as.Date("1952-7-14", format = "%F")
> format.Date(bdat, format = "%A %d %B %Y")
```

Exercise 9.

- 1. Enter your own birtday as a date. Print it using format.Date() with the format "%A %d %B %Y". Did you learn anything new?
- 2. Enter the birthday of your husband/wife/... as a date too. When will you be (were you) 100 years old together? (Hint: mean() works on vectors of dates as well.)

Working with R

3.1 Saving the work space

When exiting from R you are offered the chance of saving all the objects in your current work space. If you do so, the work space is re-instated next time you start R. It can be useful to do this, but before doing so it is worth tidying things up, because the work space can fill up with temporary objects, and it is easy to forget what these are when you resume the session.

3.2 Saving output in a file

To save the output from an R command in a file, for future use, the **sink()** command is used. For example,

```
> sink("output.txt")
> summary(births)
```

first instructs R to re-direct output away from the R terminal to the file "output.txt" and then summarizes the births data frame, the output from which goes to the sink. While a sink is open all output will go to it, replacing what is already in the file. To append output to a file, use the append=TRUE option with sink(). To close a sink, use

> sink()

Exercise 10.

- 1. Sink output to a file called "output1.txt".
- 2. Make frequency tables of hyp and sex
- 3. Make a table of mean birth weight by sex
- 4. Close the sink
- 5. From windows, have a look inside the file output1.txt and check that the output you expected is in the file.

3.3 Saving R objects in a file

The command read.table() is relatively slow because it carries out quite a lot of processing as it reads the data. To avoid doing this more than once you can save the data frame, which includes the R information, and read from this saved file in future. For example,

```
> save(births, file = "births.Rdata")
```

will save the births data frame in the file births.Rdata. By default the data frame is saved as a binary file, but the option ascii=TRUE can be used to save it as a text file. To load the object from the file use

```
> load("births.Rdata")
```

The commands **save()** and **load()** can be used with any R objects, but they are particularly useful when dealing with large data frames.

Exercise 11.

- 1. Use read.table() to read the data in the file diet.txt into a data frame called diet.
- 2. Save this data frame in the file "diet.Rdata"
- 3. Remove the data frame
- 4. Load the data frame from the file "diet.Rdata".

3.4 Using a text editor with R

When working with R it is best to use a text editor to prepare a batch file (or script) which contains R commands and then to run them from the script. This means you can use the cut and paste facilities of the editor to cut down on typing. For Windows we recommend using the text editor Tinn-R, but you can use your favourite text editor instead if you prefer. Start up the editor and enter the following lines:

```
> births <- transform(births, lowbw = factor(lowbw, labels = c("normal",
+ "low")), hyp = factor(hyp, labels = c("normal", "hyper")),
```

+ sex = factor(sex, labels = c("male", "female")))



Now save the script as mygetbirths.R and run it. One major advantage of running all your R commands from a script is that you end up with a record of exactly what you did which can be repeated at any time.

This will also help you redo the analysis in the (highly likely) event that your data changes before you have finished all analyses.

Exercise 12.

1. Create a script called mytab.R which includes the lines

```
> stat.table(hyp, data = births)
> stat.table(sex, data = births)
```

and run just these two lines.

2. Edit the script to include the lines

```
> stat.table(sex, mean(bweight), data = births)
> stat.table(hyp, mean(bweight), data = births)
```

and run these two lines.

- 3. Edit the script to create a factor cutting matage at 20, 30, 35, 40, 45 years, and run just this part of the script.
- 4. Edit the script to create a factor cutting gestwks at 20, 35, 37, 39, 45 weeks, and run just this part of the script.
- 5. Save and run the entire script.

3.5 The search path

R organizes objects in different positions on a search path. The command

> search()

shows these positions. The first is the work space, or global environment, the second is the Epi package, the third is a package of commands called methods, the fourth is a package called stats, and so on. To see what is in the work space try

> objects()

You should see just the objects births and diet. The command objects(1) does the same as objects(). To see what is in the Epi package, try

> objects(2)

There are 29 functions in this package.

When you type the name of an object R looks for it in the order of the search path and will return the first object with this name that it finds. This is why it is best to start your session with a clean workspace, otherwise you might have an object in your workspace that masks another one later in the search path.

3.6 Attaching a data frame

The function objects(1) shows that the only objects in the workspace are births and diet. To refer to variables in the births data frame by name it is necessary to specify the name of the data frame, as in births\$hyp. This is quite cumbersome, and provided you are working primarily with one data frame, it can help to put a copy of the variables from a data frame in their own position on the search path. This is done with the function

> attach(births)

which places a copy of the variables in the births data frame in position 2. You can verify this with

```
> objects(2)
```

which shows the objects in this position are the variables from the **births** data frame. Note that the methods package has now been moved up to position 3, as shown by the **search()** function.

When you type the command:

> hyp

R will look in the first position where it fails to find hyp, then the second position where it finds hyp, which now gets printed.

Although convenient, attaching a data frame can give rise to confusion. For example, when you create a new object from the variables in an attached data frame, as in

```
> subgrp <- bweight[hyp == 1]</pre>
```

the object subgrp will be in your workspace (position 1 on the search path) not in position 2. To demonstrate this, try

```
> objects(1)
> objects(2)
```

Similarly, if you modify the data frame in the workspace the changes will not carry through to the attached version of the data frame. The best advice is to regard any operation on an attached data frame as temporary, intended only to produce output such as summaries and tabulations.

Beware of attaching a data frame more than once - the second attached copy will be attached in position 2 of the search path, while the first copy will be moved up to position 3. You can see this with

```
> attach(births)
```

```
> search()
```

Having several copies of the same data set can lead to great confusion. To detach a data frame, use the command

```
> detach(births)
```

which will detach the copy in position 2 and move everything else down one position. To detach the second copy repeat the command detach(births).

Exercise 13.

- 1. Use search() to make sure you have no data frames attached.
- 2. Use objects(1) to check that you have the data frame births in your work space.
- 3. Verify that typing births\$hyp will print the data on the variable hyp but typing hyp will not.
- 4. Attach the **births** data frame in position 2 and check that the variables from this data frame are now in position 2.
- 5. Verify that typing hyp will now print the data on the the variable hyp.
- 6. Summarize the variable bweight for hypertensive women.
- > setwd(sweave.wd)

Graphs in R

There are three kinds of plotting functions in R:

- 1. Functions that generate a new plot, e.g. hist() and plot().
- 2. Functions that add extra things to an existing plot, e.g. lines() and text().
- 3. Functions that allow you to interact with the plot, e.g. locator() and identify().

The normal procedure for making a graph in R is to make a fairly simple initial plot and then add on points, lines, text etc., preferably in a script.

4.1 Simple plot on the screen

Load the births data and get an overview of the variables:

```
> library(Epi)
> data(births)
> str(births)
```

Now attach the dataframe and look at the birthweight distribution with

```
> attach(births)
> hist(bweight)
```

The histogram can be refined – take a look at the possible options with

```
> "?"(hist)
```

and try some of the options, for example:

> hist(bweight, col = "gray", border = "white")

To look at the relationship between birthweight and gestational weeks, try

```
> plot(gestwks, bweight)
```

You can change the plot-symbol by the option pch=. If you want to see all the plot symbols try:

```
> plot(1:25, pch = 1:25)
```

Exercise 14.

- 1. Make a plot of the birth weight versus maternal age with
 - > plot(matage, bweight)
- 2. Label the axes with
 - > plot(matage, bweight, xlab = "Maternal age", ylab = "Birth weight (g)")

4.2 Colours

There are many colours recognized by R. You can list them all by colours() or, equivalently, colors() (R allows you to use British or American spelling). To colour the points of birthweight versus gestational weeks, try

```
> plot(gestwks, bweight, pch = 16, col = "green")
```

This creates a solid mass of colour in the centre of the cluster of points and it is no longer possible to see individual points. You can recover this information by overwriting the points with black circles using the points() function.

```
> points(gestwks, bweight)
```

4.3 Adding to a plot

The points() function is one of several functions that add elements to an existing plot. By using these functions, you can create quite complex graphs in small steps.

Suppose we wish to recreate the plot of birthweight vs gestational weeks using different colours for male and female babies. To start with an empty plot, try

> plot(gestwks, bweight, type = "n")

Then add the points with the **points** function.

```
> points(gestwks[sex == 1], bweight[sex == 1], col = "blue")
> points(gestwks[sex == 2], bweight[sex == 2], col = "red")
```

To add a legend explaining the colours, try

```
> legend("topleft", pch = 1, legend = c("Boys", "Girls"), col = c("blue",
+ "red"))
```

which puts the legend in the top left hand corner. Finally we can add a title to the plot with

> title("Birth weight vs gestational weeks in 500 singleton births")

Using indexing for plot elements

One of the most powerful features of R is the possibility to index vectors, not only to get subsets of them, but also for repeating their elements in complex sequences.

Putting separate colours on males and female as above would become very clumsy if we had a 5 level factor instead.

Instead of specifying one color for all points, we may specify a vector of colours of the same length as the gestwks and bweight vectors. This is rather tedious to do directly, but R allows you to specify an expression anywhere, so we can use the fact that sex takes the values 1 and 2, as follows:

First create a colour vector with two colours, and take look at sex:

```
> c("blue", "red")
> sex
```

Now see what happens if you index the colour vector by sex:

> c("blue", "red")[sex]

For every occurrence of a 1 in sex you get "blue", and for every occurrence of 2 you get "red", so the result is a long vector of "blue"s and "red"s corresponding to the males and females. This can now be used in the plot:

> plot(gestwks, bweight, pch = 16, col = c("blue", "red")[sex])

The same trick can be used if we want to have a separate symbol for mothers over 40 say. We first generate the indexing variable:

> oldmum <- (matage >= 40) + 1

Note we add 1 because ($matage \geq 40$) generates a logic variable, so by adding 1 we get a numeric variable with values 1 and 2, suitable for indexing:

> plot(gestwks, bweight, pch = c(16, 3)[oldmum], col = c("blue", + "red")[sex])

so where oldmum is 1 we get pch=16 (a dot) and where oldmum is 2 we get pch=3 (a cross).

R will accept any kind of complexity in the indexing as long as the result is a valid index, so you don't need to create the variable oldmum, you can create it on the fly:

```
> plot(gestwks, bweight, pch = c(16, 3)[(matage >= 40) + 1], col = c("blue",
+ "red")[sex])
```

Exercise 15.

- 1. Make a three level factor for maternal age with cutpoints at 30 and 40 years.
- 2. Use this to make the plot of gestational weeks with three different plotting symbols. (Hint: Indexing with a factor automatically gives indexes 1,2,3 etc.).

Generating colours

R has functions that generate a vector of colours for you. For example,

> rainbow(4)

produces a vector with 4 colours (not immediately human readable, though). There are a few other functions that generates other sequences of colours, type **?rainbow** to see them.

Gray-tones are produced by the function gray (or grey), which takes a numerical argument between 0 and 1; gray(0) is black and gray(1) is white. Try:

> plot(0:10, pch = 16, cex = 3, col = gray(0:10/10))
> points(0:10, pch = 1, cex = 3)

4.4 Interacting with a plot

The locator() function allows you to interact with the plot using the mouse. Typing locator(1) shifts you to the graphics window and waits for one click of the left mouse button. When you click, it will return the corresponding coordinates.

You can use locator() inside other graphics functions to position graphical elements exactly where you want them. Recreate the birth-weight plot,

```
> plot(gestwks, bweight, pch = c(16, 3)[(matage >= 40) + 1], col = c("blue",
+ "red")[sex])
```

> print(1:10)

and then add the legend where you wish it to appear by typing

```
> legend(locator(1), pch = 1, legend = c("Boys", "Girls"), col = c("blue",
+ "red"))
```

The identify() function allows you to find out which records in the data correspond to points on the graph. Try

> identify(gestwks, bweight)

When you click the left mouse button, a label will appear on the graph identifying the row number of the nearest point in the data frame births. If there is no point nearby, R will print a warning message on the console instead. To end the interaction with the graphics window, right click the mouse: the identify function returns a vector of identified points.

Exercise 16.

- 1. Use identify() to find which records correspond to the smallest and largest number of gestational weeks.
- 2. View all the variables corresponding to these records with

births[identify(gestwks, bweight),]

4.5 Saving your graphs for use in other documents

Once you have a graph on the screen you can click on $\boxed{\text{File}} \rightarrow \boxed{\text{Save as}}$, and choose the format you want your graph in. The PDF (Acrobat reader) format is normally the most economical, and Acrobat reader has good options for viewing in more detail on the screen. The Metafile format will give you an enhanced metafile .emf, which can be imported into a Word document by $\boxed{\text{Insert}} \rightarrow \boxed{\text{Picture}} \rightarrow \boxed{}$

From File. Metafiles can be resized and edited inside Word.

If you want exact control of the size of your plot-file you can start a graphics device *before* doing the plot. Instead of appearing on the screen, the plot will be written directly to a file. After the plot has been completed you will need to close the device again in order to be able to access the file. Try:

```
> win.metafile(file = "plot1.emf", height = 3, width = 4)
> plot(gestwks, bweight)
> dev.off()
```

This will give you a enhanced metafile plot1.emf with a graph which is 3 inches tall and 4 inches wide.

4.6 The par() command

It is possible to manipulate any element in a graph, by using the graphics options. These are collected on the help page of **par()**. For example, if you want axis labels always to be horizontal, use the command **par(las=1)**. This will be in effect until a new graphics device is opened.

Look at the typewriter-version of the help-page with

```
> "?"(par)
```





It is a good idea to take a print of this (having set the text size to "smallest" because it is long) and carry it with you at any time to read in buses, cinema queues, during boring lectures etc. Don't despair, few R-users can understand what all the options are for. par() can also be used to ask about the current plot, for example par("usr") will give you the exact extent of the axes in the current plot.

If you want more plots on a single page you can use the command

> par(mfrow = c(2, 3))

This will give you a layout of 2 rows by 3 columns for the next 6 graphs you produce. The plots will appear by row, i.e. in the top row first. If you want the plots to appear columnwise, use par(mfcol=c(2,3)) (you still get 2 rows by 3 columns). To restore the layout to a single plot per page use

> par(mfrow = c(1, 1))

4.7 The Lexis diagram

If you load the Epi package you can draw a Lexis diagram by:

```
> Lexis.diagram()
```

You can draw the lifelines of the members of the diet study in the diagram by:

```
> data(diet)
> Lexis.lines(entry.date = diet$doe, exit.date = diet$dox, birth.date = diet$dob,
+ fail = diet$chd)
```

Lexis.lines recognizes variables of format Date and automatically converts them to fractional years when plotting them. The age and date axes are in years, so if you want to add text or other things to the diagram you must give coordinates in years.

You would probably want to adjust the axes of the Lexis diagram, for example by:

```
> Lexis.diagram(age = c(30, 75), date = c(1950, 1995))
```

Additional exercises

Exercise 17.

- 1. Use R to calculate 27/481.
- 2. Find the probability below 1.5 in a Gaussian (normal) distribution.
- 3. What is the probability between -1.64 and +1.64 in a Gaussian distribution?
- 4. What is the probability above 10 in a chi-squared distribution on 5 df?
- 5. Create a vector v with components $(0, 5, 10, 15, \dots 65, 70, 80, 100)$ and find its length.

Exercise 18.

- 1. Load the diet data using data(diet. You must have the Epi package loaded to do this.
- 2. Get the names of the variables in diet, and obtain a summary of diet. The length of follow-up is recorded in the variable y and the outcome at the end of follow-up is recorded in the variable chd, coded 1 for coronary heart disease and 0 otherwise.
- 3. Convert hieng to a factor and label its levels "low" and "high".
- 4. Create a table showing the frequencies of the two levels of hieng.
- 5. Convert job to a factor and label its levels "driver", "conductor" and "bank".
- 6. Create a table showing the frequencies of the three levels of job.
- 7. Create a table showing a two-way table of frequencies for hieng and job.
- 8. Use stat.table(hieng,ratio(chd,y,1000),data=diet) to make a table of rates per 1000 by hieng.
- 9. Use stat.table to make a table of rates per 1000 by hieng and job.
- 10. Create a histogram of energy.
- 11. Create a boxplot of energy.
- 12. Create a boxplot of energy by job.
- 13. Generate a new variable measuring body mass index (height/weight²) and call it bmi.
- 14. Plot the cumulative distribution funcyion (cdf) of bmi and use this to select 5 possible break points when grouping the values of bmi.
- 15. Use cut to create a factor with the 5 break points selected above, and call this bmigrp.
- 16. Use stat.table to make a table of rates per 1000 by bmigrp.
- 17. Use stat.table to make a table of rates per 1000 by hieng and bmigrp.

R commands

This R Reference Card is written by Tom Short, EPRI PEAC, tshort@epri-peac.com, 2004-10-21 and granted to the public domain. See www.Rpad.org for the source and latest version. Includes material from R for Beginners by Emmanuel Paradis (with permission).

It is also available separately as a 4-page landscape document from the R-hompage www.r-project.org, Manuals \rightarrow contributed documentation.

Getting help

Most R functions have online documentation.

help(topic) documentation on topic

?topic — the same.

help.search("topic") search the help system

apropos("topic") the names of all objects in the search list matching the regular expression "topic"

help.start() start the HTML version of help
str(a) display the internal *str*ucture of an R object
summary(a) gives a "summary" of a, usually a

statistical summary but it is *generic* meaning it has different operations for different classes of **a**

ls() show objects in the search path; specify
 pat="pat" to search on a pattern

ls.str() str() for each variable in the search path

dir() show files in the current directory

methods(a) shows S3 methods of a

methods(class=class(a)) lists all the methods to handle objects of class a.

Input and output

separator sep="" is any whitespace; use header=TRUE to read the first line as a header of column names; use as.is=TRUE to prevent character vectors from being converted to factors; use comment.char="" to prevent "#" from being interpreted as a comment; use skip=n to skip n lines before reading data; see the help for options on row naming, NA treatment, and others

read.csv("filename",header=TRUE) id. but with
 defaults set for reading comma-delimited files
read.delim("filename",header=TRUE) id. but with

defaults set for reading tab-delimited files

read.fwf(file,widths,header=FALSE,sep="",as.is=FALSE)
read a table of fixed width formatted data into
a 'data.frame'; widths is an integer vector, giving
the widths of the fixed-width fields

save(file,...) saves the specified objects (...) in the XDR platform-independent binary format save.image(file) saves all objects

cat(..., file="", sep=" ") prints the arguments
 after coercing to character; sep is the character
 separator between arguments

print(a, ...) prints its arguments; generic, meaning it can have different methods for different objects

format(x,...) format an R object for pretty printing
write.table(x,file="",row.names=TRUE,col.names=TRUE,
 sep=" ") prints x after converting to a data
 frame; if quote is TRUE, character or factor
 columns are surrounded by quotes ("); sep is the

field separator; eol is the end-of-line separator; na is the string for missing values; use col.names=NA to add a blank column header to get the column headers aligned correctly for spreadsheet input

sink(file) output to file, until sink()

Most of the I/O functions have a file argument. This can often be a character string naming a file or a connection. file="" means the standard input or output. Connections can include files, pipes, zipped files, and R variables.

On windows, the file connection can also be used with description = "clipboard". To read a table copied from Excel, use

x <- read.delim("clipboard")</pre>

To write a table to the clipboard for Excel, use write.table(x,"clipboard",sep="\t",col.names=NA) For database interaction, see packages RODBC, DBI, RMySQL, RPgSQL, and ROracle. See packages XML, hdf5, netCDF for reading other file formats.

Data creation

- c(...) generic function to combine arguments with
 the default forming a vector; with
 recursive=TRUE descends through lists
 combining all elements into one vector
- from:to generates a sequence; ":" has operator
 priority; 1:4 + 1 is "2,3,4,5"
- seq(from,to) generates a sequence by= specifies
 increment; length= specifies desired length
- seq(along=x) generates 1, 2, ..., length(along);
 useful for for loops
- data.frame(...) create
 a data frame of the named or unnamed arguments;
 data.frame(v=1:4,ch=c("a","B","c","d"),n=10);
 shorter vectors are recycled to the length of the
 longest
- list(...) create a list of the named or unnamed arguments; list(a=c(1,2),b="hi",c=3i);
- array(x,dim=) array with data x; specify dimensions like dim=c(3,4,2); elements of x recycle if x is not long enough
- matrix(x,nrow=,ncol=) matrix; elements of x recycle
 factor(x.levels=) encodes a vector x as a factor
- gl(n,k,length=n*k,labels=1:n) generate levels
- (factors) by specifying the pattern of their levels (factors) by specifying the pattern of their levels; k is the number of levels, and n is the number of replications
- expand.grid() a data frame from all combinations of the supplied vectors or factors

cbind(...) id. by columns

Slicing and extracting data

| Indexing vectors | | | | | | |
|--|---------------------------------------|--|--|--|--|--|
| x[n] | \mathbf{n}^{th} element | | | | | |
| x[-n] | all but the \mathbf{n}^{th} element | | | | | |
| x[1:n] | first n elements | | | | | |
| x[-(1:n)] | elements from $\tt n+1$ to the end | | | | | |
| x[c(1,4,2)] | specific elements | | | | | |
| x["name"] | element named "name" | | | | | |
| x[x > 3] | all elements greater than 3 | | | | | |
| x[x > 3 & x < 5] | all elements between 3 and 5 $$ | | | | | |
| x[x %in% c("a","and","the")] elements in the given set | | | | | | |
| Indexing lists | | | | | | |
| x[n] list with elements n | | | | | | |
| $x[[n]]$ n^{th} element of the list | | | | | | |
| x[["name"]] element of the list named "name" | | | | | | |
| x\$name id. | | | | | | |
| Indexing matrices | | | | | | |
| x[i,j] element at row i, column j | | | | | | |
| x[i,] row i | | | | | | |
| x[,j] column j | | | | | | |
| x[,c(1,3)] columns 1 and 3 | | | | | | |
| x["name",] row named "name" | | | | | | |
| Indexing data frames (matrix indexing plus the | | | | | | |
| following) | | | | | | |
| x[["name"]] column named "name" | | | | | | |
| x\$name id. | | | | | | |

Variable conversion

as.array(x), as.data.frame(x), as.numeric(x), as.logical(x), as.complex(x), as.character(x), ... convert type; for a complete list, use methods(as)

Variable information

- is.na(x), is.null(x), is.array(x), is.data.frame(x), is.numeric(x), is.complex(x), is.character(x), ... test for type; for a complete list, use methods(is) length(x) number of elements in x dim(x) Retrieve or set the dimension of an object; dim(x) <- c(3,2)dimnames(x) Retrieve or set the dimension names of an object nrow(x) number of rows; NROW(x) is the same but treats a vector as a one-row matrix ncol(x) and NCOL(x) id. for columns class(x) get or set the class of x; class(x) <-</pre> "mvclass" unclass(x) remove the class attribute of x attr(x,which) get or set the attribute which of x

Data selection and manipulation

- which.max(x) returns the index of the greatest
 element of x
- which.min(x) returns the index of the smallest
 element of x
- $\mathtt{rev}(\mathtt{x})$ reverses the elements of \mathtt{x}
- sort(x) sorts the elements of x in increasing order; to sort in decreasing order: rev(sort(x))
- cut(x,breaks) divides x into intervals (factors); breaks is the number of cut intervals or a vector of cut points
- which(x == a) returns a vector of the indices of x if
 the comparison operation is true (TRUE), in this
 example the values of i for which x[i] == a (the
 argument of this function must be a variable of
 mode logical)
- choose(n, k) computes the combinations of k events among n repetitions = n!/[(n-k)!k!]
- na.omit(x) suppresses the observations with missing data (NA) (suppresses the corresponding line if x is a matrix or a data frame)
- <code>na.fail(x)</code> returns an error message if x contains at least one NA
- unique(x) if x is a vector or a data frame, returns a similar object but with the duplicate elements suppressed
- table(x) returns a table with the numbers of the differents values of x (typically for integers or factors)
- subset(x, ...) returns a selection of x with respect
 to criteria (..., typically comparisons: x\$V1 <
 10); if x is a data frame, the option select gives
 the variables to be kept or dropped using a
 minus sign</pre>

- sample(x, size) resample randomly and without
 replacement size elements in the vector x, the
 option replace = TRUE allows to resample with
 replacement
- prop.table(x,margin=) table entries as fraction of marginal table

Math

- sin,cos,tan,asin,acos,atan,atan2,log,log10,exp
- max(x) maximum of the elements of x
- min(x) minimum of the elements of ${\tt x}$
- range(x) id. then c(min(x), max(x))
- sum(x) sum of the elements of x
- $\mathtt{diff}(\mathtt{x})$ lagged and iterated differences of vector \mathtt{x}
- prod(x) product of the elements of x
- mean(x) mean of the elements of x
 median(x) median of the elements of x
- quantile(x,probs=) sample quantiles corresponding
 to the given probabilities (defaults to
- 0,25,5,75,1) weighted.mean(x, w) mean of x with weights w
- rank(x) ranks of the elements of x
- var(x) or cov(x) variance of the elements of x
- (calculated on n-1); if **x** is a matrix or a data frame, the variance-covariance matrix is calculated
- sd(x) standard deviation of x
- var(x, y) or cov(x, y) covariance between x and y, or between the columns of x and those of y if they are matrices or data frames
- cor(x, y) linear correlation between x and y, or correlation matrix if they are matrices or data frames
- round(x, n) rounds the elements of x to n decimals
- $\log(\mathtt{x}, \ \mathtt{base})$ computes the logarithm of \mathtt{x} with base \mathtt{base}
- scale(x) if x is a matrix, centers and reduces the
 data; to center only use the option
 center=FALSE, to reduce only scale=FALSE (by
 default center=TRUE, scale=TRUE)
- pmin(x,y,...) a vector which ith element is the minimum of x[i], y[i], ...
- pmax(x,y,...) id. for the maximum
- cumsum(x) a vector which ith element is the sum
 from x[1] to x[i]
- cumprod(x) id. for the product
- cummin(x) id. for the minimum
- cummax(x) id. for the maximum
- union(x,y), intersect(x,y), setdiff(x,y), setequal(x,y), is.element(el,set) "set" functions
- Re(x) real part of a complex number
- Im(x) imaginary part
- Mod(x) modulus; abs(x) is the same
- Arg(x) angle in radians of the complex number
- Conj(x) complex conjugate
- fft(x) Fast Fourier Transform of an array
- mvfft(x) FFT of each column of a matrix
- filter(x,filter) applies linear filtering to a
 univariate time series or to each series separately
 of a multivariate time series
- Many math functions have a logical parameter
- $\tt na.rm=FALSE$ to specify missing data (NA) removal.

Matrices

t(x) transpose diag(x) diagonal %*% matrix multiplication solve(a,b) solves a %*% x = b for x solve(a) matrix inverse of a rowsum(x) sum of rows for a matrix-like object; rowSums(x) is a faster version colsum(x), colSums(x) id. for columns rowMeans(x) fast version of row means colMeans(x) id. for columns

Advanced data processing

- apply(X,INDEX,FUN=) a vector or array or list of values obtained by applying a function FUN to margins (INDEX) of X
- lapply(X,FUN) apply FUN to each element of the list X
 tapply(X,INDEX,FUN=) apply FUN to each cell of a
- ragged array given by X with indexes INDEX by(data,INDEX,FUN) apply FUN to data frame data subsetted by INDEX

- aggregate(x,by,FUN) splits the data frame x into subsets, computes summary statistics for each, and returns the result in a convenient form; by is a list of grouping elements, each as long as the variables in x
- stack(x, ...) transform data available as separate
 columns in a data frame or list into a single
 column
- unstack(x, ...) inverse of stack()
- reshape(x, ...) reshapes a data frame between
 'wide' format with repeated measurements in
 separate columns of the same record and 'long'
 format with the repeated measurements in
 separate records; use (direction="wide") or
 (direction="long")

Strings

- paste(...) concatenate vectors after converting to character; sep= is the string to separate terms (a single space is the default); collapse= is an optional string to separate "collapsed" results
- substr(x,start,stop) substrings in a character vector; can also assign, as substr(x, start, stop) <- value</pre>
- strsplit(x,split) split x according to the substring
 split
- grep(pattern,x) searches for matches to pattern
 within x; see ?regex
- gsub(pattern,replacement,x) replacement of matches determined by regular expression matching sub() is the same but only replaces the first occurrence.
- tolower(x) convert to lowercase
- toupper(x) convert to uppercase
- match(x,table) a vector of the positions of first
 matches for the elements of x among table
 x %in% table id. but returns a logical vector

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Dates and Times

The class Date has dates without times. POSIXct has dates and times, including time zones. Comparisons (e.g. >), seq(), and difftime() are useful. Date also allows + and -. ?DateTimeClasses gives more information. See also package chron. as.Date(s) and as.POSIXct(s) convert to the respective class. format(dt) converts to a string representation. The default string format is "2001-02-21". These accept a second argument to specify a format for conversion. Some common formats are:

- %a, %A Abbreviated and full weekday name.
- $\ensuremath{\sc k}\ensuremath{\mathsf{b}}$, $\ensuremath{\sc k}\ensuremath{\mathsf{B}}$ Abbreviated and full month name.
- % d Day of the month (01–31).
- %H Hours (00-23).
- %I Hours (01-12).
- %j Day of year (001–366).
- %m Month (01-12).
- %M Minute (00-59).
- %p AM/PM indicator.
- **%S** Second as decimal number (00-61).
- U Week (00–53); the first Sunday as day 1 of week 1.
- **%** Weekday (0–6, Sunday is 0).
- % Week (00-53); the first Monday as day 1 of week 1. % Year without century (00-99). Don't use.
- XY Year with century (00
- %z (output only.) Offset from Greenwich; -0800 is 8 hours west of.
- %Z (output only.) Time zone as a character string (empty if not available).

Where leading zeros are shown they will be used on output but are optional on input. See ?strftime.

Plotting

- plot(x, y) bivariate plot of x (on the x-axis) and y
 (on the y-axis)
- hist(x) histogram of the frequencies of x
- barplot(x) histogram of the values of x; use horiz=FALSE for horizontal bars
- dotplot(x) if x is a data frame, plots a Cleveland dot
 plot (stacked plots line-by-line and
 column-by-column)
- piechart(x) circular pie-chart
- boxplot(x) "box-and-whiskers" plot
- sunflowerplot(x, y) id. than plot() but the points
 with similar coordinates are drawn as flowers
 which petal number represents the number of
 points
- $\texttt{coplot}(\texttt{x}^{\neg}\texttt{y} ~|~ \texttt{z})$ bivariate plot of x and y for each value or interval of values of z
- interaction.plot (f1, f2, y) if f1 and f2 are
 factors, plots the means of y (on the y-axis) with
 respect to the values of f1 (on the x-axis) and of
 f2 (different curves); the option fun allows to
 choose the summary statistic of y (by default
 fun=mean)

- matplot(x,y) bivariate plot of the first column of x
 vs. the first one of y, the second one of x vs. the
 second one of y, etc.
- fourfoldplot(x) visualizes, with quarters of circles, the association between two dichotomous variables for different populations (x must be an array with dim=c(2, 2, k), or a matrix with dim=c(2, 2) if k = 1)
- assocplot(x) Cohen-Friendly graph showing the deviations from independence of rows and columns in a two dimensional contingency table
- mosaicplot(x) 'mosaic' graph of the residuals from a
 log-linear regression of a contingency table. Also
 useful for graphical display of contingency tables.
- pairs(x) if x is a matrix or a data frame, draws all
 possible bivariate plots between the columns of x
- plot.ts(x) if x is an object of class "ts", plot of x
 with respect to time, x may be multivariate but
 the series must have the same frequency and
 dates
- ts.plot(x) id. but if x is multivariate the series may have different dates and must have the same frequency
- qqnorm(x) quantiles of x with respect to the values expected under a normal law
- qqplot(x, y) quantiles of y with respect to the quantiles of x
- contour(x, y, z) contour plot (data are interpolated to draw the curves), x and y must be vectors and z must be a matrix so that dim(z)=c(length(x), length(y)) (x and y may be omitted)
- filled.contour(x, y, z) id. but the areas between
 the contours are coloured, and a legend of the
 colours is drawn as well

- stars(x) if x is a matrix or a data frame, draws a
 graph with segments or a star where each row of
 x is represented by a star and the columns are
 the lengths of the segments
- symbols(x, y, ...) draws, at the coordinates given
 by x and y, symbols (circles, squares, rectangles,
 stars, thermometres or "boxplots") which sizes,
 colours ... are specified by supplementary
 arguments
- termplot(mod.obj) plot of the (partial) effects of a
 regression model (mod.obj)
- The following parameters are common to many plotting functions:
- add=FALSE if TRUE superposes the plot on the previous one (if it exists)
- $\verb+axes=TRUE$ if <code>FALSE</code> does not draw the axes and the box
- type="p" specifies the type of plot, "p": points, "1": lines, "b": points connected by lines, "o": id. but the lines are over the points, "h": vertical lines, "s": steps, the data are represented by the top of the vertical lines, "S": id. but the data are represented by the bottom of the vertical lines
- xlim=, ylim= specifies the lower and upper limits of the axes, for example with xlim=c(1, 10) or xlim=range(x)
- xlab=, ylab= annotates the axes, must be variables
 of mode character
- main= main title, must be a variable of mode
 character

sub= sub-title (written in a smaller font)

Low-level plotting commands

- lines(x, y) id. but with lines
- text(x, y, labels, ...) adds text given by labels
 at coordinates (x,y); a typical use is: plot(x, y,
 type="n"); text(x, y, names)
- mtext(text, side=3, line=0, ...) adds text given
 by text in the margin specified by side (see
 axis() below); line specifies the line from the
 plotting area
- segments(x0, y0, x1, y1) draws lines from points
 (x0,y0) to points (x1,y1)
- arrows(x0, y0, x1, y1, angle= 30, code=2) id. with arrows at points (x0,y0) if code=2, at points (x1,y1) if code=1, or both if code=3; angle controls the angle from the shaft of the arrow to the edge of the arrow head
- abline(a,b) draws a line of slope b and intercept a
- abline(h=y) draws a horizontal line at ordinate y
- abline(v=x) draws a vertical line at abcissa x
- abline(lm.obj) draws the regression line given by lm.obj
- rect(x1, y1, x2, y2) draws a rectangle which left, right, bottom, and top limits are x1, x2, y1, and y2, respectively
- polygon(x, y) draws a polygon linking the points
 with coordinates given by x and y
- $\texttt{legend}(x,\ y,\ \texttt{legend})$ adds the legend at the point (x,y) with the symbols given by legend
- title() adds a title and optionally a sub-title
 axis(side, vect) adds an axis at the bottom
 (side=1), on the left (2), at the top (3), or on the
 right (4); vect (optional) gives the abcissa (or
 ordinates) where tick-marks are drawn
- rug(x) draws the data x on the x-axis as small
 vertical lines
- locator(n, type="n", ...) returns the coordinates
 (x, y) after the user has clicked n times on the
 plot with the mouse; also draws symbols
 (type="p") or lines (type="1") with respect to
 optional graphic parameters (...); by default
 nothing is drawn (type="n")

Graphical parameters

These can be set globally with **par(...)**; many can be passed as parameters to plotting commands.

- adj controls text justification (0 left-justified, 0.5 centred, 1 right-justified)
- bg specifies the colour of the background (ex. :
 bg="red", bg="blue", ... the list of the 657
 available colours is displayed with colors())
- bty controls the type of box drawn around the plot, allowed values are: "o", "l", "7", "c", "u" ou "]" (the box looks like the corresponding character); if bty="n" the box is not drawn
- cex a value controlling the size of texts and symbols
 with respect to the default; the following
 parameters have the same control for numbers on
 the axes, cex.axis, the axis labels, cex.lab, the
 title, cex.main, and the sub-title, cex.sub

- col controls the color of symbols and lines; use color names: "red", "blue" see colors() or as "#RRGGBB"; see rgb(), hsv(), gray(), and rainbow(); as for cex there are: col.axis, col.lab, col.main, col.sub
- font an integer which controls the style of text (1:
 normal, 2: italics, 3: bold, 4: bold italics); as for
 cex there are: font.axis, font.lab, font.main,
 font.sub
- las an integer which controls the orientation of the axis labels (0: parallel to the axes, 1: horizontal, 2: perpendicular to the axes, 3: vertical)
- lty controls the type of lines, can be an integer or string (1: "solid", 2: "dashed", 3: "dotted", 4: "dotdash", 5: "longdash", 6: "twodash", or a string of up to eight characters (between "0" and "9") which specifies alternatively the length, in points or pixels, of the drawn elements and the blanks, for example lty="44" will have the same effect than lty=2
- ${\tt lwd}$ a numeric which controls the width of lines, default 1
- mar a vector of 4 numeric values which control the space between the axes and the border of the graph of the form c(bottom, left, top, right), the default values are c(5.1, 4.1, 4.1, 2.1)
- mfcol a vector of the form c(nr,nc) which partitions
 the graphic window as a matrix of nr lines and
 nc columns, the plots are then drawn in columns
 mfrow id. but the plots are drawn by row
- pch controls the type of symbol, either an integer between 1 and 25, or a single character in "":
- ps an integer which controls the size in points of texts and symbols
- pty a character which specifies the type of the plotting region, "s": square, "m": maximal
- tck a value which specifies the length of tick-marks on the axes as a fraction of the smallest of the width or height of the plot; if tck=1 a grid is drawn
- tcl a value which specifies the length of tick-marks
 on the axes as a fraction of the height of a line of
 text (by default tcl=-0.5)
- xaxt if xaxt="n" the x-axis is set but not drawn (useful in conjonction with axis(side=1, ...)) yaxt if yaxt="n" the y-axis is set but not drawn
- (useful in conjonction with axis(side=2, ...))

Lattice (Trellis) graphics

- $\texttt{barchart}(\texttt{y^x})$ histogram of the values of y with respect to those of x
- bwplot(y~x) "box-and-whiskers" plot
- densityplot(~x) density functions plot
- histogram(~x) histogram of the frequencies of x qqmath(~x) quantiles of x with respect to the values
- expected under a theoretical distribution
 stripplot(y^x) single dimension plot, x must be
 numeric, y may be a factor
- qq(y^xx) quantiles to compare two distributions, x must be numeric, y may be numeric, character, or factor but must have two 'levels'

xyplot(y~x) bivariate plots (with many functionalities)

levelplot(z^xx*y) coloured plot of the values of z at the coordinates given by x and y (x, y and z are all of the same length)

splom(~x) matrix of bivariate plots

parallel(~x) parallel coordinates plot

Optimization and model fitting

- optim(par, fn, method = c("Nelder-Mead", "BFGS", "CG", "L-BFGS-B", "SANN") general-purpose optimization; par is initial values, fn is function to optimize (normally minimize)
- lm(formula) fit linear models; formula is typically of the form response termA + termB + ...; use I(x*y) + I(x^2) for terms made of nonlinear components
- glm(formula,family=) fit generalized linear models, specified by giving a symbolic description of the linear predictor and a description of the error distribution; family is a description of the error distribution and link function to be used in the model; see ?family
- nls(formula) nonlinear least-squares estimates of the nonlinear model parameters
- spline(x,y=) cubic spline interpolation
- loess(formula) fit a polynomial surface using local
 fitting

Many of the formula-based modeling functions have several common arguments: data= the data frame for the formala variables, subset= a subset of variables used in the fit, na.action= action for missing values: "na.fail", "na.omit", or a function. The following

generics often apply to model fitting functions:

- df.residual(fit) returns the number of residual degrees of freedom
- coef(fit) returns the estimated coefficients
 (sometimes with their standard-errors)
- residuals(fit) returns the residuals
- deviance(fit) returns the deviance
- fitted(fit) returns the fitted values

logLik(fit) computes the logarithm of the

likelihood and the number of parameters

AIC(fit) computes the Akaike information criterion or AIC

Statistics

aov(formula) analysis of variance model anova(fit,...) analysis of variance (or deviance) tables for one or more fitted model objects density(x) kernel density estimates of x

binom.test(), pairwise.t.test(), power.t.test(), prop.test(), t.test(), ... use help.search("test")

Distributions

rnorm(n, mean=0, sd=1) Gaussian (normal) rexp(n, rate=1) exponential rgamma(n, shape, scale=1) gamma rpois(n, lambda) Poisson rweibull(n, shape, scale=1) Weibull rcauchy(n, location=0, scale=1) Cauchy rbeta(n, shape1, shape2) beta rt(n, df) 'Student' (t) rf(n, df1, df2) Fisher-Snedecor (F) (χ^2) rchisq(n, df) Pearson rbinom(n, size, prob) binomial rgeom(n, prob) geometric rhyper(nn, m, n, k) hypergeometric rlogis(n, location=0, scale=1) logistic rlnorm(n, meanlog=0, sdlog=1) lognormal rnbinom(n, size, prob) negative binomial runif(n, min=0, max=1) uniform rwilcox(nn, m, n), rsignrank(nn, n) Wilcoxon's statistics All these functions can be used by replacing the

letter r with d, p or q to get, respectively, the probability density (dfunc(x, ...)), the cumulative probability density (pfunc(x, ...)), and the value of quantile (qfunc(p, ...), with 0).

Programming

arguments to be passed to it.

The Epi package

- Lexis.diagram() Draw a Lexis diagram, optionally with life lines.
- Lexis.lines() Add lines to a Lexis diagram. rateplot(rates,...) Make plots of rates from an
- age by period table.
 cal.yr(x,format) Convert x to fractional calendar
 year.
- stat.table(index,contents,...) Make tables, classified by index, of sums, ratios etc. giben in contents.
- ci.lin(obj,ctr.mat,subset,diffs,Exp) Extract
 parameters and linear functions of them from
 model objects.
- plotEst(ests,...) Make a plot of parameter
 estimates.
- twoby2(exposure,outcome,...) Analysis of a 2×2 table. Input can be either two binary variables or a matrix of counts.