# Using the Devore7 package with R

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# Contents

0	Preliminaries	7
1	Overview and Descriptive Statistics	11
	1.1 Example 1.1	. 11
	1.2 Example 1.5	
	1.3 Example 1.6	
	1.4 Example 1.9	
	1.5 Example 1.10	
	1.6 Examples 1.12 and 1.13	
	1.7 Example 1.14	
	1.8 Example 1.15	. 16
	1.9 Examples 1.17	
	1.10 Example 1.18	
	1.11 Comparative boxplots	
	1.12 Enhancing graphical displays	
<b>2</b>	Probability	21
	2.1 Example 2.23	. 21
3	Discrete Random Variables	<b>23</b>
	3.1 Example 3.31	. 24
	3.2 Example 3.32	. 24
	3.3 Example 3.35	. 25
	3.4 Example 3.36	. 25
	3.5 Example 3.38	. 25
	3.6 Example 3.39	. 25
	3.7 Example 3.40	. 26
4	Continuous Random Variables	27
	4.1 Example 4.13	. 27
	4.2 Example 4.14	
	4.3 Example 4.15	
	4.4 Example 4.16	
	4.5 Example 4.18	

4 CONTENTS

	4.6	Example 4.20	29
	4.7	Example 4.23	29
	4.8	Example 4.24	30
	4.9		30
	4.10	Example 4.25	30
			31
	4.12	Example 4.28	31
			31
	4.14	Example 4.31	33
5	Join	t Probability Distributions 3	5
	5.1	Example 5.22	35
	5.2	Normal data, like example 5.23	37
	5.3	Example 5.23	8
6	Poir	t Estimation 4	13
	6.1	Example 6.2	13
	6.2	Example 6.3	13
	6.3	Example 6.13	14
7	Stat	istical Intervals 4	15
	7.1	Example 7.2	15
	7.2	Example 7.6	15
	7.3	Example 7.8	16
	7.4	Example 7.11	17
	7.5	Example 7.15	18
8	Test	s of Hypotheses 5	1
	8.1	Example 8.8	51
	8.2	Example 8.9	53
	8.3	Example 8.10	54
	8.4	Example 8.11	54
	8.5	Example 8.13	55
9	Infe		7
	9.1	1	57
	9.2	Example 9.8	58
	9.3	1	59
	9.4	Example 9.10	60
10	The	Analysis of Variance 6	3
11	Mul	tifactor Analysis of Variance 6	5
		· ·	35
		1	6
			57
			39

CONTENTS	F
CONTENTS	.5
CONTENTS	0

	11.5	Example 11.10	1
	11.6	Example 11.11	2
A	Inst	alling R and the Devore7 package 7	5
	A.1	What is R?	5
	A.2	Obtaining and Installing R	5
	A.3	Quitting R	6
	A.4	Using data sets	6
	A.5	What is Devore 7?	7
	A.6	Installing and attaching Devore7	7
	A.7	Names of the data sets	7
	A.8	Data sets as tables	8
	A.9	Accessing individual variables	9
	A.10	Stacked data	0
	A.11	Finding help	1

6 CONTENTS

# **Preliminaries**

This document describes the use of the statistical package R as computing support in an introductory statistics course based on the text *Probability and Statistics for Engineering and the Sciences (7th edition)* by Jay Devore (Thomson Brooks-Cole, 2008). We demonstrate how R can be used to reproduce the results in many of the examples in the text.

One of the desirable features of this text is the number of examples and exercises based on real data sets the Prof. Devore has culled from the engineering literature. As they are real data, some of the data sets are large and have a complex structure. Although it is not difficult to enter these data into a computer package like R, the process is tedious and error-prone. Furthermore, it is not much of a learning experience.

We have provided copies of the data sets for the examples and the exercises in a "package", named  $\tt Devore7$ , that can be used with R. This document is also part of the  $\tt Devore7$  package.  $^1$ 

You may wish to try some of the examples in this section as you are reading it. We assume that you have both R and the Devore7 package for R installed. (See Appendix A for instructions if you need to do this first.)

#### Calculating a median

Suppose that we wish to reproduce the calculation of the median of the data on transferrin receptor concentration shown in Example 1.13 (p. 27 of the text). As there are only 12 concentrations, we could enter the data by hand. Start R and type

```
> conc = c(7.6, 8.3, 9.3, 9.4, 9.4, 9.7, 10.4, 11.5, 11.9,
+    15.2, 16.2, 20.4)
> str(conc)
num [1:12] 7.6 8.3 9.3 9.4 9.4 9.7 10.4 11.5 11.9 15.2 ...
```

<sup>&</sup>lt;sup>1</sup>For the Devore7 package, some of the data sets were not provided by the publisher. Many, but not all, missing ones were ported from the Devore6 package.

```
> median(conc)
```

[1] 10.05

The first line assigns the 12 data values as a numeric vector to the name conc, a short form of "concentration". The function named "c" concatenates a series of data values into a vector that can be assigned a name.

In the next line the "str" function is used to examine the structure of the object named "conc". The output shows that this is a numeric vector of length 12 and displays the first several data values so you can check them against the data in the text.

Finally the "median" function, applied to the conc vector returns the median.

#### Using the Devore7 package

Entering the data, as shown above, is suitable for small data sets. An alternative and preferred way to access the data for the larger data sets, is to use the Devore7 package and load the data set. The data set for Example 1.13 is called xmp01.13. In general, data sets for examples in the text are named xmpcc.nn, where cc is the two-digit chapter number and nn is the two-digit example number. Data sets for exercises are named excc.nn. (Single digit chapter or example numbers have a 0 prepended as in xmp01.13 so that the names sort in the correct order.)

You must attach the Devore7 package every time you start R if you are to have access to the data sets from the textbook like this.

To attach the package to an R session use

#### > library(Devore7)

after starting R or select Packages -> Load package -> Devore7 from the menu bar. (If this produces an error see Appendix A for instructions on installing the Devore7 package.)

After attaching the package, you can load a data set with

```
> data(xmp01.13)
> str(xmp01.13)

'data.frame': 12 obs. of 1 variable:
$ concentration: num 15.2 9.3 7.6 11.9 10.4 9.7 20.4 9.4 11.5 16.2 ...
```

The first line loads the data set into the current R session. The second line provides a description of the structure of the data set. It is a good practice always to use str on the data set after loading it.

#### Form of the data sets

The data set xmp01.13 is not a single vector like conc. It is a data table (called a "data frame" in R) with one column and 12 rows. All the data sets in the Devore7 package are data frames.

The output from str indicates that the name of the first column is concentration. To calculate the median we must give both the name of the data frame and the name of the column. We can do this in three ways, as described in Appendix A.9. We will focus on just one of these ways, which is to use the "with" function.

```
> with(xmp01.13, median(concentration))
```

```
[1] 10.05
```

The with function indicates which data set should be used to gain access to the column (or "variable") called concentration.

#### Summary

To recap:

- 1. You should have R installed on a computer and the Devore7 package for R installed. (See Appendix A for instructions if you need to do this.)
- 2. At the beginning of each session use
  - > library(Devore7)

to allow access to the data sets from the package.

3. To load the data for a specific example or a specific exercise use a name of the form xmpcc.nn or excc.nn to data() then check the structure with str().

```
> data(xmp01.13)
> str(xmp01.13)
```

In the remainder of this document we will not show these steps explicitly.

The R functions we have mentioned are shown in Table 1. See also Appendix A.11.

Function	Purpose
q()	quit R
help(name)	display help on an object (function or data set)
help.search("topic")	search for functions related to a topic
library(name)	Make data sets from a package available for loading
data(name)	Load a data set
str(name)	Display a brief description of the structure
with $(dataset,)$	Use the variables in a data set

Table 1: R functions for general use

# Overview and Descriptive Statistics

We will follow the same sequence of topics and chapter headings as in the text and will begin each chapter with a table of R functions that are used in the chapter.

Table 1.1 lists functions used in chapter 1.

Function	Description
stem(x)	stem-and-leaf display
hist(x)	histogram
boxplot(x)	boxplot
mean(x)	mean (i.e. average) value of x
median(x)	median
var(x)	sample variance
sd(x)	sample standard deviation
log(x)	natural logarithm (works on entire vectors)
log(x, 10)	common (base 10) logarithm
$\operatorname{sqrt}(x)$	square root
$x^{(1/3)}$	cube root

Table 1.1: R functions used with chapter 1

#### 1.1 Example 1.1

Example 1.1 (p. 4) lists the ambient temperatures (°F) for each test firing or actual launch of the space shuttle prior to the *Challenger* tradgedy in 1986. In Figure 1.1 (p. 4) these data are displayed as a stem-and-leaf plot and as a histogram. There are 36 data values.

A stem-and-leaf plot similar to that in Figure 1.1 (p. 4) can be produced with

> with(xmp01.01, stem(temp))

The decimal point is 1 digit(s) to the right of the |

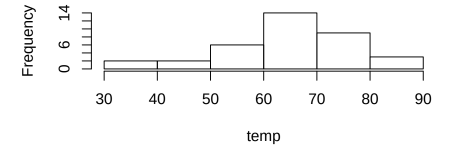
- 3 | 1
- 4 | 059
- 5 | 23788
- 6 | 01136777789
- 7 | 000023556689
- 8 | 0134

(Remember that first you must attach the Devore7 package, load the xmp01.01 data set and check its structure. We do not show those steps here.)

The hist function produces a histogram.

> with(xmp01.01, hist(temp))

#### Histogram of temp



The stem-and-leaf plot and the histogram shown here are not exactly the same as those shown in Figure 1.1 (p. 4). In section 1.12 we show how optional arguments to stem and to hist could be used to produce displays similar to those in the text.

#### 1.2 Example 1.5

The data in example 1.5 (p. 11), on the percentage of undergraduate students who are binge drinkers at 140 different campuses, are presented as a stem-and-leaf display in Figure 1.4 (p. 12).

> with(xmp01.05, stem(bingePct))

```
The decimal point is 1 digit(s) to the right of the |
  0 | 4
  0 |
  1 | 134
  1 | 5678889
  2 | 12234
  2 | 56666777889999
  3 | 0112233344
  3 | 555666677777888899999
  4 | 11122222334444
  4 | 5566666677788888999
  5 | 00111222334
  5 | 55666667777888899
  6 | 011112444
  6 | 55666778
> with(xmp01.05, stem(bingePct, scale = 0.5))
  The decimal point is 1 digit(s) to the right of the |
  0 | 4
  1 | 1345678889
  2 | 1223456666777889999
  3 | 0112233344555666677777888899999
  4 | 111222223344445566666677788888999
  5 | 00111222233455666667777888899
  6 | 01111244455666778
```

The first stem-and-leaf display is more spread out than the one in Figure 1.4 (p. 12). In the second call to stem we use the optional argument scale to shrink the scale by a factor of  $\frac{1}{2}$  so the resulting display is similar to that in Figure 1.4.

#### 1.3 Example 1.6

As in Example 1.5 a stem-and-leaf display is created, this time from data on yardages of golf courses as given in *Golf Magazine*.

```
> with(xmp01.06, stem(yardage))
The decimal point is 2 digit(s) to the right of the |
64 | 3467
65 | 1338
66 | 119
67 | 015779
```

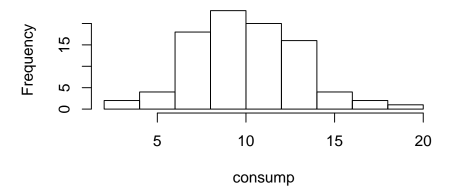
68 | 05779 69 | 0034 70 | 112455 71 | 113777 72 | 18

#### 1.4 Example 1.9

A histogram such as shown in Figure 1.8 (p. 16) is produced by the hist function.

> with(xmp01.09, hist(consump))

#### Histogram of consump

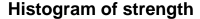


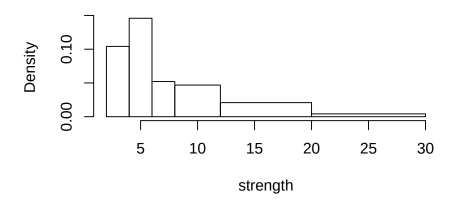
The vertical axis on this histogram is frequency. For a vertical axis on the scale of density (relative frequency divided by bin width) use the optional argument freq = FALSE.

#### 1.5 Example 1.10

Figure 1.10 (p. 28) shows an example of a histogram with unequal bin widths. The optional argument breaks to the hist function is used to set the breakpoints for the bins. When unequal bin widths are used, the vertical axis switches to the density scale.

```
> with(xmp01.10, hist(strength, breaks = c(2, 4, 6, 8, 12, + 20, 30))
```





The breaks argument is a vector created by concatenating several numbers with the  ${\tt c}$  function.

#### 1.6 Examples 1.12 and 1.13

Numeric measures of location are calculated with the functions mean and median. Another useful summary function is sum. A brief summary, including the mean, the median, the quartiles, the maximum and minimum is returned by summary.

```
> with(xmp01.12, mean(crackLength))
[1] 21.18095
> with(xmp01.12, sum(crackLength))
[1] 444.8
> with(xmp01.13, median(concentration))
[1] 10.05
> with(xmp01.12, summary(crackLength))
                Median
   Min. 1st Qu.
                           Mean 3rd Qu.
                                           Max.
   8.90
          14.00
                  21.20
                          21.18
                                  25.80
                                          45.00
> with(xmp01.13, summary(concentration))
  Min. 1st Qu.
                 Median
                           Mean 3rd Qu.
                                           Max.
  7.600
          9.375 10.050 11.610 12.730
                                         20.400
```

#### 1.7 Example 1.14

The trimmed mean, described in Example 1.14 (p. 28) is obtained by using the optional trim argument to mean.

```
> with(xmp01.14, summary(copper))
   Min. 1st Qu. Median Mean 3rd Qu. Max.
   2.000   2.700   3.300   3.667   3.875   10.100
> with(xmp01.14, mean(copper, trim = 0.1))
[1] 3.41
```

#### 1.8 Example 1.15

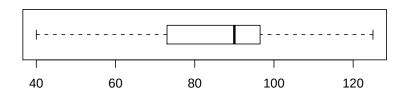
Functions var and sd provide the sample variance and sample standard deviation, respectively. The "computing formula" described on p. 34 is not used by these functions because that formula can have poor numerical properties.

```
> with(xmp01.15, var(Strength))
[1] 1.193580
> with(xmp01.15, sd(Strength))
[1] 1.092511
```

#### 1.9 Examples 1.17

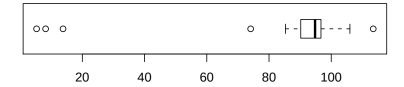
Function boxplot provides the boxplot. By default a vertical boxplot is constructed. Use the optional argument horizontal=TRUE to get a horizontal boxplot

```
> with(xmp01.17, boxplot(depth, horizontal = TRUE))
```



#### 1.10 Example 1.18

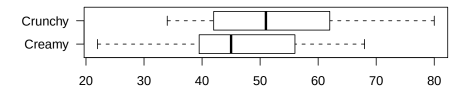
> with(xmp01.18, boxplot(C1, horizontal = TRUE))



#### 1.11 Comparative boxplots

The data for Example 1.19 (p. 38) are not available so we use the data on scores for creamy and crunchy peanut butters (exercise 1.15, p. 21) to illustrate comparative boxplots. This data set has two columns and 37 rows. The score is the first column and the indicator of "Creamy" or "Crunchy" is the second column. We say that these data are in the stacked format (see Appendix A.10 for details). In the call to boxplot we use the formula to indicate that Score is the response and Type defines the groups for the comparative boxplot.

> with(ex01.15, boxplot(Score ~ Type, horizontal = TRUE, las = 1))



#### 1.12 Enhancing graphical displays

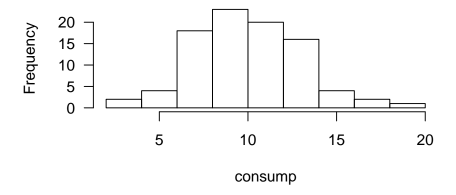
In this chapter we have used several functions, such as hist, barplot, plot, and boxplot that produce graphical displays of data. These functions all can

take optional arguments that provide more effective displays. For example, in § 1.11, we used the optional argument las=1 to boxplot to change the label style on the vertical axis so the labels are horizontal rather than vertical.

The las=1 argument can be used with other graphics functions. Compare

> with(xmp01.09, hist(consump, las = 1))

#### Histogram of consump



with the display in  $\S 1.4$ 

We have also seen how the breaks argument can be used with hist and how the scale argument can be used with stem. To reproduce a display like Figure 1.1 (p. 4) we would use

> with(xmp01.01, stem(temp, scale = 2))

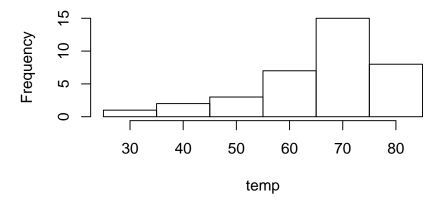
The decimal point is 1 digit(s) to the right of the |

- 3 | 1 3 | 4 | 0 4 | 59 5 | 23 5 | 788 6 | 0113 6 | 6777789
- 7 | 000023
- 7 | 556689
- 8 | 0134

and

> with(xmp01.01, hist(temp, breaks = c(25, 35, 45, 55, 65, 75, 85)))

## Histogram of temp



# Probability

Table 2.1 lists a function used in chapter 2.

Function	Description	
choose(n,k)	calculate $\binom{n}{k}$	

Table 2.1: R functions used in chapter 2

#### 2.1 Example 2.23

In this chapter on probability there is little use for R functions except for the **choose** function that evaluates the number of combinations of k objects selected from n, written  $\binom{n}{k}$  and described in section 2.3 (pp. 64–65). To calculate the first probability in example 2.23

```
> choose(15, 3) * choose(10, 3)/choose(25, 6)
```

[1] 0.3083004

# Discrete Random Variables and Probability Distributions

To quote the document "Introduction to R"

One convenient use of R is to provide a comprehensive set of statistical tables. Functions are provided to evaluate the cumulative distribution function  $P(X \le x)$ , the probability density function and the quantile function (given q, the smallest x such that  $P(X \le x) > q$ ), and to simulate from the distribution.

Distribution	R name	additional arguments
binomial	'binom'	'size, prob'
geometric	'geom'	'prob'
hypergeometric	'hyper'	'm, n, k'
negative binomial	'nbinom'	'size, prob'
Poisson	'pois'	ʻlambda'

Prefix the name given here by 'd' for the density, 'p' for the CDF, 'q' for the quantile function and 'r' for simulation (random deviates). The first argument is 'x' for 'dXXX', 'q' for 'pXXX', 'p' for 'qXXX' and 'n' for 'rXXX' (except for 'rhyper' and 'rwilcox', for which it is 'nn').

These functions are more versatile and more accurate than using probability tables.

#### 3.1 Example 3.31

```
For X having a binomial distribution with n=6 and p=0.5 we are to calculate P(X=3), P(3 \le X), and P(X \le 1).
```

```
> dbinom(3, size = 6, prob = 0.5)
[1] 0.3125
> dbinom(3:6, size = 6, prob = 0.5)
[1] 0.312500 0.234375 0.093750 0.015625
> sum(dbinom(3:6, size = 6, prob = 0.5))
[1] 0.65625
> 1 - pbinom(2, size = 6, prob = 0.5)
[1] 0.65625
> pbinom(2, size = 6, prob = 0.5, lower = FALSE)
[1] 0.65625
> pbinom(1, 6, 0.5)
```

The first call evaluates b(3;6,.5). The second call evaluates the probability function at  $3,\ldots,6$  using the ":" operator that generates the sequence from 3 to 6. If we sum this vector of probabilities we get  $P(3 \le X)$ . An alternative is to use  $P(3 \le X) = 1 - P(X \le 2)$  and evaluate  $P(X \le 2)$  with pbinom. Another alternative is to use cumulative probability in the upper tail, obtained with the optional argument lower=FALSE to pbinom. Finally pbinom is used to calculate  $P(X \le 1)$ .

#### 3.2 Example 3.32

[1] 0.109375

```
> pbinom(8, 15, 0.2)
[1] 0.999215
> dbinom(8, 15, 0.2)
[1] 0.003454764
> 1 - pbinom(7, 15, 0.2)
[1] 0.00423975
> sum(dbinom(4:7, 15, 0.2))
[1] 0.3475981
```

#### 3.3 Example 3.35

R uses a different, but equivalent, set of parameters for the hypergeometric distribution than does the text. In the text the parameters of the hypergeometric are N, the population size, n, the sample size, and M, the number of "successes" in the population. In R the sample size is called k, the parameter m corresponds to M in the text, and n is N-M.

Thus what is written in the text as h(2; 5, 12, 20) becomes

```
> dhyper(2, 12, 8, 5)
```

[1] 0.2383901

#### 3.4 Example 3.36

```
> dhyper(2, 5, 20, 10)
```

[1] 0.3853755

> phyper(2, 5, 20, 10)

[1] 0.6988142

#### 3.5 Example 3.38

The negative binomial density function, dnbinom, shown in the text as nb(10; 5, 2), has essentially the same calling sequence in R. The cumulative probability function is pnbinom.

```
> dnbinom(10, 5, 0.2)
```

[1] 0.0343941

> pnbinom(10, 5, 0.2)

[1] 0.1642337

#### 3.6 Example 3.39

```
> dpois(5, lambda = 4.5)
```

[1] 0.1708269

> ppois(5, 4.5)

[1] 0.7029304

### 3.7 Example 3.40

The Poisson distribution can be used to approximate binomial probabilities with large n and small p. However, there is no need to do so because the exact binomial probabilities can be evaluated.

```
> dbinom(1, 400, 0.005)
```

- [1] 0.2706694
- > dpois(1, 2)
- [1] 0.2706706
- > pbinom(3, 400, 0.005)
- [1] 0.8575767
- > ppois(3, 2)
- [1] 0.8571235

# Continuous Random Variables and Probability Distributions

The set of continous distributions available in R is

Distribution	R name	additional arguments
beta	'beta'	'shape1, shape2, ncp'
Cauchy	'cauchy'	'location, scale'
$\chi^2$	'chisq'	'df, ncp'
exponential	'exp'	'rate'
$\mathbf{F}$	'f'	'df1, df1, ncp'
gamma	'gamma'	'shape, scale'
log-normal	'lnorm'	'meanlog, sdlog'
logistic	'logis'	'location, scale'
normal	'norm'	'mean, sd'
Student's t	't'	'df, ncp'
uniform	'unif'	'min, max'
Weibull	'weibull'	'shape, scale'
Wilcoxon	'wilcox'	'm, n'

As with the discrete distributions, prefix the name given here by 'd' for the density, 'p' for the CDF, 'q' for the quantile function and 'r' for simulation (random deviates). The first argument is 'x' for 'dXXX', 'q' for 'pXXX', 'p' for 'qXXX' and 'n' for 'rXXX'

Not all the distributions shown above are discussed in the text.

#### 4.1 Example 4.13

Function pnorm provides the standard normal cumulative distribution function by default. The optional arguments mean and sd can be set to values other than

```
0 and 1 to provide probabilities from any normal distribution. P(Z \leq 1.25) \text{ and } P(Z \leq -1.25) \text{ are evaluated as} > pnorm(1.25) [1] 0.8943502 > pnorm(-1.25) [1] 0.1056498 P(Z > 1.25) \text{ can be evaluated in two ways} > 1 - pnorm(1.25) [1] 0.1056498
```

To evaluate probabilities of intervals, such as  $P(-0.38 \le Z \le 1.25)$ , apply pnorm to the endpoints as a vector (created with the "c" function) which returns a vector of probabilities. The "diff" function forms successive differences from which we obtain the probability in the interval.

```
> pnorm(c(-0.38, 1.25))
[1] 0.3519727 0.8943502
> diff(pnorm(c(-0.38, 1.25)))
[1] 0.5423775
```

> pnorm(1.25, lower = FALSE)

[1] 0.1056498

#### 4.2 Example 4.14

The inverse of the standard normal CDF, called the quantile function, is obtained with qnorm. Notice that the first argument to qnorm is a probability, not a percentage.

```
> qnorm(0.99)
[1] 2.326348
```

#### 4.3 Example 4.15

To obtain  $z_{\alpha}$ , use the optional argument lower=FALSE to qnorm.

```
> qnorm(0.05, lower = FALSE)
```

[1] 1.644854

#### 4.4 Example 4.16

Nonstandard normal distribution probabilities or quantiles are obtained with the optional arguments mean and sd to pnorm and qnorm. In this example  $\mu = 1.25$  and  $\sigma = 0.46$  and we wish to evaluate  $P(1.00 \le X \le 1.75)$ 

```
> diff(pnorm(c(1, 1.75), mean = 1.25, sd = 0.46))
[1] 0.5680717
```

#### 4.5 Example 4.18

```
For \mu=64 and \sigma=0.78, the 99.5th percentile is > qnorm(0.995, mean=64, sd=0.78) [1] 66.00915
```

#### 4.6 Example 4.20

The normal approximation to binomial probabilities can be calculated but, like the Poisson approximation, it is not necessary as the exact binomial probabilities can also be calculated.

```
> pnorm(10.5, mean = 12.5, sd = sqrt(12.5 * 0.75))
[1] 0.2568146
> pbinom(10, size = 50, prob = 0.25)
[1] 0.2622023
> diff(pnorm(c(4.5, 15.5), mean = 12.5, sd = sqrt(12.5 * 0.75)))
[1] 0.8319162
> diff(pbinom(c(4, 15), size = 50, prob = 0.25))
[1] 0.8348084
```

#### 4.7 Example 4.23

The parameters  $\alpha$  and  $\beta$  of the gamma distribution are named shape and scale respectively in R. In this example  $\alpha=2$  and  $\beta$  has the default value of 1.

```
> pgamma(c(3, 5), shape = 2)
[1] 0.8008517 0.9595723
```

```
> diff(pgamma(c(3, 5), shape = 2))
[1] 0.1587206
> pgamma(4, shape = 2, lower = FALSE)
```

#### 4.8 Example 4.24

[1] 0.0915782

```
> diff(pgamma(c(60, 120), shape = 8, scale = 15))
[1] 0.4959056
> pgamma(30, shape = 8, scale = 15, lower = FALSE)
[1] 0.9989033
```

#### 4.9 Examples 4.21 and 4.22

In R the parameter  $\lambda$  of the exponential distribution is called rate

```
> pexp(10, rate = 0.2)
[1] 0.8646647
> diff(pexp(c(5, 10), rate = 0.2))
[1] 0.2325442
> pexp(2, rate = 0.5, lower = FALSE)
[1] 0.3678794
```

#### 4.10 Example 4.25

The parameters  $\alpha$  and  $\beta$  of the Weibull distribution are called shape and scale.

```
> pweibull(10, shape = 2, scale = 10)
[1] 0.6321206
> qweibull(0.95, shape = 2, scale = 10)
[1] 17.30818
```

#### 4.11 Example 4.27

The lognormal distribution takes two parameters named meanlog and sdlog.

```
> diff(plnorm(c(1, 2), meanlog = 0.375, sdlog = 0.25))
[1] 0.8316108
> qlnorm(0.99, meanlog = 0.375, sdlog = 0.25)
[1] 2.602798
```

#### 4.12 Example 4.28

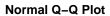
R provides probabilities and quantiles of the standard beta distribution with A=0 and B=1. The parameters  $\alpha$  and  $\beta$  are called shape1 and shape2 respectively. To use other values of A and B the scaling must be done manually. In this example  $A=2, B=5, \alpha=2$  and  $\beta=3$ . To transform to a standard beta distribution we replace x by (x-2)/(5-2)

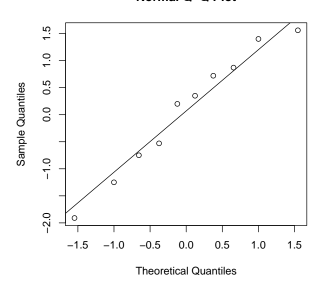
```
> pbeta((3 - 2)/(5 - 2), shape1 = 2, shape2 = 3)
[1] 0.4074074
```

#### 4.13 Examples 4.29 and 4.30

The normal probability plot is produced with qqnorm. A reference line can be added with qqline.

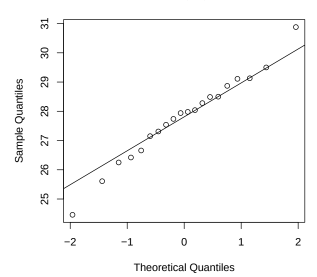
```
> with(xmp04.29, qqnorm(meas.err))
> with(xmp04.29, qqline(meas.err))
```





- > with(xmp04.30, qqnorm(Voltage))
  > with(xmp04.30, qqline(Voltage))

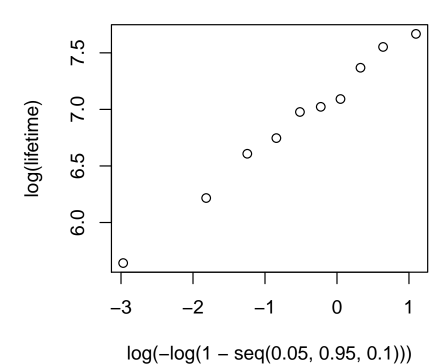
#### Normal Q-Q Plot



#### 4.14 Example 4.31

The Weibull probability plot is not available directly in R. However, the plot can be created using the formula  $\ln[-\ln(1-p)]$  for the 5th, 15th,..., and 95th percentiles as given in text. The sequence  $0.05, 0.15, \ldots, 0.95$  is generated with seq(0.05, 0.95, 0.1).

```
> with(xmp04.31, plot(log(-log(1 - seq(0.05, 0.95, 0.1))),
+ log(lifetime)))
```



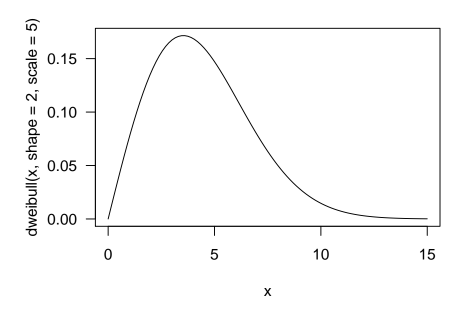
# Joint Probability Distributions and Random Samples

The main use of R in this chapter is for simulation experiments as described in section 5.3.

#### 5.1 Example 5.22

In this example six samples of size ten are drawn from a Weibull distribution with  $\alpha=2$  and  $\beta=5$ . To reproduce the plot of the density shown in Figure 5.6 we can use

> curve(dweibull(x, shape = 2, scale = 5), 0, 15, las = 1)



To get a single sample of size 10 from this Weibull distribution we use

```
> rweibull(10, shape = 2, scale = 5)
```

- [1] 7.14763 8.69608 3.16509 4.45848 0.60844 1.80645 6.17320 0.76923
- [9] 5.79083 4.47578

We could store such a sample as, say, samp, then evaluate its sample mean, sample median, and sample standard deviation.

```
> samp = rweibull(10, shape = 2, scale = 5)
> print(samp)
```

[1] 6.8219 2.0103 4.0432 1.2701 8.0921 4.9623 2.1623 4.1143 6.5781 8.0743

> mean(samp)

[1] 4.8129

> median(samp)

[1] 4.5383

> sd(samp)

[1] 2.5183

Notice that every time rweibull is called a new sample is generated.

This process of generating a sample and evaluating selected statistics on the sample could be repeated manually to get a total of 6 samples. For large simulation experiments this would quickly become tedious so we put these calculations in a loop.

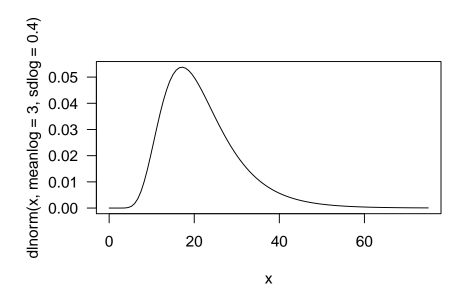
```
> means = medians = sds = numeric(6)
> for (i in 1:6) {
      samp = rweibull(10, shape = 2, scale = 5)
      print(samp)
      means[i] = mean(samp)
      medians[i] = median(samp)
      sds[i] = sd(samp)
+ }
 [1] 7.9598 3.0927 1.9392 2.8575 2.6656 6.7229 3.1295 4.0361 2.3052 7.3707
 [1] 0.97671 4.04935 4.37314 1.51435 10.54473 2.00505 5.51033
 [8] 3.39707 2.81787 1.36754
 [1] 1.8795 5.9659 2.0660 7.8419 1.1869 3.3403 5.8668 1.3481 4.3557 4.4121
 [1] 1.0304 4.1191 7.2787 4.2392 5.6721 2.1287 4.9993 4.6950 4.0448 5.8411
 [1] 3.2417 2.4885 3.4364 3.7212 3.3539 3.9331 2.9631 3.3034 7.6727 8.5520
 [1] 2.5096 7.7263 3.5716 3.7317 4.9273 6.3307 5.3553 1.0324 4.2311 3.2241
> means
[1] 4.2079 3.6556 3.8263 4.4048 4.2666 4.2640
> medians
[1] 3.1111 3.1075 3.8480 4.4671 3.3951 3.9814
> sds
[1] 2.2561 2.8296 2.2539 1.8004 2.0746 1.9223
```

In the first line we assign the names means, medians, and sds to numeric vectors of length 6. Within the loop we assign individual elements in these vectors.

## 5.2 Normal data, like example 5.23

In this example 500 samples of size n=5 are generated from a normal distribution with  $\mu=8.25$  and  $\sigma=0.75$  and the mean of each sample is calculated. We could do this in a loop, as shown above. However, it is more compact to generate a matrix with 5 rows and 500 columns then generate a histogram of the means of the columns of this matrix. Function colMeans calculates the means of the columns. Function matrix creates a matrix from a numeric vector. The user can specify the number of rows and the number of columns. If one of these is omitted, it is calculated from the length of the vector and the other dimension. We set the graphics parameter "mfrow" (multiple figures by row) to create a 2 by 2 array of plots.

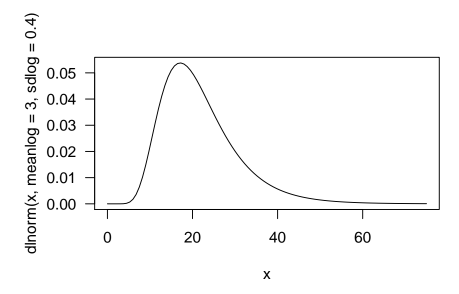
```
> par(mfrow = c(2, 2))
> samp5 = matrix(rnorm(500 * 5, mean = 8.25, sd = 0.75), ncol = 500)
> hist(colMeans(samp5), main = "Samples of size 5")
> samp10 = matrix(rnorm(500 * 10, mean = 8.25, sd = 0.75),
+ ncol = 500)
> hist(colMeans(samp10), main = "Samples of size 10")
> samp20 = matrix(rnorm(500 * 20, mean = 8.25, sd = 0.75),
+ ncol = 500)
> hist(colMeans(samp20), main = "Samples of size 20")
> samp30 = matrix(rnorm(500 * 30, mean = 8.25, sd = 0.75),
+ ncol = 500)
> hist(colMeans(samp30), main = "Samples of size 30")
```



## 5.3 Example 5.23

This example is similar to Example 5.22. To reproduce Figure 5.12 use

```
> curve(dlnorm(x, meanlog = 3, sdlog = 0.4), from = 0, to = 75,
+ las = 1)
```

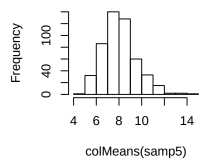


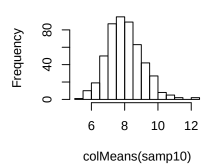
The samples and the histograms of the means are generated from

```
> par(mfrow = c(2, 2))
> samp5 = matrix(rlnorm(500 * 5, 2, 0.4), ncol = 500)
> hist(colMeans(samp5), main = "Means of samples of size 5")
> samp10 = matrix(rlnorm(500 * 10, 2, 0.4), ncol = 500)
> hist(colMeans(samp10), main = "Means of samples of size 10")
> samp20 = matrix(rlnorm(500 * 20, 2, 0.4), ncol = 500)
> hist(colMeans(samp20), main = "Means of samples of size 20")
> samp30 = matrix(rlnorm(500 * 30, 2, 0.4), ncol = 500)
> hist(colMeans(samp30), main = "Means of samples of size 30")
```

## Means of samples of size 5 Means of

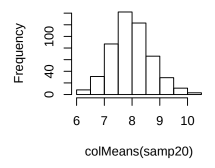
## Means of samples of size 10

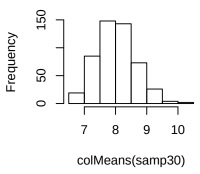




## Means of samples of size 20

## Means of samples of size 30



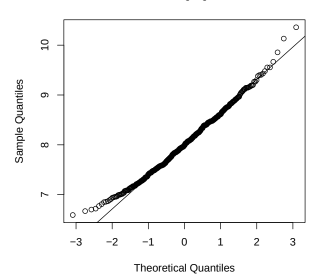


Finally the normal probability plot is generated by

<sup>&</sup>gt; qqnorm(colMeans(samp30))

<sup>&</sup>gt; qqline(colMeans(samp30))

## Normal Q-Q Plot



# Chapter 6

# **Point Estimation**

## 6.1 Example 6.2

The various estimators of location described in Example 6.2 (p. 229) can be evaluated as

```
> with(xmp06.02, mean(Voltage))
[1] 27.793
> with(xmp06.02, median(Voltage))
[1] 27.96
> with(xmp06.02, mean(range(Voltage)))
[1] 27.67
> with(xmp06.02, mean(Voltage, trim = 0.1))
[1] 27.83812
```

## **6.2** Example **6.3**

Functions  ${\tt var}$  and  ${\tt sd}$  provide  $s^2$  and s, the sample variance and standard deviation, respectively.

```
> with(xmp06.03, var(Strength))
[1] 0.25125
> with(xmp06.03, sd(Strength))
[1] 0.5012484
```

To evaluate the alternative estimator  $\hat{\sigma} = \frac{\sum (X_i - \bar{X})}{n}$  we must evaluate the formula

```
> with(xmp06.03, sum((Strength - mean(Strength))^2)/length(Strength))
[1] 0.2198438
```

Function length applied to a vector returns n, the number of elements in the vector.

## 6.3 Example 6.13

The calculation of the method of moments estimates in Example 6.13 (p. 244) can be split into stages

```
> xbar = with(xmp06.13, mean(Survival))
> xsqb = with(xmp06.13, mean(Survival^2))
> xbar
[1] 113.45
> xsqb
[1] 14087.75
> xbar^2/(xsqb - xbar^2)
[1] 10.57725
> (xsqb - xbar^2)/xbar
[1] 10.72585
```

These estimates are slightly different from those shown in the text because the intermediate results  $\bar{x}$  and  $\sum x_i^2/n$  were rounded in the text.

Maximum likelihood estimates are discussed later in chapter 6. We can evaluate the maximum likelihood estimates of  $\alpha$  and  $\beta$  for this example using the function fitdistr from the package MASS that supplements the book "Modern Applied Statistics with S (4th ed)" by Bill Venables and Brian Ripley (Springer, 2002). These estimates are determined by numerical optimization of the logarithm of the likelihood function and we must supply starting estimates. We use the method of moments estimates for this.

The MLEs are quite different from the method of moments estimates. The numbers in parentheses under the estimates are their standard errors.

# Chapter 7

# Statistical Intervals Based on a Single Sample

Table 7.1 lists functions used in chapters 7 and 8.

Function	Description
t.test(x)	Student's t test and confidence interval
prop.test(x,n)	Test and confidence interval on proportion
binom.test(x,n)	Test and confidence interval on proportion

Table 7.1: R functions used with chapters 7 and 8

## 7.1 Example 7.2

The calculation of the confidence interval for known  $\sigma$ , shown in Example 7.2, could be done in R as

```
> 80 + c(-1, 1) * 1.96 * 2/sqrt(31)
```

[1] 79.29595 80.70405

but it is probably just as easy to use a hand calculator for this.

## 7.2 Example 7.6

The calculation of the sample mean, the sample standard deviation, and the sample size can be combined into a single statement

```
> with(xmp07.06, mean(Voltage) + c(-1, 1) * 1.96 * sd(Voltage)/sqrt(length(Voltage)))
```

[1] 53.22857 56.18810

An alternative, and preferred way, of calculating the interval is to use t.test. With 48 observations the confidence interval on  $\mu$  from the t distribution is nearly identical to that from the standard normal distribution.

```
One Sample t-test

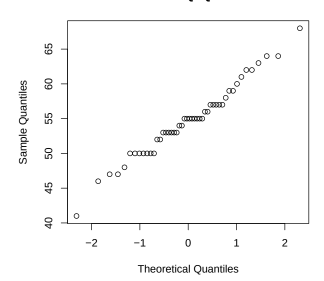
data: Voltage
t = 72.4631, df = 47, p-value < 2.2e-16
alternative hypothesis: true mean is not equal to 0
95 percent confidence interval:
53.18950 56.22716
sample estimates:
mean of x
54.70833
```

It is always a good idea to check the normal probability plot when using t.test, even with a large sample.

> with(xmp07.06, qqnorm(Voltage))

> with(xmp07.06, t.test(Voltage))

#### Normal Q-Q Plot



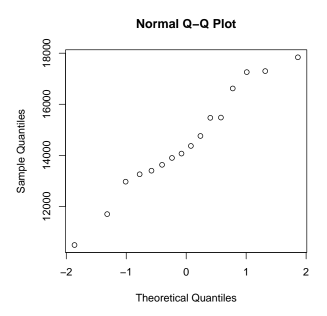
## 7.3 Example 7.8

R has two functions, binom.test and prop.test, that can be used to calculate a large-sample confidence interval on the binomial proportion, p. Neither of them corresponds exactly the the confidence interval described on p. 295.

```
> prop.test(16, 48)
        1-sample proportions test with continuity correction
data: 16 out of 48, null probability 0.5
X-squared = 4.6875, df = 1, p-value = 0.03038
alternative hypothesis: true p is not equal to 0.5
95 percent confidence interval:
0.2080794 0.4851357
sample estimates:
0.3333333
> binom.test(16, 48)
       Exact binomial test
data: 16 and 48
number of successes = 16, number of trials = 48, p-value = 0.02930
alternative hypothesis: true probability of success is not equal to 0.5
95 percent confidence interval:
0.2039597 0.4841083
sample estimates:
probability of success
```

## 7.4 Example 7.11

0.3333333

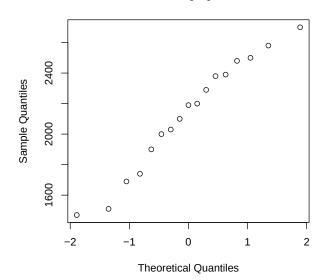


## 7.5 Example 7.15

Although there is no built-in confidence interval for  $\sigma^2$  in R, the qchisq function can be used to obtain the critical values  $\chi^2_{\alpha/2,n-1}$  and  $\chi^2_{1-\alpha/2,n-1}$  used to calculate the interval.

#### [1] 76171.31 318079.76

## Normal Q-Q Plot



# Chapter 8

# Tests of Hypotheses Based on a Single Sample

The functions described in chapter 7 are used for performing tests of hypotheses based on a single sample. Optional arguments are used to specify  $\mu_0$  or  $p_0$  and to indicate the form of the alternative hypothesis. All of these tests return a p-value, described in §8.4 (pp. 311–317). From the p-value the result of the hypothesis test for any level  $\alpha$  can be determined.

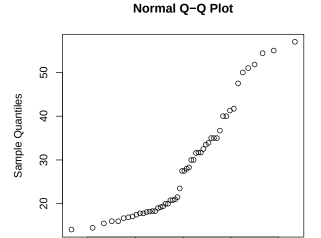
## 8.1 Example 8.8

In R the t.test function can be used with any size of data set. For large n the t test is essentially equivalent to the large-sample z test.

As the p-value of 0.2349 exceeds 0.05 we do not reject  $H_0$ :  $\mu = 30$  versus  $H_a$ :  $\mu < 30$  at level  $\alpha = 0.05$ .

Although not shown in the text book, it is of interest to examine the normal probability plot for these data

> with(xmp08.08, qqnorm(DCP))



This plot shows considerable skewness in the data. If we transform to the logarithm of the DCP measurement the skewness is diminished.

0

Theoretical Quantiles

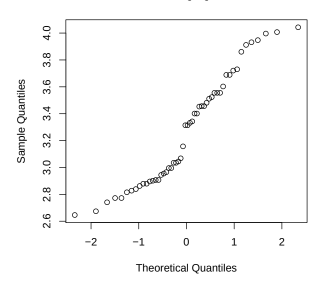
1

2

> with(xmp08.08, qqnorm(log(DCP)))

-2

### Normal Q-Q Plot



and in the logarithm scale, the hypothesis test  $H_0$ :  $\mu_{log} = \log(30)$  versus  $H_a$ :  $\mu_{log} < \log(30)$  is significant at level  $\alpha = 0.05$ .

> with(xmp08.08, t.test(log(DCP), mu = log(30), alt = "less"))

One Sample t-test

data: log(DCP)
t = -2.223, df = 51, p-value = 0.01534
alternative hypothesis: true mean is less than 3.401197
95 percent confidence interval:
 -Inf 3.370103
sample estimates:
mean of x
3.274999

## 8.2 Example 8.9

```
> with(xmp08.09, t.test(MAWL, mu = 25, alt = "greater"))
One Sample t-test
```

data: MAWL
t = 1.0382, df = 4, p-value = 0.1789
alternative hypothesis: true mean is greater than 25

```
95 percent confidence interval: 22.32433 Inf sample estimates: mean of x 27.54
```

## 8.3 Example 8.10

```
> power.t.test(n = 10, delta = 0.1, sd = 0.1, type = "one.sample",
      alt = "one.sided")
    One-sample t test power calculation
             n = 10
          delta = 0.1
            sd = 0.1
      sig.level = 0.05
         power = 0.897517
    alternative = one.sided
> power.t.test(delta = 0.1, sd = 0.1, power = 0.95, type = "one.sample",
     alt = "one.sided")
    One-sample t test power calculation
             n = 12.32052
         delta = 0.1
            sd = 0.1
      sig.level = 0.05
         power = 0.95
    alternative = one.sided
```

## 8.4 Example 8.11

The large-sample test for a population proportion is most closely related to the result of the prop.test function

```
sample estimates:
    p
0.3100851
```

The value labeled X-squared is the square of the z statistic. This version of the test uses a continuity correction. If you wish to reproduce the test statistic as given in the text book, add the optional argument correct=FALSE

In both cases the p-value is less than 0.1 so we reject  $H_0: p=0.3$  versus  $H_a: p>0.3$  at level  $\alpha=0.1$ .

## 8.5 Example 8.13

[1] 1.411737

```
The small sample test is provided by binom.test
```

> binom.test(x = 14, n = 20, p = 0.9, alt = "less")

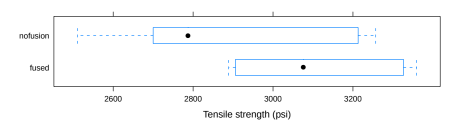
# Chapter 9

# Inference Based on Two Samples

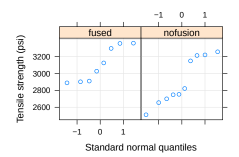
## 9.1 Example 9.7

When given vectors of numbers as arguments, the summary function returns the mean, min, max and quartiles of the given vector. The qqnorm function is used to generate a qqplot of its given argument. The simplest use of the boxplot function creates a boxplot for each vector argument passed to it.

```
> str(xmp09.07)
'data.frame': 18 obs. of 2 variables:
$ strength: int 2748 2700 2655 2822 2511 3149 3257 3213 3220 2753 ...
$ type : Factor w/ 2 levels "fused", "nofusion": 2 2 2 2 2 2 2 2 2 2 ...
> bwplot(type ~ strength, xmp09.07)
```



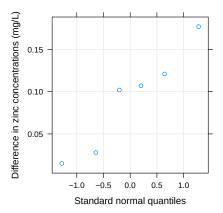
> qqmath(~strength | type, xmp09.07)



"p"), xlab = "Standard normal quantiles", aspect = 1,
ylab = "Difference in zinc concentrations (mg/L)"))

## 9.2 Example 9.8

The t.test function conducts various types of t tests. When given one vector argument, it performs a one sample test, when two vector arguments are given, and the paired option is specified as TRUE, the function can perform a paired two sample tests as well. Options can also present to specify if a test is one or two sided.



> with(xmp09.08, t.test(bottom, surface, paired = TRUE))

Paired t-test

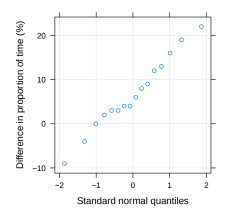
> print(qqmath(~Difference, xmp09.09, type = c("g", "p"), xlab = "Standard normal quantiles", + aspect = 1, ylab = "Difference in proportion of time (%)"))

## 9.3 Example 9.9

> str(xmp09.09)

```
'data.frame': 16 obs. of 4 variables:
$ Subject : int 1 2 3 4 5 6 7 8 9 10 ...
$ Before : int 81 87 86 82 90 86 96 73 74 75 ...
$ After : int 78 91 78 78 84 67 92 70 58 62 ...
$ Difference: int 3 -4 8 4 6 19 4 3 16 13 ...
```

> qqmath(~Difference, xmp09.09)



> with(xmp09.09, t.test(Before, After, paired = TRUE))

Paired t-test

```
data: Before and After
t = 3.2791, df = 15, p-value = 0.005072
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
    2.362371 11.137629
sample estimates:
mean of the differences
    6.75
```

## 9.4 Example 9.10

This can be done directly with the differences:

```
> Difference <- c(5, 19, 25, 10, 10, 10, 28, 46, 25, 38, 14,
+ 23, 14)
> qqnorm(Difference)
> qqline(Difference)
> t.test(Difference)

One Sample t-test

data: Difference
t = 6 1904 df = 12 n-value = 4 654e-05
```

t = 6.1904, df = 12, p-value = 4.654e-05
alternative hypothesis: true mean is not equal to 0
95 percent confidence interval:
 13.30957 27.76736
sample estimates:

mean of x 20.53846

Or, using the data frame with the paired=TRUE argument of t.test.

> with(xmp09.10, t.test(slide, digital, paired = TRUE))

Paired t-test

Chapter 10

The Analysis of Variance

## Chapter 11

# Multifactor Analysis of Variance

## 11.1 Example 11.01

An *interaction plot* shows the response versus the levels of one factor with the points joined according to the levels of the other factor. It an additive model is valid, the lines should be approximately parallel.

For a balanced data set, the order of the factors does not affect the calculations of the sums of squares nor any of the test statistics and conclusions. However, for unbalanced data the order of the factors is important. In general we put the blocking factor(s) first and the treatment factor(s) last.

The TukeyHSD function can be applied to any models with more than one factor. If we are only interested in selected factors we can use the argument which to restrict the factors being considered.

Plots are useful aids in checking if assumptions have been satisfied. To assess the constant variance assumption, residual plots are used and the normal probability plot of the residuals is used to assess the assumption of a normal distribution of the noise term. The simplest way to obtain such plots in R is to plot the fitted model object. Use which = 1 to get residuals versus fitted values, which = 2 to get the qqnorm plot of the residuals, or which = 1:2 to get both.

Before beginning, we coerce the variables brand and treatment to be factors.

```
col = 2:4, lty = 1)
> with(xmp11.01, interaction.plot(brand, treatment, strength,
      col = 2:4, lty = 1)
> anova(fm1 <- aov(strength ~ treatment + brand, xmp11.01))</pre>
Analysis of Variance Table
Response: strength
          Df Sum Sq Mean Sq F value
treatment 3 0.47969 0.15990 11.0549 0.007399
           2 0.12822 0.06411 4.4323 0.065765
Residuals 6 0.08678 0.01446
> TukeyHSD(fm1, which = "treatment")
 Tukey multiple comparisons of means
    95% family-wise confidence level
Fit: aov(formula = strength ~ treatment + brand, data = xmp11.01)
$treatment
           diff
                       lwr
                                   upr
                                           p adj
2-1 -0.46666667 -0.8065953 -0.12673802 0.0124339
3-1 -0.38000000 -0.7199286 -0.04007135 0.0315128
4-1 -0.50333333 -0.8432620 -0.16340469 0.0086330
3-2 0.08666667 -0.2532620 0.42659531 0.8141883
4-2 -0.03666667 -0.3765953 0.30326198 0.9806209
4-3 -0.12333333 -0.4632620 0.21659531 0.6185041
> plot(fm1, which = 1:2)
```

### 11.2 Example 11.05

Response: power

```
Df Sum Sq Mean Sq F value
                                      Pr(>F)
          3 116218
                     38739 278.199 2.364e-11
humid
          4 53231
                     13308 95.567 5.419e-09
brand
Residuals 12
              1671
                       139
> TukeyHSD(fm1, which = "brand")
 Tukey multiple comparisons of means
   95% family-wise confidence level
Fit: aov(formula = power ~ humid + brand, data = xmp11.05)
$brand
     diff
                 lwr
                           upr
                                   p adj
2-1 46.00 19.403556 72.59644 0.0010268
3-1 41.50 14.903556 68.09644 0.0024312
4-1 116.50 89.903556 143.09644 0.0000001
5-1 139.75 113.153556 166.34644 0.0000000
3-2 -4.50 -31.096444 22.09644 0.9812528
4-2 70.50 43.903556 97.09644 0.0000175
5-2 93.75 67.153556 120.34644 0.0000008
4-3 75.00 48.403556 101.59644 0.0000092
5-3 98.25 71.653556 124.84644 0.0000005
5-4 23.25 -3.346444 49.84644 0.0978028
> plot(TukeyHSD(fm1, which = "brand"))
> plot(TukeyHSD(fm1, which = "brand"))
11.3
        Example 11.06
> str(xmp11.06)
'data.frame':
                    24 obs. of 3 variables:
$ Resp : num 8 6.9 9.3 9.2 12 9.4 17.3 19.3 18.8 24.9 ...
$ Stimulus: Factor w/ 6 levels "L1", "L2", "T", ...: 1 2 3 4 5 6 1 2 3 4 ...
$ Subject : Factor w/ 4 levels "1","2","3","4": 1 1 1 1 1 1 2 2 2 2 ...
> anova(fm1 <- aov(Resp ~ Subject + Stimulus, data = xmp11.06))</pre>
Analysis of Variance Table
Response: Resp
         Df Sum Sq Mean Sq F value
                                       Pr(>F)
          3 13444.6 4481.5 86.1112 1.110e-09
Subject
Stimulus 5 1428.3
                      285.7 5.4888
                                      0.00455
Residuals 15 780.7
                       52.0
```

```
> with(xmp11.06, interaction.plot(Stimulus, Subject, Resp,
      col = 2:6, lty = 1))
> TukeyHSD(fm1, which = "Stimulus")
 Tukey multiple comparisons of means
    95% family-wise confidence level
Fit: aov(formula = Resp ~ Subject + Stimulus, data = xmp11.06)
$Stimulus
            diff
                         lwr
                                  upr
                                          p adj
L2-L1
           3.050 -13.5235020 19.62350 0.9895813
           4.275 -12.2985020 20.84850 0.9553756
L1+L2-L1 15.525 -1.0485020 32.09850 0.0727869
L1+T-L1 16.400 -0.1735020 32.97350 0.0532377
L2+T-L1 20.225 3.6514980 36.79850 0.0129117
T-L2
          1.225 -15.3485020 17.79850 0.9998651
L1+L2-L2 12.475 -4.0985020 29.04850 0.2019526
L1+T-L2 13.350 -3.2235020 29.92350 0.1528205
L2+T-L2 17.175 0.6014980 33.74850 0.0401628
L1+L2-T 11.250 -5.3235020 27.82350 0.2908899
L1+T-T
          12.125 -4.4485020 28.69850 0.2248769
L2+T-T
          15.950 -0.6235020 32.52350 0.0625789
L1+T-L1+L2 0.875 -15.6985020 17.44850 0.9999744
L2+T-L1+L2 4.700 -11.8735020 21.27350 0.9348445
L2+T-L1+T
           3.825 -12.7485020 20.39850 0.9719095
> plot(fm1, which = 1)
> range(xmp11.06$Resp)
[1] 6.9 96.6
> anova(fm2 <- aov(log(Resp) ~ Subject + Stimulus, data = xmp11.06))</pre>
Analysis of Variance Table
Response: log(Resp)
         Df Sum Sq Mean Sq F value
                                       Pr(>F)
Subject
          3 13.1315 4.3772 264.972 3.270e-13
Stimulus 5 1.1089 0.2218 13.425 4.331e-05
Residuals 15 0.2478 0.0165
> with(xmp11.06, interaction.plot(Stimulus, Subject, log(Resp),
      col = 2:6, lty = 1)
> plot(fm2, which = 1)
> opar <- par(pty = "s")
> plot(fm2, which = 2)
```

> par(opar)

```
> plot(TukeyHSD(fm2, which = "Stimulus"))
> with(xmp11.06, interaction.plot(Stimulus, Subject, fitted(fm2),
     col = 2:6, lty = 1)
        Example 11.07
> str(xmp11.07)
'data.frame':
                    36 obs. of 3 variables:
$ Yield : num 10.5 9.2 7.9 12.8 11.2 13.3 12.1 12.6 14 10.8 ...
$ Variety: Factor w/ 3 levels "1","2","3": 1 1 1 1 1 1 1 1 1 1 ...
$ Density: Factor w/ 4 levels "1","2","3","4": 1 1 1 2 2 2 3 3 3 4 ...
> xtabs(~Variety + Density, data = xmp11.07)
      Density
Variety 1 2 3 4
     1 3 3 3 3
     2 3 3 3 3
     3 3 3 3 3
> xtabs(Yield ~ Variety + Density, data = xmp11.07)
      Density
Variety
         1
               2
                    3
     1 27.6 37.3 38.7 32.4
     2 26.8 37.9 43.5 38.3
     3 48.9 54.3 59.8 54.5
> xtabs(Yield ~ Variety + Density, xmp11.07)/xtabs(~Variety +
     Density, xmp11.07)
      Density
Variety
                         2
                                   3
               1
     1 9.200000 12.433333 12.900000 10.800000
     2 8.933333 12.633333 14.500000 12.766667
     3 16.300000 18.100000 19.933333 18.166667
> anova(fm1 <- aov(Yield ~ Density * Variety, data = xmp11.07))</pre>
Analysis of Variance Table
Response: Yield
               Df Sum Sq Mean Sq F value
                                             Pr(>F)
Density
                3 86.69
                          28.90 18.2306 2.212e-06
Variety
                2 327.60 163.80 103.3430 1.608e-12
Density: Variety 6 8.03 1.34 0.8445
                                             0.5484
Residuals
               24 38.04
                            1.58
```

Variety

```
> with(xmp11.07, interaction.plot(Density, Variety, Yield,
      col = 2:6, lty = 1)
> anova(fm2 <- aov(Yield ~ Density + Variety, xmp11.07))</pre>
Analysis of Variance Table
Response: Yield
         Df Sum Sq Mean Sq F value
                                      Pr(>F)
Density
          3 86.69 28.90 18.816 4.690e-07
Variety
          2 327.60 163.80 106.659 2.313e-14
Residuals 30 46.07
                      1.54
> TukeyHSD(fm2)
  Tukey multiple comparisons of means
    95% family-wise confidence level
Fit: aov(formula = Yield ~ Density + Variety, data = xmp11.07)
$Density
          diff
                      lwr
                                 upr
                                         p adj
2-1 2.9111111 1.3226494 4.4995728 0.0001375
3-1 4.3000000 2.7115383 5.8884617 0.0000002
4-1 2.4333333 0.8448716 4.0217951 0.0013148
3-2 1.3888889 -0.1995728 2.9773506 0.1033936
4-2 -0.4777778 -2.0662395 1.1106839 0.8455586
4-3 -1.8666667 -3.4551284 -0.2782049 0.0163721
$Variety
                                     p adj
        diff
                   lwr
                             upr
2-1 0.875000 -0.3722268 2.122227 0.2109855
3-1 6.791667 5.5444399 8.038893 0.0000000
3-2 5.916667 4.6694399 7.163893 0.0000000
> model.tables(fm2, type = "means")
Tables of means
Grand mean
13.88889
 Density
Density
     1
            2
                   3
11.478 14.389 15.778 13.911
```

```
Variety
1 2 3
11.333 12.208 18.125
```

Coat:Strain

### 11.5 Example 11.10

```
> str(xmp11.10)
'data.frame':
                    96 obs. of 4 variables:
$ Tempr : num 3.6 3.8 3.9 3.4 3.7 3.9 2.9 2.8 2.7 2.5 ...
$ Period: int 1 1 1 1 1 1 1 1 1 ...
$ Strain: int 1 1 1 1 1 1 1 1 1 ...
$ Coat : int 1 1 1 2 2 2 3 3 3 4 ...
> xtabs(~Period + Coat + Strain, xmp11.10)
, , Strain = 1
     Coat
Period 1 2 3 4
    1 3 3 3 3
    2 3 3 3 3
    3 3 3 3 3
    4 3 3 3 3
, , Strain = 2
     Coat
Period 1 2 3 4
    1 3 3 3 3
    2 3 3 3 3
    3 3 3 3 3
    4 3 3 3 3
> anova(fm1 <- aov(Tempr ~ Period * Coat * Strain, xmp11.10))</pre>
Analysis of Variance Table
Response: Tempr
                  Df Sum Sq Mean Sq F value
                                               Pr(>F)
Period
                  1 0.257
                            0.257 2.0405
                                              0.15670
Coat
                   1 42.781 42.781 340.0850 < 2.2e-16
Strain
                   1 6.458
                             6.458 51.3409 2.265e-10
Period:Coat
                   1 0.531
                            0.531
                                    4.2214 0.04288
                   1 0.005 0.005 0.0373 0.84738
Period:Strain
```

1 0.006 0.006 0.0479 0.82733

```
Period:Coat:Strain 1 0.057
                              0.057
                                      0.4534
                                               0.50248
Residuals
                  88 11.070
                              0.126
> anova(fm2 <- update(fm1, . ~ . - Period:Strain - Period:Coat:Strain))</pre>
Analysis of Variance Table
Response: Tempr
           Df Sum Sq Mean Sq F value
                                         Pr(>F)
Period
            1 0.257
                       0.257
                               2.0753
                                        0.15317
            1 42.781 42.781 345.8856 < 2.2e-16
Coat
Strain
            1 6.458
                      6.458 52.2166 1.554e-10
Period:Coat 1 0.531
                      0.531
                               4.2935
                                        0.04112
Coat:Strain 1 0.006
                      0.006 0.0487
                                        0.82588
Residuals 90 11.132
                      0.124
        Example 11.11
11.6
> str(xmp11.11)
                    36 obs. of 4 variables:
'data.frame':
 $ abrasion: num 7.38 5.39 5.03 5.5 5.01 6.79 7.15 8.16 4.96 5.78 ...
       : Ord.factor w/ 6 levels "1"<"2"<"3"<"4"<..: 1 1 1 1 1 2 2 2 2 ...
 $ column : Ord.factor w/ 6 levels "1"<"2"<"3"<"4"<..: 1 2 3 4 5 6 1 2 3 4 ...
 $ humidity: Factor w/ 6 levels "25 percent","37 percent",..: 3 4 6 2 5 1 2 1 5 4 ...
> xtabs(as.integer(humidity) ~ row + column, xmp11.11)
   column
row 1 2 3 4 5 6
 1 3 4 6 2 5 1
 2 2 1 5 4 3 6
 3 4 6 3 5 1 2
 4 1 3 2 6 4 5
 5 6 5 1 3 2 4
 6 5 2 4 1 6 3
> xtabs(abrasion ~ row + column, xmp11.11)
   column
row 1
           2
                3
                     4
                          5
 1 7.38 5.39 5.03 5.50 5.01 6.79
 2 7.15 8.16 4.96 5.78 6.24 5.06
 3 6.75 5.64 6.34 5.31 7.81 8.05
 4 8.05 6.45 6.31 5.46 6.05 5.51
 5 5.65 5.44 7.27 6.54 7.03 5.96
 6 6.00 6.55 5.93 8.02 5.80 6.61
```

```
> anova(fm1 <- aov(abrasion ~ row + column + humidity, xmp11.11))</pre>
Analysis of Variance Table
Response: abrasion
          Df Sum Sq Mean Sq F value
                                       Pr(>F)
          5 2.1897 0.4379 2.5106 0.06407
row
          5 2.5743 0.5149 2.9516 0.03731
column
humidity 5 23.5301 4.7060 26.9789 3.03e-08
Residuals 20 3.4887 0.1744
> model.tables(fm1, cterms = "humidity", type = "mean")
Tables of means
Grand mean
6.305
humidity
humidity
25 percent 37 percent 50 percent 62 percent 75 percent 87 percent
                                     5.977
     7.683
                6.765
                          6.593
                                                 5.372
> TukeyHSD(fm1, which = "humidity")
  Tukey multiple comparisons of means
    95% family-wise confidence level
Fit: aov(formula = abrasion ~ row + column + humidity, data = xmp11.11)
$humidity
                             diff
                                         lwr
37 percent-25 percent -0.91833333 -1.6762719 -0.16039478 0.0121211
50 percent -25 percent -1.09000000 -1.8479386 -0.33206145 0.0024825
62 percent-25 percent -1.70666667 -2.4646052 -0.94872811 0.0000097
75 percent-25 percent -2.31166667 -3.0696052 -1.55372811 0.0000001
87 percent-25 percent -2.24333333 -3.0012719 -1.48539478 0.0000001
50 percent-37 percent -0.17166667 -0.9296052 0.58627189 0.9782473
62 percent-37 percent -0.78833333 -1.5462719 -0.03039478 0.0385446
75 percent-37 percent -1.39333333 -2.1512719 -0.63539478 0.0001513
87 percent-37 percent -1.32500000 -2.0829386 -0.56706145 0.0002818
62 percent-50 percent -0.61666667 -1.3746052 0.14127189 0.1545364
75 percent-50 percent -1.22166667 -1.9796052 -0.46372811 0.0007299
87 percent-50 percent -1.15333333 -1.9112719 -0.39539478 0.0013767
75 percent-62 percent -0.60500000 -1.3629386 0.15293855 0.1683501
87 percent-62 percent -0.53666667 -1.2946052 0.22127189 0.2697646
87 percent-75 percent 0.06833333 -0.6896052 0.82627189 0.9997113
```

# Appendix A

# Installing R and the Devore7 package

### A.1 What is R?

R is a freely available, open source, computer system for statistical analysis and graphics. It can be downloaded from the main R information site http://www.r-project.org, from the Comprehensive R Archive Network (CRAN) site http://cran.r-project.org, or from any of the mirrors of that site. Those in the United States, for example, are encouraged to use the U.S. mirror http://cran.us.r-project.org.

R provides facilities for data input and manipulation, for graphical and numerical summaries, for simulation and exploration of probability distributions, and for statistical analysis of data. It can be used for computing support for essentially all the topics in an introductory statistics course. This document describes how to use R for computing support in a course that uses the text *Probability and Statistics for Engineering and the Sciences (7th edition)* by Jay Devore (Duxbury, 2008).

Althought there are some limited graphical user interface (GUI) capabilites for R, it is basically a command-line system. <sup>1</sup> We will concentrate on the command-line interface, showing what the user types and what R responds. We will refer to what the user types as a "command" although, technically, we should use the term "function call".

## A.2 Obtaining and Installing R

Binary versions of R are available for various operating systems including Microsoft Windows (Windows 95 or later), the Macintosh (OS X), and several

<sup>&</sup>lt;sup>1</sup>Some GUI capabilities are provided the add-on packages Rcmdr (http://socserv.mcmaster.ca/jfox/Misc/Rcmdr/) and pmg (http://www.math.csi.cuny.edu/pmg).

Linux distributions. Complete source code for R is also available on the archives but it is quite unlikely that you will need to compile R for use with an introductory Statistics course.

As most students use R with Windows we will provide more detailed installation instructions for this operating system.

The current release of R is 2.4.1. For WIndows there is a binary installer file for R. It is approximately 30MB in size and can be found at http://cran.r-project.org/bin/windows/base/. If you have a fast network connection you should download and execute this file to install R. Without a network connection or with a slow network connection you will need to make other arrangements for obtaining a copy of this file.

The installation should provide a desktop icon or menu item for R. Use one of these to start R. The program should display a welcome message and a prompt "> ". At this point you could use it as a calculator. Try, for example,

> 2 + 2

[1] 4

## A.3 Quitting R

To quit from R you can either select File -> Exit from the menu bar or type

> q()

at the prompt, as indicated in the startup message. It is necessary to type the parentheses. That is, typing  ${\tt q}$  by itself is not sufficient.

Both of these methods will bring up a confirmation panel asking if you want to save the worksheet. In most cases you will not need to save the worksheet.

## A.4 Using data sets

A standard R installation provides several data sets that are used to demonstrate different techniques. The data command provides a list of these

> data()

You can obtain a description of a data set with the help command or with the short form for help which is ? followed by the name. Try, for example,

```
> help(pressure)
> `?`(pressure)
```

#### A.5 What is Devore7?

Notice that the description of the available data sets groups them into "packages" such as the "base" package, the "modreg" package, etc. Packages are groups of functions and data sets that extend the capabilities of R for specific purposes. The "Devore7" package provides the data sets for the 7th edition of Devore's engineering statistics text book. By installing and attaching this package you will be able to use the data sets from the examples and exercises in this text book without having to enter the data by hand.

The Devore7 package also provides a "vignette" - a document that describe particular aspects of the use of R. This document is one the vignette from the Devore7 package.

## A.6 Installing and attaching Devore7

Installing and attaching a package are two different operations. Installation involves downloading the package from a web site and installing the files on the local hard drive. It only needs to be done once. A package that has been installed can be attached to an R session after which the data sets will be available in the session.

To install the Devore7 package on a computer with access to the internet, either use the command

#### > install.packages("Devore7")

or select Packages -> Install package(s) from CRAN -> Devore7 from the menu bar.

If you do not have access to the Internet you will need to obtain a copy of the zip file whose name begins with Devore7 in the proper sub-directory http://cran.r-project.org/bin/windows/contrib/. (The exact name of the file changes as the package is updated but it will always begin with Devore7 and end with .zip.) Use the menu selection Packages -> Install package(s) from local zip files to install the package from the zip file.

We emphasize that it is only necessary to do the installation once.

To attach the package to an R session use

#### > library(Devore7)

after starting R or select Packages -> Load package -> Devore7 from the menu bar.

You must attach the package every time you start R if you are to have access to the data sets from the textbook.

### A.7 Names of the data sets

Data sets for exercises are named excc.nn where cc is two-digit chapter number and nn is the two-digit exercise number. Thus the data for exercise 27 in chapter

10 (p. 394) is called ex10.27. To provide the correct order when sorting the data set names, single-digit chapter or exercise numbers have a zero prepended. The data for exercise 1 in chapter 6 (p. 240) is called ex06.01.

Data sets for examples in the text are named xmpcc.nn.

A listing of all the data sets in the package can be obtained with

```
> data(package = "Devore7")
```

### A.8 Data sets as tables

All the data sets in the Devore7 package are in a tabular form called a *data frame* in R. Rows correspond to observations and columns correspond to "variables". We use the tabular form even when there is only one variable.

The columns have names, usually reflecting the description of the data from the exercise or the example, although names like C1 also occur frequently. (That name happens to be the default name of the first column assigned by another statistical computing system called Minitab.)

You can check the names and types of data in the data frame with str, which prints a concise summary of the structure of the data.

```
> data(xmp01.02)
> str(xmp01.02)

'data.frame': 27 obs. of 1 variable:
$ C1: num 5.9 7.2 7.3 6.3 8.1 6.8 7 7.6 6.8 6.5 ...
```

This shows that the data for example 1.2 (p. 5) consists of 27 observations of 1 variable called **strength**, which is a numeric variable. The first several data values are printed so you can check that they correspond to the values in the text.

Most of the data sets discussed in chapters 1 to 8 are univariate (i.e. only one variable), numeric data like xmp01.19. There are a few examples of univariate, categorical data such as the health complaints discussed in exercise 1.29 (p. 24)

```
> data(ex01.29)
> str(ex01.29)
'data.frame': 61 obs. of 1 variable:
$ C1: Factor w/ 8 levels "B","C","C5","F",...: 3 8 8 7 5 2 4 1 1 4 ...
```

These data are a set of observations of a variable that can take on only limited set of values named B for back pain, C for coughing, etc. In R such data are said to be a factor.

Some summary information about the variables in a data frame can be obtained with the summary function.

```
> summary(xmp01.02)
```

```
C1
Min.
        : 5.90
 1st Qu.: 7.00
Median : 7.70
Mean : 8.14
 3rd Qu.: 8.85
Max.
        :11.80
> summary(ex01.29)
       C1
 0
        :21
 J
        :10
 F
        : 9
        : 7
 N
        : 6
        : 4
 (Other): 4
```

For a numeric variable summary provides a 'five-number' summary and the mean (making a total of 6 numbers in all). For a factor summary provides a frequency table.

## A.9 Accessing individual variables

The summary function can be applied to entire data frames or to individual variables in a data frame. This is unusual. Most graphical or numerical functions apply to individual variables.

There are two ways to access a variable from within a data frame:

- 1. Use the name of the data set and the name of the variable separated by \$
- 2. attach the data frame and use the variable name by itself

For example, the two ways to obtain the stem-and-leaf plot of the space shuttle launch ambient temperature data from example 1.1 are

```
4 | 059
5 | 23788
6 | 01136777789
7 | 000023556689
8 | 0134

and

> attach(xmp01.01)
> stem(temp)

The decimal point is 1 digit(s) to the right of the |
3 | 1
4 | 059
5 | 23788
6 | 01136777789
7 | 000023556689
8 | 0134
```

### A.10 Stacked data

When storing multiple variables in an object there are two common choices. For instance suppose we had two variables

```
a = c(1, 2, 3, 4)
b = c(2, 3, 5, 7)
```

Then a data frame could have two columns, one for each

```
> df = data.frame(a, b)
> df

a b
1 1 2
2 2 3
3 3 5
4 4 7
```

This is a common storage method, but doesn't work well for independent samples, as they need not have the same number of observations, hence won't fit well in a rectangular format. As an alternative, the data can be stacked end to end, with an extra value indicating which variable the data refers to.

```
> stack(df)
```

	values	ind
1	1	a
2	2	a
3	3	a
4	4	a
5	2	b
6	3	b
7	5	b
8	7	b

This storage method also works well when model formula are used. Stacked data frames may be unstacked with unstack. For more extensive stacking options, the reshape add-on package is available from your nearest CRAN site.

## A.11 Finding help

R has an extensive amount of online documentation available through the help function.

Help on a data set Each data set in this package has a corresponding help page. For example, to view that page for the xmp01.01 data set, one would enter the command help(xmp01.01). This is more useful for data sets in other packages, as the ones in Devore7 are created generically. To view all the data sets in a package, say the Devore7 package, use the package argument: e.g., data(package="Devore7").

Help on a function name To find out more about a function, say the mean, the command help(mean) may be used. For many functions, the document writers have provided examples. The example function will execute the examples.

Seeing a vignette Many R packages have accompanying vignettes describing how the package works. This document is an example. Vignettes may be read from R with the command vignette. For instance, vignette("Devore7"). To list all available vignettes, drop the vignette name: vignette().