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Mangiafico, S.S. 2015. An R Companion for the Handbook of Biological Statistics, version 1.3.8, revised 2023.

 $\frac{r companion.org/documents/RCompanionBioStatistics.pdf}{r companion.org/r companion/).} (Web version:$

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Introduction

Purpose of This Book

This book is intended to be a supplement for *The Handbook of Biological Statistics* by John H. McDonald. It provides code for the R statistical language for some of the examples given in the *Handbook*. It does not describe the uses of, explanations for, or cautions pertaining to the analyses. For that information, you should consult the *Handbook* before using the analyses presented here.

The Handbook for Biological Statistics

This *Companion* follows the .pdf version of the third edition of the *Handbook of Biological Statistics*.

The *Handbook* provides clear explanations and examples of some the most common statistical tests used in the analysis of experiments. While the examples are taken from biology, the analyses are applicable to a variety of fields.

The *Handbook* provides examples primarily with the SAS statistical package, and with online calculators or spreadsheets for some analyses. Since SAS is a commercial package that students or researchers may not have access to, this *Companion* aims to extend the applicability of the *Handbook* by providing the examples in R, which is a free statistical package.

The .pdf version of the third edition is available at www.biostathandbook.com/HandbookBioStatThird.pdf.

Also, the *Handbook* can be accessed without cost at www.biostathandbook.com/. However, the reader should be aware that the online version may be updated since the third edition of the book.

Or, a printed copy can be purchased from www.lulu.com/shop/john-mcdonald/handbook-of-biological-statistics/paperback/product-22063985.html.

About the Author of this Companion

I have tried in this book to give the reader examples that are both as simple as possible, and that show some of the options available for the analysis. My goal for most examples is to make things comprehensible for the user without extensive R experience. The reader should realize that these goals may be partially frustrated either by the peculiarities in the R language or by the complexity required for the example.

I am neither a statistician nor an R programmer, so all advice and code in the book comes without guarantee. I'm happy to accept suggestions or corrections. Send correspondence to mangiafico@njaes.rutgers.edu.

About R

R is a free, open source, and cross-platform programming language that is well suited for statistical analyses. This means you can download R to your Windows, Mac OS, or Linux computer for free. It also means that, in theory, you can look at the code behind any of the analyses it performs to better understand the process, or to modify the code for your own purposes.

R is being used more and more in educational, academic, and commercial settings. A few advantages of working with R as a student, teacher, or researcher include:

- R functions return limited output. This helps prevent students from sorting through a lot of output they may not understand, and in essence requires the user to know what output they're asking R to produce.
- Since all functions are open source, the user has access to see how pre-defined functions are written.
- There are powerful packages written for specific type of analyses.
- There are lots of free resources available online.
- It can also be used online without installing software.

For a brief summary of some the advantages of R from the perspective of a graduate student, see thetarzan.wordpress.com/2011/07/15/why-use-r-a-grad-students-2-cents/.

It is also worth mentioning a few drawbacks with using R. New users are likely to find the code difficult to understand. Also, I think that while there is a plethora of examples for various analyses available online, it may be difficult for a beginner to adapt these examples to her own data. One goal of this book is to help alleviate these difficulties for beginners. I have some further thoughts below on avoiding pitfalls in R.

Obtaining R

This topic in SAEPER

For a more complete discussion of installing and running R, see **SAEPER**: Using R.

Standard installation

To download and install R, visit <u>cran.r-project.org/</u>. There you will find links for installation on Linux, Mac OS, and Windows operating systems.

RStudio

I also recommend using RStudio. This software is an environment for R that makes it easier to see code, output, datasets, plots, and help files together on one screen. www.rstudio.com/products/rstudio/. It is also possible to install RStudio as a portable application.

Portable application

R can be installed as a portable application. This is useful in cases where you don't want to install R on a computer but wish to run it from a portable drive. My portable installation of R with a handful of added packages is about 250 MB. The version on RStudio I have is about 400 MB. So, 1 GB of space on a usb drive is probably sufficient for the software along with additional installed packages and projects.

R Online

It is also possible to access R online, without needing to install software. One example of this is rdrr.io Snippets. This site includes common add-on packages.

A Few Notes to Get Started with R

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(dplyr)){install.packages("dplyr")}
if(!require(psych)){install.packages("psych")}
```

A cookbook approach

The examples in this book follow a "cookbook" approach as much as possible. The reader should be able to modify the examples with her own data and change the options and variable names as needed. This is more obvious with some examples than others, depending on the complexity of the code.

Color coding in this book

The text in blue in this book is R code that can be copied, pasted, and run in R. The text in red is the expected result and should not be run. In most cases I have truncated the results and included only the most relevant parts. Comments are in green. It is fine to run comments, but they have no effect on the results.

Copying and pasting code

From the website

Copying the R code pieces from the <u>website</u> version of this book should work flawlessly. Code can be copied from the webpages and pasted into the R console, the RStudio console, the RStudio editor, or a plain text file. All line breaks and formatting spaces should be preserved.

The only issue you may encounter is that if you paste code into the RStudio editor, leading spaces may be added to some lines. This is not usually a problem, but a way to avoid this is to paste the code into a plain text editor, save that file as a .R file, and open it from RStudio.

From the pdf

Copying the R code from the pdf version of this book may work less perfectly. Formatting spaces and even line breaks may be lost. Different pdf readers may behave differently.

It may help to paste the copied code in to a plain text editor to clean it up before pasting into R or saving it as a .R file. Also, if your pdf reader has a select tool that allows you to select text in a rectangle, that works better in some readers.

A sample program

The following is an example of code for R that creates a vector called *x* and a vector called *y*, performs a correlation test between *x* and *y*, and then plots *y* vs. *x*.

This code can be copied and pasted into the console area of R or RStudio, or into the editor area of RStudio, and run. You should get the output from the correlation test and the graphical output of the plot.

```
x = c(1,2,3,4,5,6,7,8,9) # create a vector of values and call it x y = c(9,7,8,6,7,5,4,3,1) # perform correlation test plot(x,y) # plot y vs. x
```

You can run fairly large chunks of code with R, though it is probably better to run smaller pieces, examining the output before proceeding to the next piece.

This kind of code can be saved as a file in the editor section of RStudio, or can be stored separately as a plain text file. By convention files for R code are saved as .R files. These files can be opened and edited with either a plain text editor or with the RStudio editor.

Assignment operators

In my examples I will use an equal sign, =, to assign a value to a variable.

```
height = 127.5
```

In examples you find elsewhere, you will more likely see a left arrow, <-, used as the assignment operator.

```
height <- 127.5
```

These are essentially equivalent, but I think the equal sign is more readable for a beginner.

Comments

Comments are indicated with a number sign, #. Comments are for human readers, and are not processed by R.

Installing and loading packages

Some of the packages used in this book do not come with R automatically but need to be installed as add-on packages. For example, if you wanted to use a function in the *psych* package to calculate the geometric mean of *x* in the sample program above:

```
x = c(1,2,3,4,5,6,7,8,9)
```

First you would need to the install the package *psych*:

```
install.packages("psych")
```

Then load the package:

```
library(psych)
```

You may then use the functions included in the package:

```
geometric.mean(x)
[1] 4.147166
```

In future sessions, you will need only to load the package; it should still be in the library from the initial installation.

If you see an error like the following, you may have misspelled the name of the package, or the package has not been installed.

```
library(psych)
```

```
Error in library(psych): there is no package called 'psych'
```

Data types

There are several data types in R. Most commonly, the functions we are using will ask for input data to be a vector, a matrix, or a data frame. Data types won't be discussed extensively here, but the examples in this book will read the data as the appropriate data type for the selected analysis.

Creating data frames from a text string of data

For certain analyses you will want to select a variable from within a data frame. In most examples using data frames, I'll create the data frame from a text string that allows us to arrange the data in columns and rows, as we normally visualize data.

A data frame can be created with the *read.table* function. Note that the text for the table is enclosed in simple double quotes and parentheses. *read.table* is pretty tolerant of extra spaces or blank lines. But if we convert a data frame to a matrix—which we will later—with *as.matrix*—I've had errors from trailing spaces at the ends of lines.

Values in the table that will have spaces or special characters can be enclosed in simple single quotes (e.g. 'Spongebob & Patrick').

```
D1 = read.table(header=TRUE, stringsAsFactors=TRUE, text="
Gender Height
        175
male
male
        176
female 162
female 165
")
D1
    Gender Height
   1 male
              175
      male
               176
   3 female
               162
   4 female
               165
```

Reading data from a file

R can also read data from a separate file. For longer data sets or complex analyses, it is helpful to keep data files and r code files separate. For example,

```
D2 = read.table("GenderHeight.dat", header=TRUE, stringsAsFactors=TRUE)
```

```
would read in data from a file called female-male.dat found in the working
   directory. In this case the file could be a space-delimited text file:
            Height
   Sex
  male
            175
            176
  male
   female
            162
   female
            165
Or, with read.csv,
  D2 = read.csv("GenderHeight.csv", header=TRUE, stringsAsFactors=TRUE)
for a comma-separated file.
   Gender, Height
   male,175
  male,176
   female, 162
   female, 165
  D2
        Gender Height
      1 male
                  175
          male
                  176
      3 female
                  162
      4 female
                  165
```

RStudio also has an easy interface in the *Tools* menu to import data from a file.

The *getwd* function will show the location of the working directory, and *setwd* can be used to set the working directory.

```
getwd()
   [1] "C:/Users/Salvatore/Documents"
setwd("C:/Users/Salvatore/Desktop")
```

Alternatively, file paths or URLs can be designated directly in the *read.table* function.

```
2 male 176
3 female 162
4 female 165
```

Variables within data frames

For the data frame *D1*created above, to look at just the variable *Gender* in this data frame:

```
D1$Gender

[1] male male female female
  Levels: female male
```

Note that *D1\$Height* is a vector of numbers.

```
D1$Height
[1] 175 176 162 165
```

So if you wanted the mean for this variable:

```
mean(D1$Height)
[1] 169.5
```

Using dplyr to create new variables in data frames

The standard method to define new variables in data frames is to use the *data.frame\$ variable* syntax. So if we wanted to add a variable to the D1 data frame above which would double *Height*:

```
D1$ Double = D1$ Height * 2
                                  # Spaces are optional
D1
     Gender Height Double
   1
       male
               175
                       350
       male
               176
                      352
   3 female
               162
                      324
   4 female
               165
                      330
```

Another method is to use the *mutate* function in the *dplyr* package:

D1

```
Gender Height Double Triple Quadruple
   male
            175
                   350
                           525
                                     700
                                     704
   male
            176
                   352
                           528
3 female
                   324
                                     648
            162
                           486
4 female
            165
                   330
                          495
                                     660
```

The *dplyr* package also has functions to select only certain columns in a data frame (*select* function) or to filter a data frame by the value of some variable (*filter* function). It can be helpful for manipulating data frames.

In the examples in this book, I will use either the \$ syntax or the *mutate* function in *dplyr*, depending on which I think makes the example more comprehensible.

Extracting elements from the output of a function

Sometimes it is useful to extract certain elements from the output of an analysis. For example, we can assign the output from a binomial test to a variable we'll call *Test*.

To see the value of *Test*:

```
Test
```

```
Exact binomial test

number of successes = 7, number of trials = 12, p-value = 0.1576

95 percent confidence interval:
    0.0000000    0.8189752
```

To see what elements are included in *Test*:

```
names(Test)

[1] "statistic" "parameter" "p.value" "conf.int" "estimate"
    "null.value" "alternative"
    [8] "method" "data.name"
```

Or with more details:

```
str(Test)
```

To view the p-value from *Test*:

```
Test$ p.value
[1] 0.1576437
```

To view the confidence interval from *Test*:

```
Test$ conf.int
[1] 0.0000000 0.8189752
[1] 0.95
```

To view the upper confidence limit from *Test*:

```
Test$ conf.int[2]
[1] 0.8189752
```

Exporting graphics

R has the ability to produce a variety of plots. Simple plots can be produced with just a few lines of code. These are useful to get a quick visualization of your data or to check on the distribution of residuals from an analysis. More in-depth coding can produce publication-quality plots.

Exporting plots from the RStudio window

In the RStudio *Plots* window, there is an *Export* icon which can be used to save the plot as image or pdf file. A method I use is to export the plot as pdf and then open this pdf with either Adobe Photoshop or the free alternative, GIMP (www.gimp.org/). These programs allow you to import the pdf at whatever resolution you need, and then crop out extra white space.

The appearance of exported plots will change depending on the size and scale of exported file. If there are elements missing from a plot, it may be because the size is not ideal. Changing the export size is also an easy way to adjust the size of the text of a plot relative to the other elements.

An additional trick in RStudio is to change the size of the plot window after the plot is produced, but before it is exported. Sometimes this can get rid of problems where, for example, words in a plot legend are cut off.

Finally, if you export a plot as a pdf, but still need to edit it further, you can open it in Inkscape, ungroup the plot elements, adjust some plot elements, and then export as a high-resolution bitmap image. Just be sure you don't change anything important, like how the data line up with the axes.

Exporting plots directly as a file

R also allows for the direct exporting of graphics as a .bmp, .jpg, .png, or .tif file. See *?png* for details. This method allows you to specify the dimensions and resolution of the outputted image.

Note that *dev.off()* is used afterwards to redirect future output to its usual channel.

```
### Optional code to set the directory where the image will be saved
setwd("C:/Users/Salvatore/Desktop")
### Create data frame
D4 = read.table(header=TRUE, stringsAsFactors=TRUE, text="
TolkienRace AvgHeight
Dwarf
            130
Hobbit
            105
            165
Man
E1f
            170
            125
Orc
")
### Output a plot as a .png file
png(filename = "TolkienPlot.png",
    width = 5,
    height = 3.75,
    units = "in",
           = 300)
    res
barplot(AvgHeight ~ TolkienRace, data=D4)
dev.off()
```

Avoiding Pitfalls in R

Grammar, spelling, and capitalization count

Probably the most common problems in programming in any language are syntax errors, for example, forgetting a comma or misspelling the name of a variable or function.

Be sure to include quotes around names requiring them; also be sure to use straight quotes (") and not the smart quotes that some word processors use automatically. It is helpful to write your R code in a plain text editor or in the editor window in RStudio.

Data types in functions

Probably the biggest cause of problems I had when I first started working with R was trying to feed functions the wrong data type. For example, if a function asks for the data as a matrix, and you give it a data frame, it won't work.

A more subtle error I've encountered is when a function is expecting a variable to be a factor vector, and it's really a character ("chr") vector.

For instance, if we create a variable in the global environment with the values of *Gender*, it will be a character vector.

```
Gender = c("male", "male", "female", "female")
str(Gender)  # What is the structure of this variable?
  chr [1:4] "male" "male" "female" "female"
```

While in the data frame, Gender was read in as a factor variable:

```
str(D1$ Gender)

Factor w/ 2 levels "female", "male": 2 2 1 1
```

One of the nice things about using RStudio is that it allows you to look at the structure of data frames and other objects in the *Environment* window.

Data types can be converted from one data type to another, but it may not be obvious how to do some conversions. Functions to convert data types include *factor*, *as.numeric*, and *as.character*.

```
Gender = c("male", "male", "female", "female")
str(Gender)
    chr [1:4] "male" "male" "female" "female"

Gender2 = factor(Gender)

Gender2
    [1] male male female female
    Levels: female male

str(Gender2)
Factor w/ 2 levels "female", "male": 2 2 1 1
```

```
Gender3 = as.numeric(Gender2)
Gender3
    [1] 2 2 1 1
str(Gender3)
    num [1:4] 2 2 1 1

Gender4 = as.character(Gender3)
Gender4
    [1] "2" "2" "1" "1"
str(Gender4)
    chr [1:4] "2" "2" "1" "1"
```

Creating data frames from vector variables

You can combine vector variables into a data frame with the *data.frame* function. However, note that the vectors need to have the same number of observations.

Style

There isn't an established style for programming in R in many respects, such as if variable names should be capitalized. In practice, people use different style conventions.

But it's helpful if you establish a convention for yourself, for example, in terms of the capitalization for variable names. For example, you could decide to always capitalize variable names and data frame names.

Some punctuation can be used variable names. For example, any of the following conventions could be used to create a fifth variable listing observations of genders.

```
Gender5 = factor(c("Female", "Male", "Nonbinary", "Female"))
Gender.5 = factor(c("Female", "Male", "Nonbinary", "Female"))
Gender_5 = factor(c("Female", "Male", "Nonbinary", "Female"))
```

Or any of the following conventions could be used to name a data frame.

```
mydata = data.frame(X=c(1,2,3), Y=c(4,5,6))

myData = data.frame(X=c(1,2,3), Y=c(4,5,6))

MyData = data.frame(X=c(1,2,3), Y=c(4,5,6))

My_Data = data.frame(X=c(1,2,3), Y=c(4,5,6))

My_Data = data.frame(X=c(1,2,3), Y=c(4,5,6))
```

Help with R

It's always a good idea to check the help information for a function before using it. Don't necessarily assume a function will perform a test as you think it will. The help information will give the options available for that function, and often those options make a difference with how the test is carried out.

Help in R

In order to see the help file for the *chisq.test* function:

```
?chisq.test
```

In order to specify the *chisq.test* function in the *stats* package, you would use:

```
?stats::chisq.test
or
help(chisq.test, package=stats)
```

In order to search all installed packages for a term:

```
??"chi-square"
```

In order to view the help for a package

```
help(package=psych)
```

CRAN documentation

Documentation for packages is also available in a .pdf format, which may be more convenient than using the help within R. Also very helpful, some packages include vignettes, which describe how a package might be used.

For a list of available packages, visit <u>cran.r-</u> <u>project.org/web/packages/available packages by name.html.</u>

And clicking on the link for the *psych* package will bring up a page with a link for the .pdf documentation, two .pdf vignettes, and other information.

Summary and Analysis of Extension Education Program Evaluation in R

Most of the analyses in this book are also presented in <u>Summary and Analysis of Extension</u> <u>Education Program Evaluation in R</u> (SAEEPER, html). Or as a pdf: <u>.pdf link</u>.

It may be useful for the reader to consult that book for additional examples and discussion.

Other online resources

Since there are many good resources for R online, an internet search for your question or analysis including the term "r" will often lead to a solution. The reader is cautioned, however, to always check the original R documentation on functions to be sure it will perform an analysis as the user desires.

A convenient tool is the *RSiteSearch* function, which will open a browser window and search for a term in functions and vignettes across a variety of sources:

```
RSiteSearch("chi-square")
```

This tool can also be accessed from: search.r-project.org/nmz.html.

R Tutorials

The descriptions of importing and manipulating data and results in this section of this book don't even scratch the surface of what is possible with R. Going beyond this very brief introduction, however, is beyond the scope of this book. I have tried to provide only enough information so that the reader unfamiliar with R will find the examples in the rest of the book comprehensible.

Luckily, there are many resources available for users wishing to better understand how to program in R, manipulate data, and perform more varied statistical analyses.

One free online resource I've found helpful is *Quick-R* (<u>www.statmethods.net/</u>).

CRAN hosts a collection of R manuals (<u>cran.r-project.org/manuals.html</u>). One that might be helpful is *An Introduction to R* by Venables.

CRAN also hosts a collection of contributed documentation (cran.r-project.org/other-docs.html), in several languages, which may prove helpful.

If readers wish to purchase a more-comprehensive and well-written textbook, *The R Book* by Michael Crawley is one option.

Formal Statistics Books

When describing a particular statistical analysis—especially one that your readers may not be familiar with—it's a good idea to cite an authoritative statistical source. A few that may be useful for this purpose:

- Biostatistical Analysis by Jerrold Zar
- Introduction to Biostatistics by Sokal and Rohlf
- Categorical Data Analysis by Alan Agresti
- Mixed-Effects Models in S and S-Plus by José Pinheiro and Douglas Bates

Tests for Nominal Variables

Exact Test of Goodness-of-Fit

The exact test goodness-of-fit can be performed with the *binom.test* function in the native *stats* package. The arguments passed to the function are: the number of successes, the number of trials, and the hypothesized probability of success. The probability can be entered as a decimal or a fraction. Other options include the confidence level for the confidence interval about the proportion, and whether the function performs a one-sided or two-sided (two-tailed) test. In most circumstances, the two-sided test is used.

Examples in Summary and Analysis of Extension Program EvaluationSAEEPER: Goodness-of-Fit Tests for Nominal Variables

Packages used in this chapter

The following commands will install these packages if they are not already installed:

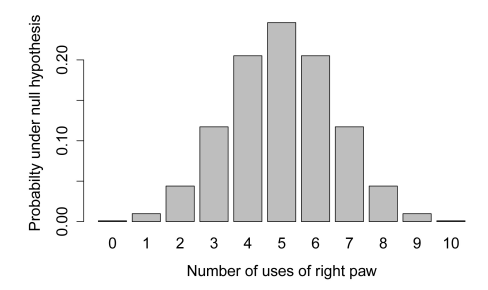
```
if(!require(XNomial)){install.packages("XNomial")}
if(!require(BSDA)){install.packages("BSDA")}
if(!require(pwr)){install.packages("pwr")}
```

Introduction When to use it Null hypothesis

See the *Handbook* for information on these topics.

How the test works Binomial test examples

Probability density plot



Comparing doubling a one-sided test and using a two-sided test

```
### -----
### Cat hair example, exact binomial test, p. 31-32
### Compares performing a one-sided test and doubling the
```

```
###
     probability, and performing a two-sided test
### -----
binom.test(7, 12, 3/4,
         alternative="less".
         conf.level=0.95)
  p-value = 0.1576
2 * Test$ p.value
                          # This extracts the p-value from the
                          # test result, we called Test
                              and multiplies it by 2
  [1] 0.3152874
binom.test(7, 12, 3/4, alternative="two.sided", conf.level=0.95)
  p-value = 0.1893
                    # Equal to the "small p values" method in the Handbook
```

Sign test

The following is an example of the two-sample dependent-samples sign test. The data are arranged as a data frame in which each row contains the values for both measurements being compared for each experimental unit. This is sometimes called "wide format" data. The SIGN.test function in the BSDA package is used. The option md=0 indicates that the expected difference in the medians is 0 (null hypothesis). This function can also perform a one-sample sign test.

```
### -----
### Tree beetle example, two-sample sign test, p. 34-35
### -----
Input =("
Row Angiosperm.feeding A.count Gymonsperm.feeding G.count
    Corthylina
                              Pityophthorus
1
                      458
                                                   200
2
    Scolytinae
                              Hylastini_Tomacini
                      5200
                                                   180
    Acanthotomicus_P 123
3
                              Orhotomicus
                                                    11
4
    Xyleborini_D
                              Ipini
                                                   195
                      1500
5
    Apion
                     1500
                              Antliarhininae
                                                    12
6
                              Allocoryninae_Oxycorinae 30
    Belinae
                       150
7
    H_Curculionidae
                              Nemonychidae
                     44002
                                                    85
                              Aseminae_Spondylinae
8
    H_Cerambycidae
                     25000
                                                    78
    Megalopodinae
                              Palophaginae
9
                     400
                                                     3
10
    H_Chrysomelidae
                     33400
                              Aulocoscelinae_Orsod
                                                    26
")
Data = read.table(textConnection(Input),header=TRUE)
```

Exact multinomial test

See example below in the "Examples" section.

Post-hoc test

Post-hoc example with manual pairwise tests

A multinomial test can be conducted with the *xmulti* function in the package *XNomial*. This can be followed with the individual binomial tests for each proportion, as post-hoc tests.

```
### Post-hoc example, multinomial and binomial test, p. 33
### ------
observed = c(72, 38, 20, 18)
expected = c(9, 3, 3, 1)
library(XNomial)
xmulti(observed,
      expected,
      detail = 2)  # 2: Reports three types of p-value
  P value (LLR) = 0.003404 # log-likelihood ratio
  P value (Prob) = 0.002255 # exact probability
  P value (Chisq) = 0.001608 # Chi-square probability
  ### Note last p-value below agrees with Handbook
successes = 72
total
        = 148
numerator = 9
denominator = 16
binom.test(successes, total, numerator/denominator,
         alternative="two.sided", conf.level=0.95)
  p-value = 0.06822
successes = 38
total = 148
```

```
numerator = 3
denominator = 16
binom.test(successes, total, numerator/denominator,
          alternative="two.sided", conf.level=0.95)
   p-value = 0.03504
successes = 20
           = 148
total
numerator = 3
denominator = 16
binom.test(successes, total, numerator/denominator,
          alternative="two.sided", conf.level=0.95)
   p-value = 0.1139
successes = 18
total
           = 148
numerator = 1
denominator = 16
binom.test(successes, total, numerator/denominator,
          alternative="two.sided", conf.level=0.95)
   p-value = 0.006057
```

Post-hoc test alternate method with custom function

When you need to do multiple similar tests, however, it is often possible to use the programming capabilities in R to do the tests more efficiently. The following example may be somewhat difficult to follow for a beginner. It creates a data frame and then adds a column called *p.Value* that contains the p-value from the *binom.test* performed on each row of the data frame.

```
### Post-hoc example, multinomial and binomial test, p. 33
### Alternate method for multiple tests
Input =("
Successes Total Numerator Denominator
72
         148
                         16
                         16
38
               3
         148
               3
20
        148
                         16
18
         148 1
                         16
")
D1 = read.table(textConnection(Input), header=TRUE)
Fun = function (x){
```

Intrinsic hypothesis

Assumptions

See the *Handbook* for information on these topics.

Examples

Binomial test examples

```
### Parasitoid examples, exact binomial test, p. 34
### -----
binom.test(10, (17+10), 0.5,
        alternative="two.sided",
        conf.level=0.95)
  p-value = 0.2478
binom.test(36, (7+36), 0.5,
        alternative="two.sided",
        conf.level=0.95)
  p-value = 8.963e-06
### Drosophila example, exact binomial test, p. 34
### -----
binom.test(140, (106+140), 0.5,
        alternative="two.sided",
        conf.level=0.95)
  p-value = 0.03516
```

```
### -----
### First Mendel example, exact binomial test, p. 35
### -----
binom.test(428, (428+152), 0.75, alternative="two.sided",
       conf.level=0.95)
 ### -----
### First Mendel example, exact binomial test, p. 35
### Alternate method with XNomial package
### -----
observed = c(428, 152)
expected = c(3, 1)
library(XNomial)
xmulti(observed,
    expected,
    detail = 2)
             # 2: reports three types of p-value
 P value (LLR) = 0.5331 # log-likelihood ratio
  P value (Prob) = 0.5022 # exact probability
 P value (Chisq) = 0.5331 # Chi-square probability
 ### Note last p-value below agrees with Handbook
```

Multinomial test example

Graphing the results

Graphing is shown in the "Chi-square Goodness-of-Fit" section.

Similar tests

The *G*–test goodness-of-fit and chi-square goodness-of-fit are presented elsewhere in this book.

How to do the test

Binomial test example where individual responses are counted

```
### Cat paw example from SAS, exact binomial test, pp. 36-37
### When responses need to be counted
Input =("
Paw
right
left
right
right
right
right
left
right
right
right
")
Gus = read.table(textConnection(Input),header=TRUE)
                                 # Note the == operator
Successes = sum(Gus$ Paw == "left")
Failures = sum(Gus$ Paw == "right")
Total = Successes + Failures
Expected = 0.5
binom.test(Successes, Total, Expected,
         alternative="less",
                                   # One-sided test!
         conf.level=0.95)
  p-value = 0.05469
conf.level=0.95)
  p-value = 0.1094
```

Other SAS examples

R code for the other SAS example is shown in the examples in previous sections.

Power analysis

Power analysis for binomial test

```
### -----
### Power analysis, binomial test, cat paw, p. 38
P0 = 0.50
P1 = 0.40
                   # This calculates effect size
H = ES.h(P0,P1)
library(pwr)
pwr.p.test(
      h=H,
      n=NULL, # NUL
sig.level=0.05, #
power=0.80, # 1 m
                            # NULL tells the function to
                                 calculate this value
                            # 1 minus Type II probability
      alternative="two.sided")
                            # Slightly different than in Handbook
  n = 193.5839
```

Power Analysis

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(pwr)){install.packages("pwr")}
```

Introduction

Parameters

How it works

See the *Handbook* for information on these topics.

Examples

Power analysis for binomial test

```
### -----
### Power analysis, binomial test, pea color, p. 43
### ------
P0 = 0.75
P1 = 0.78
H = ES.h(P0,P1) # This calculates effect size
```

```
library(pwr)

pwr.p.test(
    h=H,
    n=NULL,  # NULL tells the function to
    sig.level=0.05,  # calculate this
    power=0.90,  # 1 minus Type II probability
    alternative="two.sided")

n = 2096.953  # Somewhat different than in Handbook
```

Power analysis for unpaired t-test

```
### Power analysis, t-test, student height, pp. 43-44
M1 = 66.6

M2 = 64.6

S1 = 4.8
M1 = 66.6
                                # Mean for sample 1
                               # Mean for sample 2
# Std dev for sample 1
S2 = 3.6
                                  # Std dev for sample 2
Cohen.d = (M1 - M2)/sqrt(((S1^2) + (S2^2))/2)
library(pwr)
pwr.t.test(
       n = NULL,  # Observations in _each_ group
d = Cohen.d,
sig.level = 0.05,  # Type I probability
power = 0.80,  # 1 minus Type II probability
type = "two.sample",  # Change for one- or two-sample
alternative = "two cided")
       alternative = "two.sided")
   Two-sample t test power calculation
   n = 71.61288
   NOTE: n is number in *each* group 71.61288
```

How to do power analyses

Methods are shown in the previous examples.

Chi-square Test of Goodness-of-Fit

Examples in Summary and Analysis of Extension Program EvaluationSAEEPER: Goodness-of-Fit Tests for Nominal Variables

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(dplyr)){install.packages("dplyr")}
if(!require(ggplot2)){install.packages("ggplot2")}
if(!require(grid)){install.packages("grid")}
if(!require(pwr)){install.packages("pwr")}
```

When to use it Null hypothesis

See the *Handbook* for information on these topics.

How the test works

Chi-square goodness-of-fit example

Post-hoc test Assumptions

See the *Handbook* for information on these topics.

Examples: extrinsic hypothesis

Example: intrinsic hypothesis

Graphing the results

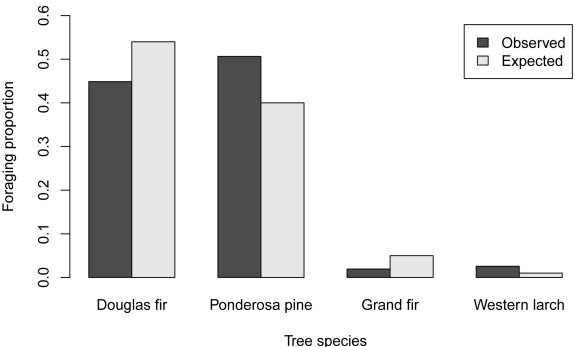
The first example below will use the *barplot* function in the native *graphics* package to produce a simple plot. First we will calculate the observed proportions and then copy those results into a matrix format for plotting. We'll call this matrix *Matriz*. See the "Chi-square Test of Independence" section for a few notes on creating matrices.

The second example uses the package *ggplot2*, and uses a data frame instead of a matrix. The data frame is named *Forage*. For this example, the code calculates confidence intervals and adds them to the data frame. This code could be skipped if those values were determined manually and put into a data frame from which the plot could be generated.

Sometimes factors will need to have the order of their levels specified for *ggplot2* to put them in the correct order on the plot, as in the second example. Otherwise R will alphabetize levels.

Simple bar plot with barplot

```
### Simple bar plot of proportions, p. 49
      Uses data in a matrix format
### ---
observed = c(70, 79, 3, 4)
expected = c(0.54, 0.40, 0.05, 0.01)
total = sum(observed)
observed.prop = observed / total
observed.prop
   [1] 0.44871795 0.50641026 0.01923077 0.02564103
### Re-enter data as a matrix
Input =("
Value
         Douglas.fir Ponderosa.pine Grand.fir
                                                  Western.larch
Observed 0.4487179
                      0.5064103
                                      0.01923077 0.02564103
Expected 0.5400000
                      0.4000000
                                      0.05000000 0.01000000
")
Matriz = as.matrix(read.table(textConnection(Input),
                  header=TRUE,
                  row.names=1))
Matriz
           Douglas fir Ponderosa pine Grand fir Western larch
                       0.5064103 0.01923077 0.02564103
   Observed 0.4487179
   Expected
             0.5400000
                            0.4000000 0.05000000
                                                    0.01000000
barplot(Matriz,
       beside=TRUE,
       legend=TRUE,
       ylim=c(0, 0.6),
       xlab="Tree species",
       ylab="Foraging proportion")
```



Bar plot with confidence intervals with ggplot2

The plot below is a bar char with confidence intervals. The code calculates confidence intervals. This code could be skipped if those values were determined manually and put in to a data frame from which the plot could be generated.

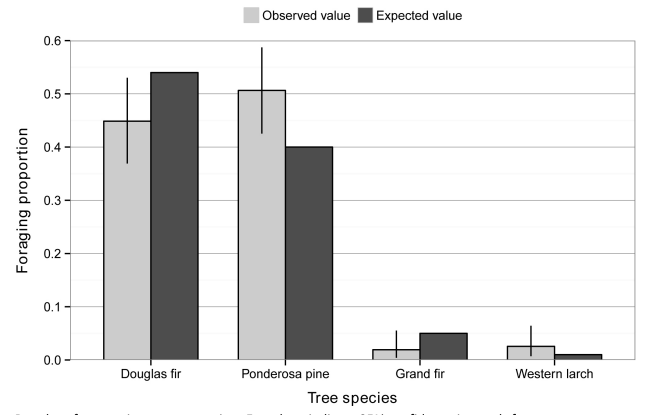
Sometimes factors will need to have the order of their levels specified for *gaplot2* to put them in the correct order on the plot. Otherwise R will alphabetize levels.

```
### -----
### Graph example, Chi-square goodness-of-fit, p. 49
      Using ggplot2
###
      Plot adapted from:
         shinyapps.stat.ubc.ca/r-graph-catalog/
###
Input =("
                 Value
                            Count
                                    Total Proportion Expected
Tree
'Douglas fir'
                 Observed
                                                     0.54
                            70
                                    156
                                          0.4487
'Douglas fir'
                            54
                                    100
                                                     0.54
                 Expected
                                         0.54
'Ponderosa pine'
                 Observed
                            79
                                    156
                                         0.5064
                                                     0.40
'Ponderosa pine'
                 Expected
                            40
                                    100
                                         0.40
                                                     0.40
'Grand fir'
                 Observed
                             3
                                    156
                                         0.0192
                                                     0.05
'Grand fir'
                             5
                                    100
                                         0.05
                                                     0.05
                 Expected
'Western larch'
                 Observed
                             4
                                    156
                                         0.0256
                                                     0.01
'Western larch'
                 Expected
                             1
                                    100
                                         0.01
                                                     0.01
")
```

Forage = read.table(textConnection(Input), header=TRUE)

```
### Specify the order of factor levels. Otherwise R will alphabetize them.
library(dplyr)
Forage =
mutate(Forage,
      Tree = factor(Tree, levels=unique(Tree)),
      value = factor(value, levels=unique(value)))
### Add confidence intervals
Forage =
mutate(Forage,
      low.ci = apply(Forage[c("Count", "Total", "Expected")],
                      function(x)
                      binom.test(x["Count"], x["Total"], x["Expected"]
                               ) $ conf.int[1]),
       upper.ci = apply(Forage[c("Count", "Total", "Expected")],
                        1,
                        function(x)
                        binom.test(x["Count"], x["Total"], x["Expected"]
                                  )$ conf.int[2]))
Forage$ low.ci [Forage$ Value == "Expected"] = 0
Forage$ upper.ci [Forage$ Value == "Expected"] = 0
Forage
                    Value Count Total Proportion Expected
             Tree
                                                         low.ci
                                                                 upper.ci
       Douglas fir Observed 70 156
                                       0.4487
                                                0.54 0.369115906 0.53030534
       Douglas fir Expected
                            54
                                100
                                       0.5400
                                                0.54 0.000000000 0.00000000
  3 Ponderosa pine Observed 79
                                156
                                       0.5064
                                                0.40 0.425290653 0.58728175
  4 Ponderosa pine Expected
                           40
                                       0.4000
                                100
                                                0.40 0.000000000 0.00000000
                          3 5
                                       0.0192
        Grand fir Observed
                                                0.05 0.003983542 0.05516994
                                156
                                100
156
100
                                       Grand fir Expected
  7 Western larch Observed
8 Western larch Expected
                            4
### Plot adapted from:
     shinyapps.stat.ubc.ca/r-graph-catalog/
library(ggplot2)
library(grid)
ggplot(Forage,
  aes(x = Tree, y = Proportion, fill = Value,
      ymax=upper.ci, ymin=low.ci)) +
      show_quide = FALSE) +
      scale_y_continuous(breaks = seq(0, 0.60, 0.1),
```

```
limits = c(0, 0.60),
         expand = c(0, 0) +
scale_fill_manual(name = "Count type"
          values = c('grey80', 'grey30'),
          labels = c("Observed value",
                     "Expected value"))
geom_errorbar(position=position_dodge(width=0.7),
              width=0.0, size=0.5, color="black") +
labs(x = "Tree species",
     y = "Foraging proportion") +
## ggtitle("Main title") +
theme_bw() +
theme(panel.grid.major.x = element_blank(),
      panel.grid.major.y = element_line(colour = "grey50"),
      plot.title = element_text(size = rel(1.5),
      face = "bold", vjust = 1.5),
      axis.title = element_text(face = "bold"),
      legend.position = "top",
      legend.title = element_blank(),
      legend.key.size = unit(0.4, "cm"),
      legend.key = element_rect(fill = "black"),
      axis.title.y = element_text(vjust= 1.8),
      axis.title.x = element_text(vjust= -0.5))
```



Bar plot of proportions vs. categories. Error bars indicate 95% confidence intervals for each observed proportion.

Similar tests

Chi-square vs. G-test

See the *Handbook* for information on these topics. The *exact test of goodness-of-fit*, the *G-test of goodness-of-fit*, and the *exact test of goodness-of-fit* tests are described elsewhere in this book.

How to do the test

Chi-square goodness-of-fit example

Power analysis

Power analysis for chi-square goodness-of-fit

G-test of Goodness-of-Fit

The G-test goodness-of-fit test can be performed with the *G.test* function in the package *RVAideMemoire*, the *GTest* function in *DescTools*. As another alternative, you can use R to calculate the statistic and p-value manually.

Examples in Summary and Analysis of Extension Program EvaluationSAEEPER: Goodness-of-Fit Tests for Nominal Variables

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(DescTools)){install.packages("DescTools")}
if(!require(RVAideMemoire)){install.packages("RVAideMemoire")}
```

When to use it
Null hypothesis
How the test works
Post-hoc test
Assumptions

See the *Handbook* for information on these topics.

Examples: extrinsic hypothesis

G-test goodness-of-fit test with DescTools and RVAideMemoire

G-test goodness-of-fit test by manual calculation

Examples of G-test goodness-of-fit test with DescTools and RVAideMemoire

Example: intrinsic hypothesis

```
### Intrinsic example, G-test goodness-of-fit, amphipod, p. 56
### -----
observed = c(1203, 2919, 1678)
expected.prop = c(.21073, 0.49665, 0.29262)
   ### Note: These are recalculated for more precision
           In this case, low precision probabilities
             change the results
   ###
expected.count = sum(observed)*expected.prop
G = 2 * sum(observed * log(observed / expected.count))
  [1] 1.032653
pchisq(G,
     df=1.
     lower.tail=FALSE)
  [1] 0.3095363
```

Graphing the results

Graphing would be the same as in the "Chi-square Test of Goodness-of-Fit" section.

Similar tests

Chi-square vs. G-test

See the *Handbook* for information on these topics. The *exact test of goodness-of-fit* and the *chisquare test of goodness-of-fit* tests are described elsewhere in this book.

How to do the test

These examples are shown above.

Power analysis

Power analysis would be the same as in the "Chi-square Test of Goodness-of-Fit" section.

Chi-square Test of Independence

The Chi-square test of independence can be performed with the *chisq.test* function in the native *stats* package in R. For this test, the function requires the contingency table to be in the form of matrix. Depending on the form of the data to begin with, this can require an extra step, either combing vectors into a matrix, or cross-tabulating the counts among factors in a data frame. None of this is too difficult, but it requires following the correct example depending on the initial form of the data.

When using *read.table* and *as.matrix* to read a table directly as a matrix, be careful of extra spaces at the end of lines or extraneous characters in the table, as these can cause errors.

Examples in Summary and Analysis of Extension Program EvaluationSAEEPER: Association Tests for Nominal Variables

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(rcompanion)){install.packages("rcompanion")}
if(!require(dplyr)){install.packages("dplyr")}
if(!require(ggplot2)){install.packages("ggplot2")}
if(!require(grid)){install.packages("grid")}
if(!require(pwr)){install.packages("pwr")}
```

When to use it

Example of chi-square test with matrix created with read.table

```
### -----
Input =("
Injection.area No.severe Severe
Thigh 4788 30
Arm 8916 76
")
Matriz = as.matrix(read.table(textConnection(Input),
                header=TRUE,
                row.names=1))
Matriz
          No.severe Severe
         4788 30
  Thigh
  Arm
          8916 76
chisq.test(Matriz,
         correct=TRUE) # Continuity correction for 2 x 2
                         # table
  Pearson's Chi-squared test with Yates' continuity correction
  X-squared = 1.7579, df = 1, p-value = 0.1849
chisq.test(Matriz,
         correct=FALSE) # No continuity correction for 2 x 2
                          # table
  Pearson's Chi-squared test
  X-squared = 2.0396, df = 1, p-value = 0.1533
```

Example of chi-square test with matrix created by combining vectors

Matriz

```
No.severe Severe
  Thigh 4788 30
  Arm
            8916
                     76
chisq.test(Matriz,
                            # Continuity correction for 2 x 2
          correct=TRUE)
                                   table
   Pearson's Chi-squared test with Yates' continuity correction
  X-squared = 1.7579, df = 1, p-value = 0.1849
chisq.test(Matriz,
                             # No continuity correction for 2 x 2
          correct=FALSE)
  Pearson's Chi-squared test
  X-squared = 2.0396, df = 1, p-value = 0.1533
```

Null hypothesis How the test works

See the *Handbook* for information on these topics.

Post-hoc tests

For the following example of post-hoc pairwise testing, we'll use the *pairwiseNominalIndependence* function from the package *rcompanion* to make the task easier. Then we'll use *pairwise.table* in the native *stats* package as an alternative.

Post-hoc pairwise chi-square tests with rcompanion

```
### Post-hoc example, Chi-square independence, pp. 60-61
### ------
Input =("
Supplement
           No.cancer Cancer
'Selenium'
           8177 575
'Vitamin E'
                    620
           8117
           8147
8167
'Selenium+E' 8147
                    555
'Placebo'
                    529
")
Matriz = as.matrix(read.table(textConnection(Input),
               header=TRUE.
               row.names=1))
Matriz
```

```
chisq.test(Matriz)
  X-squared = 7.7832, df = 3, p-value = 0.05071
library(rcompanion)
pairwiseNominalIndependence(Matriz,
                            fisher = FALSE,
                            gtest = FALSE,
                            chisq = TRUE,
                            method = "fdr")
                 Comparison p.Chisq p.adj.Chisq
      Selenium : Vitamin E 0.17700
                                        0.2960
   2 Selenium : Selenium+E 0.62800
                                         0.6280
                                      0.6280
0.2960
0.1880
0.0463
         Selenium : Placebo 0.19700
   4 Vitamin E : Selenium+E 0.06260
       Vitamin E : Placebo 0.00771
      Selenium+E : Placebo 0.44000
   6
                                       0.5280
```

Post-hoc pairwise chi-square tests with pairwise.table

```
### -----
### Post-hoc example, Chi-square independence, pp. 60-61
### As is, this code works on a matrix with two columns.
### and compares rows
### -----
Input =("
Supplement No.cancer Cancer 'Selenium' 8177 575 'Vitamin E' 8117 620 'Selenium+E' 8147 555 'Placebo' 8167 529
")
Matriz = as.matrix(read.table(textConnection(Input),
                  header=TRUE.
                  row.names=1))
Matriz
chisq.test(Matriz)
  X-squared = 7.7832, df = 3, p-value = 0.05071
FUN = function(i,j){
     chisq.test(matrix(c(Matriz[i,1], Matriz[i,2],
                        Matriz[j,1], Matriz[j,2]),
                nrow=2,
                byrow=TRUE))$ p.value
```

Assumptions

See the *Handbook* for information on this topic.

Examples

Chi-square test of independence with continuity correction and without correction

```
### Helmet example, Chi-square independence, p. 63
### -----
Input =("
PSE Head.injury Other.injury
         372
                     4715
Helemt
No.helmet 267
                    1391
Matriz = as.matrix(read.table(textConnection(Input),
                header=TRUE,
                row.names=1))
Matriz
chisq.test(Matriz,
                           # Continuity correction for 2 x 2
         correct=TRUE)
                                 table
  Pearson's Chi-squared test with Yates' continuity correction
  X-squared = 111.6569, df = 1, p-value < 2.2e-16
chisq.test(Matriz,
         correct=FALSE)
                           # No continuity correction for 2 x 2
                               table
       Pearson's Chi-squared test
       X-squared = 112.6796, df = 1, p-value < 2.2e-16
```

Chi-square test of independence

Graphing the results

The first plot below is a bar char with confidence intervals, with a style typical of the *ggplot2* package. The second plot is somewhat more similar to the style of the plot in the *Handbook*.

For each example, the code calculates proportions or confidence intervals. This code could be skipped if those values were determined manually and put in to a data frame from which the plot could be generated.

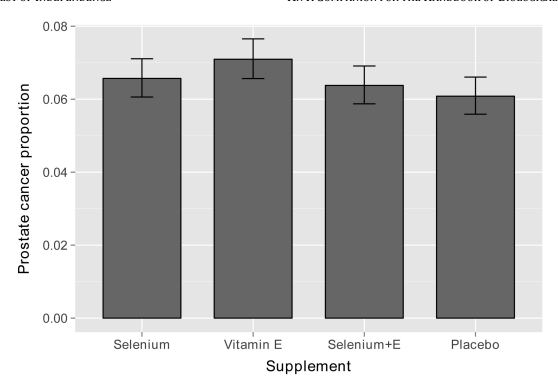
Sometimes factors will need to have the order of their levels specified for *ggplot2* to put them in the correct order on the plot. Otherwise R will alphabetize levels.

Simple bar plot with error bars showing confidence intervals

```
### ------
### Plot example, herons and egrets, Chi-square test of association,
### pp. 63-64
### ------

Input =("
Supplement No.cancer Cancer
'Selenium' 8177 575
'Vitamin E' 8117 620
'Selenium+E' 8147 555
'Placebo' 8167 529
")
```

```
Prostate = read.table(textConnection(Input),header=TRUE)
### Add sums and confidence intervals
library(dplyr)
Prostate =
mutate(Prostate.
      Sum = No.cancer + Cancer)
Prostate =
mutate(Prostate,
      Prop = Cancer / Sum,
      low.ci = apply(Prostate[c("Cancer", "Sum")], 1,
               function(y) binom.test(y['Cancer'], y['Sum'])$ conf.int[1]),
      Prostate
                                                  low.ci
    Supplement No.cancer Cancer Sum
                                                            high.ci
                                          Prop
     Selenium
                   8177 575 8752 0.06569927 0.06059677 0.07109314
  2 Vitamin E
                    8117 620 8737 0.07096257 0.06566518 0.07654816
                   8147 555 8702 0.06377844 0.05873360 0.06911770
8167 529 8696 0.06083257 0.05589912 0.06606271
  3 Selenium+E
       Placebo
### Plot (Bar chart plot)
library(ggplot2)
ggplot(Prostate,
aes(x=Supplement, y=Prop)) +
width=0.7) +
 geom_errorbar(aes(ymax=high.ci, ymin=low.ci),
                  width=0.2, size=0.5, color="black") +
xlab("Supplement") +
 ylab("Prostate cancer proportion") +
 scale_x_discrete(labels=c("Selenium", "Vitamin E",
                          "Selenium+E", "Placebo")) +
 ## ggtitle("Main title") +
 theme(axis.title=element_text(size=14, color="black",
                             face="bold", vjust=3)) +
 theme(axis.text = element_text(size=12, color = "gray25",
                              face="bold")) +
 theme(axis.title.y = element_text(vjust= 1.8)) +
 theme(axis.title.x = element_text(vjust= -0.5))
```

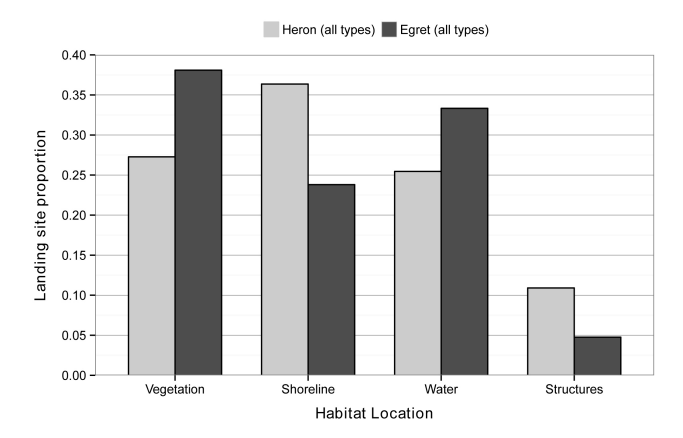


Bar plot of proportions vs. categories. Error bars indicate 95% confidence intervals for observed proportion.

Bar plot with categories and no error bars

```
### Plot example, herons and egrets, Chi-square independence,
###
     p. 64
Input =("
Habitat
             Bird
                    Count
Vegetation
            Heron
                     15
Shoreline
            Heron
                     20
Water
            Heron
                     14
Structures
            Heron
                      6
Vegetation
                      8
            Egret
                      5
Shoreline
             Egret
                      7
Water
             Egret
                      1
Structures
             Egret
")
Birds = read.table(textConnection(Input), header=TRUE)
### Specify the order of factor levels
library(dplyr)
Birds=
```

```
mutate(Birds,
        Habitat = factor(Habitat,levels=unique(Habitat)),
        Bird = factor(Bird,levels=unique(Bird)))
### Add sums and proportions
Birds$ Sum[Birds$ Bird == 'Heron'] =
        sum(Birds$ Count[Birds$ Bird == 'Heron'])
Birds$ Sum[Birds$ Bird == 'Egret'] =
        sum(Birds$ Count[Birds$ Bird == 'Egret'])
Birds=
mutate(Birds.
        prop = Count / Sum)
Birds
         Habitat Bird Count Sum
                                             prop
   1 Vegetation Heron 15 55 0.27272727
2 Shoreline Heron 20 55 0.36363636
           Water Heron 14 55 0.25454545
   3
   4 Structures Heron 6 55 0.10909091
5 Vegetation Egret 8 21 0.38095238
6 Shoreline Egret 5 21 0.23809524
7 Water Egret 7 21 0.33333333
8 Structures Egret 1 21 0.04761905
### Plot adapted from:
### shinyapps.stat.ubc.ca/r-graph-catalog/
library(ggplot2)
library(grid)
gaplot(Birds.
  aes(x = Habitat, y = prop, fill = Bird, ymax=0.40, ymin=0)) +
  geom_bar(stat="identity", position = "dodge", width = 0.7) +
geom_bar(stat="identity", position = "dodge", colour = "black",
             width = 0.7, show_guide = FALSE) +
  scale_v_continuous(breaks = seq(0, 0.40, 0.05),
                         limits = c(0, 0.40),
                         expand = c(0, 0) +
  scale_fill_manual(name = "Bird type"
                        values = c('grey80', 'grey30'),
                       labels = c("Heron (all types)",
                                     "Egret (all types)")) +
  ## geom_errorbar(position=position_dodge(width=0.7).
                      width=0.0, size=0.5, color="black") +
  labs(x = "Habitat Location", y = "Landing site proportion") +
  ## ggtitle("Main title") +
  theme_bw() +
  theme(panel.grid.major.x = element_blank(),
         panel.grid.major.y = element_line(colour = "grey50"),
```



Similar tests

Chi-square vs. G-test

See the Handbook for information on these topics. *Fisher's exact test, G-test,* and *McNemar's test* are discussed elsewhere in this book.

How to do the test

Chi-square test of independence with data as a data frame

In the following example for the chi-square test of independence, the data is read in as a data frame, not as a matrix as in previous examples. This allows more flexibility with how data are entered. For example you could have counts for same *genotype* and *health* distributed among several lines, or have a count of 1 for each row, with a separate row for each individual observation. The *xtabs* function is used to tabulate the data and convert them to a contingency table.

```
### -----
### Gardemann apolipoprotein example, Chi-square independence,
         SAS example, pp. 65-66
         Example using cross-tabulation
Input =("
Genotype Health
                       Count
ins-ins no_disease 268
ins-ins disease
ins-del no_disease
ins-del disease
                       807
                        199
                       759
del-del no_disease
                        42
del-del disease
                        184
")
Data.frame = read.table(textConnection(Input), header=TRUE)
### Cross-tabulate the data
Data.xtabs = xtabs(Count ~ Genotype + Health,
                   data=Data.frame)
Data.xtabs
          Health
   Genotype disease no_disease

    del-del
    184
    42

    ins-del
    759
    199

    ins-ins
    807
    268

summary(Data.xtabs)
                                   # includes N and factors
   Number of cases in table: 2259
   Number of factors: 2
### Chi-square test of independence
chisq.test(Data.xtabs)
   X-squared = 7.2594, df = 2, p-value = 0.02652
```

Power analysis

Power analysis for chi-square test of independence

```
### -----
### Power analysis, chi-square independence, pp. 66-67
### ------
# This example assumes you are using a Chi-square test of
```

```
independence. The example in the Handbook appears to use
   a Chi-square goodness-of-fit test
# In the pwr package, for the Chi-square test of independence,
   the table probabilities should sum to 1
Input =("
Genotype No.cancer Cancer
         0.18
                   0.165
GG
         0.24
                   0.225
GΑ
         0.08
                   0.110
AA
")
P = as.matrix(read.table(textConnection(Input),
             header=TRUE,
             row.names=1))
Р
     No.cancer Cancer
  GG
          0.18 0.165
          0.24 0.225
  GΑ
          0.08 0.110
  AA
sum(P) # Sum of values in the P matrix
  \lceil 1 \rceil 1
library(pwr)
effect.size = ES.w2(P)
degrees = (nrow(P)-1)*(ncol(P)-1) # Calculate degrees of freedom
pwr.chisq.test(
      w=effect.size,
                        # Total number of observations
      N=NULL.
      df=degrees,
      power=0.80, # 1 minus Type II probability
      sig.level=0.05) # Type I probability
          w = 0.07663476 # Answer differs significantly
          N = 1640.537 # from Handbook
         df = 2
                          # Total observations
  sig.level = 0.05
      power = 0.8
```

G-test of Independence

There are a few different options for performing G-tests of independence in R. One is the *G.test* function in the package *RVAideMemoire*. Another is the *GTest* function in the package *DescTools*.

Examples in Summary and Analysis of Extension Program EvaluationSAEEPER: Association Tests for Nominal Variables

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(DescTools)){install.packages("DescTools")}
if(!require(RVAideMemoire)){install.packages("RVAideMemoire")}
```

When to use it

G-test example with functions in DescTools and RVAideMemoire

```
### Vaccination example, G-test of independence, pp. 68-69
Input =("
Injection.area No.severe Severe
Thigh 4788 30
Arm 8916 76
")
Matriz = as.matrix(read.table(textConnection(Input),
                 header=TRUE.
                 row.names=1))
Matriz
library(DescTools)
GTest(Matriz,
     correct="none") # "none" "williams" "yates"
  Log likelihood ratio (G-test) test of independence without correction
  G = 2.1087, X-squared df = 1, p-value = 0.1465
library(RVAideMemoire)
G.test(Matriz)
  G = 2.1087, df = 1, p-value = 0.1465 # Note values differ from
                                     # the Handbook
                                     # for this example
```

Null hypothesis How the test works

See the *Handbook* for information on these topics.

Post-hoc tests

For the following example of post-hoc pairwise testing, we'll use the *pairwise.G.test* function from the package *RVAideMemoire* to make the task easier. Then we'll use *pairwise.table* in the native *stats* package as an alternative.

Post-hoc pairwise G-tests with RVAideMemoire

```
### Post-hoc example, G-test of independence, pp. 69-70
Input =("
Supplement
              No.cancer Cancer
'Selenium'
              8177
                         575
'Vitamin E' 8117
'Selenium+E' 8147
              8117
                          620
                         555
'Placebo'
               8167
                          529
")
Matriz = as.matrix(read.table(textConnection(Input),
                   header=TRUE,
                   row.names=1))
Matriz
library(RVAideMemoire)
G.test(Matriz)
   G = 7.7325, df = 3, p-value = 0.05188
library(RVAideMemoire)
pairwise.G.test(Matriz,
                p.method = "none")
                                             # Can adjust p-values:
                                             # see ?p.adjust for options
              Selenium Vitamin E Selenium+E
   Vitamin E 0.168
   Selenium+E 0.606
                       0.058
   Placebo 0.187
                      0.007
                                 0.422
```

Post-hoc pairwise G-tests with pairwise.table

As is, this function works on a matrix with two columns, and compares rows.

```
Input =("
Supplement
               No.cancer Cancer
'Selenium'
               8177
                          575
'Vitamin E'
               8117
                          620
'Selenium+E'
               8147
                          555
'Placebo'
               8167
                          529
")
Matriz = as.matrix(read.table(textConnection(Input),
                   header=TRUE,
                   row.names=1))
Matriz
library(DescTools)
GTest(Matriz,
      correct="none")
   Log likelihood ratio (G-test) test of independence without correction
  G = 7.7325, X-squared df = 3, p-value = 0.05188
FUN = function(i,j){
      GTest(matrix(c(Matriz[i,1], Matriz[i,2],
                      Matriz[j,1], Matriz[j,2]),
             nrow=2.
             byrow=TRUE),
             correct="none")$ p.value # "none" "williams" "yates"
pairwise.table(FUN,
               rownames(Matriz),
               p.adjust.method="none")
                                             # Can adjust p-values
                                             # See ?p.adjust for options
               Selenium Vitamin E Selenium+E
   Vitamin E 0.1677388
                               NA
   Selenium+E 0.6060951 0.058385135
           0.1866826 0.007004601 0.4215013
   Placebo
```

Assumptions

See the *Handbook* for information on this topic.

Examples

G-tests with DescTools and RVAideMemoire

```
### ------
### Helmet example, G-test of independence, p. 72
### ------
Input =("
PSE Head.injury Other.injury
```

```
4715
Helemt 372
No.helmet 267
                    1391
")
Matriz = as.matrix(read.table(textConnection(Input),
                header=TRUE,
                row.names=1))
Matriz
library(DescTools)
GTest(Matriz,
                     # "none" "williams" "yates"
     correct="none")
  Log likelihood ratio (G-test) test of independence without correction
  G = 101.54, X-squared df = 1, p-value < 2.2e-16
library(RVAideMemoire)
G.test(Matriz)
  G = 101.5437, df = 1, p-value < 2.2e-16
### -----
### Gardemann apolipoprotein example, G-test of independence,
### -----
Input =("
Genotype No.disease Coronary.disease
ins.ins 268
                  807
ins.del 199
del.del 42
                  759
                  184
")
Matriz = as.matrix(read.table(textConnection(Input),
                header=TRUE,
                row.names=1))
Matriz
library(DescTools)
GTest(Matriz,
                    # "none" "williams" "yates"
     correct="none")
  Log likelihood ratio (G-test) test of independence without correction
  G = 7.3008, X-squared df = 2, p-value = 0.02598
```

```
library(RVAideMemoire)

G.test(Matriz)

G = 7.3008, df = 2, p-value = 0.02598
```

Graphing the results

Graphing is discussed above in the "Chi-square Test of Independence" section.

Similar tests

Chi-square vs. G-test

See the *Handbook* for information on these topics. *Fisher's exact test, chi-square test,* and *McNemar's test* are discussed elsewhere in this book.

How to do the test

G-test of independence with data as a data frame

In the following example, the data is read in as a data frame, and the *xtabs* function is used to tabulate the data and convert them to a contingency table.

```
### Gardemann apolipoprotein example, G-test of independence,
        SAS example, pp. 74-75
        Example using cross-tabulation
### -----
Input =("
Genotype Health
                      Count
ins-ins no_disease
ins-ins disease
ins-del no_disease
                      268
                      807
                      199
ins-del disease
                      759
del-del no_disease
                      42
del-del
         disease
                      184
")
Data.frame = read.table(textConnection(Input), header=TRUE)
### Cross-tabulate the data
Data.xtabs = xtabs(Count ~ Genotype + Health,
                   data=Data.frame)
Data.xtabs
           Health
   Genotype disease no_disease
     del-del 184
                           42
     ins-del
                759
                           199
     ins-ins
                807
                          268
```

Power analysis

To calculate power or required samples, follow examples in the "Chi-square Test of Independence" section.

Fisher's Exact Test of Independence

Examples in Summary and Analysis of Extension Program EvaluationSAEEPER: Association Tests for Nominal Variables

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(rcompanion)){install.packages("rcompanion")}
```

When to use it Null hypothesis How the test works

See the *Handbook* for information on these topics.

Post-hoc tests

For the following example of post-hoc pairwise testing, we'll use the *pairwiseNominalIndependence* function from the package *rcompanion* to make the task easier.

Post-hoc pairwise Fisher's exact tests with RVAideMemoire

```
### Post-hoc example, Fisher's exact test, p. 79
Input =("
Frequency Damaged Undamaged
Daily 1
                       24
           5
Weekly
                       20
Monthly 14
                       11
Quarterly 11
                       14
Matriz = as.matrix(read.table(textConnection(Input),
                     header=TRUE,
                     row.names=1))
Matriz
fisher.test(Matriz,
             alternative="two.sided")
   p-value = 0.0001228
   alternative hypothesis: two.sided
library(rcompanion)
PT = pairwiseNominalIndependence(Matriz,
                                     fisher = TRUE,
                                     gtest = FALSE,
                                     chisq = FALSE,
                                     digits = 3)
PT
               Comparison p.Fisher p.adj.Fisher
           Daily: Weekly 0.189000 0.227000
   1
   2 Daily: Monthly 0.000102 0.000612
3 Daily: Quarterly 0.001920 0.005760
4 Weekly: Monthly 0.018600 0.037200
5 Weekly: Quarterly 0.128000 0.192000
6 Monthly: Quarterly 0.572000 0.572000
library(rcompanion)
cldList(comparison = PT$Comparison,
         p.value = PT$p.adj.Fisher,
         threshold = 0.05)
          Group Letter MonoLetter
   1
          Daily a a
```

```
2
    weekly
                        ab
               ab
3
   Monthly
              C
                         C
4 Quarterly
                         bc
               bc
Summary of results
Frequency Damaged Letter
Daily
          4%
                  a
Weekly
          20%
                  ab
Quarterly 44%
                  bc
Monthly
          56%
                    C
```

Groups sharing a letter are not significantlt different (alpha = 0.05)

Assumptions

See the *Handbook* for information on this topic.

Examples

Examples of Fisher's exact test with data in a matrix

```
### Chipmunk example, Fisher's exact test, p. 80
Input =("
Distance
          Trill No.trill
10m
                 8
          16
          3
                 18
100m
")
Matriz = as.matrix(read.table(textConnection(Input),
                   header=TRUE,
                   row.names=1))
Matriz
fisher.test(Matriz,
            alternative="two.sided")
   p-value = 0.0006862
### Drosophila example, Fisher's exact test, p. 81
Input =("
Variation
                     Synonymous Replacement
Variation Syn
'Polymorphisms' 43
'Fixed differences' 17
                                 7
")
```

```
Matriz = as.matrix(read.table(textConnection(Input),
               header=TRUE,
               row.names=1))
Matriz
fisher.test(Matriz,
         alternative="two.sided")
  p-value = 0.006653
### -----
### King penguin example, Fisher's exact test, p. 81
### -----
Input =("
Site
       Alive Dead
Lower
       43
Middle 44 6
Upper 49 1
")
Matriz = as.matrix(read.table(textConnection(Input),
               header=TRUE,
               row.names=1))
Matriz
fisher.test(Matriz,
         alternative="two.sided")
  p-value = 0.08963
  alternative hypothesis: two.sided
### -----
### Moray eel example, Fisher's exact test, pp. 81-82
Input =("
Site G.moringa G.vicinus
Grass
       127
                116
Sand
       99
                67
Border 264
                161
")
Matriz = as.matrix(read.table(textConnection(Input),
               header=TRUE,
               row.names=1))
Matriz
```

```
fisher.test(Matriz,
           alternative="two.sided")
   p-value = 0.04438
   alternative hypothesis: two.sided
### Herons example, Fisher's exact test, p. 82
### -----
Input =("
Site
             Heron Egret
Vegetation 15
Shoreline 20
             15
                   8
                 5
                  7
            14
Water
Structures 6 1
")
Matriz = as.matrix(read.table(textConnection(Input),
                  header=TRUE,
                  row.names=1))
Matriz
fisher.test(Matriz,
           alternative="two.sided")
    p-value = 0.5491
    alternative hypothesis: two.sided
```

Graphing the results

Graphing is discussed above in the "Chi-square Test of Independence" section.

Similar tests - McNemar's test

Care is needed in setting up the data for McNemar's test. For a before-and-after test, the contingency table is set-up as before and after as row and column headings, or vice-versa. Note that the total observations in the contingency table is equal to the number of experimental units. That is, in the following example there are 62 men, and the sum of the counts in the contingency table is 62. If you set up the table incorrectly, you might end with double this number, and this will not yield the correct results.

McNemar's test with data in a matrix

McNemar's test with data in a data frame

```
### Dysfunction example, McNemar test, pp. 82-83
### Example using cross-tabulation
### -----
Input =("
ED.before ED.after Count
         no
no
                   46
                  10
no
        yes
                   0
yes
        no
                   6
yes
         yes
")
Data = read.table(textConnection(Input), header=TRUE)
Data.xtabs = xtabs(Count ~ ED.before + ED.after, data=Data)
Data.xtabs
             ED.after
  ED.before no yes
             46
                  10
         yes 0
mcnemar.test(Data.xtabs, correct=FALSE)
  McNemar's chi-squared = 10, df = 1, p-value = 0.001565
```

How to do the test

Fisher's exact test with data as a data frame

```
Distance
          Sound
                  Count
          trill
10m
                  16
10m
          notrill 8
100m
         trill
                   3
100m
          notrill 18
")
Data = read.table(textConnection(Input), header=TRUE)
Data.xtabs = xtabs(Count ~ Distance + Sound, data=Data)
Data.xtabs
          Sound
  Distance notrill trill
      100m 18
      10m
                8
                      16
summary(Data.xtabs)
### Fisher's exact test of independence
fisher.test(Data.xtabs,
            alternative="two.sided")
  p-value = 0.0006862
### Bird example, Fisher's exact test, SAS example, p. 84
### Example using cross-tabulation
Input =("
Bird Substrate
                  Count
heron vegetation 15
heron shoreline 20
heron water
                  14
heron structures 6
egret vegetation 8
egret shoreline
                  5
egret water
                   7
                   1
egret structures
")
Data = read.table(textConnection(Input), header=TRUE)
Data.xtabs = xtabs(Count ~ Bird + Substrate, data=Data)
Data.xtabs
         Substrate
  Bird shoreline structures vegetation water
                            1
    egret
```

heron

20

Power analysis

To calculate power or required samples, follow examples in the "Chi-square Test of Independence" section.

6

15

14

There, the result was

```
N = 1640.537 # Total observations
```

compared with the value in the *Handbook* of N_{total} = 1523 for this section.

Small Numbers in Chi-square and G-tests

The problem with small numbers

See the *Handbook* for information on these topics.

Yates' and William's corrections in R

The following table lists the continuity corrections available for the Chi-square tests and G-tests discussed in this book.

Test	Function	Package	Correction	Option	Default	Notes
Chi-square	chisq.test	stats	Yates	correct=TRUE	TRUE	2 x 2 table only
G	G.test	RVAide Memoire	(none)			
G	GTest	DescTools	Yates	correct= "yates"	"none"	
			Williams	correct= "williams"		

Pooling

Recommendation

See the *Handbook* for information on these topics.

Repeated G-tests of Goodness-of-Fit

These examples use the *G.test* function in the *RVAideMemoire* package, but the *GTest* function in the *DescTools* package could be used in the same manner.

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(dplyr)){install.packages("dplyr")}
if(!require(RVAideMemoire)){install.packages("RVAideMemoire")}
```

When to use it Null hypothesis

See the *Handbook* for information on these topics.

How to do the test

Repeated G-tests of goodness-of-fit example

Data = read.table(textConnection(Input), header=TRUE)

Individual G-tests

```
}
   Fun.df = function (0){
               G.test(x=c(Q["R"], Q["L"]),
                      p=c(0.5, 0.5)
                      )$parameter
                   }
   Fun.p = function (Q){
              G.test(x=c(Q["R"], Q["L"]),
                     p=c(0.5, 0.5)
                     )$p.value
                   }
   library(dplyr)
   Data=
   mutate(Data,
           Prop.R = R / (R + L),
                                                             # Calculate proportion
                                                                   of right arms
           G = apply(Data[c("R", "L")], 1, Fun.G),
df = apply(Data[c("R", "L")], 1, Fun.df),
p.value = apply(Data[c("R", "L")], 1, Fun.p))
   Data
     Ethnic.group R L
                              Prop.R
                                                G df
                                                          p.Value
             Yemen 168 174 0.4912281 0.1052686 1 0.745596489
   1
   2
            Djerba 132 195 0.4036697 12.2138397 1 0.000474363
        Kurdistan 167 204 0.4501348 3.6961684 1 0.054537574
   3
            Libya 162 212 0.4331551 6.7045477 1 0.009616732
   4
   5
            Berber 143 194 0.4243323 7.7478346 1 0.005377698
           Cochin 153 174 0.4678899 1.3495524 1 0.245356383
Heterogeneity G-test
   Data.matrix = as.matrix(Data[c("R", "L")])
                                                       # We need a data matrix
                                                           to run G-test
   Data.matrix
                                                            for heterogeneity
              R
                L
      [1,] 168 174
      [2,] 132 195
      [3,] 167 204
      [4,] 162 212
      [5,] 143 194
      [6,] 153 174
   G.test(Data.matrix)
                                                       # Heterogeneity
      G-test
      G = 6.7504, df = 5, p-value = 0.2399
```

Pooled G-test

```
Total.R = sum(Data\$R)
                                                   # Set up data for pooled
  Total.L = sum(Data$L)
                                                   # G-test
   observed = c(Total.R, Total.L)
   expected = c(0.5, 0.5)
  G.test(x=observed,
         p=expected)
     G-test for given probabilities
     G = 25.0668, df = 1, p-value = 5.538e-07
Total G-test
                                                   # Set up data for total
  Total.G = sum(Data$G)
                                                   # G-test
  Total.df = sum(Data$df)
  Total.G
                                                    # Total
      [1] 31.81721
  Total.df
      [1] 6
   pchisq(Total.G,
          df= Total.df,
          lower.tail=FALSE)
      [1] 1.768815e-05
```

Example

Repeated G-tests of goodness-of-fit example

Data = read.table(textConnection(Input),header=TRUE)

Individual G-tests

```
library(RVAideMemoire)
   Fun.G = function (Q){
                                                     # Functions
             G.test(x=c(Q["D"], Q["S"]),
                                                         to calculate
                                                         individual G's and
                     p=c(0.5, 0.5)
                     )$statistic
                                                         p-values
                  }
   Fun.df = function (Q){
              G.test(x=c(Q["D"], Q["S"]),
                      p=c(0.5, 0.5)
                      )$parameter
                   }
   Fun.p = function (Q){
             G.test(x=c(Q["D"], Q["S"]),
                     p=c(0.5, 0.5)
                     )$p.value
   library(dplyr)
   Data =
   mutate(Data,
                    apply(Data[c("D", "S")], 1, Fun.G),
apply(Data[c("D", "S")], 1, Fun.df),
          G =
          df =
          p.Value = apply(Data[c("D", "S")], 1, Fun.p))
   Data
       Trial D S
                             G df
                                     p.Value
   1 Trial 1 296 366 7.415668 1 0.00646583
   2 Trial 2 78 72 0.240064 1 0.62415986
   3 Trial 3 417 467 2.829564 1 0.09254347
Heterogeneity G-test
                                                     # We need a data matrix
   Data.matrix = as.matrix(Data[c("D", "S")])
                                                         to run G-test
   Data.matrix
                                                         for heterogeneity
          D
   [1,] 296 366
   [2,] 78 72
   [3,] 417 467
```

```
G.test(Data.matrix)
                                                   # Heterogeneity
     G-test
     G = 2.8168, df = 2, p-value = 0.2445
Pooled G-test
                                                   # Set up data for pooled
   Total.D = sum(Data$D)
  Total.S = sum(Data$S)
                                                   # G-test
   observed = c(Total.D, Total.S)
   expected = c(0.5, 0.5)
   G.test(x=observed,
                                                    # Pooled
          p=expected)
     G-test for given probabilities
     G = 7.6685, df = 1, p-value = 0.005619
Total G-test
  Total.G = sum(Data$G)
                                                   # Set up data for total
                                                   # G-test
  degrees = 3
  Total.G = sum(Data$G)
                                                   # Set up data for total
                                                   # G-test
  Total.df = sum(Data$df)
                                                   # Total
  Total.G
      [1] 10.4853
  Total.df
      [1] 3
   pchisq(Total.G,
          df=Total.df,
          lower.tail=FALSE)
```

Similar tests

[1] 0.01486097

See the *Handbook* for information on these topics.

Cochran-Mantel-Haenszel Test for Repeated Tests of Independence

The Cochran–Mantel–Haenszel test can be performed in R with the *mantelhaen.test* function in the native *stats* package. A few other useful functions come from the package *vcd*. One is *woolf_test*, which performs the Woolf test for homogeneity of the odds ratio across strata levels. This has a similar function to the Breslow-Day test mentioned in the *Handbook*. If this test is significant, the C-M-H test may not be appropriate. The Breslow-Day test itself can be performed with a function in the package *DescTools*. For cautions about using this test, see the documentation for this function, or other appropriate sources.

```
library(DescTools); ?BreslowDayTest
```

There are a couple of different ways to generate the three-way contingency table. The table can be read in with the *read.ftable* function. Note that the columns are the stratum variable.

Caution should be used with the formatting, since *read.ftable* can be fussy. I've noticed that it doesn't like leading spaces in the rows. Certain editors, such as the one in RStudio, may add leading spaces when this code is pasted in. To alleviate this, delete those spaces manually, or paste the code into a plain text editor, save the file as a .R file, and then open that file with RStudio.

Another way to generate the contingency table is beginning with a data frame and tabulating the data using the *xtabs* function. The second example uses this method.

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Cochran-Mantel-Haenszel Test for 3-Dimensional Tables

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(dplyr)){install.packages("dplyr")}
if(!require(DescTools)){install.packages("DescTools")}
if(!require(ggplot2)){install.packages("ggplot2")}
if(!require(grid)){install.packages("grid")}
if(!require(vcd)){install.packages("vcd")}
```

When to use it
Null hypothesis
How the test works
Assumptions

See the *Handbook* for information on these topics.

Examples

Cochran-Mantel-Haenszel Test with data read by read.ftable

```
### Handedness example, Cochran-Mantel-Haenszel test, p. 97-98
  ### Example using read.ftable
  ### -----
        # Note no spaces on lines before row names.
        # read.ftable can be fussy about leading spaces.
  Input =(
                     Group W.Child B.adult PA.white W.men G.soldier
  Whorl
             Handed
  Clockwise Right
                             708
                                     136
                                              106
                                                    109
                                                             801
             Left
                             50
                                      24
                                              32
                                                     22
                                                             102
  CounterCl Right
                             169
                                      73
                                               17
                                                     16
                                                             180
                                      14
                                                     26
                              13
                                               4
                                                              25
             Left
  ")
  Tabla = as.table(read.ftable(textConnection(Input)))
                                      # Display a flattened table
  ftable(Tabla)
Cochran-Mantel-Haenszel test
  mantelhaen.test(Tabla)
     Mantel-Haenszel X-squared = 5.9421, df = 1, p-value = 0.01478
Woolf test
  library(vcd)
  oddsratio(Tabla, log=TRUE) # Show log odds for each 2x2
                                             W.men G.soldier
        w.Child
                    B.adult PA.white
     0.08547173 0.08319894 -0.24921579 2.08581324 0.08680711
  library(vcd)
  woolf_test(Tabla)
                                  # Woolf test for homogeneity of
                                  # odds ratios across strata.
                                      If significant, C-M-H test
                                      is not appropriate
     Woolf-test on Homogeneity of Odds Ratios (no 3-Way assoc.)
     X-squared = 22.8165, df = 4, p-value = 0.0001378
```

Breslow-Day test

```
library(DescTools)
BreslowDayTest(Tabla)
Breslow-Day Test for Homogeneity of the Odds Ratios
X-squared = 24.7309, df = 4, p-value = 5.698e-05
```

Individual Fisher exact tests

```
n = dim(Tabla)[3]
for(i in 1:n){
  Name = dimnames(Tabla)[3]$Group[i]
   P.value = fisher.test(Tabla[,,i])$p.value
   cat(Name, "\n")
   cat("Fisher test p-value: ", P.value, "\n")
  cat("\n")
  }
 ### Note: "Group" must be the name of the stratum variable
  w.child
   Fisher test p-value: 0.7435918
   B.adult
   Fisher test p-value: 0.8545009
   PA.white
   Fisher test p-value: 0.7859788
  W.men
   Fisher test p-value: 6.225227e-08
  G.soldier
   Fisher test p-value: 0.7160507
```

Cochran-Mantel-Haenszel Test with data entered as a data frame

```
### -----
### Mussel example, Cochran-Mantel-Haenszel test, pp. 98-99
### Example using cross-tabulation of a data frame
### -----

Input =("
Location Habitat Allele Count
Tillamook marine 94 56
Tillamook estuarine 94 69
```

```
Tillamook marine
                       non-94
                                  40
Tillamook estuarine
                       non-94
                                  77
Yaquina
           marine
                           94
                                  61
Yaquina
           estuarine
                           94
                                 257
Yaquina
                       non-94
           marine
                                  57
Yaquina
           estuarine
                       non-94
                                 301
Alsea
           marine
                           94
                                  73
Alsea
           estuarine
                           94
                                  65
Alsea
           marine
                       non-94
                                  71
                                  79
Alsea
           estuarine
                       non-94
Umpqua
           marine
                           94
                                  71
Umpqua
           estuarine
                           94
                                  48
                                  55
Umpqua
           marine
                       non-94
Umpqua
           estuarine
                       non-94
                                  48
")
Data = read.table(textConnection(Input), header=TRUE)
### Specify the order of factor levels
### Otherwise, R will alphabetize them
library(dplyr)
Data =
mutate(Data,
       Location = factor(Location, levels=unique(Location)),
       Habitat = factor(Habitat, levels=unique(Habitat)),
       Allele = factor(Allele, levels=unique(Allele)))
### Cross-tabulate the data
      Note here, Location is stratum variable (is last)
###
###
                 Habitat x Allele are 2 x 2 tables
Data.xtabs = xtabs(Count ~ Allele + Habitat + Location,
                   data=Data)
ftable(Data.xtabs)
                                        # Display a flattened table
                    Location Tillamook Yaquina Alsea Umpqua
   Allele Habitat
   94
          marine
                                    56
                                            61
                                                   73
                                                          71
                                            257
                                                   65
                                                          48
          estuarine
                                    69
   non-94 marine
                                    40
                                            57
                                                   71
                                                          55
          estuarine
                                    77
                                            301
                                                   79
                                                          48
```

Cochran-Mantel-Haenszel test

```
mantelhaen.test(Data.xtabs)
```

```
Mantel-Haenszel X-squared = 5.0497, df = 1, p-value = 0.02463
```

```
Woolf test
```

```
library(vcd)
   oddsratio(Data.xtabs, log=TRUE) # Show log odds for each 2x2
      Tillamook
                  Yaquina
                              Alsea Umpqua
      0.4461712 0.2258568 0.2228401 0.2553467
   library(vcd)
   woolf_test(Data.xtabs)
                                        # Woolf test for homogeneity of
                                            odds ratios across strata.
                                            If significant, C-M-H test
                                            is not appropriate
     Woolf-test on Homogeneity of Odds Ratios (no 3-Way assoc.)
      X-squared = 0.5292, df = 3, p-value = 0.9124
Breslow-Day test
   library(DescTools)
   BreslowDayTest(Data.xtabs)
      Breslow-Day Test for Homogeneity of the Odds Ratios
      X-squared = 0.5295, df = 3, p-value = 0.9124
Individual Fisher exact tests
   n = dim(Data.xtabs)[3]
   for(i in 1:n){
      Name = dimnames(Data.xtabs)[3]$Location[i]
      P.value = fisher.test(Data.xtabs[,,i])$p.value
      cat(Name, "\n")
      cat("Fisher test p-value: ", P.value, "\n")
      cat("\n")
     }
    ### Note: "Location" must be the name of the stratum variable
     Tillamook
      Fisher test p-value: 0.1145223
      Yaquina
```

```
Fisher test p-value: 0.2665712
     Alsea
     Fisher test p-value: 0.4090355
     Umpqua
     Fisher test p-value: 0.4151874
Cochran-Mantel-Haenszel Test with data read by read.ftable
  ### ------
  ### Niacin example, Cochran-Mantel-Haenszel test, p. 99
  ### Example using read.ftable
  ### -----
       # Note no spaces on lines before row names.
       # read.ftable can be fussy about leading spaces.
  Input =(
                  Study FATS AFREGS ARBITER.2 HATS CLAS.1
  Supplement Revasc
                                  1 1
86 37
  Niacin
                      4
46 67
11 12
41 60
                       2
                                               2
            Yes
                                               92
            No
  Placebo
           Yes
                                  4
                                           6
                                               1
                                  76 32 93
            No
  ")
  Tabla = as.table(read.ftable(textConnection(Input)))
  ftable(Tabla)
                                   # Display a flattened table
Cochran-Mantel-Haenszel test
  mantelhaen.test(Tabla)
     Mantel-Haenszel X-squared = 12.7457, df = 1, p-value = 0.0003568
Woolf test
  library(vcd)
  oddsratio(Tabla, log=TRUE) # Show log odds for each 2x2
                 AFREGS ARBITER.2
                                       HATS
                                               CLAS.1
     -1.8198174 -1.2089603 -1.5099083 -1.9369415 0.7039581
  library(vcd)
```

woolf test for homogeneity of
odds ratios across strata.

woolf_test(Tabla)

```
If significant, C-M-H test
                                        # is not appropriate
     Woolf-test on Homogeneity of Odds Ratios (no 3-Way assoc.)
     X-squared = 3.4512, df = 4, p-value = 0.4853
Breslow-Day test
   library(DescTools)
   BreslowDayTest(Tabla)
      Breslow-Day Test for Homogeneity of the Odds Ratios
     X-squared = 4.4517, df = 4, p-value = 0.3483
Individual Fisher exact tests
   n = dim(Tabla)[3]
   for(i in 1:n){
      Name = dimnames(Tabla)[3]$Study[i]
      P.value = fisher.test(Tabla[,,i])$p.value
      cat(Name, "\n")
      cat("Fisher test p-value: ", P.value, "\n")
      cat("\n")
    ### Note: "Study" must be the name of the stratum variable
      Fisher test p-value: 0.01581505
      AFREGS
      Fisher test p-value: 0.0607213
     ARBITER.2
      Fisher test p-value: 0.1948915
     HATS
      Fisher test p-value: 0.1075169
     CLAS.1
```

Graphing the results

Simple bar plot with categories and no error bars

Fisher test p-value: 1

```
### ------### Simple bar plot of proportions, p. 99
```

```
###
         Uses data in a matrix format
### --
Input =("
          Tillamook Yaquina Alsea
Habitat
                                        Umpqua
Marine
          0.5833
                      0.5169
                                0.5069 0.5635
                                0.4514
                      0.4606
                                        0.5000
Estuarine 0.4726
")
Matriz = as.matrix(read.table(textConnection(Input),
                    header=TRUE,
                    row.names=1))
Matriz
barplot(Matriz,
        beside=TRUE,
        legend=TRUE,
        ylim=c(0, 0.9),
        xlab="Location",
        ylab="Lap94 proportion")
       0.8
                                                        Marine
                                                    Estuarine
   Lap94 proportion
       9
       Ö
       4
       \sim
```

Bar plot with categories and error bars

Tillamook

Ö

0

This example includes code to calculate the confidence intervals for the error bars and add them to the data frame. This code could be excluded if these values were calculated manually and added to the data frame.

Location

Alsea

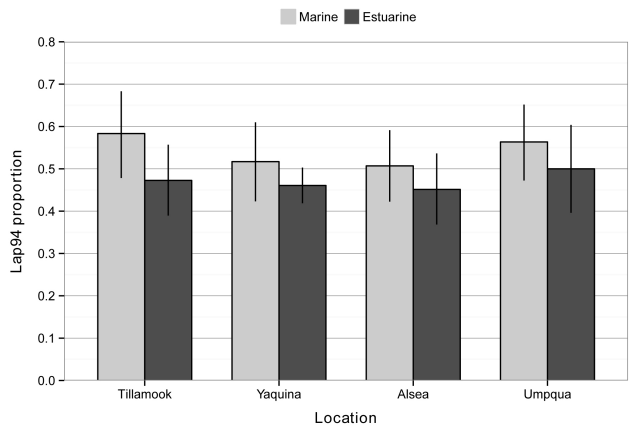
Umpqua

```
### -----
### Graph example, bar plot of proportions, p. 99
### Using ggplot2
### Plot adapted from:
### shinyapps.stat.ubc.ca/r-graph-catalog/
```

Yaquina

```
### -----
Input =("
Location Habitat Allele Count Total Lap.94.Proportion
Tillamook Marine 94 56 96 0.5833
Tillamook Estuarine 94 69 146 0.4726
Yaquina Marine 94 61 118 0.5169
Yaquina Estuarine 94 257 558 0.4606
Alsea Marine 94 73 144 0.5069
Alsea Estuarine 94 65 144 0.4514
Umpqua Marine 94 71 126 0.5635
Umpqua Estuarine 94 48 96 0.5000
")
Data = read.table(textConnection(Input),header=TRUE)
### Specify the order of factor levels
### Otherwise, R will alphabetize them
library(dplyr)
Data =
mutate(Data,
         Location = factor(Location, levels=unique(Location)),
         Habitat = factor(Habitat, levels=unique(Habitat)),
         Allele = factor(Allele, levels=unique(Data$ Allele)))
### Add confidence intervals
Fun.low = function (x){
             binom.test(x["Count"], x["Total"],
             0.5)$ conf.int[1]
            }
Fun.up = function (x){
               binom.test(x["Count"], x["Total"],
               0.5)$ conf.int[2]
Data =
mutate(Data,
             low.ci = apply(Data[c("Count", "Total")], 1, Fun.low),
             upper.ci = apply(Data[c("Count", "Total")], 1, Fun.up))
Data
       Location Habitat Allele Count Total Lap.94.Proportion
                                                                              low.ci upper.ci
    1 Tillamook Marine 94 56 96 0.5833 0.4782322 0.6831506
    2 TITIAMOON ESCUALTINE 94 69 146 0.4726 0.3894970 0.5568427 3 Yaquina Marine 94 61 118 0.5169 0.4231343 0.6098931 4 Yaquina Estuarine 94 257 558 0.4606 0.4186243 0.5029422 5 Alsea Marine 94 73 144 0.5069 0.4224208 0.5911766 6 Alsea Estuarine 94 65 144 0.4514 0.3684040 0.5364149
                                 94 69 146
                                                                  0.4726 0.3894970 0.5568427
```

```
7
                         94
                               71
                                   126
                                                0.5635 0.4723096 0.6516209
       Umpqua
               Marine
  8
       Umpqua Estuarine
                         94
                               48
                                    96
                                                0.5000 0.3961779 0.6038221
### Plot adapted from:
     shinyapps.stat.ubc.ca/r-graph-catalog/
library(ggplot2)
library(grid)
ggplot(Data,
  aes(x = Location, y = Lap.94.Proportion, fill = Habitat,
      ymax=upper.ci, ymin=low.ci)) +
      show_quide = FALSE) +
      scale_y_continuous(breaks = seq(0, 0.80, 0.1),
               limits = c(0, 0.80),
               expand = c(0, 0)) +
      scale_fill_manual(name = "Count type"
                values = c('grey80', 'grey30'),
labels = c("Marine",
                           "Estuarine")) +
      geom_errorbar(position=position_dodge(width=0.7),
                    width=0.0, size=0.5, color="black") +
      labs(x = "Location",
           y = "Lap94 proportion") +
      ## ggtitle("Main title") +
      theme_bw() +
      theme(panel.grid.major.x = element_blank(),
            panel.grid.major.y = element_line(colour = "grey50"),
            plot.title = element_text(size = rel(1.5),
            face = "bold", vjust = 1.5),
            axis.title = element_text(face = "bold"),
            legend.position = "top",
            legend.title = element_blank(),
            legend.key.size = unit(0.4, "cm"),
            legend.key = element_rect(fill = "black"),
            axis.title.y = element_text(vjust= 1.8),
            axis.title.x = element_text(vjust= -0.5))
```



Bar plot of proportions vs. categories. Error bars indicate 95% confidence intervals for proportion.

Similar tests

See the *Handbook* for information on this topic.

How to do the test

R code for the SAS example is shown in the "Examples" section above.

Descriptive Statistics

Statistics of Central Tendency

Most common statistics of central tendency can be calculated with functions in the native *stats* package. The *psych* and *DescTools* packages add functions for the geometric mean and the harmonic mean. The *describe* function in the *psych* package includes the mean, median, and trimmed mean along with other common statistics. In the native *stats* package, *summary* is a quick way to see the mean, median, and quantiles for numeric variables in a data frame. The mode is not commonly calculated, but can be found in *DescTools*.

Many functions which determine common statistics of central tendency or dispersion will return an NA if there are any missing values (NA's) in the analyzed data. In most cases this behavior can be changed with the na.rm=TRUE option, which will simply exclude any NA's in the data. The functions shown here either exclude NA's by default or use the na.rm=TRUE option.

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Descriptive Statistics

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(psych)){install.packages("psych")}
if(!require(DescTools)){install.packages("DescTools")}
```

Introduction

The normal distribution

See the *Handbook* for information on these topics.

Different measures of central tendency

Methods are described in the "Example" section below.

Example

```
### Central tendency example, pp. 105-106
Input =("
Stream
                            Fish
Mill_Creek_1
                            76
Mill_Creek_2
                           102
North_Branch_Rock_Creek_1
                            12
North_Branch_Rock_Creek_2
                            39
                            55
Rock_Creek_1
Rock_Creek_2
                            93
                            98
Rock_Creek_3
Rock_Creek_4
                             53
```

```
Turkey_Branch 102
")
Data = read.table(textConnection(Input), header=TRUE)
```

Arithmetic mean

```
mean(Data$ Fish, na.rm=TRUE)
[1] 70
```

Geometric mean

```
library(psych)
geometric.mean(Data$ Fish)
   [1] 59.83515
library(DescTools)
Gmean(Data$ Fish)
   [1] 59.83515
```

Harmonic mean

```
library(psych)
harmonic.mean(Data$ Fish)
  [1] 45.05709
library(DescTools)
Hmean(Data$ Fish)
  [1] 45.05709
```

Median

```
median(Data$ Fish, na.rm=TRUE)
[1] 76
```

Mode

```
library(DescTools)
```

```
Mode(Data$ Fish)
[1] 102
```

Summary and describe functions for means, medians, and other statistics

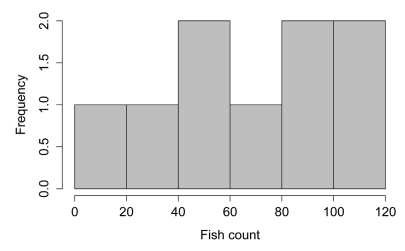
The interquartile range (IQR) is 3^{rd} Qu. minus 1^{st} Qu.

```
summary(Data$ Fish)
                           # Also works on whole data frames
                           # Will also report count of NA's
     Min. 1st Qu.
                  Median
                            Mean 3rd Qu.
                                           Max.
       12
               53
                      76
                              70
                                     98
                                            10
library(psych)
              type=2)
describe(Data$ Fish,
                          # Also works on whole data frames
                        # Type of skew and kurtosis
                  sd median trimmed mad min max range skew kurtosis
  1
     1 9 70 32.09
                      76
                                 70 34.1 12 102
                                                   90 -0.65
                                                              -0.69 10.7
```

Histogram

```
hist(Data$ Fish,
    col="gray",
    main="Maryland Biological Stream Survey",
    xlab="Fish count")
```

Maryland Biological Stream Survey



DescTools to produce summary statistics and plots

The *Desc* function in the package *DescTools* produces summary information for individual variables or whole data frames. It has custom output for factor, numeric, integer, and date variables.

```
### -----
### Central tendency example, pp. 105-106
### ------
Input =("
Stream
                           Fish
Mill_Creek_1
                           76
Mill_Creek_2
                          102
North_Branch_Rock_Creek_1 12
North_Branch_Rock_Creek_2
                           39
                           55
Rock_Creek_1
Rock_Creek_2
                           93
Rock_Creek_3
                           98
                           53
Rock_Creek_4
Turkey_Branch
                           102
")
Data = read.table(textConnection(Input),header=TRUE)
### Add a numeric variable with the same values as Fish
Data$Fish.num = as.numeric(Data$Fish)
### Produce summary statistics and plots
library(DescTools)
Desc(Data,
    plotit=TRUE)
   ______
   1 - Stream (factor)
     length n NAs levels unique dupes
          9
               9
                     0 9 9 n
                         level freq perc cumfreq cumperc
                  Mill_Creek_1 1 .111 1 .111
   1
  Mill_Creek_1 1 .111 1 .111

Mill_Creek_2 1 .111 2 .222

North_Branch_Rock_Creek_1 1 .111 3 .333

North_Branch_Rock_Creek_2 1 .111 4 .444

Rock_Creek_1 1 .111 5 .556

Rock_Creek_2 1 .111 6 .667

Rock_Creek_3 1 .111 7 .778

Rock_Creek_4 1 .111 8 .889

Turkey_Branch 1 .111 9 1.000
```

. < results snipped > .

.

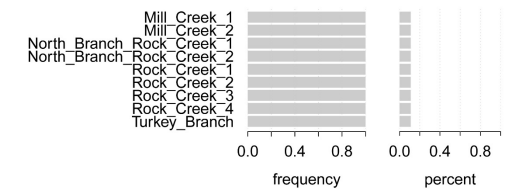
3 - Fish.num (numeric)

length	n	NAS	unique	0s		meanSE
9	9	0	8	0		10.695
.05	.10	.25	median	.75	.90	.95
22.800	33.600	53	76	98	102	102
rng	sd		mad	IQR	skew	kurt
90	32.086		34.100	45	-0.448	-1.389

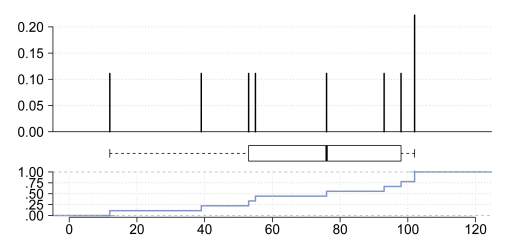
lowest: 12, 39, 53, 55, 76 highest: 55, 76, 93, 98, 102 (2)

Shapiro-Wilks normality test p.value: 0.23393

1 - Stream (factor)



3 - Fish.num (numeric)



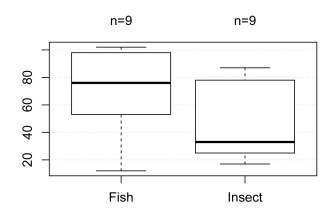
DescTools with grouped data

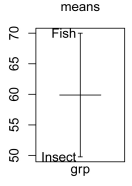
```
Summary statistics with grouped data, hypothetical data
Input =("
Stream
                            Animal
                                     Count
Mill_Creek_1
                            Fish
                                      76
Mill_Creek_2
                            Fish
                                     102
North_Branch_Rock_Creek_1
                            Fish
                                      12
North_Branch_Rock_Creek_2
                            Fish
                                      39
Rock_Creek_1
                            Fish
                                      55
Rock_Creek_2
                            Fish
                                      93
Rock_Creek_3
                            Fish
                                      98
Rock_Creek_4
                            Fish
                                      53
Turkey_Branch
                            Fish
                                     102
Mill_Creek_1
                            Insect
                                      28
Mill_Creek_2
                            Insect
                                      85
North_Branch_Rock_Creek_1
                            Insect
                                      17
North_Branch_Rock_Creek_2
                            Insect
                                      20
Rock_Creek_1
                                      33
                            Insect
Rock_Creek_2
                                      75
                            Insect
Rock_Creek_3
                                      78
                            Insect
Rock_Creek_4
                            Insect
                                      25
Turkey_Branch
                            Insect
                                      87
")
D2 = read.table(textConnection(Input),header=TRUE)
library(DescTools)
Desc(Count ~ Animal,
     D2,
```

```
digits=1,
  plotit=TRUE)
Count ~ Animal
Summary:
n pairs: 18, valid: 18 (100%), missings: 0 (0%), groups: 2
          Fish Insect
          70.0"
                   49.8'
mean
           76.0"
                   33.0'
median
           32.1
                   30.4
sd
IQR
          45.0
                   53.0
              9
                      9
         0.500
                  0.500
np
NAS
0s
              0
                      0
' min, " max
Kruskal-Wallis rank sum test:
```

Count ~ Animal

Kruskal-wallis chi-squared = 2.125, df = 1, p-value = 0.1449





How to calculate the statistics

Methods are described in the "Example" section above.

Statistics of Dispersion

Measures of dispersion—such as range, variance, standard deviation, and coefficient of variation—can be calculated with standard functions in the native *stats* package. In addition, a function, here called *summary.list*, can be defined to output whichever statistics are of interest.

Introduction

See the *Handbook* for information on this topic.

Example

Statistics of dispersion example

```
### ------
### Statistics of dispersion example, p. 111
### -----
Input =("
Stream
                    Fish
Mill_Creek_1
                     76
Mill_Creek_2
                    102
North_Branch_Rock_Creek_1
                     12
North_Branch_Rock_Creek_2
                     39
Rock_Creek_1
                     55
Rock_Creek_2
                     93
Rock_Creek_3
                     98
Rock_Creek_4
                     53
Turkey_Branch
                    102
")
Data = read.table(textConnection(Input), header=TRUE)
```

Range

```
range(Data$ Fish, na.rm=TRUE)

[1] 12 102  # Min and max

max(Data$ Fish, na.rm=TRUE) - min(Data$ Fish, na.rm=TRUE)

[1] 90
```

Sum of squares

Not included here.

Parametric variance

Not included here.

Sample variance

```
var(Data$ Fish, na.rm=TRUE)
[1] 1029.5
```

Standard deviation

```
sd(Data$ Fish, na.rm=TRUE)
[1] 32.08582
```

Coefficient of variation, as percent

```
sd(Data$ Fish, na.rm=TRUE)/
mean(Data$ Fish, na.rm=TRUE)*100
[1] 45.83689
```

Custom function of desired measures of central tendency and dispersion

```
### Note NA's removed in the following function
summary.list = function(x)list(
N.with.NA.removed= length(x[!is.na(x)]),
Count.of.NA= length(x[is.na(x)]),
Mean=mean(x, na.rm=TRUE),
Median=median(x, na.rm=TRUE),
Max.Min=range(x, na.rm=TRUE),
 Range=max(Data$ Fish, na.rm=TRUE) - min(Data$ Fish, na.rm=TRUE),
Variance=var(x, na.rm=TRUE),
 Std.Dev=sd(x, na.rm=TRUE),
Coeff.Variation.Prcnt=sd(x, na.rm=TRUE)/mean(x, na.rm=TRUE)*100,
 Std.Error=sd(x, na.rm=TRUE)/sqrt(length(x[!is.na(x)])),
Quantile=quantile(x, na.rm=TRUE)
)
summary.list(Data$ Fish)
   $N.with.NA.removed
   Γ11 9
   $Count.of.NA
   [1] 0
   $Mean
   [1] 70
   $Median
   [1] 76
   $Range
   [1] 12 102
   $Variance
   [1] 1029.5
   $Std.Dev
```

```
[1] 32.08582
$Coeff.Variation.Prcnt
[1] 45.83689
$Std.Error
[1] 10.69527
$Quantile
    0% 25% 50% 75% 100%
    12 53 76 98 102
```

How to calculate the statistics

Methods are described in the "Example" section above.

Standard Error of the Mean

The standard error of the mean can be calculated with standard functions in the native *stats* package. The *describe* function in the *psych* package includes the standard error of the mean along with other descriptive statistics. This function is useful to summarize multiple variables in a data frame.

Introduction Similar statistics

See the *Handbook* for information on these topics.

Example

Standard error example

```
### Standard error example, p. 115
Input =("
Stream
                           Fish
Mill_Creek_1
                            76
Mill_Creek_2
                           102
North_Branch_Rock_Creek_1
                            12
North_Branch_Rock_Creek_2
                            39
Rock_Creek_1
                            55
                            93
Rock_Creek_2
Rock_Creek_3
                            98
Rock_Creek_4
                            53
Turkey_Branch
                           102
")
Data = read.table(textConnection(Input), header=TRUE)
```

```
### Calculate standard error manually
sd(Data$ Fish, na.rm=TRUE) /
   sqrt(length(Data$Fish[!is.na(Data$ Fish)]))
                                             # Standard error
   [1] 10.69527
### Use describe function from psych package for standard error
### Also works on whole data frames
library(psych)
describe(Data$ Fish,
                             # Type of skew and kurtosis
              type=2)
                   sd median trimmed mad min max range skew kurtosis
             70 32.09
                          76
                                  70 34.1 12 102
                                                  90 -0.65
```

How to calculate the standard error

Methods are described in the "Example" section above.

Confidence Limits

Introduction

See the *Handbook* for information on this topic.

Confidence limits for measurement variables

Methods are described in the "How to calculate confidence limits" section below.

Confidence limits for nominal variables

Examples are given in the "How to calculate confidence limits" section below.

Statistical testing with confidence intervals Similar statistics

Examples

See the *Handbook* for information on these topics.

How to calculate confidence limits

The confidence limits about the mean—calculated using the *t*-value discussed in the *Handbook*—can be determined with variety of functions. One is *t.test* in the native *stats* package. Another is the *CI* function in the *Rmisc* package, which also has the function *summarySE* that presents the mean, standard deviation, standard error, and confidence interval for data designated as groups.

The bootstrap method noted in the *Handbook* can be achieved with the *boot* and *boot.ci* functions in the *boot* package.

Confidence intervals for mean with t.test, Rmisc, and DescTools

```
### Confidence interval for measurement data, blacknose fish , p. 120
### -----
Input =("
Stream
                         Fish
Mill_Creek_1
                        76
Mill_Creek_2
                        102
North_Branch_Rock_Creek_1
                         12
North_Branch_Rock_Creek_2 39
                         55
Rock_Creek_1
Rock_Creek_2
                         93
Rock_Creek_3
                         98
                         53
Rock_Creek_4
Turkey_Branch
                        102
")
Data = read.table(textConnection(Input), header=TRUE)
### Use t.test to produce confidence interval
t.test(Data$ Fish,
      conf.level=0.95) # Confidence interval of the mean
  95 percent confidence interval:
   45.33665 94.66335
### Use CI in Rmisc package to produce confidence interval
library(Rmisc)
CI(Data$ Fish,
                             # Confidence interval of the mean
  ci=0.95)
     upper mean lower
  94.66335 70.00000 45.33665
### Use MeanCI in DescTools package to produce confidence interval
library(DescTools)
MeanCI(Data$ Fish,
      conf.level=0.95) # Confidence interval of the mean
      mean lwr.ci upr.ci
  70.00000 45.33665 94.66335
```

Confidence intervals for means for grouped data

```
### -----
### Confidence interval for grouped data, hypothetical data
Input =("
Stream
                       Animal Count
Mill_Creek_1
                      Fish
                              76
Mill_Creek_2
North_Branch_Rock_Creek_1 Fish 12
North_Branch_Rock_Creek_2 Fish 39
Fish 55
Mill_Creek_2
                      Fish
                              102
                     Fish 93
Fish 98
Rock_Creek_2
                    Fish
Rock_Creek_3
                              53
Rock_Creek_4
                      Fish
Turkey_Branch
                      Fish 102
                      Insect 76
Mill_Creek_1
Mill_Creek_2
                      Insect 102
North_Branch_Rock_Creek_1 Insect
                               12
North_Branch_Rock_Creek_2 Insect
                               39
")
D2 = read.table(textConnection(Input),header=TRUE)
library(Rmisc)
summarySE(data=D2,
                            # Will produce confidence intervals
        measurevar="Count",
                          # for groups defined by a variable
        groupvars="Animal",
        conf.interval = 0.95)
    Animal N Count
                     sd
                             se
  1 Fish 9 70.00 32.08582 10.69527 24.66335
  2 Insect 4 57.25 39.72719 19.86360 63.21483
```

Confidence intervals for mean by bootstrap

```
### -----
### Confidence interval for measurement data, blacknose fish , p. 120
### -----
Input =("
                  Fish
Stream
Mill_Creek_1
                  76
Mill_Creek_2
                  102
North_Branch_Rock_Creek_1 12
North_Branch_Rock_Creek_2 39
                  55
Rock_Creek_1
                  93
Rock_Creek_2
```

```
Rock_Creek_3
                               98
   Rock_Creek_4
                               53
   Turkey_Branch
                              102
  Data = read.table(textConnection(Input),header=TRUE)
Confidence intervals for mean by bootstrap with DescTools
   MeanCI(Data$Fish, method="boot", type="norm", R=10000)
               lwr.ci
                          upr.ci
      70.00000 50.17986 89.84836
      # May be different for different iterations
  MeanCI(Data$Fish, method="boot", type="basic", R=10000)
         mean lwr.ci
                          upr.ci
      70.00000 51.44444 90.66667
      # May be different for different iterations
Confidence intervals for mean by bootstrap with boot package
   library(boot)
   Fun = function(x, index) {
                     return(c(mean(x[index]),
                              var(x[index]) / length(index)))
                       }
   Boot = boot(data=Data$Fish,
               statistic=Fun,
               R=10000)
  mean(Boot$t[,1])
      [1] 70.01229
                          # Mean by bootstrap
                          # May be different for different iterations
   boot.ci(Boot,
           conf=0.95)
      Intervals :
                 Normal
                                     Basic
                                                       Studentized
      Level
            (50.22, 89.76) (51.11, 90.44) (38.85, 91.72)
      95%
                Percentile
      Level
                                      вса
           (49.56, 88.89) (47.44, 87.22)
      Calculations and Intervals on Original Scale
```

Note that the bootstrapped confidence limits vary from
the calculated ones above because the original data set has
few values and is not necessarily normally distributed.

Confidence interval for proportions

The confidence interval for a proportion can be determined with the *binom.test* function, and more options are available in the *BinomCI* function and *MultinomCI* function in the *DescTools* package. More advanced techniques for confidence intervals on proportions and differences in proportions can be found in the *PropCIs* package.

```
### -----
### Confidence interval for nominal data, colorblind example, p. 118
### -----
binom.test(2, 20, 0.5,
        alternative="two.sided",
        conf.level=0.95)
  95 percent confidence interval:
   0.01234853 0.31698271
### ------
### Confidence interval for nominal data, Gus data, p. 121
Input =("
Paw
right
left
right
right
right
riaht
left
right
right
right
")
Gus = read.table(textConnection(Input),header=TRUE)
Successes = sum(Gus$ Paw == "left")  # Note the == operator
Failures = sum(Gus$ Paw == "right")
Total = Successes + Failures
Expected = 0.5
binom.test(Successes, Total, Expected,
        alternative="two.sided",
        conf.level=0.95)
```

```
95 percent confidence interval:
0.02521073 0.55609546
### Agrees with exact confidence interval from SAS
```

Confidence interval for proportions using DescTools

Confidence interval for single proportion

Confidence interval for multinomial proportion

Tests for One Measurement Variable

Student's t-test for One Sample

Introduction When to use it Null hypothesis How the test works Assumptions

See the *Handbook* for information on these topics.

Example

One sample t-test with observations as vector

Graphing the results

See the *Handbook* for information on this topic.

Similar tests

The *paired t-test* and *two-sample t-test* are presented elsewhere in this book.

How to do the test

One sample t-test with observations in data frame

```
### -----
### One-sample t-test, SAS example, pp. 125
### ------

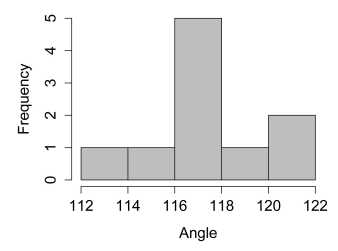
Input =("
Angle
120.6
116.4
117.2
118.1
114.1
```

```
116.9
113.3
121.1
116.9
117.0
")
Data = read.table(textConnection(Input), header=TRUE)
            = Data$ Angle
observed
theoretical = 50
t.test(observed,
       mu = theoretical,
       conf.int=0.95)
  One Sample t-test
   t = 87.3166, df = 9, p-value = 1.718e-14
   ### Does not agree with Handbook. The Handbook results are incorrect.
   ### The SAS code produces the following result.
                     T-Tests
     variable
                   DF
                         t Value
                                    Pr > |t|
     angle
                    9
                           87.32
                                      <.0001
```

Histogram

```
hist(Data$ Angle,
    col="gray",
    main="Histogram of values",
    xlab="Angle")
```

Histogram of values



Histogram of data in a single population from a one-sample t-test. Distribution of these values should be approximately normal.

Power analysis

Power analysis for one-sample t-test

```
### -----
### Power analysis, t-test, one-sample,
### hip joint example, pp. 125-126
M1 = 70
                               # Theoretical mean
M2 = 71
                               # Mean to detect
S1 = 2.4
                               # Standard deviation
S2 = 2.4
                               # Standard deviation
Cohen.d = (M1 - M2)/sqrt(((S1^2) + (S2^2))/2)
library(pwr)
pwr.t.test(
                                # Observations
       n = NULL
       d = Cohen.d,
       sig.level = 0.05, # Type I probability
power = 0.90, # 1 minus Type II probability
type = "one.sample", # Change for one- or two-sample
       alternative = "two.sided")
   One-sample t test power calculation
     n = 62.47518
```

Student's t-test for Two Samples

Introduction
When to use it
Null hypothesis
How the test works
Assumptions

See the *Handbook* for information on these topics.

Example

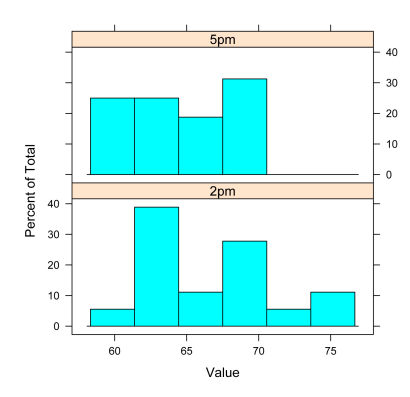
Two-sample t-test, independent (unpaired) observations

```
### ------
### Two-sample t-test, biological data analysis class, pp. 128-129
### ------
```

```
Input =("
Group Value
2pm
       69
       70
2pm
       66
2pm
2pm
       63
2pm
       68
2pm
       70
2pm
       69
       67
2pm
2pm
       62
2pm
       63
       76
2pm
       59
2pm
       62
2pm
       62
2pm
2pm
       75
       62
2pm
       72
2pm
2pm
       63
       68
5pm
5pm
       62
5pm
       67
5pm
       68
5pm
       69
       67
5pm
       61
5pm
5pm
       59
5pm
       62
       61
5pm
5pm
       69
       66
5pm
5pm
       62
5pm
       62
       61
5pm
       70
5pm
")
Data = read.table(textConnection(Input), header=TRUE)
bartlett.test(Value ~ Group, data=Data)
### If p-value >= 0.05, use var.equal=TRUE below
   Bartlett's K-squared = 1.2465, df = 1, p-value = 0.2642
t.test(Value ~ Group, data=Data,
       var.equal=TRUE,
       conf.level=0.95)
   Two Sample t-test
   t = 1.2888, df = 32, p-value = 0.2067
```

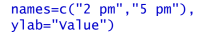
Plot of histograms

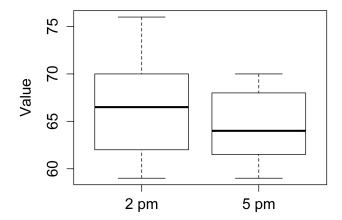
```
library(lattice)
```



Histograms for each population in a two-sample t-test. For the t-test to be valid, the data in each population should be approximately normal. If the distributions are different, minimally Welch's t-test should be used. If the data are not normal or the distributions are different, a non-parametric test like Mann-Whitney U-test or permutation test may be appropriate.

Box plots





Box plots of two populations from a two-sample t-test.

Similar tests

Welch's t-test is discussed below. The *paired t-test* and *signed-rank test* are discussed in this book in their own chapters. *Analysis of variance* (anova) is discussed in several subsequent chapters.

As non-parametric alternatives, the *Mann–Whitney U-test* and the *permutation test* for two independent samples are discussed in the chapter *Mann–Whitney and Two-sample Permutation Test*.

Welch's t-test

Welch's t-test is shown above in the "Example" section ("Two sample unpaired t-test"). It is invoked with the *var.equal=FALSE* option in the *t.test* function.

How to do the test

The SAS example from the *Handbook* is shown above in the "Example" section.

Power analysis

Power analysis for t-test

Mann-Whitney and Two-sample Permutation Test

The Mann–Whitney U-test is a nonparametric test, also called the Mann–Whitney–Wilcoxon test. It tests for a difference in central tendency of two groups, or, with certain assumptions, for the difference in medians. It is conducted with the *wilcox.test* function in the native *stats* package. It can be used with continuous or ordinal measurements.

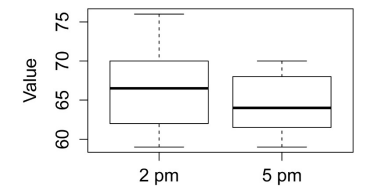
As another non-parametric alternative to t-tests, a permutation test can be used. An example is shown in the "Permutation test for independent samples" section of this chapter.

Mann-Whitney U-test

```
### Mann-Whitney U-test, biological data analysis class, pp. 128-129
Input =("
Group Value
2pm
       69
2pm
       70
2pm
       66
       63
2pm
2pm
       68
2pm
       70
2pm
       69
2pm
       67
       62
2pm
2pm
       63
2pm
       76
       59
2pm
2pm
       62
2pm
       62
       75
2pm
2pm
       62
       72
2pm
```

```
2pm
        63
5pm
        68
        62
5pm
5pm
        67
        68
5pm
        69
5pm
        67
5pm
        61
5pm
5pm
        59
5pm
        62
5pm
        61
        69
5pm
5pm
        66
5pm
        62
        62
5pm
        61
5pm
        70
5pm
")
Data = read.table(textConnection(Input), header=TRUE)
```

Box plots



```
wilcox.test(Value ~ Group, data=Data)
    wilcoxon rank sum test with continuity correction
    w = 186, p-value = 0.1485
```

Permutation test for independent samples

Permutation tests are nonparametric tests, and can be performed with the *coin* package. The permutation test compares values across groups, and can also be used to compare ranks or counts. This test is analogous to a nonparametric t-test. Normality is not assumed but the test

may require that distributions have similar variance or shape to be interpreted as a test of means.

```
### -----
### Two-sample permutation test, biological data analysis class,
    pp. 128-129
### -----
Input =("
Group Value
2pm
     69
2pm
     70
2pm
     66
2pm
      63
2pm
      68
2pm
      70
2pm
      69
2pm
      67
2pm
      62
2pm
      63
2pm
      76
2pm
      59
      62
2pm
2pm
      62
2pm
      75
      62
2pm
      72
2pm
      63
2pm
5pm
      68
5pm
      62
5pm
      67
5pm
      68
5pm
      69
5pm
      67
5pm
      61
5pm
      59
5pm
      62
      61
5pm
5pm
      69
5pm
      66
5pm
      62
5pm
      62
      61
5pm
5pm
      70
"<u>`</u>
Data = read.table(textConnection(Input), header=TRUE)
library(coin)
independence_test(Value ~ Group,
               data = Data)
  Asymptotic General Independence Test
```

Z = 1.2761, p-value = 0.2019

Chapters Not Covered in This Book

Introduction
Step-by-step analysis of biological data
Types of biological variables
Probability
Basic concepts of hypothesis testing
Confounding variables
Independence
Normality
Data transformations

See the *Handbook* for information on these topics.

Homoscedasticity and heteroscedasticity

Bartlett's test is performed with the *bartlett.test* function. Levene's test can be invoked with the *leveneTest* function in the *car* package. This test can also be used for a model with two independent variables. They are used in the chapter on *One-way anova*.

Type I, II, and III Sums of Squares

An in-depth discussion of Type I, II, and III sum of squares is beyond the scope of this book, but readers should at least be aware of them. They come into play in analysis of variance (anova) tables, when calculating sum of squares, F-values, and p-values.

Perhaps the most salient point for beginners is that SAS tends to use Type III by default whereas R will use Type I with the *anova* function. In R, Type II and Type III tests are accessed through *Anova* in the *car* package, as well as through some other functions for other types of analyses. However, for Type III tests to be correct, the way R codes factors has to be changed from its default with the *options(contrasts =...)* function. Changing this will not affect Type I or Type II tests.

Type I sum of squares are "sequential." In essence the factors are tested in the order they are listed in the model.

Type III sum of squares are "partial." In essence, every term in the model is tested in light of every other term in the model. That means that main effects are tested in light of interaction terms as well as in light of other main effects.

Type II sum of squares are similar to Type III, except that they preserve the principle of marginality. This means that main factors are tested in light of one another, but not in light of the interaction term.

When data are balanced and the design is simple, types I, II, and III will give the same results. But readers should be aware that results may differ for unbalanced data or more complex designs. The code below gives an example of this.

There are disagreements as to which type should be used routinely in analysis of variance. In reality, the user should understand what hypothesis she wants to test, and then choose the appropriate tests. As general advice, I would recommend not using Type I except in cases where you intend to have the effects assessed sequentially. Beyond that, probably a majority of those in the R community recommend Type II tests, while SAS users are more likely to consider Type III tests.

Some experimental designs will call for using a specified type of sum of squares, for example when you see "/SS1" or "HTYPE=1" in SAS code.

As a final note, readers should not confuse these sums of squares with "Type I error", which refers to rejecting a null hypothesis when it is actually true (a false positive), and "Type II error", which is failing to reject null hypothesis when it actually false (a false negative).

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(car)){install.packages("car")}
```

Setting contrasts for different sums of squares

```
options(contrasts = c("contr.sum", "contr.poly"))
  ### needed for type III tests

options(contrasts = c("contr.treatment", "contr.poly"))
  ### returns the default contrasts in R
```

Contrasts can also be set within a model.

Example for Type I, II, II sums of squares

Type I tests

```
model.1 = lm(Y ~ A + B + A:B)

anova(model)

Analysis of Variance Table

Df Sum Sq Mean Sq F value Pr(>F)
A 1 1488.37 1488.37 18.3892 0.002658 **
B 1 18.22 18.22 0.2252 0.647809
A:B 1 390.15 390.15 4.8204 0.059406 .
Residuals 8 647.50 80.94
```

Type II tests

Type III tests

```
model.3 = lm(Y \sim A + B + A:B,
             contrasts=list(A="contr.sum", B="contr.sum"))
library(car)
Anova(model.3, type="III")
   Anova Table (Type III tests)
                Sum Sq Df F value
                                      Pr(>F)
   (Intercept) 11343.8 1 140.1544 2.377e-06 ***
                1135.3 1 14.0275 0.005662 **
                  66.1 1
   В
                           0.8173
                                   0.392381
                 390.1 1
                           4.8204
                                   0.059406 .
   A:B
   Residuals
                647.5
```

One-way Anova

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Introduction to Parametric Tests

SAEEPER: One-way ANOVA

SAEEPER: What are Least Square Means?

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(dplyr)){install.packages("dplyr")}
if(!require(FSA)){install.packages("FSA")}
if(!require(car)){install.packages("car")}
if(!require(agricolae)){install.packages("agricolae")}
if(!require(multcomp)){install.packages("multcomp")}
if(!require(DescTools)){install.packages("DescTools")}
if(!require(lsmeans)){install.packages("lsmeans")}
if(!require(multcompView)){install.packages("multcompView")}
if(!require(Rmisc)){install.packages("Rmisc")}
if(!require(ggplot2)){install.packages("ggplot2")}
if(!require(pwr)){install.packages("pwr")}
```

When to use it

Analysis for this example is described below in the "How to do the test" section below.

Null hypothesis How the test works Assumptions Additional analyses

See the *Handbook* for information on these topics.

Tukey-Kramer test

The Tukey mean separation tests and others are shown below in the "How to do the test" section.

Partitioning variance

This topic is not covered here.

Example

Code for this example is not included here. An example is covered below in the "How to do the test" section.

Graphing the results

Graphing of the results is shown below in the "How to do the test" section.

Similar tests

Two-sample t-test, Two-way anova, Nested anova, Welch's anova, and Kruskal-Wallis are presented elsewhere in this book.

A *permutation test*, presented in the *One-way Analysis with Permutation Test* chapter, can also be employed as a nonparametric alternative.

How to do the test

The *lm* function in the native *stats* package fits a linear model by least squares, and can be used for a variety of analyses such as regression, analysis of variance, and analysis of covariance. The analysis of variance is then conducted either with the *Anova* function in the *car* package for Type II or Type III sum of squares, or with the *anova* function in the native *stats* package for Type I sum of squares.

If the analysis of variance indicates a significant effect of the independent variable, multiple comparisons among the levels of this factor can be conducted using Tukey or Least Significant Difference (LSD) procedures. The problem of inflating the Type I Error Rate when making multiple comparisons is discussed in the *Multiple Comparisons* chapter in the *Handbook*. R functions which make multiple comparisons usually allow for adjusting p-values. In R, the "BH", or "fdr", procedure is the Benjamini–Hochberg procedure discussed in the *Handbook*. See *?p.adjust* for more information.

One-way anova example

```
### One-way anova, SAS example, pp. 155-156
Input =("
Location Aam
Tillamook 0.0571
Tillamook 0.0813
Tillamook 0.0831
Tillamook 0.0976
Tillamook 0.0817
Tillamook 0.0859
Tillamook 0.0735
Tillamook 0.0659
Tillamook 0.0923
Tillamook 0.0836
Newport
          0.0873
          0.0662
Newport
          0.0672
Newport
          0.0819
Newport
Newport
          0.0749
Newport
          0.0649
          0.0835
Newport
Newport
          0.0725
Petersburg 0.0974
Petersburg 0.1352
Petersburg 0.0817
```

```
Petersburg 0.1016
Petersburg 0.0968
Petersburg 0.1064
Petersburg 0.1050
Magadan
        0.1033
Magadan
        0.0915
Magadan
         0.0781
Magadan
         0.0685
Magadan
         0.0677
Magadan
          0.0697
Magadan
         0.0764
Magadan
        0.0689
Tvarminne 0.0703
Tvarminne 0.1026
Tvarminne 0.0956
Tvarminne 0.0973
Tvarminne 0.1039
Tvarminne 0.1045
")
Data = read.table(textConnection(Input),header=TRUE)
```

Specify the order of factor levels for plots and Dunnett comparison

Produce summary statistics

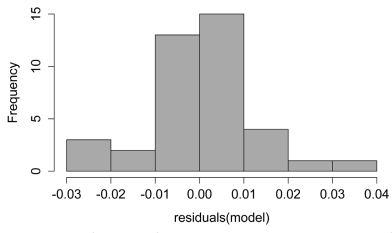
Fit the linear model and conduct ANOVA

```
### If you use type="III", you need the following line before the analysi
  ### options(contrasts = c("contr.sum", "contr.poly"))
               Sum Sq Df F value
  Location 0.0045197
                          7.121 0.0002812 ***
                      4
  Residuals 0.0053949 34
anova(model)
                                          # Produces type I sum of squares
                  Sum Sq
                            Mean Sq F value
            4 0.0045197 0.00112992
                                      7.121 0.0002812 ***
  Residuals 34 0.0053949 0.00015867
summary(model)
                  # Produces r-square, overall p-value, parameter estimates
  Multiple R-squared: 0.4559, Adjusted R-squared: 0.3918
   F-statistic: 7.121 on 4 and 34 DF, p-value: 0.0002812
```

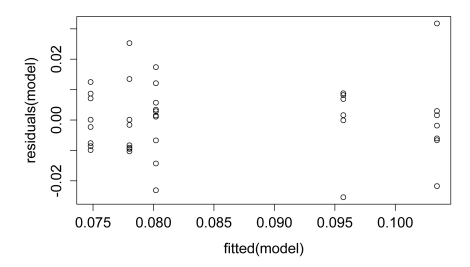
Checking assumptions of the model

```
hist(residuals(model),
      col="darkgray")
```

Histogram of residuals(model)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

```
### additional model checking plots with: plot(model)
### alternative: library(FSA); residPlot(model)
```

Tukey and Least Significant Difference mean separation tests (pairwise comparisons)

Tukey and other multiple comparison tests can be performed with a handful of functions. The functions *TukeyHSD*, *HSD.test*, and *LSD.test* are probably not appropriate for cases where there are unbalanced data or unequal variances among levels of the factor, though *TukeyHSD* does make an adjustment for mildly unbalanced data. It is my understanding that the *multcomp* and *Ismeans* packages are more appropriate for unbalanced data. Another alternative is the *DTK* package that performs mean separation tests on data with unequal sample sizes and no assumption of equal variances.

Tukey comparisons in agricolae package

```
library(agricolae)

(HSD.test(model, "Location"))  # outer parentheses print result

trt means M

1 Petersburg 0.1034429 a

2 Tvarminne 0.0957000 ab

3 Tillamook 0.0802000 bc

4 Magadan 0.0780125 bc

5 Newport 0.0748000 c
```

Means sharing the same letter are not significantly different

LSD comparisons in agricolae package

```
library(agricolae)
(LSD.test(model, "Location",
                               # outer parentheses print result
           alpha = 0.05,
           p.adj="none"))
                               # see ?p.adjust for options
            trt
                    means M
   1 Petersburg 0.1034429 a
   2 Tvarminne 0.0957000 a
   3 Tillamook 0.0802000 b
   4 Magadan
                0.0780125 b
   5 Newport
                0.0748000 b
   # Means sharing the same letter are not significantly different
```

Multiple comparisons in *multcomp* package

Note that "Tukey" here does not mean Tukey-adjusted comparisons. It just sets up a matrix to compare each mean to each other mean.

```
library(multcomp)
mc = glht(model,
         mcp(Location = "Tukey"))
mcs = summary(mc, test=adjusted("single-step"))
mcs
   ### Adjustment options: "none", "single-step", "Shaffer",
                           "westfall", "free", "holm", "hochberg",
   ###
                           "hommel", "bonferroni", "BH", "BY", "fdr"
   ###
Linear Hypotheses:
                             Estimate Std. Error t value Pr(>|t|)
Newport - Tillamook == 0
                            -0.005400
                                        0.005975 -0.904 0.89303
Petersburg - Tillamook == 0 0.023243
                                        0.006208
                                                 3.744 0.00555 **
Magadan - Tillamook == 0
                            -0.002188
                                        0.005975 -0.366 0.99596
Tvarminne - Tillamook == 0
                                                   2.383
                             0.015500
                                        0.006505
                                                         0.14413
Petersburg - Newport == 0
                             0.028643
                                        0.006519
                                                  4.394
                                                         < 0.001 ***
Magadan - Newport == 0
                             0.003213
                                        0.006298
                                                  0.510
                                                         0.98573
Tvarminne - Newport == 0
                             0.020900
                                        0.006803
                                                  3.072
                                                         0.03153 *
Magadan - Petersburg == 0
                            -0.025430
                                        0.006519 -3.901 0.00376 **
Tvarminne - Petersburg == 0 -0.007743
                                        0.007008
                                                 -1.105 0.80211
Tvarminne - Magadan == 0
                            0.017688
                                        0.006803
                                                 2.600 0.09254 .
cld(mcs.
    level=0.05.
    decreasing=TRUE)
                Newport Petersburg
   Tillamook
                                      Magadan Tvarminne
                     "c"
                                "a"
                                          "bc"
         "bc"
                                                     "ab"
```

Means sharing a letter are not significantly different

Multiple comparisons to a control in *multcomp* package

```
### Control is the first level of the factor
library(multcomp)
mc = glht(model,
          mcp(Location = "Dunnett"))
summary(mc, test=adjusted("single-step"))
   ### Adjustment options: "none", "single-step", "Shaffer",
                           "Westfall", "free", "holm", "hochberg",
   ###
                           "hommel", "bonferroni", "BH", "BY", "fdr"
   ###
   Linear Hypotheses:
                                Estimate Std. Error t value Pr(>|t|)
                               -0.005400 0.005975 -0.904 0.79587
   Newport - Tillamook == 0
                                                     3.744 0.00252 **
   Petersburg - Tillamook == 0 0.023243
                                           0.006208
   Magadan - Tillamook == 0 -0.002188 0.005975 -0.366 0.98989
   Tvarminne - Tillamook == 0 \quad 0.015500 \quad 0.006505 \quad 2.383 \quad 0.07794.
```

Multiple comparisons to a control with Dunnett Test

```
### The control group can be specified with the control option,
     or will be the first level of the factor
library(DescTools)
DunnettTest(Aam ~ Location,
           data = Data
    Dunnett's test for comparing several treatments with a control :
      95% family-wise confidence level
                              diff
                                         lwr.ci
                                                    upr.ci
                                                             pval
  Newport-Tillamook
                       -0.00540000 -0.020830113 0.01003011 0.7958
  Petersburg-Tillamook 0.02324286 0.007212127 0.03927359 0.0026 **
  Magadan-Tillamook -0.00218750 -0.017617613 0.01324261 0.9899
                        0.01550000 -0.001298180 0.03229818 0.0778 .
  Tvarminne-Tillamook
```

Multiple comparisons with least square means

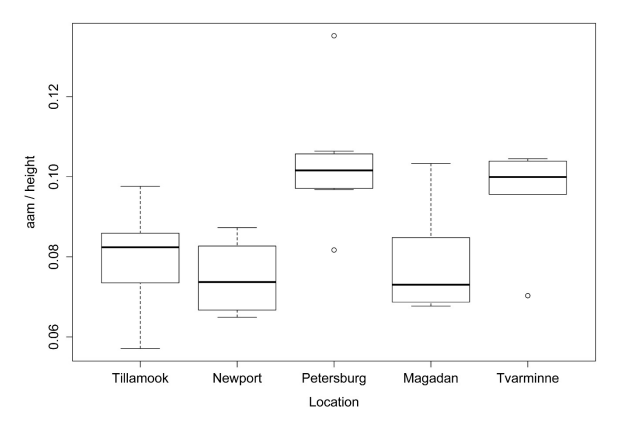
Least square means can be calculated for each group. Here a Tukey adjustment is applied for multiple comparisons among group least square means. The multiple comparisons can be displayed as a compact letter display.

```
library(lsmeans)
library(multcompView)
```

```
leastsquare = lsmeans(model,
                     pairwise ~ Location.
                     adjust = "tukey")
   $contrasts
                                                SE df t.ratio p.value
   contrast
                              estimate
   Tillamook - Newport
                           0.005400000 0.005975080 34
                                                        0.904 0.8935
   Tillamook - Petersburg -0.023242857 0.006207660 34
                                                       -3.744 0.0057
   Tillamook - Magadan
                           0.002187500 0.005975080 34
                                                        0.366 0.9960
   Tillamook - Tvarminne -0.015500000 0.006504843 34
                                                       -2.383 0.1447
   Newport - Petersburg -0.028642857 0.006519347 34 -4.394 0.0009
                          -0.003212500 0.006298288 34
                                                      -0.510 0.9858
   Newport - Magadan
   Newport - Tvarminne
                                                      -3.072 0.0317
                          -0.020900000 0.006802928 34
   Petersburg - Magadan 0.025430357 0.006519347 34
                                                        3.901 0.0037
   Petersburg - Tvarminne 0.007742857 0.007008087 34
                                                        1.105 0.8028
                          -0.017687500 0.006802928 34 -2.600 0.0929
   Magadan - Tvarminne
   P value adjustment: tukey method for comparing a family of 5 estimates
cld(leastsquare,
   alpha
          = 0.05.
   Letters = letters,
   adjust="tukey")
   Location
                lsmean
                                SE df
                                        lower.CL
                                                   upper.CL .group
   Newport
              0.0748000 0.004453562 34 0.06268565 0.08691435
   Magadan
              0.0780125 0.004453562 34 0.06589815 0.09012685
   Tillamook 0.0802000 0.003983387 34 0.06936459 0.09103541
   Tvarminne 0.0957000 0.005142530 34 0.08171155 0.10968845
                                                               bc
   Petersburg 0.1034429 0.004761058 34 0.09049207 0.11639365
  Confidence level used: 0.95
  Conf-level adjustment: sidak method for 5 estimates
   P value adjustment: tukey method for comparing a family of 5 estimates
   significance level used: alpha = 0.05
```

Graphing the results

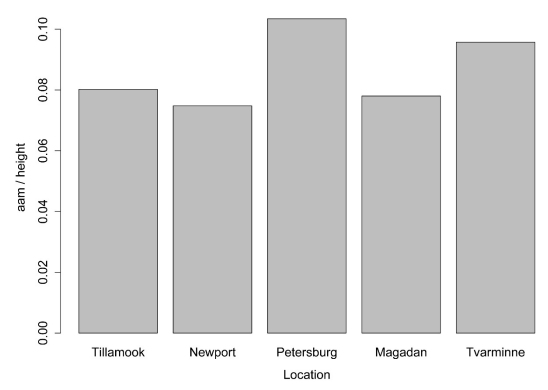
Simple box plots of values across groups



Box plots of values for each level of the independent variable for a one-way analysis of variance (ANOVA).

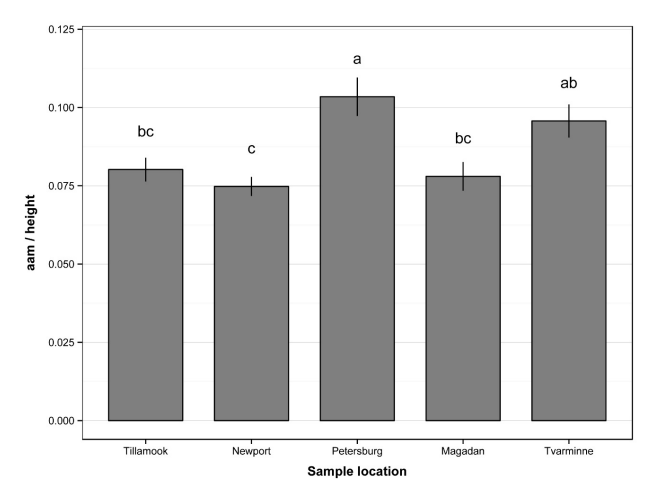
Simple bar plot of means across groups

```
### Summarize the data frame (Data) into a table
library(Rmisc)
Data2 = summarySE(data=Data,
          "Aam",
          groupvars="Location",
          conf.interval = 0.95)
Tabla = as.table(Data2$Aam)
rownames(Tabla) = Data2$Location
Tabla
   Tillamook
                                      Magadan Tvarminne
                Newport Petersburg
    0.0802000 0.0748000 0.1034429
                                     0.0780125 0.0957000
barplot(Tabla,
        ylab="aam / height",
        xlab="Location")
```



Bar plot of means for each level of the independent variable for a one-way analysis of variance (ANOVA).

Bar plot of means with error bars across groups



Bar plot of means for each level of the independent variable of a one-way analysis of variance (ANOVA). Error indicates standard error of the mean. Bars sharing the same letter are not significantly different according to Tukey's HSD test.

Welch's anova

Bartlett's test and Levene's test can be used to check the homoscedasticity of groups from a one-way anova. A significant result for these tests (p < 0.05) suggests that groups are heteroscedastic. One approach with heteroscedastic data in a one way anova is to use the Welch correction with the *oneway.test* function in the native *stats* package. A more versatile approach is to use the *white.adjust=TRUE* option in the *Anova* function from the *car* package.

```
### Levene test for homogeneity of variance
library(car)
leveneTest(Aam ~ Location,
           data = Data
   Levene's Test for Homogeneity of Variance (center = median)
         Df F value Pr(>F)
   group 4 0.12 0.9744
         34
### Welch's anova for unequal variances
oneway.test(Aam ~ Location,
            data=Data,
            var.equal=FALSE)
  One-way analysis of means (not assuming equal variances)
   F = 5.6645, num df = 4.000, denom df = 15.695, p-value = 0.00508
### white-adjusted anova for heteroscedasticity
model = lm(Aam \sim Location,
           data=Data)
library(car)
Anova(model, Type="II",
      white.adjust=TRUE)
                  F Pr(>F)
   Location 4 5.4617 0.001659 **
   Residuals 34
```

Power analysis

Power analysis for one-way anova

```
### ----
### Power analysis for anova, pp. 157
### ----
library(pwr)

groups = 5
means = c(10, 10, 15, 15, 15)
sd = 12

grand.mean = mean(means)
Cohen.f = sqrt( sum( (1/groups) * (means-grand.mean)^2) ) /sd
```

Kruskal-Wallis Test

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Kruskal-Wallis Test

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(dplyr)){install.packages("dplyr")}
if(!require(FSA)){install.packages("FSA")}
if(!require(DescTools)){install.packages("DescTools")}
if(!require(rcompanion)){install.packages("rcompanion")}
if(!require(multcompView)){install.packages("multcompView")}
```

When to use it

See the *Handbook* for information on this topic.

Null hypothesis

This example shows just summary statistics, histograms by group, and the Kruskal–Wallis test. An example with plots, post-hoc tests, and alternative tests is shown in the "Example" section below.

Kruskal-Wallis test example

```
### -----
### Kruskal-wallis test, hypothetical example, p. 159
### -----

Input =("
Group Value
Group.1 1
Group.1 2
Group.1 3
Group.1 4
```

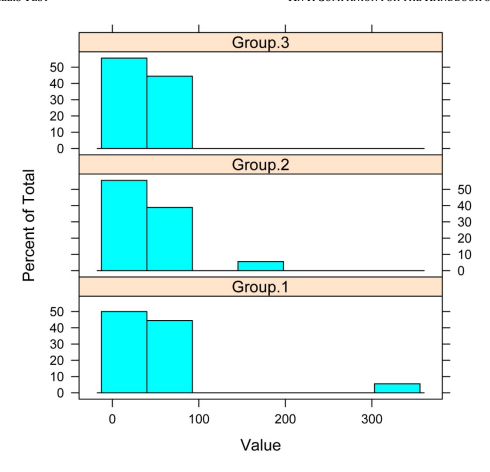
```
Group.1
              5
              6
Group.1
              7
Group.1
              8
Group.1
              9
Group.1
             46
Group.1
Group.1
             47
Group.1
             48
Group.1
             49
             50
Group.1
Group.1
             51
             52
Group.1
Group.1
             53
Group.1
            342
Group.2
             10
Group.2
             11
Group.2
             12
Group.2
             13
Group.2
             14
Group.2
             15
Group.2
             16
Group.2
             17
Group.2
             18
Group.2
             37
             58
Group.2
             59
Group.2
Group.2
             60
Group.2
             61
Group.2
             62
Group.2
             63
Group.2
             64
            193
Group.2
Group.3
             19
Group.3
             20
Group.3
             21
             22
Group.3
Group.3
             23
Group.3
             24
Group.3
             25
             26
Group.3
Group.3
             27
Group.3
             28
Group.3
             65
Group.3
             66
Group.3
             67
Group.3
             68
Group.3
             69
Group.3
             70
Group.3
             71
Group.3
             72
")
```

Data = read.table(textConnection(Input),header=TRUE)

Medians and descriptive statistics

As noted in the *Handbook*, each group has identical medians and means.

Histograms for each group



Kruskal-Wallis test

In this case, there is a significant difference in the distributions of values among groups, as is evident both from the histograms and from the significant Kruskal–Wallis test. Only in cases where the distributions in each group are similar can a significant Kruskal–Wallis test be interpreted as a difference in medians.

How the test works Assumptions

See the *Handbook* for information on these topics.

Example

The Kruskal–Wallis test is performed on a data frame with the *kruskal.test* function in the native *stats* package. Shown first is a complete example with plots, post-hoc tests, and alternative methods, for the example used in R help. It is data measuring if the mucociliary efficiency in the rate of dust removal is different among normal subjects, subjects with obstructive airway disease, and subjects with asbestosis. For the original citation, use the *?kruskal.test* command.

For both the submissive dog example and the oyster DNA example from the *Handbook*, a Kruskal–Wallis test is shown later in this chapter.

Kruskal-Wallis test example

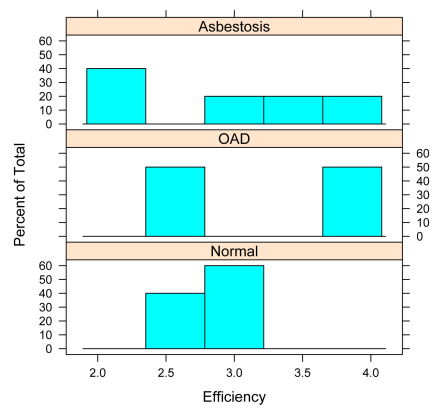
```
### -----
### Kruskal-wallis test, asbestosis example from R help for
### kruskal.test
Input =("
   Health Efficiency
Normal 2.9
Obs Health
1
             3.0
2
   Normal
3 Normal 2.5
4 Normal 2.6
5 Normal 3.2
6 OAD 3.8
   OAD 3.0
OAD 2.7
OAD 4.0
OAD 2.4
7
8
9
10 Asbestosis 2.8
11 Asbestosis 3.4
12 Asbestosis 3.7
13 Asbestosis 2.2
14 Asbestosis 2.0
")
Data = read.table(textConnection(Input), header=TRUE)
### Specify the order of factor levels
library(dplyr)
Data =
mutate(Data,
      Health = factor(Health, levels=unique(Health)))
```

Medians and descriptive statistics

Graphing the results

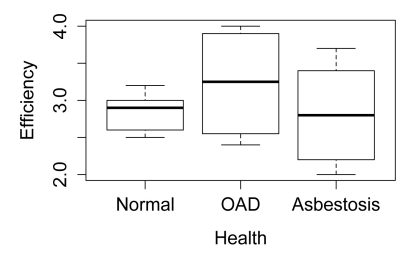
Stacked histograms of values across groups

```
library(lattice)
```



Stacked histograms for each group in a Kruskal–Wallis test. If the distributions are similar, then the Kruskal–Wallis test will test for a difference in medians.

Simple box plots of values across groups



Kruskal-Wallis test

Dunn test for multiple comparisons

If the Kruskal–Wallis test is significant, a post-hoc analysis can be performed to determine which levels of the independent variable differ from each other level. Probably the most popular test for this is the Dunn test, which is performed with the *dunnTest* function in the *FSA* package. Adjustments to the p-values could be made using the *method* option to control the familywise error rate or to control the false discovery rate. See *?p.adjust* for details.

Zar (2010) states that the Dunn test is appropriate for groups with unequal numbers of observations.

If there are several values to compare, it can be beneficial to have R convert this table to a compact letter display for you. The *cldList* function in the *rcompanion* package can do this.

Nemenyi test for multiple comparisons

Zar (2010) suggests that the Nemenyi test is not appropriate for groups with unequal numbers of observations.

```
library(DescTools)
PT = NemenyiTest(x = Data$Efficiency,
                 g = Data$Health,
                 dist="tukey")
PT
  Nemenyi's test of multiple comparisons for independent samples (tukey)
                    mean.rank.diff
                                      pval
  OAD-Normal
                               1.8 0.7972
   Asbestosis-Normal
                              -0.6 0.9720
   Asbestosis-OAD
                              -2.4 0.6686
library(rcompanion)
cldList(comparison = PT$Comparison,
        p.value = PT$P.adi.
        threshold = 0.05)
   Error: No significant differences.
```

Pairwise Mann-Whitney U-tests

Another post-hoc approach is to use pairwise Mann–Whitney U-tests. To prevent the inflation of type I error rates, adjustments to the p-values can be made using the *p.adjust.method* option to control the familywise error rate or to control the false discovery rate. See ?*p.adjust* for details.

If there are several values to compare, it can be beneficial to have R convert this table to a compact letter display for you. The *multcompLetters* function in the *multcompView* package can do this, but first the table of p-values must be converted to a full table.

```
PT = pairwise.wilcox.test(Data$Efficiency,
                           Data$Health,
                           p.adjust.method="none")
                               # Can adjust p-values;
                               # See ?p.adjust for options
PT
   Pairwise comparisons using Wilcoxon rank sum test
              Normal OAD
   OAD
              0.73
   Asbestosis 1.00
                     0.41
PT = PT$p.value
library(rcompanion)
PT1 = fullPTable(PT)
PT1
                 Normal
                               OAD Asbestosis
   Normal
              1.0000000 0.7301587 1.0000000
              0.7301587 1.0000000 0.4126984
   Asbestosis 1.0000000 0.4126984 1.0000000
library(multcompView)
multcompLetters(PT1,
                compare="<",</pre>
                threshold=0.05,
                Letters=letters.
                reversed = FALSE)
       Normal
                     OAD Asbestosis
                                 "a"
          "a"
                     "a"
       ### Values sharing the same letter are not significantly different
```

Kruskal-Wallis test example

```
### ------
### Kruskal-Wallis test, submissive dog example, pp. 161-162
Input =("
Dog
            Sex
                     Rank
Merlino Male
                      1
                      2
Gastone
            Male
        Male
Male
Male
                      3
 Pippo
 Leon
                      4
                      5
 Golia
 Lancillotto Male
                      6
           Female 7
Female 8
Female 9
Female 10
Male 11
                      7
Mamy
             Female
Nanà
 Isotta
Diana
 Simba
 Pongo
             Male
                     12
 Semola
             Male
                     13
 Kimba
             Male
                     14
Morgana Female 15
Stella Female 16
Hansel
            Male
                     17
Cucciola
Mammolo
             Male
                     18
Mammolo
             Male
                     19
             Male 20
Dotto
Gongolo Male 21
Gretel Female 22
Brontolo Female 23
 Eolo
             Female 24
           Female 25
Female 26
Mag
 Emy
 Pisola
             Female 27
 ")
Data = read.table(textConnection(Input), header=TRUE)
kruskal.test(Rank ~ Sex,
             data = Data
   Kruskal-Wallis chi-squared = 4.6095, df = 1, p-value = 0.03179
```

Graphing the results

Graphing of the results is shown above in the "Example" section.

Similar tests

One-way anova is presented elsewhere in this book.

How to do the test Kruskal-Wallis test example

```
### Kruskal-wallis test, oyster DNA example, pp. 163-164
Input =("
   Markername Markertype fst

        CVB1
        DNA
        -0.005

        CVB2m
        DNA
        0.116

        CVJ5
        DNA
        -0.006

        CVJ6
        DNA
        0.095

        CVL1
        DNA
        0.053

        CVL3
        DNA
        0.003

        6Pgd
        protein
        -0.005

        Aat-2
        protein
        0.016

        Acp-3
        protein
        0.041

        Adk-1
        protein
        0.066

        Est-1
        protein
        0.066

        Est-3
        protein
        0.004

        Lap-1
        protein
        0.049

        Lap-2
        protein
        0.058

        Pgi
        protein
        -0.002

        Pgm-1
        protein
        0.015

        Pgm-2
        protein
        0.044

        Sdh
        protein
        0.024

   CVB1
                                       DNA -0.005
   Sdh
                                                                             0.024
                                        protein
")
Data = read.table(textConnection(Input),header=TRUE)
kruskal.test(fst ~ Markertype,
                                        data = Data
         Kruskal-wallis chi-squared = 0.0426, df = 1, p-value = 0.8365
```

Power Analysis

See the *Handbook* for information on this topic.

References

Zar, J.H. 2010. Biostatistical Analysis, 5th ed. Pearson Prentice Hall: Upper Saddle River, NJ.

One-way Analysis with Permutation Test

Permutation tests are non-parametric tests that do not assume normally-distributed errors. However, these tests may assume that distributions have similar variance or shape to be

interpreted as a test of means.

A one-way anova using permutation tests can be performed with the *coin* package. A post-hoc analysis can be conducted with pairwise permutation tests analagous to pairwise t-tests. This can be accomplished with the functions *pairwisePermutationTest* and *pairwisePermutationMatrix* in the *rcompanion* package, which rely on the *independence_test* function in the *coin* package.

For more information on permutation tests available in the *coin* package, see:

```
help(package="coin")
```

Consult the chapters on *One-way Anova* and *Kruskal–Wallis Test* for general consideration about conducting analysis of variance.

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Permutation Test of Independence

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(coin)){install.packages("coin")}
if(!require(FSA)){install.packages("FSA")}
if(!require(rcompanion)){install.packages("rcompanion")}
if(!require(multcompView)){install.packages("multcompView")}
```

Permutation test for one-way analysis

```
### One-way permutation test, hypothetical data
Input =("
Factor Response
  Α
        4.6
  Α
        5.5
        3.4
  Α
        5.0
  Α
  Α
        3.9
        4.5
  Α
        3.6
  В
        4.5
  В
        2.4
  В
        4.0
  В
  В
        2.9
        3.5
  В
  C
        2.6
        3.5
  C
  C
        1.4
  C
        3.0
  C
        1.9
  C
        2.5
```

```
4.7
  D
  D
        5.6
  D
        3.5
  D
        5.1
        4.0
  D
  D
        4.6
Data = read.table(textConnection(Input), header=TRUE)
Data$Factor = factor(Data$Factor,
                      ordered=FALSE,
                      levels=unique(Data$Factor))
    # Order factors, otherwise R will alphabetize them
boxplot(Response ~ Factor,
        data = Data,
        ylab ="Response",
        xlab ="Factor")
       2
   Response
       4
       က
       2
                 Α
                                                    D
                             В
                                        C
```

Permutation test

Pairwise permutation tests

Pairwise permutation tests could be used as a post-hoc test for a significant permutation test. If no p-value adjustment is made, then the type I error rate may be inflated due to multiple comparisons. Here, the "fdr" p-value adjustment method is used to control the false discovery

Factor

rate.

<u>Table output with pairwisePermutationTest</u>

```
### Order groups by median
Data$Factor = factor(Data$Factor,
             levels = c("D", "A", "B", "C"))
library(FSA)
headtail(Data)
### Pairwise tests
library(rcompanion)
PT = pairwisePermutationTest(Response ~ Factor,
                           data = Data,
                           method="fdr")
PT
    Comparison Stat p.value p.adjust
  1 D - A = 0 -0.2409 0.8096 0.80960
  2 D - B = 0 -2.074 0.03812 0.06106
  3 D - C = 0 -2.776 0.005505 0.01876
  4 A - B = 0 1.952 0.05088 0.06106
  5 A - C = 0 2.734 0.006253 0.01876
  6 B - C = 0 1.952 0.05088 0.06106
library(rcompanion)
cldList(p.adjust ~ Comparison,
       data = PT,
       threshold = 0.05)
      Group Letter MonoLetter
  1
      D
             a
  2
       Α
             a
                        a
  3
        В
                        ab
              ab
  4
        C
             b
                         b
```

<u>Compact letter display output with pairwisePermutationMatrix</u>

```
### Pairwise tests
library(rcompanion)
PM = pairwisePermutationMatrix(Response ~ Factor,
                                data = Data.
                                method="fdr")
PM
   $Unadjusted
                     В
   D NA 0.8096 0.03812 0.005505
            NA 0.05088 0.006253
   A NA
   B NA
                    NA 0.050880
   C NA
            NA
                    NA
   $Method
   [1] "fdr"
   $Adjusted
  D 1.00000 0.80960 0.06106 0.01876
   A 0.80960 1.00000 0.06106 0.01876
   в 0.06106 0.06106 1.00000 0.06106
  C 0.01876 0.01876 0.06106 1.00000
library(multcompView)
multcompLetters(PM$Adjusted,
                compare="<",</pre>
                threshold=0.05,
                Letters=letters,
                reversed = FALSE)
    D A B C "a" "ab" "b"
```

Nested Anova

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Using Random Effects in Models SAEEPER: What are Least Square Means?

SAEEPER: One-way ANOVA with Random Blocks

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(nlme)){install.packages("nlme")}
if(!require(multcomp)){install.packages("multcomp")}
if(!require(multcompview)){install.packages("multcompview")}
if(!require(lsmeans)){install.packages("lsmeans")}
if(!require(lme4)){install.packages("lme4")}
if(!require(lmerTest)){install.packages("lmerTest")}
if(!require(TukeyC)){install.packages("TukeyC")}
```

When to use it
Null hypotheses
How the test works
Partitioning variance and optimal allocation of resources
Unequal sample sizes
Assumptions
Example
Graphing the results
Similar tests
See the Handbook for information on these topics.

How to do the test

Nested anova example with mixed effects model (nlme)

One approach to fit a nested anova is to use a mixed effects model. Here *Tech* is being treated as a fixed effect, while *Rat* is treated as a random effect. Note that the F-value and p-value for the test on *Tech* agree with the values in the *Handbook*. The effect of *Rat* will be tested by comparing this model to a model without the *Rat* term. The model is fit using the *lme* function in *nlme*.

```
### Nested anova, SAS example, pp. 171-173
Input =("
Tech Rat Protein
 Janet 1 1.119
 Janet 1 1.2996
 Janet 1 1.5407
 Janet 1 1.5084
 Janet 1 1.6181
 Janet 1 1.5962
 Janet 1 1.2617
 Janet 1 1.2288
 Janet 1 1.3471
 Janet 1 1.0206
 Janet 2 1.045
 Janet 2 1.1418
 Janet 2 1.2569
 Janet 2 0.6191
 Janet 2 1.4823
```

```
Janet 2 0.8991
 Janet 2 0.8365
 Janet 2
          1.2898
 Janet 2
          1.1821
 Janet 2 0.9177
 Janet 3
         0.9873
 Janet 3
         0.9873
 Janet 3
          0.8714
 Janet 3
          0.9452
 Janet 3
          1.1186
          1.2909
 Janet 3
 Janet 3 1.1502
 Janet 3 1.1635
 Janet 3
          1.151
 Janet 3 0.9367
 Brad 5 1.3883
 Brad 5
          1.104
 Brad 5
          1.1581
 Brad 5 1.319
 Brad 5 1.1803
 Brad 5 0.8738
 Brad 5
         1.387
 Brad 5 1.301
 Brad 5
          1.3925
 Brad 5
          1.0832
 Brad 6 1.3952
 Brad 6 0.9714
 Brad 6
          1.3972
 Brad 6
         1.5369
 Brad 6 1.3727
 Brad 6 1.2909
 Brad 6 1.1874
 Brad 6 1.1374
 Brad 6 1.0647
 Brad 6 0.9486
 Brad 7
          1.2574
 Brad 7 1.0295
 Brad 7
         1.1941
 Brad 7
          1.0759
 Brad 7
          1.3249
 Brad 7 0.9494
 Brad 7 1.1041
 Brad 7 1.1575
 Brad
     7
          1.294
     7
 Brad
          1.4543
")
Data = read.table(textConnection(Input), header=TRUE)
### Since Rat is read in as an integer variable, convert it to factor
Data$Rat = as.factor(Data$Rat)
library(nlme)
```

Post-hoc comparison of means

Note that "Tukey" here instructs the *glht* function to compare all means, not to perform a Tukey adjustment of multiple comparisons.

```
library(multcomp)
posthoc = glht(model,
               linfct = mcp(Tech="Tukey"))
mcs = summary(posthoc,
              test=adjusted("single-step"))
mcs
   ### Adjustment options: "none", "single-step", "Shaffer",
                           "Westfall", "free", "holm", "hochberg",
   ###
                           "hommel", "bonferroni", "BH", "BY", "fdr"
   ###
   Linear Hypotheses:
                     Estimate Std. Error z value Pr(>|z|)
   Janet - Brad == 0 - 0.05060 0.09781 - 0.517
cld(mcs.
    level=0.05.
    decreasing=TRUE)
    Brad Janet
     "a"
       ### Means sharing a letter are not significantly different
```

Post-hoc comparison of least-square means

Least squares means are adjusted for other terms in the model. If the experimental design is unbalanced or there is missing data, the least square means may differ significantly from arithmetic means for treatments, but are generally more representative of the population means than the arithmetic means would be.

Note that the adjustments for multiple comparisons (adjust ="tukey") appears in both the *Ismeans* and *cld* functions.

```
library(multcompView)
library(lsmeans)
leastsquare = lsmeans(model,
                       pairwise ~ Tech,
                       adjust="tukey")
                                              ### Tukey-adjusted comparisons
leastsquare
   $1smeans
    Tech
            lsmean
                          SE df lower.CL upper.CL
    Brad 1.211023 0.06916055 5 1.0332405 1.388806
    Janet 1.160420 0.06916055 4 0.9683995 1.352440
   Confidence level used: 0.95
   $contrasts
                                     SE df t.ratio p.value
    contrast
                   estimate
    Brad - Janet 0.05060333 0.09780778 4 0.517 0.6322
cld(leastsquare,
    alpha=0.05,
    Letters=letters, ### Use lower-case letters for .group adjust="tukey") ### Tukey-adjusted comparisons
   Tech
                           SE df asymp.LCL asymp.UCL .group
           lsmean
    Janet 1.160420 0.06916018 NA 1.005745 1.315095 a
    Brad 1.211023 0.06916018 NA 1.056348 1.365698 a
       ### Means sharing a letter in .group are not significantly different
```

Test the significance of the random effect in the mixed effects model

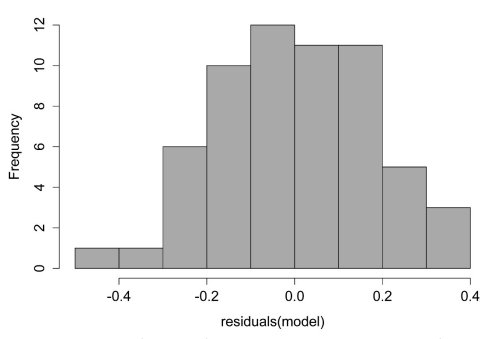
In order to the test the significance of the random effect from our model (*Rat*), we can fit a new model with only the fixed effects from the model. For this we use the *gls* function in the *nlme* package. We then compare the two models with the *anova* fuction. Note that the p-value does not agree with p-value from the *Handbook*, because the technique is different, though in this case the conclusion is the same. As a general precaution, if your models are fit with "REML" (restricted maximum likelihood) estimation, then you should compare only models with the same fixed effects. If you need to compare models with different fixed effects, use "ML" as the estimation method for all models.

```
Model df AIC BIC logLik Test L.Ratio p-value model 1 4 -7.819054 0.4227176 7.909527 model.fixed 2 3 -4.499342 1.6819872 5.249671 1 vs 2 5.319713 0.0211
```

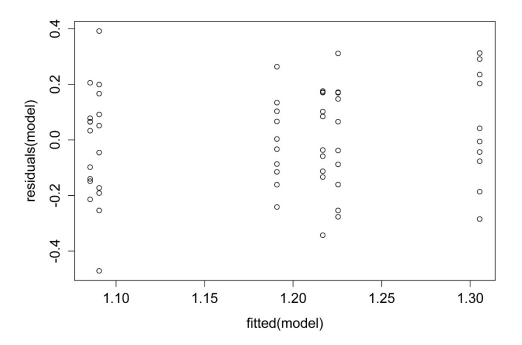
Checking assumptions of the model

```
hist(residuals(model),
      col="darkgray")
```

Histogram of residuals(model)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

additional model checking plots with: plot(model)

Mixed effects model with Imer

The following is an abbreviated example of a nested anova using the *lmer* function in the *lme4* package. See the previous example in this chapter for explanation and model-checking.

```
### Nested anova, SAS example, pp. 171-173
Input =("
 Tech
       Rat Protein
           1.119
 Janet 1
           1.2996
 Janet 1
 Janet 1
           1.5407
           1.5084
 Janet 1
 Janet 1
           1.6181
 Janet 1
           1.5962
           1.2617
 Janet 1
 Janet 1
           1.2288
 Janet 1
           1.3471
           1.0206
 Janet 1
           1.045
 Janet 2
 Janet 2
           1.1418
           1.2569
 Janet 2
 Janet 2
           0.6191
           1.4823
 Janet 2
 Janet 2
           0.8991
```

```
0.8365
 Janet 2
          1.2898
 Janet 2
 Janet 2
          1.1821
 Janet 2
          0.9177
          0.9873
 Janet 3
 Janet 3
          0.9873
          0.8714
 Janet 3
 Janet 3
          0.9452
 Janet 3
          1.1186
 Janet 3
          1.2909
 Janet 3
          1.1502
 Janet 3
          1.1635
          1.151
 Janet 3
 Janet 3
          0.9367
 Brad 5
          1.3883
 Brad 5
          1.104
 Brad 5
          1.1581
      5
          1.319
 Brad
 Brad 5
          1.1803
 Brad 5
          0.8738
          1.387
 Brad
      5
 Brad
      5
          1.301
      5
          1.3925
 Brad
 Brad
      5
          1.0832
          1.3952
 Brad
      6
          0.9714
 Brad
      6
         1.3972
 Brad 6
 Brad 6
          1.5369
 Brad
      6
          1.3727
 Brad 6 1.2909
 Brad
      6
         1.1874
 Brad
      6
          1.1374
 Brad
      6
          1.0647
 Brad
      6
          0.9486
          1.2574
 Brad
      7
          1.0295
 Brad
      7
          1.1941
 Brad
      7
 Brad 7
          1.0759
 Brad
     7
          1.3249
 Brad 7
          0.9494
 Brad 7
          1.1041
 Brad 7
          1.1575
      7
          1.294
 Brad
Brad
          1.4543
")
Data = read.table(textConnection(Input), header=TRUE)
Data$Rat = as.factor(Data$Rat)
library(lme4)
library(lmerTest)
model = lmer(Protein ~ Tech + (1|Rat),
            data=Data,
```

```
REML=TRUE)
anova(model)
   Analysis of Variance Table of type III with Satterthwaite
   approximation for degrees of freedom
          Sum Sq Mean Sq NumDF DenDF F.value Pr(>F)
  Tech 0.0096465 0.0096465 1 4 0.26768 0.6322
rand(model)
  Analysis of Random effects Table:
      Chi.sq Chi.DF p.value
   Rat 5.32 1 0.02 *
difflsmeans(model,
           test.effs="Tech")
  Differences of LSMEANS:
                  Estimate Standard Error DF t-value Lower CI Upper CI p-value
  Tech Brad - Janet 0.1 0.0978 4.0 0.52 -0.221 0.322 0.6
library(multcomp)
posthoc = glht(model,
              linfct = mcp(Tech="Tukey"))
mcs = summary(posthoc,
             test=adjusted("single-step"))
mcs
   Linear Hypotheses:
                    Estimate Std. Error z value Pr(>|z|)
   Janet - Brad = 0 - 0.05060 \quad 0.09781 \quad -0.517 \quad 0.605
   (Adjusted p values reported -- single-step method)
cld(mcs,
    level=0.05,
   decreasing=TRUE)
   Brad Janet
    "a" "a"
```

Nested anova example with the aov function

```
### ------### Nested anova, SAS example, pp. 171-173
```

```
Input =("
Tech Rat Protein
Janet 1
         1.119
         1.2996
Janet 1
Janet 1
         1.5407
Janet 1
         1.5084
Janet 1
         1.6181
Janet 1
         1.5962
Janet 1
         1.2617
         1.2288
Janet 1
         1.3471
Janet 1
Janet 1
         1.0206
Janet 2
         1.045
Janet 2
         1.1418
Janet 2
         1.2569
Janet 2
         0.6191
Janet 2
         1.4823
Janet 2
         0.8991
         0.8365
Janet 2
Janet 2
         1.2898
Janet 2
         1.1821
Janet 2
         0.9177
         0.9873
Janet 3
         0.9873
Janet 3
         0.8714
Janet 3
Janet 3
         0.9452
Janet 3
         1.1186
Janet 3
         1.2909
Janet 3
         1.1502
Janet 3
         1.1635
Janet 3
         1.151
Janet 3
         0.9367
         1.3883
Brad 5
Brad 5
         1.104
         1.1581
Brad 5
Brad 5
         1.319
Brad 5
         1.1803
Brad 5
         0.8738
Brad 5
         1.387
Brad 5
         1.301
         1.3925
Brad 5
Brad 5
         1.0832
         1.3952
Brad 6
Brad 6
         0.9714
         1.3972
Brad 6
         1.5369
Brad 6
         1.3727
Brad 6
         1.2909
Brad 6
         1.1874
Brad 6
Brad 6
         1.1374
Brad 6
         1.0647
Brad 6
         0.9486
Brad 7
         1.2574
```

```
1.0295
Brad 7
         1.1941
Brad 7
Brad 7
         1.0759
Brad 7
         1.3249
         0.9494
Brad 7
Brad 7
         1.1041
Brad 7
         1.1575
Brad 7
         1.294
Brad 7 1.4543
")
Data = read.table(textConnection(Input), header=TRUE)
### Since Rat is read in as an integer variable, convert it to factor
Data$Rat = as.factor(Data$Rat)
```

Using the aov function for a nested anova

The *aov* function in the native stats package allows you to specify an error component to the model. When formulating this model in R, the correct error is *Rat*, not *Tech/Rat* (Rat within Tech) as used in the SAS example. The SAS model will tolerate *Rat* or *Rat(Tech)*.

The summary of the *aov* will produce the correct test for *Tech*. The test for *Rat* can be performed by manually calculating the p-value for the F-test using the output for *Error:Rat* and *Error:Within*.

See the rattlesnake example in the *Two-way anova* chapter for designating an error term in a repeated-measures model.

Using Mean Sq and Df values to get p-value for H = Tech and Error = Rat

```
pf(q=0.03841/0.14349,
    df1=1,
    df2=4,
    lower.tail=FALSE)

[1] 0.6321845

### Note: This is same test as summary(fit)
```

<u>Using Mean Sq and Df values to get p-value for H = Rat and Error = Within</u>

Post-hoc comparison of means with Tukey

The *aov* function with an *Error* component produces an object of *aovlist* type, which unfortunately isn't handled by many post-hoc testing functions. However, in the *TukeyC* package, you can specify a model and error term. For unbalanced data, the *dispersion* parameter may need to be modified.

Two-way Anova

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Two-way ANOVA
SAEEPER: Using Random Eff

SAEEPER: Using Random Effects in Models SAEEPER: What are Least Square Means?

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(FSA)){install.packages("FSA")}
if(!require(ggplot2)){install.packages("ggplot2")}
if(!require(car)){install.packages("car")}
if(!require(multcompView)){install.packages("multcompView")}
if(!require(lsmeans)){install.packages("lsmeans")}
if(!require(grid)){install.packages("grid")}
if(!require(nlme)){install.packages("nlme")}
if(!require(lme4)){install.packages("lme4")}
if(!require(lmerTest)){install.packages("lmerTest")}
if(!require(rcompanion)){install.packages("rcompanion")}
```

When to use it Null hypotheses How the test works Assumptions

See the *Handbook* for information on these topics.

Examples

The rattlesnake example is shown at the end of the "How to do the test" section.

How to do the test

For notes on linear models and conducting anova, see the "How to do the test" section in the *One-way anova* chapter of this book. For two-way anova with robust regression, see the chapter on *Two-way Anova with Robust Estimation*.

Two-way anova example

```
### Two-way anova, SAS example, pp. 179-180
Input = ("
id Sex
          Genotype Activity
 1 male
          ff
                   1.884
 2 male
          ff
                   2.283
 3 male
          fs
                  2.396
 4 female ff
                  2.838
 5 male
          fs
                   2.956
 6 female ff
                   4.216
 7 female ss
                   3.620
 8 female ff
                   2.889
 9 female fs
                   3.550
 10 male
                   3.105
          fs
11 female fs
                  4.556
                   3.087
12 female fs
13 male
          ff
                   4.939
14 male ff
                   3.486
 15 female ss
                   3.079
```

1 female

3 female

male

male

```
16 male
               fs
                         2.649
    17 female
               fs
                         1.943
    19 female
               ff
                         4.198
    20 female
               ff
                         2.473
    22 female
               ff
                         2.033
    24 female
               fs
                         2.200
    25 female
               fs
                         2.157
    26 male
               SS
                         2.801
    28 male
                         3.421
               SS
    29 female
               ff
                         1.811
    30 female
                         4.281
               fs
    32 female
               fs
                         4.772
    34 female
               SS
                         3.586
    36 female
               ff
                         3.944
    38 female
                         2.669
               SS
    39 female ss
                         3.050
    41 male
                         4.275
               SS
    43 female
               SS
                         2.963
    46 female
                         3.236
               SS
    48 female ss
                         3.673
   49 male
                         3.110
               SS
   ")
   Data = read.table(textConnection(Input), header=TRUE)
Means and summary statistics by group
   library(FSA)
   Sum = Summarize(Activity ~ Sex + Genotype,
                  data = Data
   Sum
           Sex Genotype n
                                                      Q1 median
                             mean
                                          sd
                                               min
                                                                    Q3
                                                                         max
                     ff 8 3.05025 0.9599032 1.811 2.363 2.864 4.008 4.216
                     ff 4 3.14800 1.3745115 1.884 2.183 2.884 3.849 4.939
          male
                     fs 8 3.31825 1.1445388 1.943 2.189 3.318 4.350 4.772
      3 female
                     fs 4 2.77650 0.3168433 2.396 2.586 2.802 2.993 3.105
          male
      5 female
                    ss 8 3.23450 0.3617754 2.669 3.028 3.158 3.594 3.673
                     ss 4 3.40175 0.6348109 2.801 3.033 3.266 3.634 4.275
          male
   ### Add standard error
   Sum$se = Sum$sd/sqrt(Sum$n)
   Sum
          Sex Genotype n
                                           min
                                                  01 median
                           mean
                                      sd
                                                              03
```

ff 8 3.05025 0.9599032 1.811 2.363 2.864 4.008 4.216 0.3393770 ff 4 3.14800 1.3745115 1.884 2.183 2.884 3.849 4.939 0.6872558

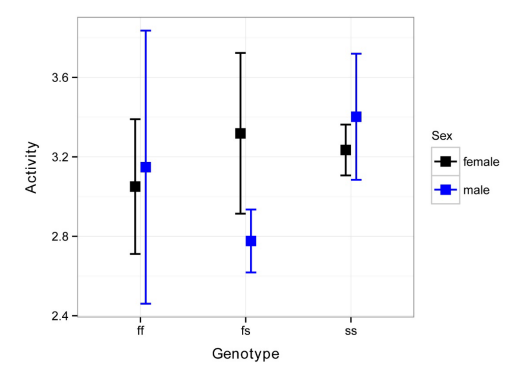
fs 8 3.31825 1.1445388 1.943 2.189 3.318 4.350 4.772 0.4046556

fs 4 2.77650 0.3168433 2.396 2.586 2.802 2.993 3.105 0.1584216

```
5 female ss 8 3.23450 0.3617754 2.669 3.028 3.158 3.594 3.673 0.1279069 6 male ss 4 3.40175 0.6348109 2.801 3.033 3.266 3.634 4.275 0.3174054
```

Interaction plot using summary statistics

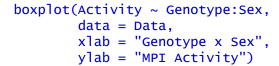
You may see an error, "ymax not defined"
In this case, it does not appear to affect anything

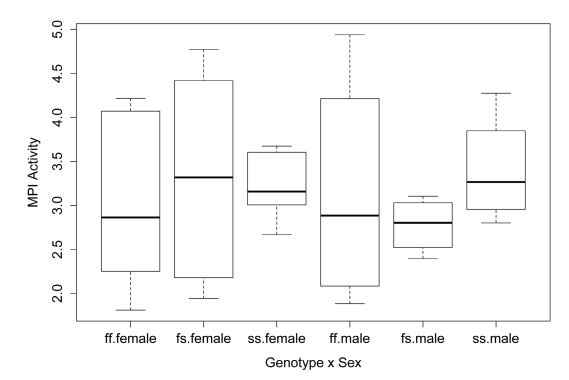


Interaction plot for a two-way anova. Square points represent means for groups, and error bars indicate standard errors of the mean.

Simple box plot of main effect and interaction

Genotype





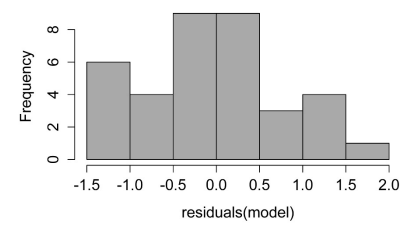
Fit the linear model and conduct ANOVA

```
model = lm(Activity ~ Sex + Genotype + Sex:Genotype,
          data=Data)
library(car)
Anova(model,
     type="II") ### Type II sum of squares
  ### If you use type="III", you need the following line before the analysi
  ### options(contrasts = c("contr.sum", "contr.poly"))
                Sum Sq Df F value Pr(>F)
                0.0681 1 0.0861 0.7712
  Sex
                0.2772 2 0.1754 0.8400
  Genotype
  Sex:Genotype 0.8146 2 0.5153 0.6025
  Residuals 23.7138 30
anova(model)
               # Produces type I sum of squares
               Df Sum Sq Mean Sq F value Pr(>F)
               1 0.0681 0.06808 0.0861 0.7712
  Sex
               2 0.2772 0.13862 0.1754 0.8400
  Genotype
  Sex:Genotype 2 0.8146 0.40732 0.5153 0.6025
              30 23.7138 0.79046
  Residuals
summary(model) # Produces r-square, overall p-value, parameter estimates
  Multiple R-squared: 0.04663, Adjusted R-squared: -0.1123
   F-statistic: 0.2935 on 5 and 30 DF, p-value: 0.9128
```

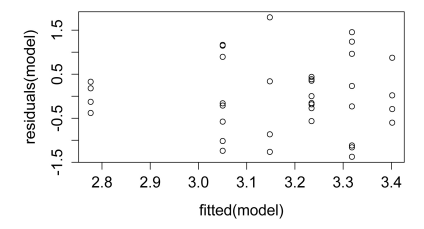
Checking assumptions of the model

```
hist(residuals(model),
      col="darkgray")
```

Histogram of residuals(model)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

additional model checking plots with: plot(model)
alternative: library(FSA); residPlot(model)

Post-hoc comparison of least-square means

For notes on least-square means, see the "Post-hoc comparison of least-square means" section in the *Nested anova* chapter in this book.

One advantage of the using the *Ismeans* package for post-hoc tests is that it can produce comparisons for interaction effects.

In general, if the interaction effect is significant, you will want to look at comparisons of means for the interactions. If the interaction effect is not significant but a main effect is, it is appropriate to look at comparisons among the means for that main effect. In this case, because no effect of *Sex, Genotype*, or *Sex:Genotype* was significant, we would not actually perform any mean separation test.

Mean separations for main factor with Ismeans

```
library(multcompview)
library(lsmeans)
lsmeans = lsmeans::lsmeans ### Uses the lsmeans function
                           ### from the 1smeans package,
                           ### not from the lmerTest package
leastsquare = lsmeans(model,
                      pairwise ~ Genotype,
                      adjust="tukey")
cld(leastsquare,
    alpha=.05,
    Letters=letters.
    adjust="tukey")
   Genotype lsmean
                            SE df lower.CL upper.CL .group
    fs
             3.047375 0.2722236 30 2.359065 3.735685 a
    ff
             3.099125 0.2722236 30 2.410815 3.787435 a
             3.318125 0.2722236 30 2.629815 4.006435 a
    SS
       ### Means sharing a letter in .group are not significantly different
```

Mean separations for interaction effect with *Ismeans*

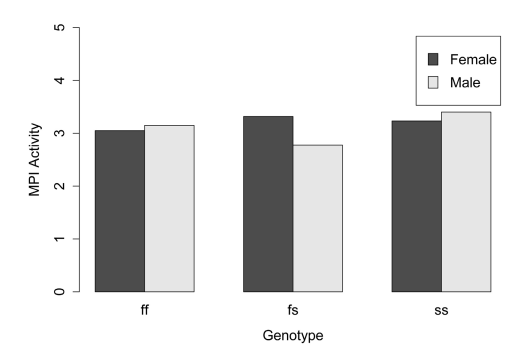
```
library(multcompView)
library(lsmeans)
lsmeans = lsmeans::lsmeans ### Uses the lsmeans function
                           ### from the 1smeans package,
                           ### not from the lmerTest package
leastsquare = lsmeans(model,
                      pairwise ~ Sex:Genotype,
                      adjust="tukey")
cld(leastsquare,
    alpha=.05,
    Letters=letters,
    adjust="tukey")
   Sex
         Genotype lsmean
                                  SE df lower.CL upper.CL .group
   male
                   2.77650 0.4445393 30 1.524666 4.028334 a
```

```
female ff 3.05025 0.3143368 30 2.165069 3.935431 a male ff 3.14800 0.4445393 30 1.896166 4.399834 a female ss 3.23450 0.3143368 30 2.349319 4.119681 a female fs 3.31825 0.3143368 30 2.433069 4.203431 a male ss 3.40175 0.4445393 30 2.149916 4.653584 a ### Note that means are listed from low to high, ### not in the same order as Summarize
```

Graphing the results

Simple bar plot with categories and no error bars

```
### Re-enter data as matrix
Input =("
        ff
Sex
                 fs
                          SS
Female 3.05025 3.31825 3.23450
Male
        3.14800 2.77650 3.40175
")
Matriz = as.matrix(read.table(textConnection(Input),
                   header=TRUE,
                   row.names=1))
Matriz
               ff
                       fs
   Female 3.05025 3.31825 3.23450
   Male
         3.14800 2.77650 3.40175
barplot(Matriz,
        beside=TRUE,
        legend=TRUE,
        ylim=c(0, 5),
        xlab="Genotype",
        ylab="MPI Activity")
```

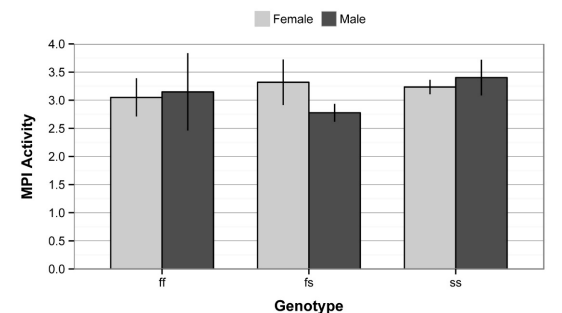


Bar plot with error bars with ggplot2

This plot uses the data frame created by *Summarize* in *FSA*. Error bars indicate standard error of the means (*se* in the data frame).

```
library(FSA)
Sum = Summarize(Activity ~ Sex + Genotype,
                data = Data
Sum
### Add standard error
Sum$se = Sum$sd/sqrt(Sum$n)
Sum
        Sex Genotype n
                                     sd
                                         min
                                                Q1 median
                                                             Q3
   1 female
                ff 8 3.05025 0.9599032 1.811 2.363 2.864 4.008 4.216 0.3393770
      male
                 ff 4 3.14800 1.3745115 1.884 2.183 2.884 3.849 4.939 0.6872558
                 fs 8 3.31825 1.1445388 1.943 2.189 3.318 4.350 4.772 0.4046556
   3 female
      male
                 fs 4 2.77650 0.3168433 2.396 2.586 2.802 2.993 3.105 0.1584216
   5 female
                 ss 8 3.23450 0.3617754 2.669 3.028 3.158 3.594 3.673 0.1279069
                 ss 4 3.40175 0.6348109 2.801 3.033 3.266 3.634 4.275 0.3174054
      male
### Plot adapted from:
      shinyapps.stat.ubc.ca/r-graph-catalog/
library(ggplot2)
library(grid)
```

```
ggplot(Sum,
  aes(x = Genotype,
      y = mean.
      fill = Sex,
      ymax=mean+se,
      ymin=mean-se)) +
      show.legend = FALSE) +
      scale_y\_continuous(breaks = seq(0, 4, 0.5),
               limits = c(0, 4),
               expand = c(0, 0)) +
      scale_fill_manual(name = "Count type"
                values = c('grey80', 'grey30'),
                labels = c("Female"
                          "Male")) +
      geom_errorbar(position=position_dodge(width=0.7),
                   width=0.0, size=0.5, color="black")
      labs(x = "Genotype",
           y = "MPI Activity") +
      ## ggtitle("Main title") +
      theme_bw() +
      theme(panel.grid.major.x = element_blank(),
            panel.grid.major.y = element_line(colour = "grey50"),
            plot.title = element_text(size = rel(1.5),
            face = "bold", vjust = 1.5),
            axis.title = element_text(face = "bold"),
            legend.position = "top",
            legend.title = element_blank(),
            legend.key.size = unit(0.4, "cm"),
            legend.key = element_rect(fill = "black"),
            axis.title.y = element_text(vjust= 1.8),
            axis.title.x = element_text(vjust= -0.5)
          )
```



Bar plot for a two-way anova. Bar heights represent means for groups, and error bars indicate standard errors of the mean.

Rattlesnake example - two-way anova without replication, repeated measures

This example could be interpreted as two-way anova without replication or as a one-way repeated measures experiment. Below it is analyzed as a two-way fixed effects model using the *lm* function, and as a mixed effects model using the *nlme* package and *lme4* packages.

```
### Two-way anova, rattlesnake example, pp. 177-178
Input = ("
 Day Snake Openings
      D1
                85
               107
 1
      D3
 1
      D5
                 61
 1
                 22
      D8
 1
      D11
                40
                 65
 1
      D12
 2
      D1
                 58
 2
      D3
                 51
 2
      D5
                 60
 2
      D8
                41
 2
      D11
                45
 2
                 27
      D12
 3
      D1
                15
 3
                 30
      D3
 3
      D5
                 68
 3
      D8
                 63
 3
                 28
      D11
 3
      D12
                 3
 4
                 57
      D1
 4
                12
      D3
 4
      D5
                36
 4
      D8
                 21
 4
      D11
                 10
      D12
                 16
")
Data = read.table(textConnection(Input), header=TRUE)
### Treat Day as a factor variable
Data$Day = as.factor(Data$Day)
```

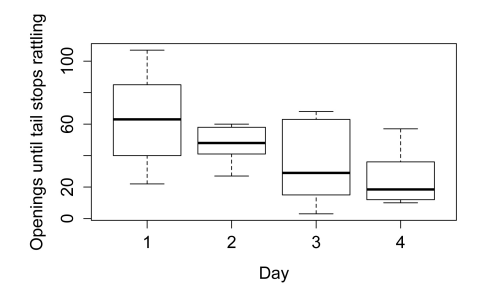
Using two-way fixed effects model

Means and summary statistics by group

```
library(FSA)
```

```
Sum = Summarize(Openings ~ Day,
                data = Data)
Sum
    Day n
               mean
                          sd min
                                    Q1 median
                                                  Q3 max
      1 6 63.33333 30.45434
                             22 45.25
                                         63.0 80.00 107
       2 6 47.00000 12.21475
                              27 42.00
                                         48.0 56.25
                              3 18.25
       3 6 34.50000 25.95958
                                         29.0 54.75
                                                      68
       4 6 25.33333 18.08498
                             10 13.00
                                         18.5 32.25
                                                      57
```

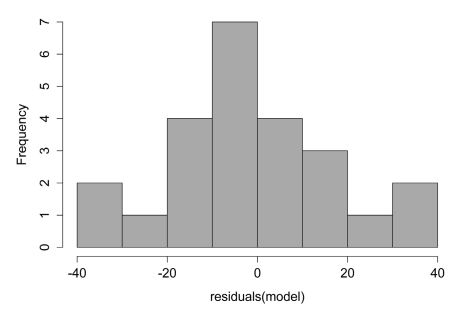
Simple box plots



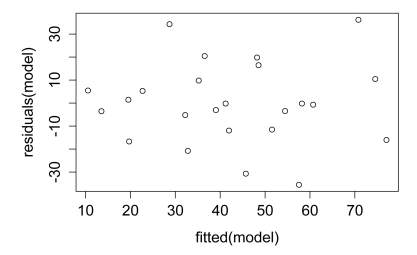
Fit the linear model and conduct ANOVA

Checking assumptions of the model

Histogram of residuals(model)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

```
### additional model checking plots with: plot(model)
### alternative: library(FSA); residPlot(model)
```

Mean separations for main factor with *Ismeans*

For notes on least-square means, see the "Post-hoc comparison of least-square" means section in the *Nested anova* chapter in this book. For other mean separation techniques for a main factor in anova, see "Tukey and Least Significant Difference mean separation tests (pairwise comparisons)" section in the *One-way anova* chapter.

```
library(multcompView)
library(lsmeans)
lsmeans = lsmeans::lsmeans ### Uses the lsmeans function
                                from the 1smeans package.
                           ###
                           ###
                                not from the lmerTest package
leastsquare = lsmeans(model,
                      pairwise ~ Day,
                      adjust="tukey")
cld(leastsquare,
    alpha=.05,
    Letters=letters,
    adjust="tukey")
   Day
         1smean
                      SE df
                              lower.CL upper.CL .group
        25.33333 9.034476 15 -0.2085871 50.87525
    3
        34.50000 9.034476 15
                             8.9580796 60.04192
    2
        47.00000 9.034476 15 21.4580796 72.54192
                                                   ab
        63.33333 9.034476 15 37.7914129 88.87525
```

Means sharing a letter in .group are not significantly different

Using mixed effects model with nlme

This is an abbreviated example using the *lme* function in the *nlme* package.

```
library(nlme)
model = lme(Openings ~ Day, random=~1|Snake,
           data=Data.
           method="REML")
anova.lme(model,
          type="sequential",
          adjustSigma = FALSE)
              numDF denDF F-value p-value
   (Intercept) 1 15 71.38736 <.0001
                  3
                       15 3.32005 0.0487
   Day
library(multcompView)
library(lsmeans)
lsmeans = lsmeans::lsmeans ### Uses the lsmeans function
                          ### from the 1smeans package,
                          ### not from the lmerTest package
leastsquare = lsmeans(model,
                     pairwise ~ Day,
                     alpha=.05,
                     adjust="tukey")
cld(leastsquare,
    alpha=.05,
    Letters=letters,
    adjust="tukey")
                    SE df lower.CL upper.CL .group
   Day
        lsmean
       25.33333 9.304196 5 -9.9416542 60.60832 a
    3
       34.50000 9.304196 5 -0.7749876 69.77499 ab
      47.00000 9.304196 5 11.7250124 82.27499 ab
       63.33333 9.304196 5 28.0583458 98.60832
     ### Means sharing a letter in .group are not significantly different
```

Using mixed effects model with Imer

This is an abbreviated example using the *lmer* function in the *lme4* package.

```
data=Data,
              REML=TRUE)
  anova(model)
     Analysis of Variance Table of type III with Satterthwaite
     approximation for degrees of freedom
         Sum Sq Mean Sq NumDF DenDF F.value Pr(>F)
     Day 4877.8 1625.9 3 15 3.3201 0.04866 *
  rand(model)
     Analysis of Random effects Table:
           Chi.sq Chi.DF p.value
     Snake 0.0915
                       1
                             0.8
Least square means with the Ismeans package
  library(multcompView)
  library(lsmeans)
  lsmeans = lsmeans::lsmeans ### Uses the lsmeans function
                             ### from the Ismeans package.
                             ### not from the lmerTest package
  leastsquare = lsmeans(model,
                       pairwise ~ Day,
                       alpha=.05,
                       adjust="tukey")
  cld(leastsquare,
      alpha=.05.
      Letters=letters,
      adjust="tukey")
                        SE df
                                   lower.CL upper.CL .group
     Day
          lsmean
          25.33333 9.304196 19.81 -0.1441779 50.81084 a
      4
          34.50000 9.304196 19.81 9.0224887 59.97751 ab
          47.00000 9.304196 19.81 21.5224887 72.47751 ab
          63.33333 9.304196 19.81 37.8558221 88.81084
     Degrees-of-freedom method: satterthwaite
     Confidence level used: 0.95
     Conf-level adjustment: sidak method for 4 estimates
      P value adjustment: tukey method for comparing a family of 4 estimates
     significance level used: alpha = 0.05
```

Means sharing a letter in .group are not significantly different

Least square means using the *lmerTest* package

```
lsmeans = lmerTest::lsmeans ### Uses the lsmeans function
                           ### from the lmerTest package,
                           ###
                                not from the 1smeans package
LT = lsmeans(model,
            test.effs = "Day")
LT
  Least Squares Means table:
         Day Estimate Standard Error DF t-value Lower CI Upper CI p-value
               63.33 9.30 19.8
                                                         82.8 <2e-16 ***
   Day 1 1.0
                                         6.81
                                                43.91
  Day 2 2.0
                                                         66.4 1e-04 ***
               47.00
                             9.30 19.8
                                         5.05
                                                27.58
  Day 3 3.0
Day 4 4.0
                                                         53.9 0.001 **
               34.50
                             9.30 19.8 3.71
                                                15.08
                                         2.72
                                                5.91
                                                         44.8 0.013 *
               25.33
                             9.30 19.8
PT = difflsmeans(model,
                test.effs="Day")
PT
  Differences of LSMEANS:
            Estimate Standard Error DF t-value Lower CI Upper CI p-value
  Day 1 - 2
                              12.78 15.0 1.28 -10.90
                16.3
                                                             43.6
                                                                    0.220
  Day 1 - 3
                28.8
                              12.78 15.0
                                           2.26
                                                   1.60
                                                             56.1
                                                                    0.039 *
  Day 1 - 4
Day 2 - 3
Day 2 - 4
                             12.78 15.0 2.97 10.77
12.78 15.0 0.98 -14.73
                                                                    0.009 **
                38.0
                                                             65.2
                12.5
                                                             39.7
                                                                    0.343
                              12.78 15.0 1.70 -5.57
                                                             48.9 0.111
                21.7
  Day 3 - 4
                9.2
                              12.78 15.0 0.72 -18.07
                                                             36.4 0.484
### Extract lsmeans table
Sum = PT$diffs.lsmeans.table
### Extract comparisons and p-values
Comparison = rownames(Sum)
P.value = Sum\"p-value"
### Adjust p-values
P.value.adj = p.adjust(P.value,
                      method = "none")
### Fix names of comparisons
Comparison = gsub("-", "- Day", Comparison)
```

```
### Produce compact letter display
library(rcompanion)
cldList(comparison = Comparison,
       p.value = P.value.adi,
       threshold = 0.05)
    Group Letter MonoLetter
              a
  1 Day1
  2 Day2
              ab
                         ab
          b
b
   3 Day3
                         b
  4 Day4
                         b
```

Two-way Anova with Robust Estimation

A two-way anova using robust estimators can be performed with the *WRS2* package. Options for estimators are M-estimators, trimmed means, and medians. This type of analysis is resistant to deviations from the assumptions of the traditional ordinary-least-squares anova, and are robust to outliers. However, it may not be appropriate for data that deviate too widely from parametric assumptions.

The main analysis using M-estimators for a two-way anova is conducted with the *pbad2way* function in the *WRS2* package. Post-hoc tests can be performed with the *mcp2a* function in the *WRS2* package or with my custom functions *pairwiseRobustTest* and *pairwiseRobustMatrix*, which rely on the *pb2gen* function in *WRS2*.

My custom function *groupwiseHuber* uses *the HuberM* function in the *DescTools* package to determine the Huber M-estimators across groups in a data frame.

For more information on robust tests available in the *WRS2* package, see:

```
help(package="WRS2")
```

Consult the chapter on *Two-way Anova* for general consideration about conducting analysis of variance.

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(rcompanion)){install.packages("rcompanion")}
if(!require(ggplot2)){install.packages("ggplot2")}
if(!require(WRS2)){install.packages("WRS2")}
if(!require(multcompView)){install.packages("multcompView")}
if(!require(psych)){install.packages("psych")}
```

Example

```
### -----
### Two-way anova with robust estimators, hypothetical data
### Using WRS2 package
Input = ("
Factor.A Factor.B Response
        x 0.9
1
        y
               1.4
1
        X
               1.3
1
               2.0
       У
1
               1.6
        X
1
        y
               2.6
        X
               2.4
m
                3.6
        y
m
                2.8
        X
                3.7
m
        У
                3.2
        X
m
               3.0
m
        У
       X
               1.6
n
   x
y
x
y
               1.2
n
               2.0
n
               1.9
n
               2.7
n
                0.9
")
```

Data = read.table(textConnection(Input), header=TRUE)

Produce Huber M-estimators and confidence intervals by group

Factor.B

Interaction plot using summary statistics

```
library(ggplot2)
pd = position_dodge(.2)
ggplot(Sum, aes(x=Factor.A,
                y=M.Huber,
                color=Factor.B)) +
    geom_errorbar(aes(ymin=lower.ci,
                      ymax=upper.ci),
                   width=.2, size=0.7, position=pd) +
    geom_point(shape=15, size=4, position=pd) +
    theme_bw() +
    theme(
          axis.title.y = element_text(vjust= 1.8),
          axis.title.x = element_text(vjust= -0.5),
          axis.title = element_text(face = "bold")) +
    scale_color_manual(values = c("black", "blue"))
      3
   M.Huber
```

Two-way analysis of variance for M-estimators

The est = "mom" option uses a modified M-estimator for the analysis. To analyze using medians, use the est = "median" option in the pbad2way function in the WRS2 package. To analyze using trimmed means, use the t2way function in the WRS2 package.

m Factor.A

Produce post-hoc tests for main effects with mcp2a

```
post = mcp2a(Response ~ Factor.A + Factor.B + Factor.A:Factor.B,
           data = Data
           est = "mom",
                          # M-estimator
           nboot = 5000) # number of bootstrap samples
post$contrasts
post
                                       Factor.A1: Factor.A2: Factor.A3:
     Factor.A1 Factor.A2 Factor.A3 Factor.B1 Factor.B1 Factor.B1 Factor.B1
  1_x 1 1 0 1
                                         1
                                                      1
          1 1 0
-1 0 1
-1 0 1
0 -1 -1
0 -1 -1
                                    -1
                                             -1
                                                       -1
                                                                  0
  1_y
                                    1
                                            -1
                                                       0
                                                                 1
  m_x
  m_y
                                    -1
                                                       0
                                            1
                                                                 -1
                                    1
                                             0
                                                       -1
                                                                 -1
  n_x
  n_y
                                             0
                                                       1
                                    -1
                          V1 ci.lower ci.upper p-value
                   -3.18333 -4.20000 -1.60000 0.00000
  Factor.A1
                    -0.16667 -1.70000 1.36667 0.40233
  Factor.A2
                     3.01667 1.40000 4.05000 0.00000
  Factor.A3
                    -0.81667 -2.28333 1.00000 0.22233
  Factor.B1
  Factor.A1:Factor.B1 0.11667 -1.50000 1.16667 0.48033
  Factor.A2:Factor.B1 -1.50000 -3.10000 0.00000 0.01767
  Factor.A3:Factor.B1 -1.61667 -2.80000 0.00000 0.01433
  ### The Factor.A1 contrast compares 1 to m; since it is significant,
      l is significantly different than m.
  ### The Factor.A2 contrast compares 1 to n; since it is not significant,
```

Produce post-hoc tests for main effects with pairwiseRobustTest or pairwiseRobustMatrix

Table and compact letter display with pairwiseRobustTest

l is not significantly different than n.

```
### Pairwise robust test
library(rcompanion)
nboot = 5000.
                        method = "fdr")
                          # adjust p-values; see ?p.adjust for options
РТ
    Comparison Statistic p.value p.adjust
   1 \quad n - 1 = 0 \quad 0.08333 \quad 0.7204 \quad 0.7204
   2 \quad n - m = 0 -1.4 \quad 0.0014 \quad 0.0021
   3 \quad 1 \quad - \quad m = 0 \quad -1.483 \quad 6e-04 \quad 0.0018
   ### p-values may differ
### Produce compact letter display
library(rcompanion)
cldList(comparison = PT$Comparison,
        p.value = PT$p.adjust,
        threshold = 0.05)
     Group Letter MonoLetter
   1
         n
           a
   2
         1
                a
                          a
   3
         m
                b
                           b
```

Compact letter display output with pairwiseRobustMatrix

```
PM$Adjusted
```

Produce post-hoc tests for interaction effect

```
### Create a factor which is the interaction of Factor.A and Factor.B
Data$Factor.int = interaction (Data$Factor.A, Data$Factor.B)
### Order groups by median
Data$Factor.int = factor(Data$Factor.int,
                levels = c("m.y", "m.x", "n.x", "l.y", "n.y", "l.x"))
### Check data frame
library(psych)
headTail(Data)
      Factor.A Factor.B Response Factor.int
  8
                         3.6
                    У
                                     m.y
  10
                           3.7
            m
                     У
                                     m.y
  12
            m
                  y
x
                           3
                                     m.y
                          2.4
           m
                                     m.x
          <NA>
                 <NA>
                           . . .
                                     <NA>
  . . .
                  У
                           0.9
  18
           n
                                      n.y
  1
            1
                           0.9
                                     1.x
                   X
  3
            1
                           1.3
                                     1.x
                   X
                           1.6
                                      1.x
                     X
```

Table and compact letter display with pairwiseRobustTest

```
library(rcompanion)
PT = pairwiseRobustTest(x = Data$Response,
                       g = Data$Fa
est = "mom",
                              = Data$Factor.int,
                       nboot = 5000,
                       method = "fdr")
                          # adjust p-values; see ?p.adjust for options
PT
        Comparison Statistic p.value p.adjust
  1 \quad \text{m.y} - \text{m.x} = 0 \quad -0.85 \quad 0.1348
                                      0.1615
  2 \quad m.y - n.x = 0 -1.55 3 \quad m.y - 1.y = 0 -1.65
                                       0.0000
                                     0.0000
                                  0.0000
  4 \text{ m.y} - \text{n.y} = 0 -2.317
  5 \text{ m.y} - 1.x = 0 -2.383
                                 0.0000
                      -0.7 0.1312 0.1615
  6 \text{ m.x} - \text{n.x} = 0
  0.8355
                                     0.1615
  12 \text{ n.x} - 1.x = 0   0.8333   0.0664   0.1423
  13 l.y - n.y = 0 -0.6667
                              0.14 0.1615
  14 \ 1.y - 1.x = 0 \quad -0.7333 \quad 0.1296 \quad 0.1615
  15 \text{ n.y} - 1.x = 0 -0.06667 0.944
                                      0.9440
  ### p-values may differ
### Produce compact letter display
library(rcompanion)
cldList(comparison = PT$Comparison,
       p.value = PT$p.adjust,
       threshold = 0.05)
    Group Letter MonoLetter
  1 m.y a
                        а
     m.x ab
n.x bc
1.y bc
  2
                        ab
  3
                        bc
  4
                         bc
  5
             C
                         C
      1.x
              C
                          C
```

<u>Compact letter display output with pairwiseRobustMatrix</u>

```
### Order groups by median
Data$Factor.int = factor(Data$Factor.int,
```

```
levels = c("m.y", "m.x", "n.x", "l.y", "n.y", "l.x"))
### Pairwise robust tests
library(rcompanion)
PM = pairwiseRobustMatrix(x = Data$Response,
                         g
                                = Data$Factor.int.
                          est = "mom",
                         nboot = 5000,
                         method = "fdr")
                           # adjust p-values; see ?p.adjust for options
PM
   $Unadjusted
      m.y
            m.x
                   n.x 1.y n.y 1.x
  m.y NA 0.1312 0.000 0.0000 0.0000 0.0000
              NA 0.126 0.1320 0.0000 0.0000
  m.x NA
                    NA 0.7638 0.1328 0.0680
   n.x NA
              NA
                           NA 0.1304 0.1408
   l.y NA
              NA
                    NA
                                  NA 0.9318
   n.y NA
              NA
                    NA
                           NA
   1.x NA
              NA
                    NA
                           NA
                                  NA
                                         NA
   $Method
   [1] "fdr"
   $Adjusted
                      n.x
                             1.y n.y
         m.y
                m.x
  m.y 1.0000 0.1625 0.0000 0.0000 0.0000 0.0000
  m.x 0.1625 1.0000 0.1625 0.1625 0.0000 0.0000
   n.x 0.0000 0.1625 1.0000 0.8184 0.1625 0.1457
   1.y 0.0000 0.1625 0.8184 1.0000 0.1625 0.1625
   n.y 0.0000 0.0000 0.1625 0.1625 1.0000 0.9318
   1.x 0.0000 0.0000 0.1457 0.1625 0.9318 1.0000
   ### p-values may differ
library(multcompView)
multcompLetters(PM$Adjusted,
               compare="<",
               threshold=0.05,
               Letters=letters,
               reversed = FALSE)
  m.y m.x n.x l.y n.y l.x "a" "ab" "bc" "bc" "c" "c"
   ### Note, means are not ordered from largest to smallest
```

Paired t-test

Paired t-tests can be conducted with the *t.test* function in the native *stats* package using the *paired=TRUE* option. Data can be in long format or short format. Examples of each are shown in this chapter.

As a non-parametric alternative to paired t-tests, a permutation test can be used. An example is shown in the "Permutation test for dependent samples" section of this chapter.

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Paired t-test

SAEEPER: One-way Permutation Test of Symmetry for Paired Ordinal Data

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(ggplot2)){install.packages("ggplot2")}
if(!require(coin)){install.packages("coin")}
if(!require(pwr)){install.packages("pwr")}
```

When to use it

The horseshoe crab example is shown at the end of the "How to do the test" section.

Null hypothesis

Assumption

How the test works

See the *Handbook* for information on these topics.

Examples

The flicker feather example is shown in the "How to do the test" section.

Graphing the results

Plots are shown in the "How to do the test" section.

How to do the test

Paired t-test, data in wide format, flicker feather example

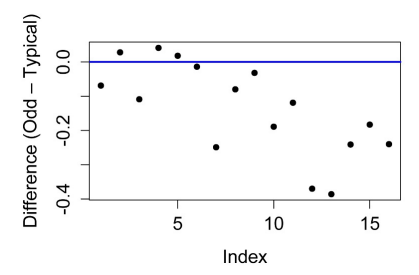
```
### ------
### Paired t-test, Flicker feather example, p. 185
### ------

Input = ("
Bird Typical Odd
A -0.255 -0.324
B -0.213 -0.185
C -0.190 -0.299
```

```
-0.185
              -0.144
D
 Е
      -0.045
              -0.027
       -0.025
               -0.039
 F
 G
      -0.015
               -0.264
       0.003
               -0.077
Н
               -0.017
 Ι
       0.015
               -0.169
       0.020
 J
 K
       0.023
              -0.096
L
       0.040
              -0.330
       0.040
               -0.346
M
               -0.191
Ν
       0.050
       0.055
               -0.128
0
P
       0.058
               -0.182
")
Data = read.table(textConnection(Input), header=TRUE)
```

Paired t-test

Simple plot of differences

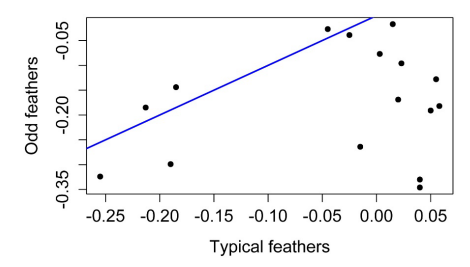


A simple plot of differences between one sample and the other. Points below the blue line indicate observations where Typical is greater than Odd, that is where (Odd - Typical) is negative.

Simple 1-to-1 plot of values

```
plot(Data$Typical, Data$Odd,
    pch = 16,
    xlab="Typical feathers",
    ylab="Odd feathers")
```

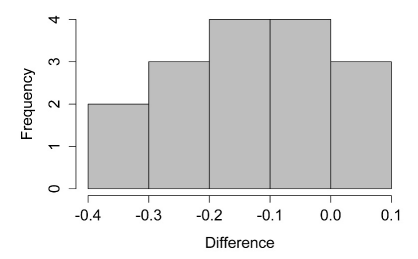
abline(0,1, col="blue", lwd=2)



Plot of paired samples from a paired t-test. Circles below or to the right of the blue one-to-one line indicate observations with a higher value for *Typical* than for *Odd*.

Checking assumptions of the model

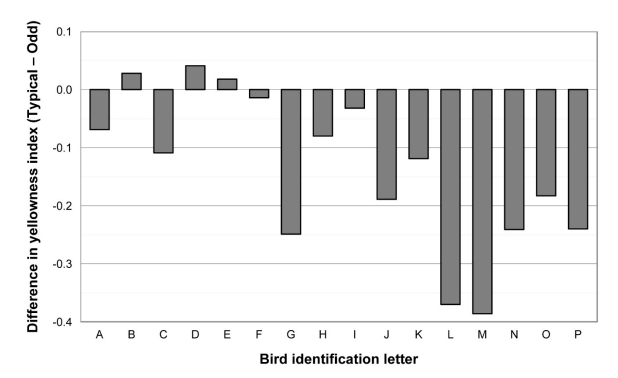
Histogram of differences



Histogram of differences of two populations from a paired t-test. Distribution of differences should be approximately normal. Bins with negative values indicate observations with a higher value for Typical than for Odd.

Graphing the results

```
Data$Difference = Data$Odd - Data$Typical
library(ggplot2)
ggplot(Data,
       aes(x = Bird,
           y = Difference)) +
  geom_bar(stat = "identity",
           fill = "grey50"
           colour = "black",
           width = 0.6) +
  scale_y_continuous(breaks = seq(-0.4, 0.1, 0.1),
           limits = c(-0.4, 0.1),
           expand = c(0, 0)) +
  #ggtitle("Chart title") +
  labs(x = "Bird identification letter",
       y = "Difference in yellowness index (Typical - Odd)") +
  theme_bw() +
  theme(panel.grid.major.x = element_blank(),
```



Paired t-test, data in wide format, horseshoe crab example

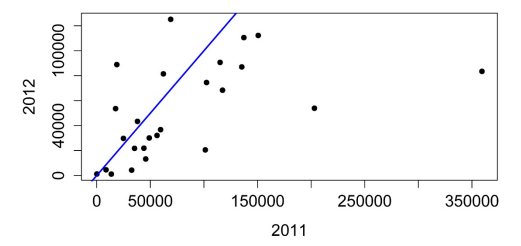
```
### Paired t-test, Horseshoe crab example, pp. 181-182
   # Note, if you use "2011" as a variable name,
       the read.table function will convert it to "X2011"
Input = ("
Beach
                     Year.2011
                                   Year.2012
'Bennetts Pier'
                      35282
                                    21814
'Big Stone'
                                    83500
                     359350
'Broadkill'
                      45705
                                    13290
'Cape Henlopen'
                      49005
                                    30150
'Fortescue'
                      68978
                                   125190
'Fowler'
                       8700
                                     4620
'Gandys'
                                    88926
                      18780
'Higbees'
                      13622
                                     1205
'Highs'
                                    29800
                      24936
```

```
'Kimbles'
                      17620
                                    53640
'Kitts Hummock'
                     117360
                                    68400
'Norburys Landing'
                     102425
                                    74552
'North Bowers'
                      59566
                                    36790
'North Cape May'
                                     4350
                      32610
'Pickering'
                     137250
                                   110550
'Pierces Point'
                      38003
                                    43435
'Primehook'
                     101300
                                    20580
'Reeds'
                      62179
                                    81503
'Slaughter'
                     203070
                                    53940
'South Bowers'
                     135309
                                    87055
'South CSL'
                                   112266
                     150656
'Ted Harvey'
                     115090
                                    90670
'Townbank'
                                    21942
                      44022
'villas'
                                    32140
                      56260
'woodland'
                        125
                                     1260
")
```

Data = read.table(textConnection(Input), header=TRUE)

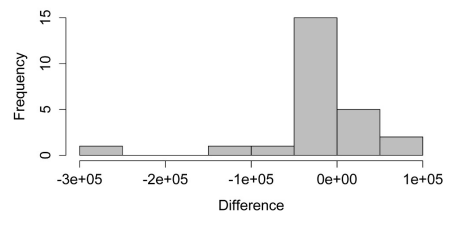
Paired t-test

Simple 1-to-1 plot of values



Plot of paired samples from a paired t-test. Circles below and to the right of the blue one-to-one line indicate observations with a higher value for 2011 than for 2012.

Histogram of differences



Histogram of differences in two populations from paired t-test. Distribution of differences should be approximately normal. Bins with negative values indicate observations with a higher score for 2011 than for 2012.

Paired t-test, data in long format

```
### ------
### Paired t-test, long format data, Flicker feather example, p. 185
### ------
```

```
Input = ("
 Bird
         Feather
                    Length
         Typical
                    -0.255
 Α
                    -0.213
 В
         Typical
 C
         Typical
                    -0.19
                    -0.185
 D
         Typical
 Ē
         Typical
                   -0.045
 F
         Typical
                    -0.025
                    -0.015
 G
         Typical
 н
         Typical
                     0.003
                     0.015
 Ι
         Typical
 J
                     0.02
         Typical
 K
         Typical
                     0.023
                     0.04
         Typical
 L
                     0.04
 Μ
         Typical
 N
         Typical
                     0.05
 0
         Typical
                     0.055
 Р
         Typical
                     0.058
         odd
                    -0.324
 Α
                    -0.185
         odd
 В
 C
         odd
                    -0.299
         Odd
                    -0.144
 D
 Ε
         odd
                    -0.027
         odd
 F
                    -0.039
 G
         odd
                    -0.264
         odd
                    -0.077
 н
 Ι
         odd
                    -0.017
                    -0.169
 J
         odd
         Odd
                    -0.096
 K
 L
         odd
                    -0.33
                    -0.346
         odd
 Μ
                    -0.191
 Ν
         odd
 0
         Odd
                    -0.128
         odd
                    -0.182
 Р
")
Data = read.table(textConnection(Input), header=TRUE)
      ### Note: data must be ordered so that the first observation of Group 1
            is the same subject as the first observation of Group 2
t.test(Length ~ Feather,
       data=Data,
       paired=TRUE,
       conf.level=0.95)
   t = -4.0647, df = 15, p-value = 0.001017
   mean of the differences
                  -0.137125
```

Permutation test for dependent samples

This permutation test is analogous to a paired t-test.

```
### ------
### Paired two-sample permutation test, long format data
### Flicker feather example, p. 185
Input = ("
Bird
       Feather Length
       Typical -0.255
       Typical -0.213
В
C
       Typical -0.19
D
       Typical
                -0.185
                -0.045
       Typical
Е
F
       Typical -0.025
               -0.015
G
       Typical
                0.003
Н
       Typical
Ι
       Typical
               0.015
                0.02
J
       Typical
K
       Typical
                 0.023
                 0.04
L
       Typical
M
       Typical
                0.04
       Typical
                0.05
N
       Typical
                0.055
0
                0.058
Р
       Typical
       odd
                -0.324
Α
В
       odd
                -0.185
C
       odd
                -0.299
D
       odd
                -0.144
Е
       Odd
                -0.027
F
       odd
                -0.039
              -0.264
-0.077
G
       odd
н
       odd
Ι
       odd
               -0.017
J
       odd
                -0.169
                -0.096
K
       odd
L
       odd
                -0.33
М
       Odd
                -0.346
       odd
                -0.191
N
       odd
0
                -0.128
       odd
                -0.182
Р
")
Data = read.table(textConnection(Input), header=TRUE)
library(coin)
independence_test(Length ~ Feather | Bird,
                data = Data
  Asymptotic General Independence Test
  Z = -2.8959, p-value = 0.003781
```

Power analysis

Power analysis for paired t-test

```
### Power analysis, paired t-test, pp. 185-186
Detect = 0.1
                                # Difference in means to detect
SD = 0.135
                                # Standard deviation of differences
Cohen.d = Detect/SD
library(pwr)
pwr.t.test(
                                 # Number of _pairs_ of observations
      n = NULL
      d = Cohen.d,
      sig.level = 0.05, # Type I probability
      power = 0.90,
                                 # 1 minus Type II probability
      type = "paired",
                                # paired t-test
      alternative = "two.sided")
   Paired t test power calculation
  n = 21.16434
  NOTE: n is number of *pairs*
```

Wilcoxon Signed-rank Test

Examples in Summary and Analysis of Extension Program Evaluation

SAEEPER: Two-sample Paired Rank-sum Test SAEEPER: Sign Test for Two-sample Paired Data

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(BSDA)){install.packages("BSDA")}
```

When to use it

The poplar example is shown below in the "How to do the test" section.

Null hypothesis How it works Examples Graphing the results

See the *Handbook* for information on these topics.

Similar tests

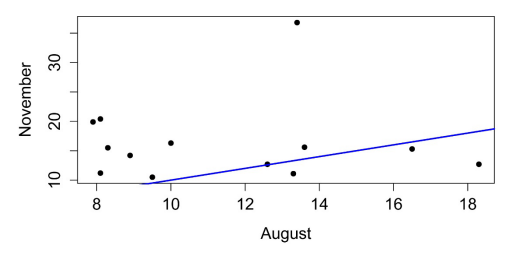
Paired t-test and permutation test are described in the *Paired t-test* chapter. The sign test is described below.

How to do the test

Wilcoxon signed-rank test example

```
### ------
### Wilcoxon signed-rank test, poplar example, p. 189
### -----
Input = ("
            August November
Clone
 Balsam_Spire 8.1 11.2
Beaupre 10.0 16.3
Hazendans 16.5 15.3
Hoogvorst 13.6 15.6
Raspalje 9.5 10.5
Unal
             8.3
                   15.5
 Columbia_River 18.3 12.7
 Fritzi_Pauley 13.3
                    11.1
             7.9
Trichobel
                    19.9
              8.1
                    20.4
Gaver
Gibecq
            8.9
12.6
                    14.2
                   12.7
Wolterson 13.4 36.8
")
Data = read.table(textConnection(Input), header=TRUE)
wilcox.test(Data$August,
          Data$November,
          paired=TRUE)
  Wilcoxon signed rank test
  V = 16, p-value = 0.03979
     ### Matches "Signed Rank" p-value in SAS output
```

Simple 1-to-1 plot of values



Plot of paired samples from a Wilcoxon signed-rank test. Circles above and to the left of the blue one-to-one line indicate observations with a higher value for November than for August.

Sign test example

The following is an example of the two-sample dependent-samples sign test. The data are arranged as a data frame in which each row contains the values for both measurements being compared for each experimental unit. This is sometimes called "wide format" data. The SIGN.test function in the BSDA package is used. The option md=0 indicates that the expected difference in the medians is 0 (null hypothesis). This function can also perform a one-sample sign test.

```
### Two-sample sign test, poplar example, p. 189
Input = ("
 Clone
                         November
                August
                 8.1
 Balsam_Spire
                         11.2
                10.0
                         16.3
 Beaupre
                16.5
                         15.3
 Hazendans
 Hoogvorst
                13.6
                         15.6
 Raspalje
                  9.5
                         10.5
                  8.3
                         15.5
 Unal
 Columbia_River
                  18.3
                         12.7
 Fritzi_Pauley
                  13.3
                         11.1
 Trichobel
                  7.9
                         19.9
                   8.1
                         20.4
 Gaver
                  8.9
 Gibecq
                         14.2
 Primo
                         12.7
                  12.6
 Wolterson
                  13.4
                         36.8
")
Data = read.table(textConnection(Input), header=TRUE)
library(BSDA)
```

Regressions

Correlation and Linear Regression

Introduction

The amphipod egg example is shown below in the "How to do the test" section.

When to use them Correlation versus linear regression Correlation and causation Null hypothesis Independent vs. dependent variables How the test works Assumptions

See the *Handbook* for information on these topics.

Examples

The species diversity example is shown below in the "How to do the test" section.

Graphing the results Similar tests

How to do the test

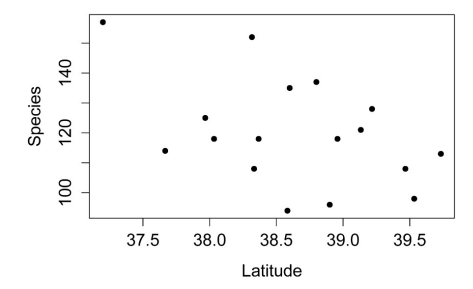
Correlation and linear regression example

```
### -----
### Correlation and linear regression, species diversity example
### pp. 207-208
Input = ("
Town State Latitude Species 'Bombay Hook' DE 39.217 128 'Cape Henlopen' DE 38.800 137 'Middletown' DE 39.467 108 'Milford' DE 38.958 118 'Rehoboth' DE 38.600 135
                                39.467 108
38.958 118
38.600 135
'Rehoboth'
'Seaford-Nanticoke'
DE
                                   38.583
                                               94
                                            113
                                   39.733
'Crisfield'
                                   38.033
                                               118
                          MD
'Denton'
                                   38.900
                                              96
                         MD
'Elkton'
                          MD
                                   39.533
                                               98
'Lower Kent County'
'Ocean City'
'Salisbury'
                                   39.133
                                               121
                          MD
                          MD
                                   38.317
                                               152
                                   38.333
                          MD
                                               108
'S Dorchester County' MD
                                   38.367
                                            118
```

```
'Cape Charles' VA 37.200 157
'Chincoteague' VA 37.967 125
'Wachapreague' VA 37.667 114
")
```

Data = read.table(textConnection(Input),header=TRUE)

Simple plot of the data



Correlation

Correlation can be performed with the *cor.test* function in the native *stats* package. It can perform Pearson, Kendall, and Spearman correlation procedures. Methods for multiple correlation of several variables simultaneously are discussed in the *Multiple regression* chapter.

Pearson correlation

Pearson correlation is the most common form of correlation. It is a parametric test, and assumes that the data are linearly related and that the residuals are normally distributed.

-0.4628844

Kendall correlation

Kendall rank correlation is a non-parametric test that does not assume a distribution of the data or that the data are linearly related. It ranks the data to determine the degree of correlation.

Spearman correlation

Spearman rank correlation is a non-parametric test that does not assume a distribution of the data or that the data are linearly related. It ranks the data to determine the degree of correlation, and is appropriate for ordinal measurements.

Linear regression

Linear regression can be performed with the *lm* function in the native *stats* package. A robust regression can be performed with the *lmrob* function in the *robustbase* package.

```
Multiple R-squared: 0.2143, Adjusted R-squared: 0.1619
F-statistic: 4.09 on 1 and 15 DF, p-value: 0.06134

library(car)

Anova(model, type="II")  # shows p-value for effects in model

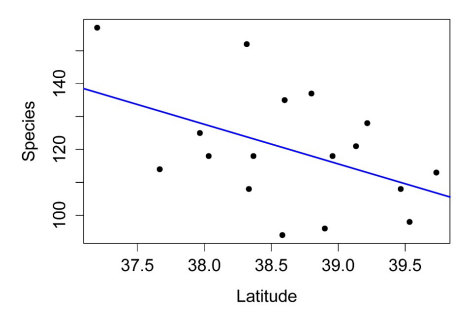
Response: Species
Sum Sq Df F value Pr(>F)
Latitude 1096.6 1 4.0903 0.06134 .
Residuals 4021.4 15
```

Plot linear regression

```
int = model$coefficient["(Intercept)"]
slope =model$coefficient["Latitude"]

plot(Species ~ Latitude,
    data = Data,
    pch=16,
    xlab = "Latitude",
    ylab = "Species")

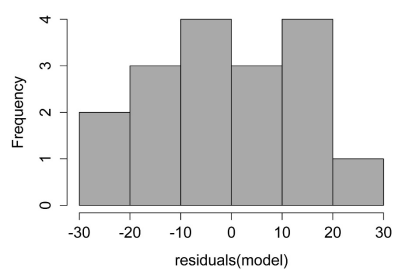
abline(int, slope,
        lty=1, lwd=2, col="blue")  # style and color of line
```



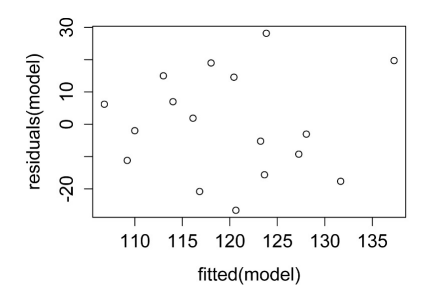
Checking assumptions of the model

```
hist(residuals(model),
     col="darkgray")
```

Histogram of residuals(model)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

additional model checking plots with: plot(model)
alternative: library(FSA); residPlot(model)

Robust regression

The *lmrob* function in the *robustbase* package produces a linear regression which is not sensitive to outliers in the response variable. It uses MM-estimation.

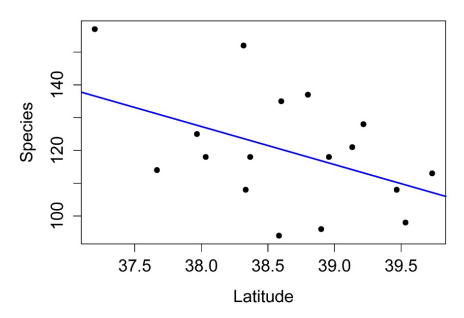
```
library(robustbase)
model = lmrob(Species ~ Latitude,
             data = Data
summary(model)
                                 # shows parameter estimates, r-square
              Estimate Std. Error t value Pr(>|t|)
                                           0.0259 *
   (Intercept) 568.830
                          230.203 2.471
   Latitude
               -11.619
                            5.912 -1.966
                                            0.0681 .
  Multiple R-squared: 0.1846, Adjusted R-squared: 0.1302
model.null = lmrob(Species ~ 1,
                  data = Data
anova(model, model.null)
                                # shows p-value for model
    pseudoDf Test.Stat Df Pr(>chisq)
   1
          15
   2
          16
                3.8634 1 0.04935 *
```

Plot the model

```
int = model$coefficient["(Intercept)"]
slope =model$coefficient["Latitude"]

plot(Species ~ Latitude,
    data = Data,
    pch=16,
    xlab = "Latitude",
    ylab = "Species")

abline(int, slope,
    lty=1, lwd=2, col="blue")  # style and color of line
```



Linear regression example

```
### Linear regression, amphipod eggs example
### pp. 191-193
### -----
Input = ("
Weight Eggs
 5.38
         29
 7.36
         23
 6.13
         22
 4.75
         20
 8.10
         25
 8.62
         25
 6.30
         17
 7.44
         24
 7.26
         20
 7.17
         27
 7.78
         24
 6.23
         21
 5.42
         22
 7.87
         22
 5.25
         23
         35
 7.37
 8.01
         27
 4.92
         23
 7.03
         25
 6.45
         24
 5.06
         19
 6.72
         21
 7.00
         20
 9.39
         33
 6.49
         17
 6.34
         21
```

```
6.16
        25
 5.74
        22
")
Data = read.table(textConnection(Input), header=TRUE)
model = lm(Eggs ~ Weight,
          data = Data
summary(model)
                              # shows parameter estimates.
                               # p-value for model, r-square
  Coefficients:
             Estimate Std. Error t value Pr(>|t|)
  (Intercept) 12.6890 4.2009 3.021 0.0056 **
            Weight
  Multiple R-squared: 0.2055, Adjusted R-squared: 0.175
  F-statistic: 6.726 on 1 and 26 DF, p-value: 0.0154
  ### Neither the r-squared nor the p-value agrees with what is reported
  ###
         in the Handbook.
library(car)
Anova(model, type="II") # shows p-value for effects in model
           Sum Sq Df F value Pr(>F)
            93.89 1 6.7258 0.0154 *
  Weight
  Residuals 362.96 26
```

Power analysis

Power analysis for correlation

Spearman Rank Correlation

When to use it Null hypothesis Assumption How the test works

See the *Handbook* for information on these topics.

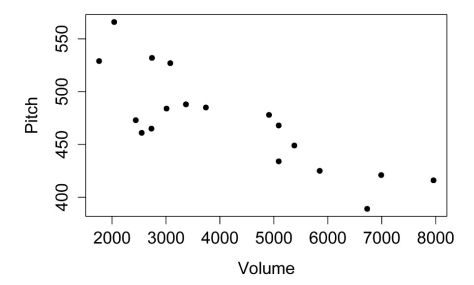
Example

Example of Spearman rank correlation

```
### Spearman rank correlation, frigatebird example
### p. 212
### -----
Input = ("
Volume Pitch
 1760
         529
 2040
         566
 2440
         473
 2550
         461
 2730
         465
 2740
         532
 3010
         484
 3080
         527
 3370
         488
         485
 3740
 4910
         478
 5090
         434
 5090
         468
 5380
         449
 5850
         425
 6730
         389
 6990
         421
7960
         416
")
Data = read.table(textConnection(Input), header=TRUE)
cor.test( ~ Pitch + Volume,
         data=Data,
         method = "spearman",
         continuity = FALSE,
         conf.level = 0.95)
   Spearman's rank correlation rho
   S = 1708.382, p-value = 0.0002302
   sample estimates:
```

rho -0.7630357

Simple plot of the data



Graphing the results

See the *Handbook* for information on this topic.

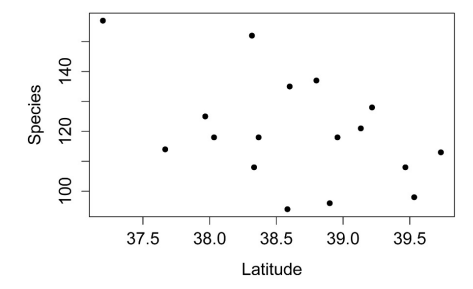
How to do the test

Example of Spearman rank correlation

```
### Spearman rank correlation, species diversity example
### p. 214
### ----
Input = ("
Town
                      State
                              Latitude
                                         Species
'Bombay Hook'
                                39.217
                                           128
                         DE
'Cape Henlopen'
                                38.800
                                           137
                         DE
'Middletown'
                                39.467
                                           108
                         DE
'Milford'
                                38.958
                                           118
                         DE
'Rehoboth'
                         DE
                                38.600
                                           135
'Seaford-Nanticoke'
                                38.583
                                            94
                         DE
'Wilmington'
                                39.733
                                           113
                         DE
'Crisfield'
                         MD
                                38.033
                                           118
'Denton'
                                38.900
                                            96
                         MD
'Elkton'
                                39.533
                                            98
                         MD
'Lower Kent County'
                                39.133
                                           121
                         MD
'Ocean City'
                                38.317
                                           152
                         MD
```

```
'Salisbury'
                               38.333
                                          108
                        MD
'S Dorchester County'
                               38.367
                                          118
                        MD
'Cape Charles'
                               37.200
                                          157
                        VA
'Chincoteague'
                        VA
                               37.967
                                          125
'Wachapreague'
                               37.667
                                          114
                        VA
")
Data = read.table(textConnection(Input), header=TRUE)
cor.test( ~ Species + Latitude,
         data=Data,
         method = "spearman",
         continuity = FALSE,
         conf.level = 0.95)
   Spearman's rank correlation rho
   S = 1111.908, p-value = 0.1526
   -0.3626323
```

Simple plot of the data



Curvilinear Regression

When to use it

Null hypotheses
Assumptions
How the test works
Examples
Graphing the results
Similar tests

See the *Handbook* for information on these topics.

How to do the test

This chapter will fit models to curvilinear data using three methods: 1) Polynomial regression; 2) B-spline regression with polynomial splines; and 3) Nonlinear regression with the *nls* function. In this example, each of these three will find essentially the same best-fit curve with very similar p-values and R-squared values.

Polynomial regression

Polynomial regression is really just a special case of multiple regression, which is covered in the *Multiple regression* chapter. In this example we will fit a few models, as the *Handbook* does, and then compare the models with the extra sum of squares test, the Akaike information criterion (AIC), and the adjusted R-squared as model fit criteria.

For a linear model (*lm*), the adjusted R-squared is included with the output of the *summary(model)* statement. The AIC is produced with its own function call, *AIC(model)*. The extra sum of squares test is conducted with the *anova* function applied to two models.

For AIC, smaller is better. For adjusted R-squared, larger is better. A non-significant p-value for the extra sum of squares test comparing model *a* to model *b* indicates that the model with the extra terms does not significantly reduce the error sum of squares over the reduced model. Which is to say, a non-significant p-value suggests the model with the additional terms is not better than the reduced model.

```
### Polynomial regression, turtle carapace example
### pp. 220-221
Input = ("
Length Clutch
 284
          3
          2
 290
          7
 290
          7
 290
 298
         11
 299
         12
 302
         10
 306
          8
 306
          8
 309
          9
 310
         10
 311
         13
```

```
7
 317
 317
          9
 320
          6
 323
         13
 334
          2
 334
          8
")
Data = read.table(textConnection(Input),header=TRUE)
### Change Length from integer to numeric variable
      otherwise, we will get an integer overflow error on big numbers
Data$Length = as.numeric(Data$Length)
### Create quadratic, cubic, quartic variables
library(dplyr)
Data =
mutate(Data,
       Length2 = Length*Length,
       Length3 = Length*Length,
       Length4 = Length*Length*Length)
library(FSA)
headtail(Data)
      Length Clutch Length2 Length3
                                           Length4
               3 80656 22906304 6505390336
   1
         284
   2
         290
                 2 84100 24389000 7072810000
         290
                 7 84100 24389000 7072810000
         323 13 104329 33698267 100013.32
334 2 111556 37259704 12444741136
8 111556 37259704 12444741136
   16
   17
   18
```

Define the models to compare

194

Generate the model selection criteria statistics for these models

```
0.0276
                           0.0563
                                     0.49
                                              0.63
  Length
  Multiple R-squared: 0.0148, Adjusted R-squared:
                                                     -0.0468
  F-statistic: 0.24 on 1 and 16 DF, p-value: 0.631
AIC(model.1)
   [1] 99.133
summary(model.2)
  Coefficients:
               Estimate Std. Error t value Pr(>|t|)
                          2.70e+02
                                             0.0046 **
   (Intercept) -9.00e+02
                                     -3.33
  Length
               5.86e+00
                          1.75e+00
                                      3.35
                                             0.0044 **
                          2.83e-03
                                     -3.33
                                             0.0045 **
  Length2
              -9.42e-03
  Multiple R-squared: 0.434, Adjusted R-squared: 0.358
   F-statistic: 5.75 on 2 and 15 DF, p-value: 0.014
AIC(model.2)
   [1] 91.16157
anova(model.1, model.2)
  Analysis of Variance Table
    Res.Df
              RSS Df Sum of Sq
                                 F Pr(>F)
  1
        16 186.15
        15 106.97 1
                        79.178 11.102 0.00455 **
```

Model selection criteria for four polynomial models. Model 2 has the lowest AIC, suggesting it is the best model from this list for these data. Likewise model 2 shows the largest adjusted R-squared. Finally, the extra SS test shows model 2 to be better than model 1, but that model 3 is not better than model 2. All this evidence indicates selecting model 2.

Continue this process for the remainder of the models

Model	AIC	Adjusted R- squared	p-value for extra SS from previous model
1	99.1	- 0.047	
2	91.2	0.36	0.0045
3	92.7	0.33	0.55
4	94.4	0.29	0.64

Compare models with *compareLM* and *anova*

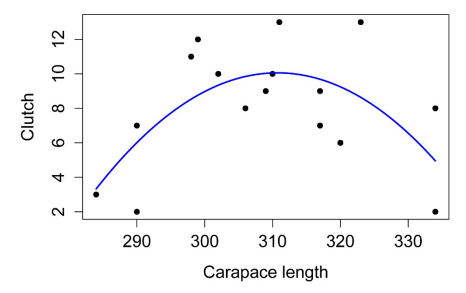
This process can be automated somewhat by using my *compareLM* function and by passing multiple models to the *anova* function. Any of AIC, AICc, or BIC can be minimized to select the best model. If you have no preference, I might recommend using AICc.

```
model.1 = lm (Clutch \sim Length,
                                                                   data=Data)
model.2 = 1m (Clutch ~ Length + Length2,
                                                                   data=Data)
model.3 = lm (Clutch ~ Length + Length2 + Length3,
                                                                   data=Data)
model.4 = lm (Clutch ~ Length + Length2 + Length3 + Length4, data=Data)
library(rcompanion)
compareLM(model.1, model.2, model.3, model.4)
   $Fit.criteria
     Rank Df.res
                 AIC AICC
                                BIC R.squared Adj.R.sq p.value Shapiro.W Shapiro.p
            16 99.13 100.80 101.80  0.01478  -0.0468 0.63080
                                                                  0.9559
                                                                            0.5253
              15 91.16 94.24 94.72
                                      0.43380
                                               0.3583 0.01403
                                                                  0.9605
                                                                            0.6116
             14 92.68 97.68 97.14 0.44860
13 94.37 102.00 99.71 0.45810
                                               0.3305 0.03496
                                                                  0.9762
                                                                            0.9025
                                               0.2914 0.07413
                                                                  0.9797
                                                                            0.9474
anova(model.1, model.2, model.3, model.4)
                RSS Df Sum of Sq F Pr(>F)
     Res.Df
   1
         16 186.15
   2
          15 106.97 1
                           79.178 10.0535 0.007372 ** ## Compares m.2 to m.1
         14 104.18 1 2.797 0.3551 0.561448 ## Compares m.3 to m.2 13 102.38 1 1.792 0.2276 0.641254 ## Compares m.4 to m.3
   3
```

Investigate the final model

```
model.final = lm (Clutch ~ Length + Length2,
                 data=Data)
summary(model.final)
                                    # Shows coefficients,
                                    # overall p-value for model, R-squared
  Coefficients:
               Estimate Std. Error t value Pr(>|t|)
   (Intercept) -9.00e+02 2.70e+02 -3.33 0.0046 **
               5.86e+00
                                     3.35
                         1.75e+00
                                            0.0044 **
  Lenath
  Length2
              -9.42e-03 2.83e-03 -3.33
                                            0.0045 **
  Multiple R-squared: 0.434, Adjusted R-squared: 0.358
  F-statistic: 5.75 on 2 and 15 DF, p-value: 0.014
library(car)
Anova(model.final, type="II")
                                     # Shows p-values for individual terms
                                   196
```

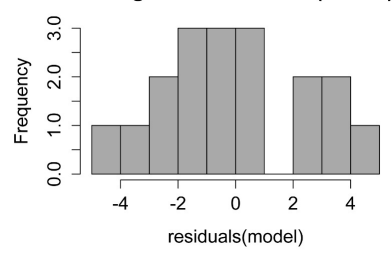
Simple plot of model



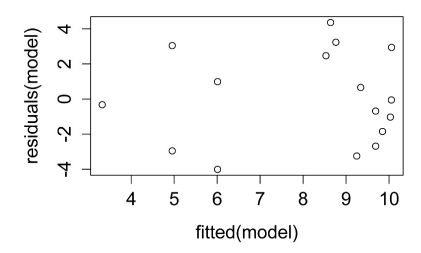
Checking assumptions of the model

```
hist(residuals(model.final),
     col="darkgray")
```

Histogram of residuals(model)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

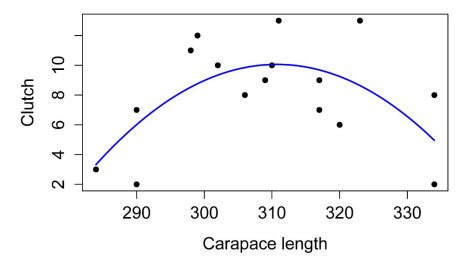
additional model checking plots with: plot(model.final)

B-spline regression with polynomial splines

B-spline regression uses smaller segments of linear or polynomial regression which are stitched together to make a single model. It is useful to fit a curve to data when you don't have a theoretical model to use (e.g. neither linear, nor polynomial, nor nonlinear). It does not assume a

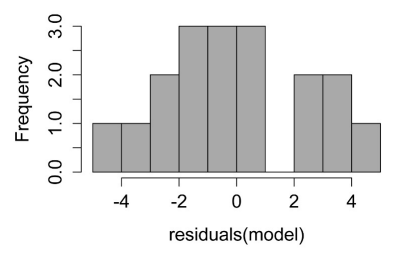
linear relationship between the variables, but the residuals should still be normal and independent. The model may be influenced by outliers.

```
### B-spline regression, turtle carapace example
   ### pp. 220-221
   Input = ("
   Length Clutch
    284 3
    290
             2
    290
            7
            7
    290
    298
            11
    299
            12
    302
            10
    306
             8
            8
    306
            9
    309
    310
            10
    311
            13
    317
            7
    317
            9
    320
             6
    323
            13
    334
            2
    334
             8
   ")
   Data = read.table(textConnection(Input), header=TRUE)
   library(splines)
   model = lm(Clutch ~ bs(Length,
                            knots = 5,  # How many internal segment nodes?
degree = 2),  # 1=local linear fits, 2=quadratic
              data = Data
   summary(model)
                                           # Display p-value and R-squared
      Residual standard error: 2.671 on 15 degrees of freedom
      Multiple R-squared: 0.4338, Adjusted R-squared: 0.3583
      F-statistic: 5.747 on 2 and 15 DF, p-value: 0.01403
Simple plot of model
   plot(Clutch ~ Length,
        data = Data.
        pch=16,
        xlab = "Carapace length",
        ylab = "Clutch")
   i = seq(min(Data$Length), max(Data$Length), len=100) # x-values for line
```

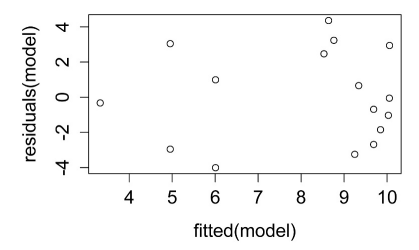


Checking assumptions of the model

Histogram of residuals(model)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

additional model checking plots with: plot(model)

Nonlinear regression

Nonlinear regression can fit various nonlinear models to a data set. These model might include exponential models, logarithmic models, decay curves, or growth curves. The *nls* function works by an iterative process, starting with user supplied estimates for the parameters in the model, and finding successively better parameter estimates until certain convergence criteria are met.

In this example, we assume that we want to fit a parabola to our data, but we'll use the vertex form of the equation $(y = a \cdot (x-h) + k)$. This form is handy because the point (h, k) indicates the vertex of the parabola.

Note in the formula in the *nls* call below, that there are variables from our data (*Clutch* and *Length*), and parameters we want to estimate (*Lcenter*, *Cmax*, and *a*).

There's no set process for choosing starting estimates for the parameters. Often, the parameters will be meaningful. For example, here, *Lcenter* is the *x*-coordinate of the vertex and *Cmax* is the *y*-coordinate of the vertex. So we can guess at reasonable values for these. The parameter *a* would be difficult to guess at, though we know it should be negative because the parabola opens downward.

Because *nls* uses an iterative process based on initial estimates of the parameters, it fails to find a solution if the estimates are too far off, or it may return a set of parameter estimates that don't fit the data well. It is important to plot the solution and make sure it is reasonable. I have seen *nls* have difficulty with models that have more than three parameters. The package *nlmrt* uses a different process for determining the iterations, and may be better to fit difficult models.

If you wish to have an overall p-value for the model and a pseudo-R-squared for the model, the model will need to be compared with a null model. Technically for this to be valid, the null model must be nested within the fitted model. That means that the null model is a special case of the fitted model. In our example, if we were to force a to be zero, that would leave a model $Clutch \sim constant$, where constant would be a parameter that estimates the mean of the Clutch variable. Many theoretical models do not have this property; that is, they don't have a constant or linear term. They are therefore considered nonlinear models. In these cases, nls can still be used to fit the model, but the extra steps determining the model's overall p-value and pseudo-R-squared are technically not valid. In these cases, models could be compared with the Akaike information criterion (AIC).

The p-value for the model, relative to the null model, is determined with the extra SS (F) test (*anova* function) or likelihood ratio test (*lrtest* in the package *lmtest*).

There are various pseudo-R-squared values that have been developed for models without r-squared defined. My function *nagelkerke* calculates the McFadden, the Cox and Snell, and the Nagelkereke pseudo-R-squared. For *nls* models, a null model must be explicitly defined and passed to the function. The Nagelkereke is a modification of the Cox and Snell so that it has a maximum of 1. I find the Nagelkereke to usually be satisfactory for *nls*, *lme*, and *gls* models. As a technical note, for *gls* and *lme* models, my function uses the likelihood for the model with ML fitting (REML = FALSE).

Pseudo-R-squared values are not directly comparable to multiple R-squared values, though in the examples in this chapter, the Nagelkereke is reasonably close to the multiple R-squared for the quadratic parabola model.

```
### ------
### Nonlinear regression, turtle carapace example
### pp. 220-221
### -----
Input = ("
Length Clutch
284
      3
      2
290
      7
290
      7
290
298
      11
299
      12
302
      10
306
      8
306
      8
309
      9
310
      10
311
      13
317
      7
      9
317
320
      6
323
      13
334
      2
334
      8
```

```
")
Data = read.table(textConnection(Input),header=TRUE)
model = nls(Clutch \sim a * (Length - Lcenter)^2 + Cmax,
           data
                = Data,
           start = c(Lcenter = 310,
                      cmax =
                                 12,
                                 -1).
            trace = FALSE,
            nls.control(maxiter = 1000)
            )
summary(model)
   Parameters:
           Estimate Std. Error t value Pr(>|t|)
   Lcenter 310.72865 2.37976 130.57 < 2e-16 ***
   Cmax 10.05879
                       0.86359 11.65 6.5e-09 ***
           -0.00942
                       0.00283 -3.33 0.0045 **
```

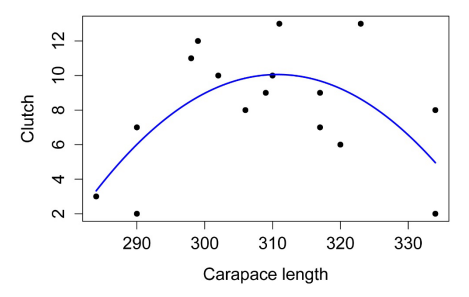
Determine overall p-value and pseudo-R-squared

```
model.null = nls(Clutch ~ I,
           data = Data,
           start = c(I = 8),
           trace = FALSE)
anova(model, model.null)
    Res.Df Res.Sum Sq Df Sum Sq F value Pr(>F)
  1
        15
               106.97
        17
               188.94 -2 -81.971 5.747 0.01403 *
library(rcompanion)
nagelkerke(fit = model,
          null = model.null)
  $Pseudo.R.squared.for.model.vs.null
                               Pseudo.R.squared
  McFadden
                                       0.109631
  Cox and Snell (ML)
                                       0.433836
  Nagelkerke (Cragg and Uhler)
                                     0.436269
```

<u>Determine confidence intervals for parameters</u>

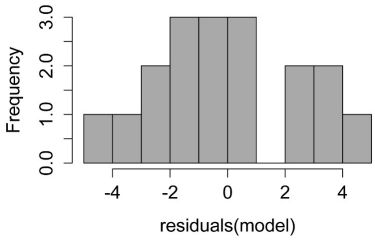
```
2.5 %
                          97.5 %
  Lcenter 305.6563154 315.800988774
  Cmax 8.2180886 11.899483768
          -0.0154538 -0.003395949
Boot=nlsBoot(model)
summary(Boot)
  _____
  Bootstrap statistics
            Estimate Std. error
  Lcenter 311.07998936 2.872859816
  Cmax 10.13306941 0.764154661
          -0.00938236 0.002599385
  a
  Median of bootstrap estimates and percentile confidence intervals
                Median 2.5%
                                     97.5%
  Lcenter 310.770796703 306.78718266 316.153528168
  Cmax 10.157560932 8.58974408 11.583719723
  a
          -0.009402318 -0.01432593 -0.004265714
```

Simple plot of model

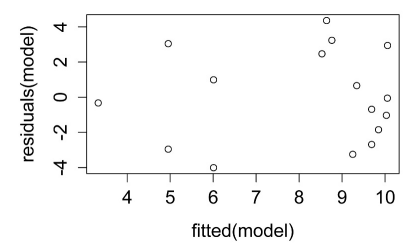


Checking assumptions of the model

Histogram of residuals(model)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

Analysis of Covariance

When to use it

The cricket example is shown in the "How to do the test" section.

Null hypotheses
Assumptions
How the test works
Examples
Graphing the results
Similar tests

See the *Handbook* for information on these topics.

How to do the test

Analysis of covariance example with two categories and type II sum of squares

This example uses type II sum of squares, but otherwise follows the example in the *Handbook*. The parameter estimates are calculated differently in R, so the calculation of the intercepts of the lines is slightly different.

```
### ------
### Analysis of covariance, cricket example
### pp. 228-229
### ------

Input = ("
Species Temp Pulse
    ex    20.8   67.9
    ex    20.8   65.1
```

```
24
                 77.3
 ex
          24
                 78.7
 ex
          24
                 79.4
 ex
 ex
          24
                 80.4
          26.2
                 85.8
 ex
          26.2
                 86.6
 ex
          26.2
                 87.5
 ex
          26.2
                 89.1
 ex
          28.4
               98.6
 ex
          29
                100.8
 ex
          30.4
               99.3
 ex
          30.4 101.7
 ex
          17.2
                44.3
 niv
          18.3
                47.2
 niv
          18.3
 niv
                47.6
          18.3
                49.6
 niv
 niv
         18.9
                 50.3
          18.9
 niv
                 51.8
          20.4
                 60
 niv
 niv
          21
                 58.5
          21
                 58.9
 niv
         22.1
                 60.7
 niv
         23.5
 niv
                 69.8
 niv
         24.2
                 70.9
         25.9
                 76.2
 niv
         26.5
 niv
                 76.1
         26.5
 niv
                 77
                 77.7
niv
          26.5
niv
          28.6
                 84.7
")
```

Data = read.table(textConnection(Input),header=TRUE)

Simple plot

```
plot(x = Data$Temp,
    y = Data$Pulse,
    col = Data$Species,
    pch = 16,
    xlab = "Temperature",
    ylab = "Pulse")

legend('bottomright',
    legend = levels(Data$Species),
    col = 1:2,
    cex = 1,
    pch = 16)
```

Analysis of covariance

```
options(contrasts = c("contr.treatment", "contr.poly"))
### These are the default contrasts in R
```

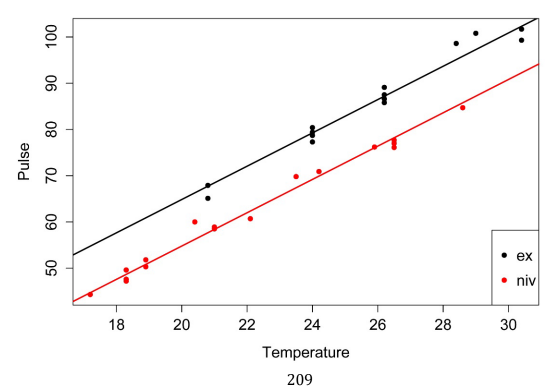
```
model.1 = lm (Pulse ~ Temp + Species + Temp:Species,
              data = Data
library(car)
Anova(model.1, type="II")
   Anova Table (Type II tests)
                Sum Sq Df F value
                                      Pr(>F)
                4376.1 1 1388.839 < 2.2e-16 ***
   Temp
                 598.0 1 189.789 9.907e-14 ***
   Species
                   4.3 1
   Temp:Species
                             1.357
                                      0.2542
   ### Interaction is not significant, so the slope across groups
   ### is not different.
model.2 = lm (Pulse ~ Temp + Species,
              data = Data
library(car)
Anova(model.2, type="II")
   Anova Table (Type II tests)
             Sum Sq Df F value
                                  Pr(>F)
             4376.1 1 1371.4 < 2.2e-16 ***
   Temp
              598.0 1
                       187.4 6.272e-14 ***
   Species
   ### The category variable (Species) is significant,
   ### so the intercepts among groups are different
summary(model.2)
   Coefficients:
                Estimate Std. Error t value Pr(>|t|)
                           2.55094 -2.827 0.00858 **
               -7.21091
   (Intercept)
                            0.09729 37.032 < 2e-16 ***
   Temp
                 3.60275
   Speciesniv -10.06529
                            0.73526 -13.689 6.27e-14 ***
   ### Note that these estimates are different than in the Handbook,
         but the calculated results will be identical.
   ### The slope estimate is the same.
   ### The intercept for species 1 (ex) is (intercept).
   ### The intercept for species 2 (niv) is (intercept) + Speciesniv.
   ### This is determined from the contrast coding of the Species
   ### variable shown below, and the fact that Speciesniv is shown in
   ### coefficient table above.
```

contrasts(Data\$Species)

```
niv
ex 0
niv 1
```

Simple plot with fitted lines

```
I.nought = -7.21091
I1 = I.nought + 0
I2 = I.nought + -10.06529
B = 3.60275
plot(x
         = Data$Temp,
     y = Data$Pulse,
     col = Data$Species,
     pch = 16,
     xlab = "Temperature",
     ylab = "Pulse")
legend('bottomright',
       legend = levels(Data$Species),
       col = 1:2,
       cex = 1,
       pch = 16)
abline(I1, B,
       lty=1, lwd=2, col = 1)
abline(I2, B,
       lty=1, lwd=2, col = 2)
```



p-value and R-squared of combined model

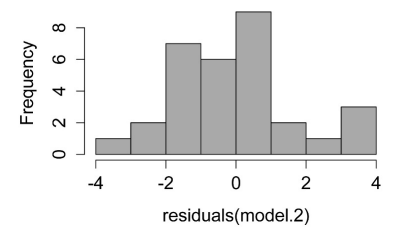
```
summary(model.2)

Multiple R-squared: 0.9896, Adjusted R-squared: 0.9888
F-statistic: 1331 on 2 and 28 DF, p-value: < 2.2e-16</pre>
```

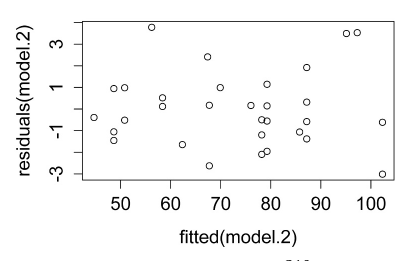
Checking assumptions of the model

```
hist(residuals(model.2),
     col="darkgray")
```

Histogram of residuals(model.2)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

```
### additional model checking plots with: plot(model.2)
### alternative: library(FSA); residPlot(model.2)
```

*Analysis of covariance example with three categories and type II sum of squares*This example uses type II sum of squares, and considers a case with three groups.

```
### Analysis of covariance, hypothetical data
Input = ("
Species Temp
                Pulse
          20.8
                 67.9
ex
          20.8
                 65.1
 ex
          24
                 77.3
 ex
         24
                 78.7
 ex
          24
                 79.4
 ex
          24
                 80.4
 ex
          26.2
                 85.8
 ex
          26.2
                 86.6
 ex
          26.2
                 87.5
 ex
 ex
          26.2
                 89.1
          28.4
                 98.6
 ex
          29
                100.8
 ex
          30.4
                99.3
 ex
          30.4 101.7
 ex
                 44.3
 niv
          17.2
 niv
          18.3
                 47.2
          18.3
                 47.6
 niv
          18.3
                 49.6
 niv
          18.9
 niv
                 50.3
 niv
          18.9
                 51.8
 niv
          20.4
                 60
          21
                 58.5
 niv
          21
                 58.9
 niv
          22.1
                 60.7
 niv
          23.5
 niv
                 69.8
          24.2
                 70.9
 niv
          25.9
                 76.2
 niv
          26.5
                 76.1
 niv
          26.5
                 77
 niv
 niv
          26.5
                 77.7
          28.6
                 84.7
 niv
          17.2
 fake
                 74.3
 fake
          18.3
                 77.2
 fake
          18.3
                 77.6
 fake
         18.3
                 79.6
```

```
fake
            18.9
                   80.3
   fake
           18.9
                   81.8
    fake
            20.4
                   90
    fake
            21
                   88.5
            21
    fake
                   88.9
            22.1
   fake
                   90.7
            23.5
                   99.8
    fake
   fake
           24.2
                   100.9
   fake
           25.9
                   106.2
   fake
           26.5
                   106.1
         26.5
   fake
                   107
   fake
           26.5
                   107.7
   fake
            28.6
                   114.7
   ")
  Data = read.table(textConnection(Input), header=TRUE)
Simple plot
  plot(x = Data$Temp,
       y = Data$Pulse,
       col = Data$Species,
       pch = 16,
       xlab = "Temperature",
       ylab = "Pulse")
  legend('bottomright',
          legend = levels(Data$Species),
         col = 1:3,
         cex = 1,
         pch = 16
Analysis of covariance
  options(contrasts = c("contr.treatment", "contr.poly"))
     ### These are the default contrasts in R
  model.1 = lm (Pulse ~ Temp + Species + Temp:Species,
                data = Data
  library(car)
  Anova(model.1, type="II")
                  Sum Sq Df F value Pr(>F)
                  7026.0 1 2452.4187 <2e-16 ***
     Temp
                  7835.7 2 1367.5377 <2e-16 ***
     Species
                     5.2 2 0.9126 0.4093
     Temp:Species
     ### Interaction is not significant, so the slope among groups
     ### is not different.
```

```
model.2 = lm (Pulse ~ Temp + Species,
             data = Data
library(car)
Anova(model.2, type="II")
            Sum Sq Df F value Pr(>F)
            7026.0 1 2462.2 < 2.2e-16 ***
            7835.7 2 1373.0 < 2.2e-16 ***
  Species
  Residuals 125.6 44
  ### The category variable (Species) is significant,
  ### so the intercepts among groups are different
summary(model.2)
  Coefficients:
               Estimate Std. Error t value Pr(>|t|)
   (Intercept) -6.35729 1.90713 -3.333 0.00175 **
  Temp
                3.56961
                           0.07194 49.621 < 2e-16 ***
  Speciesfake 19.81429 0.66333 29.871 < 2e-16 ***
  Speciesniv -10.18571 0.66333 -15.355 < 2e-16 ***
  ### The slope estimate is the Temp coefficient.
  ### The intercept for species 1 (ex) is (intercept).
  ### The intercept for species 2 (fake) is (intercept) + Speciesfake.
  ### The intercept for species 3 (niv) is (intercept) + Speciesniv.
  ### This is determined from the contrast coding of the Species
  ### variable shown below.
contrasts(Data$Species)
       fake niv
          0 0
  ex
              0
  fake
          1
  niv
          0
            1
```

Simple plot with fitted lines

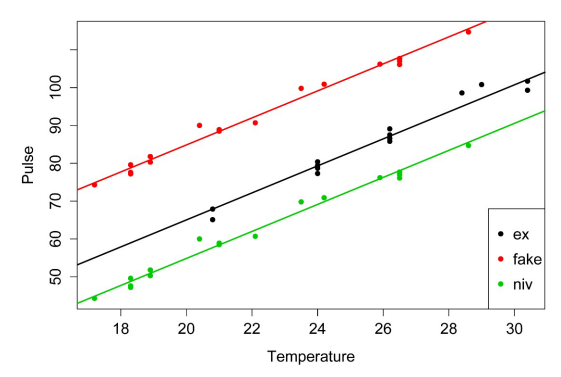
```
xlab = "Temperature",
   ylab = "Pulse")

legend('bottomright',
        legend = levels(Data$species),
        col = 1:3,
        cex = 1,
        pch = 16)

abline(I1, B,
        lty=1, lwd=2, col = 1)

abline(I2, B,
        lty=1, lwd=2, col = 2)

abline(I3, B,
        lty=1, lwd=2, col = 3)
```



p-value and R-squared of combined model

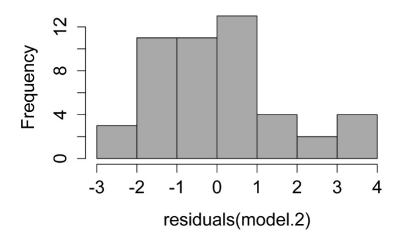
```
summary(model.2)

Multiple R-squared: 0.9919, Adjusted R-squared: 0.9913
F-statistic: 1791 on 3 and 44 DF, p-value: < 2.2e-16</pre>
```

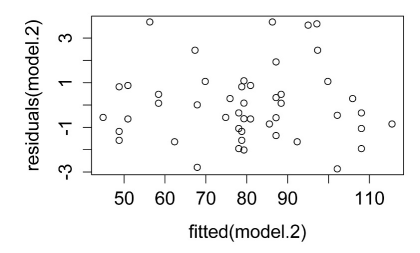
Checking assumptions of the model

```
hist(residuals(model.2),
     col="darkgray")
```

Histogram of residuals(model.2)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

additional model checking plots with: plot(model.2)
alternative: library(FSA); residPlot(model.2)

Power analysis

See the *Handbook* for information on this topic.

Multiple Regression

When to use it
Null hypothesis
How it works
Using nominal variables in a multiple regression
Selecting variables in multiple regression
Assumptions

See the *Handbook* for information on these topics.

Example

The Maryland Biological Stream Survey example is shown in the "How to do the multiple regression" section.

Graphing the results Similar tests

See the *Handbook* for information on these topics.

How to do multiple regression

Multiple correlation

Whenever you have a dataset with multiple numeric variables, it is a good idea to look at the correlations among these variables. One reason is that if you have a dependent variable, you can easily see which independent variables correlate with that dependent variable. A second reason is that if you will be constructing a multiple regression model, adding an independent variable that is strongly correlated with an independent variable already in the model is unlikely to improve the model much, and you may have a good reason to chose one variable over another.

Finally, it is worthwhile to look at the distribution of the numeric variables. If the distributions differ greatly, using Kendall or Spearman correlations may be more appropriate. Also, if independent variables differ in distribution from the dependent variable, the independent variables may need to be transformed. In this example, *Longnose*, *Acreage*, *Maxdepth*, *NO3*, and *SO4* are relatively log-normally distributed, while *DO2* and *Temp* are relatively normal in distribution. It may be advisable in this case to transform these variable so that they all have similar distributions (not shown here).

With the *corr.test* function in the *psych package*, the "Correlation matrix" shows r-values and the "Probability values" table shows p-values. The *PerformanceAnalytics* plot shows r-values, with asterisks indicating significance, as well as a histogram of the individual variables. Either of these indicates that *Longnose* is significantly correlated with *Acreage*, *Maxdepth*, and *NO3*.

```
### ------### Multiple correlation and regression, stream survey example ### pp. 236-237
```

<pre>Input = ("</pre>							
Stream	Longnose	Acerage	DO2	Maxdepth	NO3	S04	Temp
BASIN_RUN	13	2528	9.6	80	2.28	16.75	15.3
BEAR_BR	12	3333	8.5	83	5.34	7.74	19.4
BEAR_CR	54	19611	8.3	96	0.99	10.92	19.5
BEAVER_DAM_CR	19	3570	9.2	56	5.44	16.53	17
BEAVER_RUN	37	1722	8.1	43	5.66	5.91	19.3
BENNETT_CR	2	583	9.2	51	2.26	8.81	12.9
BIG_BR	72	4790	9.4	91	4.1	5.65	16.7
BIG_ELK_CR	164	35971	10.2	81	3.2	17.53	13.8
BIG_PIPE_CR	18	25440	7.5	120	3.53	8.2	13.7
BLUE_LICK_RUN	1	2217	8.5	46	1.2	10.85	14.3
BROAD_RUN	53	1971	11.9	56	3.25	11.12	22.2
BUFFALO_RUN	16	12620	8.3	37	0.61	18.87	16.8
BUSH_CR	32	19046	8.3	120	2.93	11.31	18
CABIN_JOHN_CR	21	8612	8.2	103	1.57	16.09	15
CARROLL_BR	23	3896	10.4	105	2.77	12.79	18.4
COLLIER_RUN	18	6298	8.6	42	0.26	17.63	18.2
CONOWINGO_CR	112	27350	8.5	65	6.95	14.94	24.1
DEAD_RUN	25	4145	8.7	51	0.34	44.93	23
DEEP_RUN	5	1175	7.7	57	1.3	21.68	21.8
DEER_CR	26	8297	9.9	60	5.26	6.36	19.1
DORSEY_RUN	8	7814	6.8	160	0.44	20.24	22.6
FALLS_RUN	15	1745	9.4		2.19	10.27	14.3
FISHING_CR	11	5046	7.6	109	0.73	7.1	19
FLINTSTONE_CR	11	18943	9.2	50	0.25	14.21	18.5
GREAT_SENECA_CR	87	8624	8.6	78	3.37	7.51	21.3
GREENE_BR	33	2225	9.1		2.3	9.72	20.5
GUNPOWDER_FALLS	22	12659	9.7	65	3.3	5.98	18
HAINES_BR	98	1967	8.6	50	7.71	26.44	16.8
HAWLINGS_R	1	1172	8.3	73	2.62	4.64	20.5
HAY_MEADOW_BR	5	639	9.5	26	3.53	4.46	20.1
HERRINGTON_RUN	1	7056	6.4	60	0.25	9.82	24.5
HOLLANDS_BR	38	1934	10.5	85	2.34	11.44	12
ISRAEL_CR	30	6260	9.5	133	2.41	13.77	21
LIBERTY_RES	12	424	8.3	62	3.49	5.82	20.2
LITTLE_ANTIETAM_CR	24	3488	9.3	44	2.11	13.37	24
LITTLE_BEAR_CR	6	3330	9.1	67	0.81	8.16	14.9
	15	2227	6.8	54	0.33	7.6	24
LITTLE_CONOCOCHEAGUE_CR LITTLE_DEER_CR	38	8115	9.6	110	3.4	9.22	20.5
	84	1600	10.2	56	3.54	5.69	19.5
LITTLE_FALLS	3	15305	9.7			6.96	17.5
LITTLE_GUNPOWDER_R LITTLE_HUNTING_CR	18	7121	9.5	58	2.6 0.51	7.41	16
LITTLE_PAINT_BR	63	5794	9.4		1.19	12.27	17.5
MAINSTEM_PATUXENT_R	239	8636	8.4		3.31	5.95	18.1
MEADOW_BR	234	4803	8.5	93	5.01	10.98	24.3
MILL_CR	6	1097	8.3	53	1.71	15.77	13.1
	76	9765	9.3		4.38	5.74	16.9
MORGAN_RUN	25	4266	8.9		2.05	12.77	17
MUDDY_BR	8	1507	7.4		0.84	16.3	21
MUDLICK_RUN	23	3836	8.3	121	1.32	7.36	18.5
NORTH_BR	16		7.4		0.29	2.5	18.3
NORTH_BR_CASSELMAN_R		17419					
NORTHWEST_BR ANACOSTTA B	6	8735	8.2	63	1.56	13.22	20.8
NORTHWEST_BR_ANACOSTIA_R	100	22550	8.4		1.41	14.45	23
OWENS_CR	80	9961	8.6	79 61	1.02	9.07	21.8
PATAPSCO_R	28	4706	8.9	61	4.06	9.9	19.7
PINEY_BR	48	4011	8.3	52 100	4.7	5.38	18.9
PINEY_CR	18	6949	9.3		4.57	17.84	18.6
PINEY_RUN	36 19	11405 904	9.2	70 39	2.17	10.17	23.6
PRETTYBOY_BR	13	904	9.8	צכ	6.81	9.2	19.2

```
RED_RUN
                        32
                                  3332
                                         8.4 73
                                                       2.09
                                                             5.5
                                                                    17.7
                                         6.8 33
                                                       2.47
ROCK_CR
                        3
                                   575
                                                             7.61
                                                                    18
                                         7.7 73
                       106
                                 29708
                                                       0.63 12.28
                                                                    21.4
SAVAGE_R
SECOND_MINE_BR
                        62
                                  2511
                                        10.2 60
                                                       4.17 10.75
                                                                    17.7
                                 18422
SENECA_CR
                        23
                                         9.9 45
                                                       1.58
                                                             8.37
                                                                    20.1
                                         7.6 46
                        2
SOUTH_BR_CASSELMAN_R
                                  6311
                                                       0.64 21.16
                                                                    18.5
                        26
SOUTH_BR_PATAPSCO
                                  1450
                                        7.9 60
                                                       2.96
                                                             8.84
                                                                    18.6
SOUTH_FORK_LINGANORE_CR
                        20
                                  4106
                                        10.0 96
                                                       2.62
                                                             5.45
                                                                    15.4
                        38
                                 10274
                                         9.3 90
                                                       5.45 24.76
TUSCARORA_CR
                                                                    15
                                                       5.25 14.19
WATTS_BR
                        19
                                   510
                                         6.7 82
                                                                    26.5
")
Data = read.table(textConnection(Input),header=TRUE)
### Create a new data frame with only the numeric variables.
### This is required for corr.test and chart.Correlation
library(dplyr)
Data.num =
   select(Data,
          Longnose,
          Acerage,
          DO2,
          Maxdepth,
          NO3,
          SO4,
          Temp)
library(FSA)
headtail(Data.num)
      Longnose Acerage DO2 Maxdepth NO3
                                            SO4 Temp
   1
            13
                  2528 9.6
                                 80 2.28 16.75 15.3
   2
            12
                  3333 8.5
                                  83 5.34 7.74 19.4
   3
            54
                 19611 8.3
                                  96 0.99 10.92 19.5
   66
            20
                 4106 10.0
                                  96 2.62 5.45 15.4
                                  90 5.45 24.76 15.0
   67
            38
                10274 9.3
                                  82 5.25 14.19 26.5
            19
                   510 6.7
   68
library(psych)
corr.test(Data.num,
          use = "pairwise",
          method="pearson",
          adjust="none",  # Can adjust p-values; see ?p.adjust for options
          alpha=.05)
   Correlation matrix
            Longnose Acerage DO2 Maxdepth
                                              NO3
                                                    SO4 Temp
                1.00
                        0.35 0.14 0.30 0.31 -0.02 0.14
   Lonanose
   Acerage
                0.35
                        1.00 -0.02
                                       0.26 -0.10 0.05 0.00
                                      -0.06 0.27 -0.07 -0.32
                0.14
                       -0.02 1.00
   DO2
```

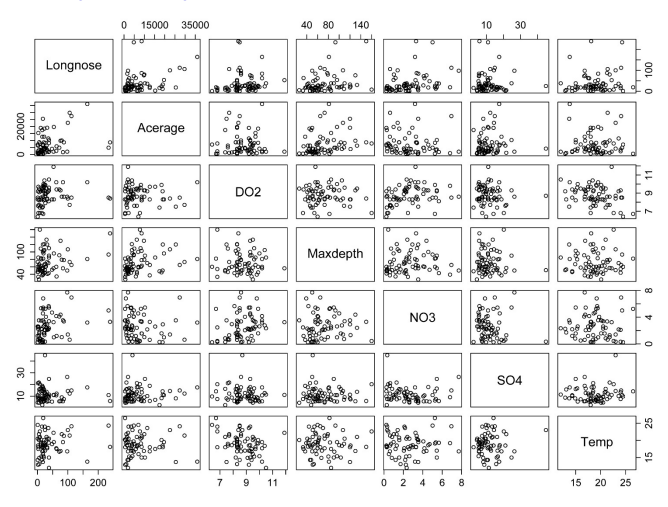
```
Maxdepth
             0.30
                    0.26 - 0.06
                                   1.00
                                         0.04 -0.05 0.00
NO3
             0.31
                    -0.10 0.27
                                   0.04
                                         1.00 -0.09 0.00
            -0.02
                    0.05 -0.07
                                   -0.05 -0.09 1.00 0.08
S04
Temp
             0.14
                    0.00 - 0.32
                                   0.00 0.00 0.08 1.00
Sample Size
```

Probability values (Entries above the diagonal are adjusted for multiple tests.)

```
Longnose Acerage DO2 Maxdepth NO3 SO4 Temp
             0.00
                     0.00 0.27
                                   0.01 0.01 0.89 0.26
Longnose
Acerage
             0.00
                     0.00 0.86
                                   0.03 0.42 0.69 0.98
                     0.86 0.00
             0.27
                                   0.64 0.02 0.56 0.01
DO2
Maxdepth
             0.01
                     0.03 0.64
                                   0.00 0.77 0.69 0.97
NO3
             0.01
                     0.42 0.02
                                   0.77 0.00 0.48 0.99
             0.89
                     0.69 0.56
                                   0.69 0.48 0.00 0.52
S04
                                   0.97 0.99 0.52 0.00
             0.26
                     0.98 0.01
Temp
```

pairs(data=Data,

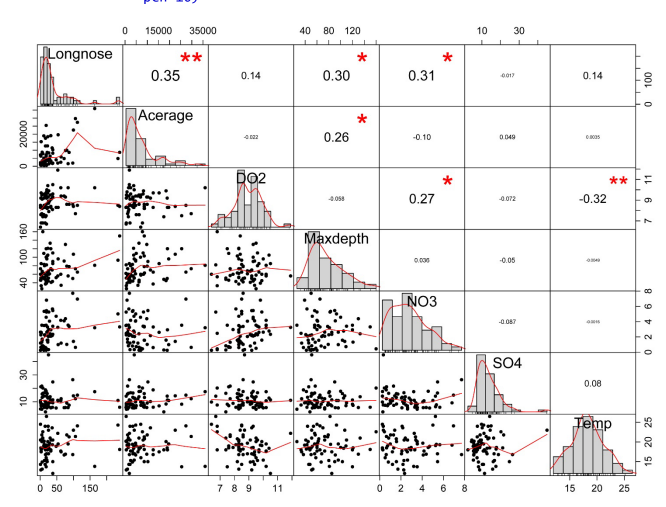
~ Longnose + Acerage + DO2 + Maxdepth + NO3 + SO4 + Temp)



library(PerformanceAnalytics)

chart.Correlation(Data.num,

method="pearson",
histogram=TRUE,
pch=16)



Multiple regression

Model selection using the *step* function

The *step* function has options to add terms to a model (*direction="forward"*), remove terms from a model (*direction="backward"*), or to use a process that both adds and removes terms (*direction="both"*). It uses AIC (Akaike information criterion) as a selection criterion. You can use the option k = log(n) to use BIC instead.

You can add the *test="F"* option to see the p-value for adding or removing terms, but the test will still follow the AIC statistic. If you use this, however, note that a significant p-value essentially argues for the term being included in the model, whether it's its addition or its removal that's being considered.

A full model and a null are defined, and then the function will follow a procedure to find the model with the lowest AIC. The final model is shown at the end of the output, with the *Call:* indication, and lists the coefficients for that model.

Stepwise procedure

```
model.null = lm(Longnose ~ 1,
                   data=Data)
   model.full = lm(Longnose ~ Acerage + DO2 + Maxdepth + NO3 + SO4 + Temp,
                   data=Data)
   step(model.null,
        scope = list(upper=model.full),
                direction="both",
                data=Data)
      Longnose ~ 1
                 Df Sum of Sq RSS
                1 17989.6 131841 518.75
      + Acerage
      + NO3
                  1
                      14327.5 135503 520.61
      + Maxdepth 1 13936.1 135894 520.81
      <none>
                              149831 525.45
     + Temp 1 2931.0 146899 526.10
+ DO2 1 2777.7 147053 526.17
+ SO4 1 45.3 149785 527.43
      < snip... more steps >
      Longnose ~ Acerage + NO3 + Maxdepth
                 Df Sum of Sq
                                RSS
                               107904 509.13
      <none>
               1 2948.0 104956 509.24
      + Temp
                       669.6 107234 510.70
      + DO2
                 1
     - Maxdepth 1 6058.4 113962 510.84
+ SO4 1 5.9 107898 511.12
      - Acerage 1 14652.0 122556 515.78
      NO3
               1 16489.3 124393 516.80
      call:
      lm(formula = Longnose ~ Acerage + NO3 + Maxdepth, data = Data)
      Coefficients:
      (Intercept) Acerage NO3
-23.829067 0.001988 8.673044
                                                 Maxdepth
                                                 0.336605
Define final model
   model.final = lm(Longnose ~ Acerage + Maxdepth + NO3,
                    data=Data)
   summary(model.final) # Show coefficients, R-squared, and overall p-value
                    Estimate Std. Error t value Pr(>|t|)
      (Intercept) -2.383e+01 1.527e+01 -1.560 0.12367
```

```
Acerage 1.988e-03 6.742e-04 2.948 0.00446 **
Maxdepth 3.366e-01 1.776e-01 1.896 0.06253 .
NO3 8.673e+00 2.773e+00 3.127 0.00265 **

Multiple R-squared: 0.2798, Adjusted R-squared: 0.2461
F-statistic: 8.289 on 3 and 64 DF, p-value: 9.717e-05
```

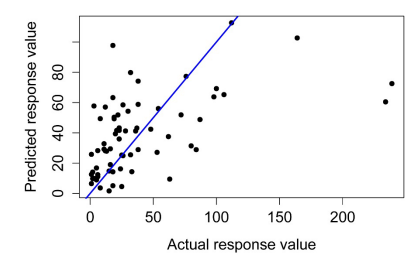
Analysis of variance for individual terms

Simple plot of predicted values with 1-to-1 line

```
Data$predy = predict(model.final)

plot(predy ~ Longnose,
    data=Data,
    pch = 16,
    xlab="Actual response value",
    ylab="Predicted response value")

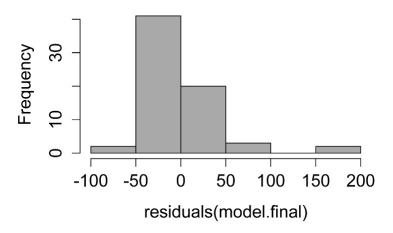
abline(0,1, col="blue", lwd=2)
```



Checking assumptions of the model

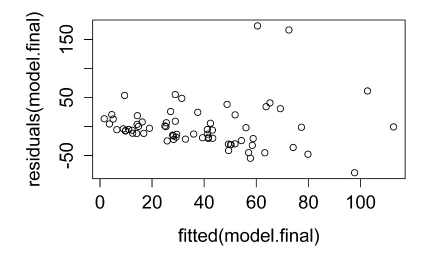
hist(residuals(model.final),
 col="darkgray")

Histogram of residuals(model.final)



A histogram of residuals from a linear model. The distribution of these residuals should be approximately normal.

plot(fitted(model.final),
 residuals(model.final))



A plot of residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

additional model checking plots with: plot(model.final)

Model fit criteria

Model fit criteria are available to decide which model is most appropriate. The step function uses AIC, or optionally BIC, but there are others. You don't want to use multiple R-squared, because it will continue to improve as more terms are added into the model. Instead, you want to use a criterion that balances the improvement in explanatory power with not adding extraneous terms to the model. Adjusted R-squared is a modification of R-squared that includes this balance. Larger is better. AIC is based on information theory and measures this balance. AICc is an adjustment to AIC that is more appropriate for data sets with relatively fewer observations. BIC is similar to AIC, but penalizes more for additional terms in the model. Smaller is better for AIC, AICc, and BIC. There are differing opinions on which model fitting criteria is best to use, but if you have no opinion, I would recommend AICc for routine use.

Using the *step* procedure to automatically find an optimal model is an option, but some people caution against using an automated procedure because it might not hone in on the best model. Instead, you can look at the model fit criteria for competing models manually. There may be reasons why you wish to include or exclude some terms in the model, and it may be useful to look at different model selection criteria simultaneously.

In my *compare.lm* function below, *Shapiro.W* and *Shapiro.p* are results from the Shapiro–Wilks test for normality on the model residuals. A higher Shapiro W and a higher Shapiro p indicate that the residuals are more normally distributed. You should be aware, however, that any model with a high number of observation may yield a significant p-value (p < 0.05) for the Shapiro–Wilks test. It is best to investigate the residuals visually.

In the following example, we'll look only at the terms that are significantly correlated with *Longnose* (*Acreage*, *Maxdepth*, and *NO3*), and then add in the other terms just to show the decrease in AICc by adding extra terms.

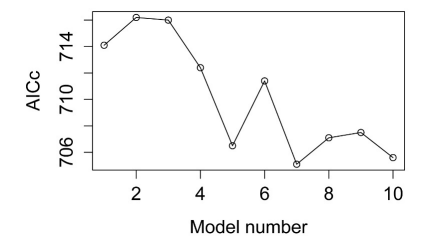
Note that AIC and BIC are calculated differently than in the *step* function.

```
model.1 = lm(Longnose ~ Acerage,
                                                           data=Data)
model.2 = lm(Longnose \sim Maxdepth,
                                                           data=Data)
model.3 = lm(Longnose \sim NO3,
                                                           data=Data)
model.4 = lm(Longnose ~ Acerage + Maxdepth,
                                                           data=Data)
model.5 = lm(Longnose ~ Acerage + NO3,
                                                           data=Data)
model.6 = lm(Longnose \sim Maxdepth + NO3,
                                                           data=Data)
model.7 = lm(Longnose ~ Acerage + Maxdepth + NO3,
                                                           data=Data)
model.8 = lm(Longnose ~ Acerage + Maxdepth + NO3 + DO2, data=Data)
model.9 = lm(Longnose ~ Acerage + Maxdepth + NO3 + SO4, data=Data)
model.10 = lm(Longnose ~ Acerage + Maxdepth + NO3 + Temp, data=Data)
library(rcompanion)
compareLM(model.1, model.2, model.3, model.4, model.5, model.6,
          model.7, model.8, model.9, model.10)
   $Models
      Formula
     "Longnose ~ Acerage"
   2 "Longnose ~ Maxdepth"
```

```
"Longnose ~ NO3"
  "Longnose ~ Acerage + Maxdepth"
   "Longnose ~ Acerage + NO3"
  "Longnose ~ Maxdepth + NO3"
7
  "Longnose ~ Acerage + Maxdepth + NO3"
  "Longnose ~ Acerage + Maxdepth + NO3 + DO2"
  "Longnose ~ Acerage + Maxdepth + NO3 + SO4"
10 "Longnose ~ Acerage + Maxdepth + NO3 + Temp"
$Fit.criteria
  Rank Df.res
                           BIC R.squared Adj.R.sq
                                                    p.value Shapiro.W Shapiro.p
                AIC AICC
           66 713.7 714.1 720.4
                                 0.12010 0.10670 3.796e-03
                                                               0.7278 6.460e-10
           66 715.8 716.2 722.4
                                 0.09301 0.07927 1.144e-02
                                                               0.7923 2.115e-08
3
     2
           66 715.6 716.0 722.2
                                 0.09562 0.08192 1.029e-02
                                                               0.7361 9.803e-10
4
     3
           65 711.8 712.4 720.6
                                 0.16980 0.14420 2.365e-03
                                                               0.7934 2.250e-08
5
     3
           65 705.8 706.5 714.7
                                 0.23940 0.21600 1.373e-04
                                                               0.7505 2.055e-09
6
     3
                                 0.18200 0.15690 1.458e-03
                                                               0.8149 8.405e-08
           65 710.8 711.4 719.6
7
     4
                                                               0.8108 6.511e-08
           64 704.1 705.1 715.2
                                 0.27980 0.24610 9.717e-05
8
           63 705.7 707.1 719.0
                                 0.28430 0.23890 2.643e-04
                                                               0.8041 4.283e-08
9
           63 706.1 707.5 719.4
                                 0.27990 0.23410 3.166e-04
                                                               0.8104 6.345e-08
10
           63 704.2 705.6 717.5
                                 0.29950 0.25500 1.409e-04
                                                               0.8225 1.371e-07
```

Model 7 is the model which minimizes AICc, which is the same model ### chosen by the step function

lines(Result\$Fit.criteria\$AICc)



A plot of AICc (modified Akaike information criterion) of several models. Model 7 minimizes AICc, and is therefore chosen as the best model out of this set.

Comparing models with likelihood ratio test

It may also be helpful to compare models with the extra sum of squares test or likelihood ratio test to see if additional terms significantly reduce the error sum of squares.

One of the compared models should be nested within the other. That is, the one model should be the same as the other, except with additional terms. For example in the set of models below, it is appropriate to compare *model.7* to *model.4*. Or to compare each of *model.8*, *model.9*, and *model.10* to *model.7*.

For a single comparison, the *anova* function can be used for the Extra SS test, or *Irtest* in *Imtest* can be used for the likelihood ratio test. For multiple comparisons, the *extraSS* and *Irt* functions in the *FSA* package can be used. The *extraSS* function works only for *Im* and *nIs* models, whereas the *Irt* function works on a wider range of model objects.

```
model.4 = lm(Longnose ~ Acerage + Maxdepth,
                                                               data=Data)
model.7 = lm(Longnose ~ Acerage + Maxdepth + NO3,
                                                               data=Data)
model.8 = lm(Longnose ~ Acerage + Maxdepth + NO3 + DO2, data=Data)
model.9 = lm(Longnose ~ Acerage + Maxdepth + NO3 + SO4, data=Data)
model.10 = lm(Longnose ~ Acerage + Maxdepth + NO3 + Temp, data=Data)
anova(model.7, model.4)
   Analysis of Variance Table
   Model 1: Longnose ~ Acerage + Maxdepth + NO3
   Model 2: Longnose ~ Acerage + Maxdepth
     Res.Df
                RSS Df Sum of Sq F Pr(>F)
   1
         64 107904
         65 124393 -1 -16489 9.7802 0.002654 **
library(lmtest)
lrtest(model.7, model.4)
   Likelihood ratio test
   Model 1: Longnose ~ Acerage + Maxdepth + NO3
   Model 2: Longnose ~ Acerage + Maxdepth
     #Df LogLik Df Chisq Pr(>Chisq)
   1 5 -347.05
       4 -351.89 -1 9.6701 0.001873 **
library(FSA)
extrass(model.8, model.9, model.10,
        com=mode1.7)
   Model 1: Longnose ~ Acerage + Maxdepth + NO3 + DO2
```

```
Model 2: Longnose ~ Acerage + Maxdepth + NO3 + SO4
  Model 3: Longnose ~ Acerage + Maxdepth + NO3 + Temp
  Model A: Longnose ~ Acerage + Maxdepth + NO3
      Dfo
               RSSO DfA
                             RSSA Df
                                            SS
                                                    F Pr(>F)
   1vA 63 107234.38 64 107903.97 -1 -669.59 0.3934 0.5328
   2VA 63 107898.06 64 107903.97 -1
                                       -5.91 0.0035 0.9533
   3vA 63 104955.97 64 107903.97 -1 -2948.00 1.7695 0.1882
lrt(model.8, model.9, model.10,
   com=mode1.7)
  Model 1: Longnose ~ Acerage + Maxdepth + NO3 + DO2
  Model 2: Longnose ~ Acerage + Maxdepth + NO3 + SO4
  Model 3: Longnose ~ Acerage + Maxdepth + NO3 + Temp
  Model A: Longnose ~ Acerage + Maxdepth + NO3
      DfO
             logLikO DfA
                            logLikA Df
                                          logLik Chisq Pr(>Chisq)
  1vA 63 -346.83881 64 -347.05045 -1
                                          0.21164 0.4233
   2vA 63 -347.04859 64 -347.05045 -1
                                          0.00186 0.0037
                                                             0.9513
   3vA 63 -346.10863 64 -347.05045 -1
                                          0.94182 1.8836
                                                            0.1699
```

Power analysis

See the *Handbook* for information on this topic.

Simple Logistic Regression

When to use it Null hypothesis How the test works Assumptions

See the *Handbook* for information on these topics.

Examples

The Mpi example is shown below in the "How to do the test" section.

Graphing the results Similar tests

See the *Handbook* for information on these topics.

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(car)){install.packages("car")}
if(!require(lmtest){install.packages("lmtest")}
if(!require(tidyr)){install.packages("tidyr")}
if(!require(rcompanion)){install.packages("rcompanion")}
if(!require(FSA){install.packages("FSA")}
if(!require(popbio)){install.packages("popbio")}
```

How to do the test

Logistic regression can be performed in R with the *glm* (generalized linear model) function. This function uses a link function to determine which kind of model to use, such as logistic, probit, or Poisson. These are indicated in the *family* and *link* options. See *?glm* and *?family* for more information.

Assumptions

Generalized linear models have fewer assumptions than most common parametric tests. Observations still need to be independent, and the correct link function needs to be specified. So, for example you should understand when to use a poisson regression, and when to use a logistic regression. However, the normal distribution of data or residuals is not required.

Specifying the counts of "successes" and "failures"

Logistic regression has a dependent variable with two levels. In R, this can be specified in three ways. 1) The dependent variable can be a factor variable where the first level is interpreted as "failure" and the other levels are interpreted as "success". (As in the second example in this chapter). 2) The dependent variable can be a vector of proportions of successes, with the caveat that the number of observations for each proportion is indicated in the *weights* option. 3) The dependent variable can be a matrix with two columns, with the first column being the number of "successes" and the second being the number of "failures". (As in the first example in this chapter).

Not all proportions or counts are appropriate for logistic regression analysis

Note that in each of these specifications, both the number of successes and the number of failures is known. You should not perform logistic regression on proportion data where you don't know (or don't tell R) how many individuals went into those proportions. In statistics, 75% is different if it means 3 out of 4 rather than 150 out of 200. As another example where logistic regression doesn't apply, the weight people lose in a diet study expressed as a proportion of initial weight cannot be interpreted as a count of "successes" and "failures". Here, you might be able to use common parametric methods, provided the model assumptions are met; log or arc-sine transformations may be appropriate. Likewise, if you count the number of people in front of you in line, you can't interpret this as a percentage of people since you don't know how many people are *not* in front of you in line. In this case with count data as the dependent variable, you might use poisson regression.

Overdispersion

One potential problem to be aware of when using generalized linear models is overdispersion. This occurs when the residual deviance of the model is high relative to the residual degrees of freedom. It is basically an indication that the model doesn't fit the data well.

It is my understanding, however, that overdispersion is technically not a problem for a simple logistic regression, that is one with a binomial dependent and a single continuous independent variable. Overdispersion is discussed in the chapter on *Multiple logistic regression*.

Pseudo-R-squared

R does not produce r-squared values for generalized linear models, and common r-squared are not defined for these models. However, a variety of pseudo r-squared measures can be reported.

My function *nagelkerke* will calculate the McFadden, Cox and Snell, and Nagelkereke pseudo r-squared for glm and other model fits. The Cox and Snell is also called the ML, and the Nagelkerke is also called the Cragg and Uhler. These pseudo r-squared values compare the maximum likelihood of the model to a nested null model fit with the same method. They should not be thought of as the same as the R-squared from an ordinary-least-squares linear (OLS) model, but instead as a relative measure among similar models. The Cox and Snell and Efron's pseudo R-squared for an OLS linear model, however, will be equivalent to R-squared for that model.

Some authors recommend McFadden pseudo-R-squared for logistic regression. (See the Cross Validated discussion in the References.) Efron's pseudo r-squared and count pseudo r-squared are also recommended (see IDRE in the References).

Testing for p-values

Note that testing p-values for a logistic or poisson regression uses Chi-square tests. This is achieved through the *test="Wald"* option in *Anova* to test the significance of each coefficient, and the *test="Chisq"* option in *anova* for the significance of the overall model. A likelihood ratio test can also be used to test the significance of the overall model.

Logistic regression example

```
### Logistic regression, amphipod example, p. 247
Input = ("
                    Latitude mpi90
                                     mpi 100
 Location
 Port_Townsend,_WA 48.1
                                47
                                      139
 Neskowin,_OR
                    45.2
                               177
                                      241
 Siuslaw_R.,_OR
                    44.0
                              1087
                                     1183
 Umpqua_R.,_OR
                    43.7
                               187
                                      175
 Coos_Bay,_OR
                                      671
                    43.5
                               397
                                40
                                       14
 San_Francisco,_CA 37.8
                    36.6
                                39
                                       17
 Carmel,_CA
 Santa_Barbara,_CA 34.3
                                30
                                        0
```

```
Data$Total = Data$mpi90 + Data$mpi100
Data$Percent = Data$mpi100 / + Data$Total
```

Model fitting

Coefficients and exponentiated cofficients

```
summary(model)
  Coefficients:
            Estimate Std. Error z value Pr(>|z|)
  Latitude 0.17864
                      0.02104 8.490 <2e-16 ***
confint(model)
                2.5 % 97.5 %
  (Intercept) -9.5003746 -5.8702453
  Latitude 0.1382141 0.2208032
exp(model$coefficients)  # exponentiated coefficients
  (Intercept) Latitude
  0.0004775391 1.1955899446
exp(confint(model))
                   # 95% CI for exponentiated coefficients
                  2.5 %
                          97.5 %
  (Intercept) 7.482379e-05 0.002822181
  Latitude 1.148221e+00 1.247077992
```

Analysis of variance for individual terms

```
Latitude 1 72.076 < 2.2e-16 ***
```

Overall p-value for model

```
anova(model,
     update(model, ~1),
                           # update here produces null model for comparison
     test="Chisq")
  Analysis of Deviance Table
  Model 1: Trials ~ Latitude
  Model 2: Trials ~ 1
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
  1
            6
                  70.333
  2
            7
                 153.633 -1 -83.301 < 2.2e-16 ***
library(lmtest)
1rtest(model)
  Likelihood ratio test
  Model 1: Trials ~ Latitude
  Model 2: Trials ~ 1
    #Df LogLik Df Chisq Pr(>Chisq)
    2 -56.293
      1 -97.944 -1 83.301 < 2.2e-16 ***
```

Pseudo r-squared

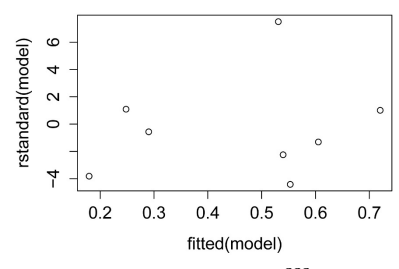
In order to calculate the pseudo r-squared for a logistic model, it's necessary to convert the data to long format, and re-fit the model. Here, I'll re-type the data in a slightly longer format, and then use *tidyr::uncount* to convert the data frame to true long format, with 4444 observations.

```
Data = read.table(header=TRUE, stringsAsFactors=TRUE, text="
                    Latitude Allele Count
 Location
 Port_Townsend,_WA
                              mpi90
                                          47
                    48.1
 Neskowin,_OR
                    45.2
                              mpi90
                                         177
 Siuslaw_R.,_OR
                    44.0
                              mpi90
                                        1087
 Umpqua_R.,_OR
                              mpi90
                                         187
                    43.7
 Coos_Bay,_OR
                    43.5
                              mpi90
                                         397
                    37.8
                              mpi90
 San_Francisco,_CA
                                          40
 Carmel,_CA
                    36.6
                              mpi90
                                          39
 Santa_Barbara,_CA
                   34.3
                              mpi90
                                          30
Port_Townsend,_WA
                    48.1
                              mpi100
                                         139
Neskowin,_OR
                    45.2
                              mpi100
                                         241
Siuslaw_R.,_OR
                    44.0
                              mpi100
                                        1183
Umpqua_R.,_OR
                    43.7
                              mpi100
                                         175
                              mpi100
Coos_Bay,_OR
                    43.5
                                         671
San_Francisco,_CA
                              mpi100
                    37.8
                                          14
San_Francisco,_CA
                    36.6
                              mpi100
                                          17
```

```
34.3 mpi100
Santa_Barbara,_CA
                                          0
")
   str(Data)
   'data.frame':16 obs. of 4 variables:
    $ Location: Factor w/ 8 levels "Carmel,_CA","Coos_Bay,_OR",..: 4 3 7 8 2 5 1 6 4 3
  $ Latitude: num 48.1 45.2 44 43.7 43.5 37.8 36.6 34.3 48.1 45.2 ... $ Allele : Factor w/ 2 levels "mpi100", "mpi90": 2 2 2 2 2 2 2 1 1 ... $ Count : int 47 177 1087 187 397 40 39 30 139 241 ...
library(tidyr)
Long = uncount(Data, Count)
str(Long)
   'data.frame':4444 obs. of 3 variables:
    $ Allele : Factor w/ 2 levels "mpi100", "mpi90": 2 2 2 2 2 2 2 2 2 2 ...
model.2 = glm(Allele ~ Latitude, data = Long,
              family = binomial())
summary(model.2)
  Coefficients:
                Estimate Std. Error z value Pr(>|z|)
   (Intercept) 7.64686 0.92487 8.268 <2e-16 ***
   Latitude -0.17864 0.02104 -8.490 <2e-16 ***
library(car)
Anova(model.2, test="Wald")
  Analysis of Deviance Table (Type II tests)
            Df Chisq Pr(>Chisq)
   Latitude 1 72.076 < 2.2e-16 ***
library(rcompanion)
nagelkerke(model.2)
   $Pseudo.R.squared.for.model.vs.null
   Pseudo.R.squared
  McFadden
                                       0.0136160
  Cox and Snell (ML)
                                       0.0185699
```

```
Nagelkerke (Cragg and Uhler)
                                        0.0248401
   $Likelihood.ratio.test
     Df.diff LogLik.diff Chisq
                                    p.value
           -41.65 83.301 7.0476e-20
library(rcompanion)
efronRSquared(model.2)
   EfronRSquared
          0.0176
library(rcompanion)
countRSquare(model.2)
   $Result
     Count.R2 Count.R2.corrected
       0.567
                          0.0389
   $Confusion.matrix
             Predicted
   Actual
                0
                     1
                       Sum
   0
             2409
                    31 2440
    1
             1895
                   109 2004
             4304
                  140 4444
    Sum
```

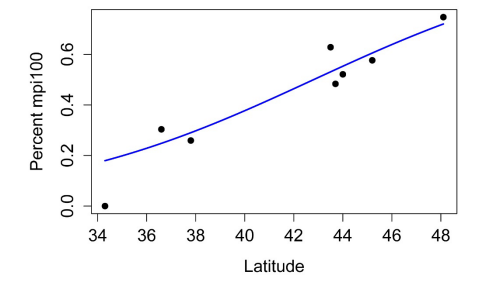
Plot of standardized residuals



A plot of standardized residuals vs. predicted values. The residuals should be unbiased and homoscedastic. For an illustration of these properties, see this diagram by Steve Jost at DePaul University: condor.depaul.edu/sjost/it223/documents/resid-plots.gif.

```
### additional model checking plots with: plot(model)
```

Plotting the model



Logistic regression example

```
### Logistic regression, favorite insect example, p. 248
Data = read.table(header=TRUE, stringsAsFactors=TRUE, text="
Height Insect
62
         beetle
 66
         other
 61
         beetle
 67
         other
 62
         other
 76
         other
         other
 66
```

```
70
         beetle
 67
         other
 66
         other
 70
         other
 70
         other
 77
         beetle
 76
         other
 72
         beetle
 76
         beetle
 72
         other
 70
         other
 65
         other
 63
         other
 63
         other
 70
         other
72
         other
70
         beetle
         other
74
")
```

Model fitting

Coefficients and exponentiated cofficients

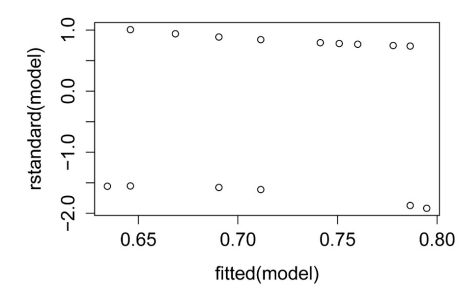
```
summary(model)
  Coefficients:
             Estimate Std. Error z value Pr(>|z|)
  (Intercept) 4.41379 6.66190 0.663 0.508
  Height
             -0.05016
                         0.09577 -0.524
                                          0.600
confint(model)
                  2.5 %
                           97.5 %
  (Intercept) -8.4723648 18.4667731
  Height
             -0.2498133 0.1374819
exp(model$coefficients) # exponentiated coefficients
  (Intercept)
                  Height
   82.5821122
               0.9510757
exp(confint(model))
                       # 95% CI for exponentiated coefficients
                    2.5 %
                               97.5 %
  (Intercept) 0.0002091697 1.047171e+08
           0.7789461738 1.147381e+0
  Height
```

Analysis of variance for individual terms

```
library(car)
  Anova(model, type="II", test="Wald")
     Analysis of Deviance Table (Type II tests)
     Response: Insect
              Df Chisq Pr(>Chisq)
              1 0.2743 0.6004
     Heiaht
     Residuals 23
Overall p-value for model
  anova(model,
        update(model, ~1),  # update here produces null model for comparison
        test="Chisq")
     Analysis of Deviance Table
     Model 1: Insect ~ Height
     Model 2: Insect ~ 1
       Resid. Df Resid. Dev Df Deviance Pr(>Chi)
     1
         23 29.370
     2
          24 29.648 -1 -0.27779 0.5982
  library(lmtest)
  1rtest(model)
     Likelihood ratio test
     Model 1: Insect ~ Height
     Model 2: Insect ~ 1
       #Df LogLik Df Chisq Pr(>Chisq)
     1 2 -14.685
     2 1 -14.824 -1 0.2778 0.5982
Pseudo-R-squared
  library(rcompanion)
```

```
nagelkerke(model)
   $Pseudo.R.squared.for.model.vs.null
                                Pseudo.R.squared
  McFadden
                                      0.00936978
   Cox and Snell (ML)
                                      0.01105020
  Nagelkerke (Cragg and Uhler)
                                      0.01591030
                                     236
```

Plot of standardized residuals



Plotting the model

```
### Convert Insect to a numeric variable, levels 0 and 1
Data$Insect.num=as.numeric(Data$Insect)-1
library(FSA)
headtail(Data)
    Height Insect Insect.num
```

```
1
             62 beetle
                                  0
      2
             66 other
                                  1
      3
             61 beetle
                                  0
      23
             72 other
                                  1
                                  0
      24
             70 beetle
      25
             74 other
                                  1
   plot(Insect.num ~ Height,
        data = Data,
        xlab="Height"
        ylab="Insect",
        pch=19)
   curve(predict(model,data.frame(Height=x),type="response"),
         lty=1, lwd=2, col="blue",
         add=TRUE)
    0.8
Insect
                                  70
                    65
                                                75
```

Convert Insect to a logical variable, levels TRUE and FALSE
Data\$Insect.log=(Data\$Insect=="other")

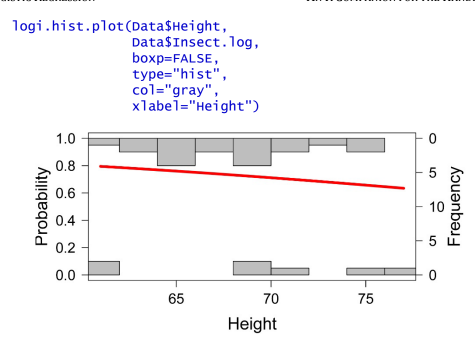
Height

library(FSA)

headtail(Data)

```
Height Insect Insect.num Insect.log
       62 beetle
                           0
1
                                  FALSE
2
       66 other
                           1
                                   TRUE
3
       61 beetle
                          0
                                  FALSE
23
       72 other
                          1
                                   TRUE
24
       70 beetle
                          0
                                  FALSE
25
                          1
       74 other
                                   TRUE
```

library(popbio)



Logistic regression example with significant model and abbreviated code

```
### Logistic regression, hypothetical example
### Abbreviated code and description
Data = read.table(header=TRUE, stringsAsFactors=TRUE, text="
Continuous
             Factor
 62
              Α
 63
              Α
 64
              Α
 65
 66
 67
 68
 69
 70
 71
 72
 73
 74
 75
 72.5
 73.5
              В
 74.5
              В
 75
              В
 76
              В
 77
              В
 78
              В
 79
              В
 80
              В
 81
              В
```

```
82
             В
 83
             В
 84
             В
 85
             В
 86
             В
")
model = glm(Factor ~ Continuous,
            data=Data,
            family = binomial(link="logit"))
summary(model)
   Coefficients:
               Estimate Std. Error z value Pr(>|z|)
   (Intercept) -66.4981 32.3787 -2.054 0.0400 *
   Continuous 0.9027
                          0.4389
                                     2.056
                                             0.0397 *
library(car)
Anova(model, type="II", test="Wald")
   Analysis of Deviance Table (Type II tests)
   Response: Factor
              Df Chisq Pr(>Chisq)
   Continuous 1 4.229 0.03974 *
   Residuals 27
anova(model,
      update(model, ~1),
      test="Chisq")
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
   1
            27
                   12.148
   2
            28
                   40.168 -1
                              -28.02 1.2e-07 ***
library(rcompanion)
nagelkerke(model)
                                Pseudo.R.squared
  McFadden
                                        0.697579
   Cox and Snell (ML)
                                        0.619482
   Nagelkerke (Cragg and Uhler)
                                        0.826303
library(rcompanion)
efronRSquared(model)
```

```
EfronRSquared 0.71
```

library(rcompanion)

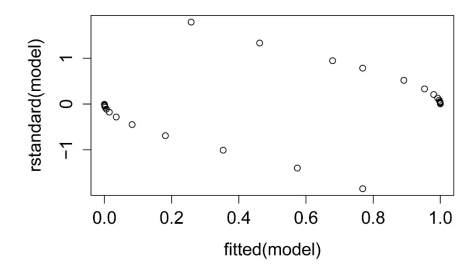
countRSquare(model)

\$Result

```
Count.R2 Count.R2.corrected 0.862 0.714
```

\$Confusion.matrix

Predicted Actual 0 1 Sum 0 12 2 14 1 2 13 15 Sum 14 15 29



Convert Factor to a numeric variable, levels 0 and 1

Data\$Factor.num=as.numeric(Data\$Factor)-1

library(FSA)

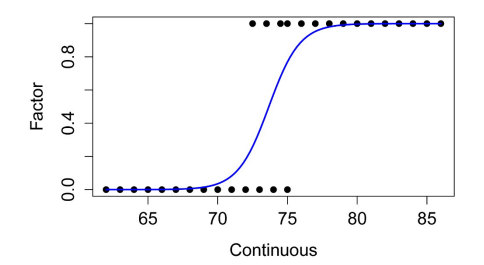
headtail(Data)

	Continuous	Factor	Factor.num
1	62	Α	0
2	63	Α	0
3	64	Α	0

```
27 84 B 1
28 85 B 1
29 86 B 1
```

```
plot(Factor.num ~ Continuous,
    data = Data,
    xlab="Continuous",
    ylab="Factor",
    pch=19)

curve(predict(model,data.frame(Continuous=x),type="response"),
    lty=1, lwd=2, col="blue",
    add=TRUE)
```



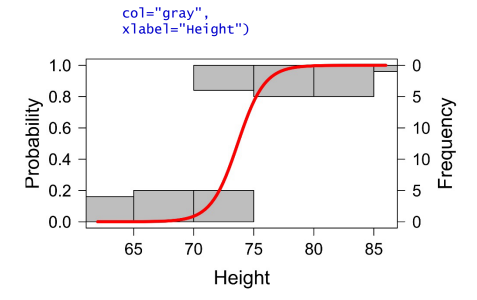
Convert Factor to a logical variable, levels TRUE and FALSE
Data\$Factor.log=(Data\$Factor=="B")

library(FSA)

headtail(Data)

	Continuous	Factor	Factor.num	Factor.log
1	62	Α	0	FALSE
2	63	Α	0	FALSE
3	64	Α	0	FALSE
27	84	В	1	TRUE
28	85	В	1	TRUE
29	86	В	1	TRUE

library(popbio)



Power analysis

See the *Handbook* for information on this topic.

References

IDRE . 2011. FAQ: What are pseudo R-squareds? UCLA. <u>stats.oarc.ucla.edu/other/mult-pkg/faq/general/faq-what-are-pseudo-r-squareds/</u>.

Cross Validated. 2014. McFadden's Pseudo-R2 Interpretation. stats.stackexchange.com/questions/82105/mcfaddens-pseudo-r2-interpretation.

Multiple Logistic Regression

When to use it

The bird example is shown in the "How to do multiple logistic regression" section.

Null hypothesis How it works Selecting variables in multiple logistic regression

See the *Handbook* for information on these topics.

Assumptions

See the *Handbook* and the "How to do multiple logistic regression" section below for information on this topic.

Example Graphing the results Similar tests

See the *Handbook* for information on these topics.

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(dplyr)){install.packages("dplyr")}
if(!require(FSA){install.packages("FSA")}
if(!require(psych)){install.packages("psych")}
if(!require(rcompanion)){install.packages("rcompanion")}
if(!require(car){install.packages("car")}
if(!require(lmtest)){install.packages("lmtest")}
if(!require(PerformanceAnalytics)){install.packages("PerformanceAnalytics")}
```

How to do multiple logistic regression

Don't use stepwise procedures

In general, stepwise procedures are not recommended to determine an appropriate model for multiple regression. It is better to use knowledge of the measured variables, or to use penalized models like ridge regression or lasso regression.

However, a stepwise procedure will be used here to highlight some methods to compare models, and to compare with the example in the *Handbook*.

Stepwise regression in R

Multiple logistic regression can be determined by a stepwise procedure using the *step* function. This function selects models to minimize AIC, not according to p-values as does the SAS example in the *Handbook*. Note, also, that in this example the *step* function found a different model than did the procedure in the *Handbook*.

It is often advised to not blindly follow a stepwise procedure, but to also compare competing models using fit statistics (AIC, AICc, BIC), or to build a model from available variables that are biologically or scientifically sensible.

Multiple correlation

Multiple correlation is one tool for investigating the relationship among potential independent variables. For example, if two independent variables are correlated to one another, likely both won't be needed in a final model, but there may be reasons why you would choose one variable over the other.

```
### ------
### Multiple logistic regression, bird example, p. 254-256
### -----
```

when using read.table, the column headings need to be on the ### same line. If the headings will spill over to the next line, ### be sure to not put an enter or return at the end of the top ### line. The same holds for each line of data.

D	ata = r	read.ta	able(h	neade	r=TRI	JE, s	string	sasf	actors	=TRUE	, te	xt="			
	Species	Status	Length	Mass	Range	Migr	Insect	Diet	Clutch	Broods	Wood	Upland	Water	Release	Indiv
	Cyg_olor	1	1520		1.21	1	12	2	6	1	0	0	1	6	29
	Cyg_atra	1	1250		0.56	1	0	1	6	1	0	0	1	10	85
	Cer_nova	1	870		0.07	1	0	1	4	1	0	0	1	3	8
	Ans_caer	0	720	2517		3	12	2	3.8	1	0	0	1	1	10
	Ans_anse	0	820		3.45	3	0	1	5.9	1	0	0	1	2	7
	Bra_cana	1	770		2.96	2	0	1	5.9	1	0	0	1	10	60
	Bra_sand	0	50		0.01	1	0	1	4	2	0	0	0	1	2
	Alo_aegy	0	680		2.71	1	NA	2	8.5	1	0	0	1	1	8
	Ana_plat	1	570		9.01	2	6	2	12.6	1	0	0	1	17	1539
	Ana_acut	0	580		7.9	3	6	2	8.3	1	0	0	1	3	102
	Ana_pene	0 0	480 470		4.33 1.04	3	0 12	1 2	8.7 13.5	1	0 1	0	1	5 5	32 10
	Aix_spon	0	450		2.17	3	12	2	9.5	1	0	0	1	3	9
	Ayt_feri Ayt_fuli	0	435		4.81	3	12	2	10.1	1	0	0	1	2	5
	Ore_pict	0	275		0.31	1	3	1	9.5	1	1	1	0	9	398
	Lop_cali	1	256		0.24	1	3	1	14.2	2	0	0	0	15	1420
	Col_virg	1	230		0.77	1	3	1	13.7	1	0	0	0	17	1156
	Ale_grae	1	330		2.23	1	3	1	15.5	1	0	1	0	15	362
	Ale_rufa	0	330		0.22	1	3	2	11.2	2	0	0	0	2	20
	Per_perd	0	300		2.4	1	3	1	14.6	1	0	1	0	24	676
	Cot_pect	0	182		0.33	3	NA	2	7.5	1	0	0	0	3	NA
	Cot_aust		180		0.69	2	12	2	11	1	Ö	0	1	11	601
	Lop_nyct	0	800		0.28	1	12	2	5	1	1	1	0	4	6
	Pha_colc	1	710		1.25	1	12	2	11.8	1	1	0	0	27	244
	Syr_reev	0	750	949	0.2	1	12	2	9.5	1	1	1	0	2	9
	Tet_tetr	0	470	900	4.17	1	3	1	7.9	1	1	1	0	2	13
	Lag_lago	0	390	517	7.29	1	0	1	7.5	1	1	1	0	2	4
	Ped_phas	0	440	815	1.83	1	3	1	12.3	1	1	0	0	1	22
	Tym_cupi	0	435	770	0.26	1	4	1	12	1	0	0	0	3	57
	Van_vane	0	300	226	3.93	2	12	3	3.8	1	0	0	0	8	124
	Plu_squa	0	285	318	1.67	3	12	3	4	1	0	0	1	2	3
	Pte_alch	0	350	225	1.21	2	0	1	2.5	2	0	0	0	1	8
	Pha_chal	0	320		0.6	1	12	2	2	2	1	0	0	8	42
	Ocy_loph	0	330	205	0.76	1	0	1	2	7	1	0	1	4	23
	Leu_mela	0	372	NA	0.07	1	12	2	2	1	1	0	0	6	34
	Ath_noct		220		4.84	1	12	3	3.6	1	1	0	0	7	221
	Tyt_alba	0	340		8.9	2	0	3	5.7	2	1	0	0	1	7
	Dac_nova	1	460		0.34	1	12	3	2	1	1	0	0	7	21
	Lul_arbo	0	150		1.78	2	4	2	3.9	2	1	0	0	1	5
	Ala_arve	1	185		5.19	2	12	2	3.7	3	0	0	0	11	391
	Pru_modu	1	145		1.95	2	12 12	2	3.4	2	1	0	0	14 11	245 123
	Eri_rebe	0 0	140		2.31	2	12	2	5 4.7	2	1	0	0	4	
	Lus_mega Tur_meru	1	161 255	82.6	1.88	2	12	2	3.8	3	1	0	0	16	7 596
	Tur_phil	1	230		4.84	2	12	2	4.7	2	1	0	0	12	343
	Syl_comm	0	140		3.39	3	12	2	4.6	2	1	0	0	1	2
	Syl_atri	0	140		2.43	2	5	2	4.6	1	1	0	0	1	5
	Man_mela	0	180	NA	0.04	1	12	3	1.9	5	1	0	0	1	2
	Man_mela	0	265		0.25	1	12	2	2.6	NA	1	0	0	1	80
	Gra_cyan	0	275		0.83	1	12	3	3	2	1	0	1	1	NA
	Gym_tibi	1	400		0.82	1	12	3	4	1	1	0	0	15	448
	Cor_mone	0	335		3.4	2	12	2	4.5	1	1	0	0	2	3
	Cor_fruq	1	400		3.73	1	12	2	3.6	1	1	0	0	10	182
	Stu_vulg		222		3.33	2	6	2	4.8	2	1	0	Ö	14	653
			_				-		-						

```
Acr_tris 1
                    230 111.3 0.56 1
                                          12
                                                                                                    88
                                                        3 7
                                                  2
                                                                                          12
 Pas_dome 1
                    149 28.8 6.5
                                                        3.9
                                                               3
                                                                      1
                                                                           0
                                                                                   0
                                                                                                   416
Pas_mont 0
                                                        4.7
                                                                                                    14
                                                  2
                                                                                   0
                                                                                           3
                   133
                           22 6.8
                                     1
                                           6
                                                               3
                                                                      1
                                                                           0
Aeg_temp 0
                   120 NA 0.17 1
                                           6
                                                  2
                                                        4.7
                                                               3
                                                                      1
                                                                                   0
                                                                                           3
                                                                                                    14
 Emb_gutt 0
                   120
                           19 0.15 1
                                           4
                                                  1
                                                               3
                                                                      0
                                                                           0
                                                                                   0
                                                                                           4
                                                                                                   112
Poe_gutt 0
                   100 12.4 0.75
                                           4
                                                               3
                                                                      0
                                                                           0
                                                                                   0
                                                  1
                                                        4.7
                                                                                           1
                                                                                                    12
Lon_punc 0
Lon_cast 0
Pad_oryz 0
                                           0
                                                                           0
                                                                                   0
                                                                                           1
                    110 13.5 1.06
                                     1
                                                  1
                                                        5
                                                               3
                                                                      0
                                                                                                     8
                    100
                         NA
                              0.13
                                     1
                                           4
                                                  1
                                                        5
                                                               NA
                                                                      0
                                                                           0
                                                                                   1
                                                                                           4
                                                                                                    45
                              0.09
                                           0
                                                                           0
                                                                                   0
                                                                                           2
                   160 NA
                                     1
                                                  1
                                                        5
                                                               NA
                                                                      0
                                                                                                     6
Fri_coel 1
                    160 23.5 2.61
                                          12
                                                        4.9
                                                                           0
                                                                                   0
                                                                                         17
                                                                                                   449
                                                                      1
                                                  2
Fri_mont 0
                    146 21.4 3.09
                                     3
                                          10
                                                        6
                                                               NA
                                                                      1
                                                                           0
                                                                                   0
                                                                                          7
                                                                                                   121
Car_chlo 1
                   147
                         29
                              2.09
                                     2
                                           7
                                                  2
                                                        4.8
                                                               2
                                                                      1
                                                                           0
                                                                                   0
                                                                                          6
                                                                                                    65
Car_spin 0
                    117
                         12
                              2.09
                                           3
                                                  1
                                                        4
                                                                      1
                                                                           0
                                                                                   0
                                                                                          3
                                                                                                    54
Car_card 1
                   120 15.5 2.85
                                     2
                                                                                                   626
                                           4
                                                  1
                                                        4.4
                                                               3
                                                                           0
                                                                                   0
                                                                                         14
                                                                      1
Aca_flam 1
                   115 11.5 5.54
                                     2
                                           6
                                                  1
                                                        5
                                                               2
                                                                      1
                                                                           0
                                                                                   0
                                                                                         10
                                                                                                   607
Aca_flavi 0
                   133 17
                              1.67
                                           0
                                                                      0
                                                                                   0
                                                                                                    61
Aca_cann 0
                   136 18.5 2.52
                                                        4.7
                                                                                         12
                                                                                                   209
                                           6
                                                  1
                                                               2
                                                                           0
                                                                                   0
                                     2
                                                                      1
Pyr_pyrr 0
Emb_citr 1
                         23.5 3.57
                                           4
                                                  1
                                                                                   0
                   142
                                                        4
                                                               3
                                                                      1
                                                                           0
                                                                                          2
                                                                                                  NA
                    160
                                                  2
                                                                                                   656
                         28.2 4.11
                                           8
                                                        3.3
                                                               3
                                                                      1
                                                                           0
                                                                                   0
                                                                                         14
Emb_hort 0
                   163 21.6 2.75
                                                  2
                                                                                   0
                                          12
                                                        5
                                                               1
                                                                      0
                                                                           0
                                                                                          1
                                                                                                     6
Emb_cirl 1
                   160 23.6 0.62
                                          12
                                                                                   0
                                                                                                    29
                                                        3.5
                                                               2
                                                                      1
                                                                           0
                                                                                           3
Emb_scho 0
                   150 20.7 5.42
                                     1
                                                  2
                                                               2
                                                                      0
                                                                                   1
                                                                                          2
                                                                                                     9
                                          12
                                                        5.1
                                                                           0
Pir_rubr 0
Age_phoe 0
Stu_negl 0
                                                                                                     2
                   170 31
                              0.55 3
                                          12
                                                  2
                                                        4
                                                                           0
                                                                                   0
                                                                                          1
                                                               NA
                                                                      1
                    210 36.9 2
                                     2
                                           8
                                                  2
                                                        3.7
                                                               1
                                                                      0
                                                                           0
                                                                                   1
                                                                                           1
                                                                                                     2
                                                                                                     2
                   225 106.5 1.2
                                                        4.8
                                     2
                                          12
                                                               2
                                                                      0
                                                                           0
                                                                                   0
                                                                                          1
")
```

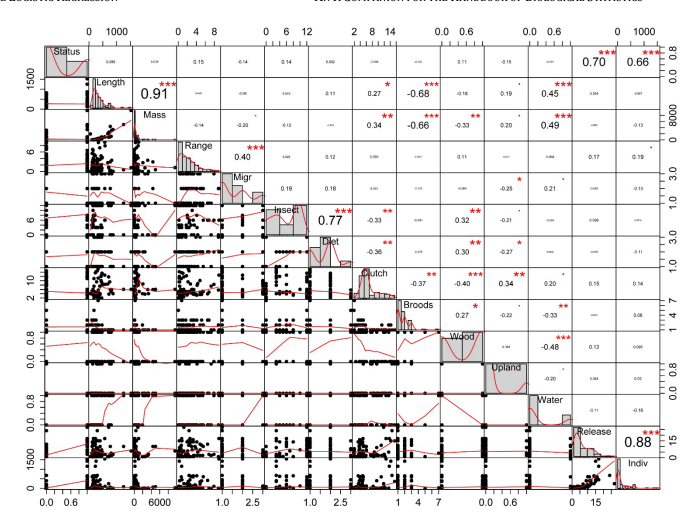
Create a data frame of numeric variables

```
### Select only those variables that are numeric or can be made numeric
library(dplyr)
Data.num =
   select(Data.
          Status,
          Length,
          Mass,
          Range,
          Migr,
          Insect.
          Diet,
          clutch,
          Broods,
          Wood,
          Upland,
          Water,
          Release,
          Indiv)
### Covert integer variables to numeric variables
Data.num$Status = as.numeric(Data.num$Status)
Data.num$Length = as.numeric(Data.num$Length)
Data.num$Migr
                 = as.numeric(Data.num$Migr)
Data.num$Insect = as.numeric(Data.num$Insect)
Data.num$Diet
                 = as.numeric(Data.num$Diet)
```

Data.num\$Broods = as.numeric(Data.num\$Broods)

```
= as.numeric(Data.num$Wood)
Data.num$Wood
Data.num$Upland = as.numeric(Data.num$Upland)
Data.num$water = as.numeric(Data.num$water)
Data.num$Release = as.numeric(Data.num$Release)
Data.num$Indiv
                  = as.numeric(Data.num$Indiv)
### Examine the new data frame
library(FSA)
headtail(Data.num)
                   Mass Range Migr Insect Diet Clutch Broods Wood Upland Water Release Indiv
      Status Length
          1 1520 9600.0 1.21
                                     12
                                               6.0
                                                       1
   2
          1
             1250 5000.0 0.56
                                1
                                      0
                                           1
                                               6.0
                                                       1
                                                            0
                                                                  0
                                                                       1
                                                                              10
                                                                                   85
   3
              870 3360.0 0.07
                                      0
                                           1
                                               4.0
                                                                  0
                                                                       1
                                                                              3
                                                                                    8
          1
                                1
                                                       1
                                                            0
   77
              170
                    31.0 0.55
                                      12
                                           2
                                               4.0
                                                      NA
                                                            1
                                                                  0
                                                                       0
                                                                              1
                                                                                    2
                                           2
   78
          0
              210
                    36.9
                         2.00
                                2
                                      8
                                               3.7
                                                            0
                                                                  0
                                                                       1
                                                                              1
                                                                                    2
                                                       1
              225 106.5 1.20
                                      12
                                               4.8
                                                       2
                                                                              1
                                                                                    2
```

Examining correlations among variables



library(psych)

Multiple logistic regression example

In this example, the data contain missing values. In SAS, missing values are indicated with a period, whereas in R missing values are indicated with NA. SAS will often deal with missing values seamlessly. While this makes things easier for the user, it may not ensure that the user understands what is being done with these missing values. In some cases, R requires that user be explicit with how missing values are handled.

One method to handle missing values in a multiple regression would be to remove all observations from the data set that have any missing values. This is what we will do prior to the stepwise procedure, creating a data frame called *Data.omit*.

However, when we create our final model, we may want to exclude only those observations that have missing values in the variables that are actually included in that final model.

For testing the overall p-value of the final model, plotting the final model, or using the *glm.compare* function, we will create a data frame called *Data.final* with only those observations excluded.

There are some cautions about using the *step* procedure with certain glm fits, though models in the binomial and Poisson families should be okay. See *?stats::step* for more information.

```
Man_mela 0
                           0.04 1
                  180 NA
                                        12
                                               3
2
                                                     1 9
                         59 0.25
                                        12
Man_mela 0
                  265
                                                     2.6
                                                           NA
                                                                  1
                                                                       0
                                                                               0
                                                                                      1
                                                                                               80
                  275
                        128 0.83 1
                                        12
Gra_cyan 0
                                               3
                                                     3
                                                           2
                                                                  1
                                                                       0
                                                                              1
                                                                                      1
                                                                                             NA
                                                                                              448
Gym_tibi 1
                  400
                        380 0.82 1
                                        12
                                               3
                                                                  1
                                                                               0
                                                                                     15
Cor_mone 0
                  335
                        203 3.4
                                        12
                                                     4.5
                                                           1
                                                                  1
                                                                       0
                                                                              0
                                                                                      2
                  400
                       425 3.73
                                        12
                                               2
                                                                       0
                                                                              0
                                                                                     10
Cor_frug 1
                                                     3.6
                                                           1
                                                                                              182
                                                                  1
Stu_vulg 1
Acr_tris 1
                  222 79.8 3.33
                                               2
                                                           2
                                                                              0
                                                                                              653
                                        6
                                                     4.8
                                                                  1
                                                                       0
                                                                                     14
                                               2
                  230 111.3 0.56 1
                                        12
                                                     3.7
                                                           1
                                                                  1
                                                                       0
                                                                              0
                                                                                      5
                                                                                               88
Pas_dome 1
                  149 28.8 6.5
                                               2
                                                     3.9
                                                           3
                                                                              0
                                                                                     12
                                                                                              416
                                  1
                                        6
                                                                  1
                                                                       0
Pas_mont 0
                  133
                         22 6.8
                                         6
                                                     4.7
                                                                              0
                                                                  1
                                                                                      3
                                                                                               14
                                               2
Aeg_temp 0
                  120 NA
                            0.17 1
                                         6
                                                     4.7
                                                           3
                                                                  1
                                                                       0
                                                                              0
                                                                                      3
                                                                                               14
Emb_gutt 0
                  120
                         19 0.15
                                  1
                                         4
                                               1
                                                           3
                                                                  0
                                                                       0
                                                                              0
                                                                                      4
                                                                                              112
                                                     5
Poe_gutt 0
Lon_punc 0
                  100 12.4 0.75
                                         4
                                               1
                                                     4.7
                                                           3
                                                                  0
                                                                       0
                                                                              0
                                                                                      1
                                                                                               12
                  110 13.5 1.06
                                         0
                                  1
                                               1
                                                           3
                                                                  0
                                                                       0
                                                                              0
                                                                                      1
                                                                                                8
                                                     5
Lon_cast 0
                  100 NA
                            0.13 1
                                         4
                                               1
                                                     5
                                                                  0
                                                                       0
                                                                              1
                                                                                               45
                                                           NA
                                                                                      2
Pad_oryz 0
                  160 NA
                            0.09
                                        0
                                                                  0
                                                                       0
                                                                              0
                                                                                                6
                                                           NA
Fri_coel 1
                  160 23.5 2.61
                                               2
                                                     4.9
                                                                              0
                                                                                              449
                                  2
                                        12
                                                                       0
                                                                                     17
                                                           2
                                                                  1
Fri_mont 0
Car_chlo 1
                  146 21.4 3.09
                                        10
                                               2
                                                                              0
                                                     6
                                                           NA
                                                                  1
                                                                       0
                                                                                      7
                                                                                              121
                  147
                                               2
                       29
                            2.09
                                         7
                                                     4.8
                                                           2
                                                                  1
                                                                       0
                                                                              0
                                                                                      6
                                                                                               65
                  117 12
Car_spin 0
                            2.09
                                  3
                                         3
                                                                              0
                                                                                      3
                                                                                               54
                                               1
                                                     4
                                                           2
                                                                  1
                                                                       0
Car_card 1
                  120 15.5 2.85
                                                     4.4
                                                                              0
                                                                                              626
                                               1
                                                                  1
                                                                       0
                                                                                     14
                  115 11.5 5.54
                                  2
                                         6
                                               1
                                                           2
                                                                       0
                                                                              0
                                                                                     10
Aca_flam 1
                                                     5
                                                                  1
                                                                                              607
                  133 17 1.67
136 18.5 2.52
Aca_flavi 0
                                  2
                                         0
                                               1
                                                     5
                                                           3
                                                                  0
                                                                       1
                                                                              0
                                                                                      3
                                                                                               61
Aca_cann 0
                                  2
                                         6
                                               1
                                                     4.7
                                                           2
                                                                  1
                                                                       0
                                                                              0
                                                                                     12
                                                                                              209
Pyr_pyrr 0
                  142 23.5 3.57
                                  1
                                         4
                                               1
                                                     4
                                                           3
                                                                  1
                                                                       0
                                                                              0
                                                                                      2
                                                                                             NA
Emb_citr 1
                  160 28.2 4.11
                                  2
                                         8
                                               2
                                                     3.3
                                                                       0
                                                                              0
                                                                                     14
                                                                                              656
                                                           3
                                                                  1
                                               2
Emb_hort 0
                  163 21.6 2.75
                                       12
                                                                  0
                                                                       0
                                                                              0
                                                                                      1
                  160 23.6 0.62 1
                                       12
                                               2
                                                     3.5
                                                           2
                                                                       0
                                                                              0
                                                                                      3
                                                                                               29
Emb_cirl 1
                                                                  1
Emb_scho 0
Pir_rubr 0
                  150
                       20.7 5.42
                                        12
                                                     5.1
                                                           2
                                                                  0
                                                                       0
                                                                              1
                                                                                      2
                                                                                                9
                  170 31 0.55 3
                                                                                                2
                                               2
                                        12
                                                     4
                                                           NA
                                                                  1
                                                                       0
                                                                              0
                                                                                      1
Age_phoe 0
                  210 36.9 2
                                                                                                2
                                        8
                                               2
                                                     3.7
                                                                  0
                                                                       0
                                                           1
                                                                              1
                                                                                      1
Stu_negl 0
                  225 106.5 1.2
                                       12
                                                                               0
                                                                                      1
                                                     4.8
```

Determining model with step procedure

```
### Create new data frame with all missing values removed (NA's)
Data.omit = na.omit(Data)
### Define full and null models and do step procedure
model.null = glm(Status ~ 1,
                 data=Data.omit,
                 family = binomial())
model.full = glm(Status ~ Length + Mass + Range + Migr + Insect + Diet +
                          Clutch + Broods + Wood + Upland + Water +
                          Release + Indiv,
                 data=Data.omit,
                 family = binomial())
step(model.null,
     scope = list(upper=model.full),
             direction="both",
             test="Chisq",
             data=Data)
   Start: AIC=92.34
```

```
Status ~ 1
         Df Deviance
                     AIC
                             LRT Pr(>Chi)
             56.130 60.130 34.213 4.940e-09 ***
+ Release 1
             60.692 64.692 29.651 5.172e-08 ***
+ Indiv
          1
+ Migr
          1
             85.704 89.704 4.639 0.03125 *
+ Upland
          1
             86.987 90.987 3.356 0.06696 .
             88.231 92.231 2.112
+ Insect
         1
                                  0.14614
             90.343 92.343
<none>
+ Mass
          1
            88.380 92.380 1.963
                                  0.16121
             88.781 92.781 1.562
+ Wood
          1
                                  0.21133
+ Diet
          1
             89.195 93.195 1.148
                                  0.28394
             89.372 93.372 0.972 0.32430
+ Length
          1
             90.104 94.104 0.240 0.62448
+ Water
         1
+ Broods
         1
             90.223 94.223 0.120
                                  0.72898
             90.255 94.255 0.088
+ Range
         1
                                  0.76676
+ Clutch
        1 90.332 94.332 0.012
                                  0.91420
< several more steps >
Step: AIC=42.03
Status ~ Upland + Migr + Mass + Indiv + Insect + Wood
         Df Deviance
                      AIC
                             LRT Pr(>Chi)
             28.031 42.031
<none>
Wood
          1
             30.710 42.710 2.679 0.101686
         1 26.960 42.960 1.071 0.300673
+ Diet
+ Length 1
             27.965 43.965 0.066 0.796641
+ Water
         1
             27.970 43.970 0.062 0.803670
+ Broods 1
            27.983 43.983 0.048 0.825974
+ Clutch 1
             28.005 44.005 0.027 0.870592
             28.009 44.009 0.022 0.881631
+ Release 1
+ Range
          1
             28.031 44.031 0.000 0.999964
Insect
          1 32.369 44.369 4.338 0.037276 *
          1 35.169 47.169 7.137 0.007550 **
- Migr
         1 38.302 50.302 10.270 0.001352 **
- Upland
- Mass
         1 43.402 55.402 15.371 8.833e-05 ***
        1 71.250 83.250 43.219 4.894e-11 ***
- Indiv
```

Final model

```
Indiv     0.0137061     0.0038703     3.541     0.000398 ***
Insect     0.2394720     0.1373456     1.744     0.081234     .
wood     1.8134445     1.3105911     1.384     0.166455
```

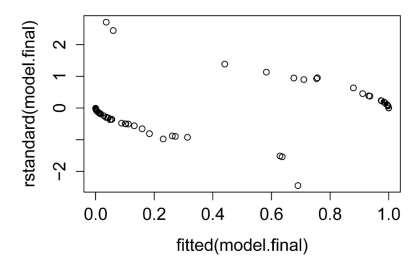
Analysis of variance for individual terms

```
library(car)
  Anova(model.final, type="II", test="Wald")
Pseudo-R-squared
   library(rcompanion)
   nagelkerke(model.final)
      $Pseudo.R.squared.for.model.vs.null
                                   Pseudo.R.squared
     McFadden
                                           0.700475
     Cox and Snell (ML)
                                           0.637732
     Nagelkerke (Cragg and Uhler)
                                          0.833284
   library(rcompanion)
   efronRSquared(model.final)
      EfronRSquared
             0.738
   library(rcompanion)
   countRSquare(model.final)
      $Result
       Count.R2 Count.R2.corrected
      1 0.914
                             0.778
      $Confusion.matrix
           Predicted
      Actual 0 1 Sum
        0 40 3 43
         1
             3 24 27
        Sum 43 27 70
```

Overall p-value for model

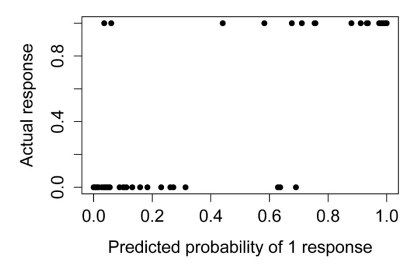
```
### Create data frame with variables in final model and NA's omitted
library(dplyr)
```

```
Data.final =
      select(Data,
             Status.
             Upland,
             Migr,
             Mass,
             Indiv,
             Insect,
             Wood)
  Data.final = na.omit(Data.final)
   ### Define null models and compare to final model
  model.null = glm(Status \sim 1,
                     data=Data.final,
                     family = binomial(link="logit")
   anova(model.final,
         model.null,
         test="Chisq")
     Analysis of Deviance Table
     Model 1: Status ~ Upland + Migr + Mass + Indiv + Insect + Wood
     Model 2: Status ~ 1
       Resid. Df Resid. Dev Df Deviance Pr(>Chi)
      1
               63
                  30.392
      2
               69
                      93.351 -6 -62.959 1.125e-11 ***
  library(lmtest)
   lrtest(model.final)
      Likelihood ratio test
       #Df LogLik Df Chisq Pr(>Chisq)
      1 7 -15.196
      2 1 -46.675 -6 62.959 1.125e-11 ***
Plot of standardized residuals
   plot(fitted(model.final),
        rstandard(model.final))
```



Simple plot of predicted values

```
### Create data frame with variables in final model and NA's omitted
library(dplyr)
Data.final =
   select(Data,
          Status,
          Upland,
          Migr,
          Mass,
          Indiv,
          Insect,
          Wood)
Data.final = na.omit(Data.final)
Data.final$predy = predict(model.final,
                            type="response")
plot(Status ~ predy,
     data = Data.final,
     pch = 16,
     xlab="Predicted probability of 1 response",
     ylab="Actual response")
```



Check for overdispersion

Overdispersion is a situation where the residual deviance of the glm is large relative to the residual degrees of freedom. These values are shown in the *summary* of the model. One guideline is that if the ratio of the residual deviance to the residual degrees of freedom exceeds 1.5, then the model is overdispersed. Overdispersion indicates that the model doesn't fit the data well: the explanatory variables may not well describe the dependent variable or the model may not be specified correctly for these data. If there is overdispersion, one potential solution is to use the quasibinomial *family* option in *glm*.

```
summary(model)

Null deviance: 93.351 on 69 degrees of freedom
Residual deviance: 30.392 on 63 degrees of freedom

summary(model.final)$deviance / summary(model.final)$df.residual

[1] 0.482417
```

Alternative to assess models: using compare.glm

An alternative to, or a supplement to, using a stepwise procedure is comparing competing models with fit statistics. My *compare.glm* function will display AIC, AICc, BIC, and pseudo R-squared for glm models. The models used should all be fit to the same data. That is, caution should be used if different variables in the data set contain missing values. If you don't have any preference on which fit statistic to use, I might recommend AICc, or BIC if you'd rather aim for having fewer terms in the final model.

A series of models can be compared with the standard *anova* function. Models should be nested within the previous model or the next model in the list in the *anova* function; and models should be fit to the same data. When comparing multiple regression models, a p-value to include a new term is often relaxed is 0.10 or 0.15.

In the following example, the models chosen with the stepwise procedure are used. Note that while model 9 minimizes AIC and AICc, model 8 minimizes BIC. The anova results suggest that model 8 is not a significant improvement to model 7. These results give support for selecting any of model 7, 8, or 9. Note that the SAS example in the *Handbook* selected model 4.

```
### Create data frame with just final terms and no NA's
library(dplyr)
Data.final =
   select(Data,
          Status.
          Upland,
          Migr,
          Mass,
          Indiv,
          Insect,
          (boow
Data.final = na.omit(Data.final)
### Define models to compare.
model.1=glm(Status \sim 1,
            data=Data.omit, family=binomial())
model.2=glm(Status ~ Release,
            data=Data.omit, family=binomial())
model.3=glm(Status ~ Release + Upland,
            data=Data.omit, family=binomial())
model.4=glm(Status ~ Release + Upland + Migr,
            data=Data.omit, family=binomial())
model.5=glm(Status ~ Release + Upland + Migr + Mass,
            data=Data.omit, family=binomial())
model.6=glm(Status ~ Release + Upland + Migr + Mass + Indiv,
            data=Data.omit, family=binomial())
model.7=glm(Status ~ Release + Upland + Migr + Mass + Indiv + Insect,
            data=Data.omit, family=binomial())
model.8=glm(Status ~ Upland + Migr + Mass + Indiv + Insect,
            data=Data.omit, family=binomial())
model.9=glm(Status ~ Upland + Migr + Mass + Indiv + Insect + Wood,
            data=Data.omit, family=binomial())
### Use compare.glm to assess fit statistics.
library(rcompanion)
compareGLM(model.1, model.2, model.3, model.4, model.5, model.6,
           model.7, model.8, model.9)
   $Models
     Formula
   1 "Status ~ 1"
   2 "Status ~ Release"
```

```
3 "Status ~ Release + Upland"
   4 "Status ~ Release + Upland + Migr"
   5 "Status ~ Release + Upland + Migr + Mass"
   6 "Status ~ Release + Upland + Migr + Mass + Indiv"
   7 "Status ~ Release + Upland + Migr + Mass + Indiv + Insect"
   8 "Status ~ Upland + Migr + Mass + Indiv + Insect"
   9 "Status ~ Upland + Migr + Mass + Indiv + Insect + Wood"
   $Fit.criteria
     Rank Df.res
                  AIC AICC
                               BIC McFadden Cox.and.Snell Nagelkerke
                                                                        p.value
   1
        1
              66 94.34 94.53 98.75
                                     0.0000
                                                   0.0000
                                                               0.0000
   2
        2
              65 62.13 62.51 68.74
                                     0.3787
                                                   0.3999
                                                               0.5401 2.538e-09
   3
        3
              64 56.02 56.67 64.84
                                     0.4684
                                                              0.6325 3.232e-10
                                                   0.4683
   4
       4
              63 51.63 52.61 62.65
                                     0.5392
                                                   0.5167
                                                              0.6979 7.363e-11
   5
       5
              62 50.64 52.04 63.87
                                                              0.7263 7.672e-11
                                     0.5723
                                                   0.5377
   6
       6
              61 49.07 50.97 64.50
                                    0.6118
                                                   0.5618
                                                              0.7588 5.434e-11
   7
       7
              60 46.42 48.90 64.05
                                     0.6633
                                                   0.5912
                                                              0.7985 2.177e-11
   8
        6
              61 44.71 46.61 60.14
                                                              0.7961 6.885e-12
                                     0.6601
                                                   0.5894
        7
              60 44.03 46.51 61.67
                                     0.6897
                                                   0.6055
                                                              0.8178 7.148e-12
### Use anova to compare each model to the previous one.
anova(model.1, model.2, model.3, model.4, model.5, model.6,
     model.7, model.8, model.9,
      test="Chisq")
  Analysis of Deviance Table
  Model 1: Status ~ 1
   Model 2: Status ~ Release
  Model 3: Status ~ Release + Upland
  Model 4: Status ~ Release + Upland + Migr
  Model 5: Status ~ Release + Upland + Migr + Mass
  Model 6: Status ~ Release + Upland + Migr + Mass + Indiv
  Model 7: Status ~ Release + Upland + Migr + Mass + Indiv + Insect
  Model 8: Status ~ Upland + Migr + Mass + Indiv + Insect
   Model 9: Status ~ Upland + Migr + Mass + Indiv + Insect + Wood
     Resid. Df Resid. Dev Df Deviance Pr(>Chi)
   1
            66
                   90.343
   2
            65
                   56.130 1
                               34.213 4.94e-09 ***
   3
                   48.024 1
                               8.106 0.004412 **
            64
   4
            63
                   41.631 1
                                6.393 0.011458 *
   5
                   38.643 1
                               2.988 0.083872 .
            62
   6
                   35.070 1
                               3.573 0.058721 .
            61
   7
                   30.415 1
                                4.655 0.030970 *
            60
   8
            61
                   30.710 -1
                               -0.295 0.587066
   9
            60
                   28.031 1
                                2.679 0.101686
```

Power analysis

See the *Handbook* for information on this topic.

Multiple Comparisons

The problem with multiple comparisons

See the *Handbook* for information on this topic. Also see sections of this book with the terms "multiple comparisons", "Tukey", "pairwise", "post-hoc", "p.adj", "p.adjust", 'p.method", or "adjust".

Controlling the familywise error rate: Bonferroni correction

Example is shown below in the "How to do the tests" section

Controlling the false discovery rate: Benjamini-Hochberg procedure

Example is shown below in the "How to do the tests" section

Assumption

When not to correct for multiple comparisons

See the *Handbook* for information on these topics.

How to do the tests

R has built in methods to adjust a series of p-values either to control the family-wise error rate or to control the false discovery rate.

The methods Holm, Hochberg, Hommel, and Bonferroni control the family-wise error rate. These methods attempt to limit the probability of even one false discovery (a type I error, incorrectly rejecting the null hypothesis when there is no real effect), and so are all relatively strong (conservative).

The methods BH (Benjamini–Hochberg, which is the same as FDR in R) and BY control the false discovery rate. These methods attempt to control the expected proportion of false discoveries.

For more information on these methods, see *?p.adjust* or other resources.

Note that these methods require only the p-values to adjust and the number of p-values that are being compared. This is different from methods such as Tukey or Dunnett that require also the variability of the underlying data. Tukey and Dunnett are considered familywise error rate methods.

To get some sense of how conservative these different adjustments are, see the two plots below in this chapter.

There is no definitive advice on which p-value adjustment measure to use. In general, you should choose a method which will be familiar to your audience or in your field of study. In addition, there may be some logic which allows you to choose how you balance the probability of making a type I error relative to a type II error. For example, in a preliminary study, you might want to

keep as many significant values as possible to not exclude potentially significant factors from future studies. On the other hand, in a medical study where people's lives are at stake and very expensive treatments are being considered, you would want to have a very high level of certainty before concluding that one treatment is better than another.

Multiple comparisons example with 25 p-values

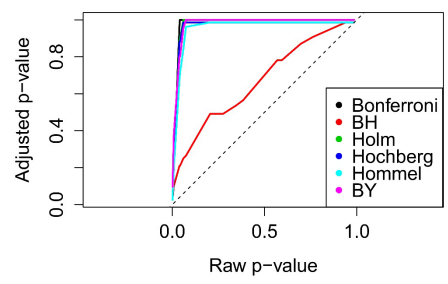
```
### -----
### Multiple comparisons example, p. 262-263
Input = ("
 Food
Blue_fish
                 Raw.p
                 . 34
 Bread .594
Butter .212
Carbohydrates .384
 Cereals_and_pasta .074
 Dairy_products .94
 Eggs
                  .275
                  . 696
 Fats
 Fruit .269
Legumes .341
                  .06
 Nuts .00
Olive_oil .008
Potatoes .569
Processed_meat .986
 Nuts
Proteins
Red meat
                   .042
Red_meat
                   .251
Semi-skimmed_milk .942
Skimmed_milk .222
Sweets
                   .762
Total_calories .001
Total_meat .975
Vegetables .216
White_fish .205
White_meat .041
Whole_milk .039
")
Data = read.table(textConnection(Input), header=TRUE)
### Order data by p-value
Data = Data[order(Data$Raw.p),]
### Check if data is ordered the way we intended
library(FSA)
headtail(Data)
```

```
Food Raw.p
                            Total_calories 0.001
             20
             12
                                                     Olive_oil 0.008
             25
                                                   Whole_milk 0.039
             17 Semi-skimmed_milk 0.942
                                                   Total_meat 0.975
             14
                                    Processed_meat 0.986
 ### Perform p-value adjustments and add to data frame
Data$Bonferroni =
                          p.adjust(Data$Raw.p,
                                                                method = "bonferroni")
 Data$BH =
                          p.adjust(Data$Raw.p,
                                                                method = "BH")
Data$Holm =
                          p.adjust(Data$ Raw.p,
                                                                method = "holm")
 Data$Hochberg =
                           p.adjust(Data$ Raw.p,
                                                                method = "hochberg")
 Data$Hommel =
                          p.adjust(Data$ Raw.p,
                                                                method = "hommel")
 Data$BY =
                          p.adjust(Data$ Raw.p,
                                                                method = "BY")
 Data
           Tood Raw.p Bonferroni BH Holm Hochberg Hommel BY 0 Total_calories 0.001 0.025 0.0250000 0.025 0.025 0.025 0.025 0.09539895 0.001 0.008 0.200 0.1000000 0.192 0.192 0.192 0.38159582 0.001 0.000000 0.897 0.882 0.682 0.80135122 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.001 0.00
                                                                           Food Raw.p Bonferroni
                                                                                                                                                                                        BH Holm Hochberg Hommel
```

```
1.000 0.9071429 1.000
19
             Sweets 0.762
                                                       0.986 0.986 1.00000000
6
     Dairy_products 0.940
                               1.000 0.9860000 1.000
                                                       0.986 0.986 1.00000000
17 Semi-skimmed_milk 0.942
                               1.000 0.9860000 1.000
                                                       0.986 0.986 1.00000000
         Total_meat 0.975
                               1.000 0.9860000 1.000
                                                       0.986 0.986 1.00000000
                                                       0.986 0.986 1.00000000
14
     Processed_meat 0.986
                               1.000 0.9860000 1.000
```

Plot

```
X = Data\$Raw.p
Y = cbind(Data$Bonferroni,
          Data$BH,
          Data$Holm,
          Data$Hochberg,
          Data$Hommel,
          Data$BY)
matplot(X, Y,
        xlab="Raw p-value",
        ylab="Adjusted p-value",
        type="1",
        asp=1,
        col=1:6,
        1ty=1,
        1wd=2)
legend('bottomright',
       legend = c("Bonferroni", "BH", "Holm", "Hochberg", "Hommel", "BY"),
       col = 1:6,
       cex = 1,
       pch = 16)
abline(0, 1,
       col=1,
       1ty=2,
       lwd=1)
```



Plot of adjusted p-values vs. raw p-values for a series of 25 p-values. The dashed line represents a one-to-one line.

Multiple comparisons example with five p-values

```
### Multiple comparisons example, hypothetical example
### ------
Input = ("
Factor
        Raw.p
       .001
Α
      .01
В
C
       .025
      .05
D
       .1
Е
")
Data = read.table(textConnection(Input),header=TRUE)
### Perform p-value adjustments and add to data frame
Data$Bonferroni =
    p.adjust(Data$Raw.p,
            method = "bonferroni")
Data$BH =
     signif(p.adjust(Data$Raw.p,
            method = "BH"),
Data$Holm =
     p.adjust(Data$ Raw.p,
            method = "holm")
Data$Hochberg =
    p.adjust(Data$ Raw.p,
            method = "hochberg")
Data$Hommel =
     p.adjust(Data$ Raw.p,
            method = "hommel")
Data$BY =
     signif(p.adjust(Data$ Raw.p,
            method = "BY"),
          4)
Data
   Factor Raw.p Bonferroni BH Holm Hochberg Hommel
  1 A 0.001 0.005 0.00500 0.005 0.005 0.005 0.01142
```

```
2 B 0.010 0.050 0.02500 0.040 0.040 0.040 0.05708
3 C 0.025 0.125 0.04167 0.075 0.075 0.075 0.09514
4 D 0.050 0.250 0.06250 0.100 0.100 0.100 0.14270
5 E 0.100 0.500 0.10000 0.100 0.100 0.22830
```

Plot

```
X = Data\$Raw.p
Y = cbind(Data$Bonferroni,
         Data$BH,
         Data$Holm,
         Data$Hochberg,
         Data$Hommel,
         Data$BY)
matplot(X, Y,
        xlab="Raw p-value",
        ylab="Adjusted p-value",
        type="1",
        asp=1,
        col=1:6,
        1ty=1,
        1wd=2)
legend('bottomright',
       legend = c("Bonferroni", "BH", "Holm", "Hochberg", "Hommel", "BY"),
       col = 1:6,
       cex = 1,
       pch = 16
abline(0, 1,
        col=1,
        1ty=2,
        lwd=1)
   Adjusted p-value
        0.4
                                               Bonferroni
                                               BH
                                               Holm
                                              Hochberg
                                               Hommel<sup>®</sup>
                                               BY
           -0.4
                     -0.2
                                0.0
                                          0.2
                                                    0.4
                             Raw p-value
```

Plot of adjusted p-values vs. raw p-values for a series of five p-values between 0 and 0.1. Note that Holm and Hochberg have the same values as Hommel, and so are hidden by Hommel. The dashed line represents a one-to-one line.

Miscellany

Chapters Not Covered in this Book

Meta-analysis
Using spreadsheets for statistics
Guide to fairly good graphs
Presenting data in tables
Getting started with SAS
Choosing a statistical test
See the Handbook for information on these topics.

Other Analyses

Post-hoc Contrasts in Models

Contrasts can be used to make specific comparisons of treatments within a model.

One common use is when a factorial design is used, but control or check treatments are used in addition to the factorial design. In the first example below, there are two treatments (D and C) each at two levels (1 and 2), and then there is a *Control* treatment. The approach used here is to analyze the experiment as a one-way analysis of variance, and then use contrasts to test various hypotheses.

Another common use is when there are several treatments that could be thought of as members of a group. In the second example below, there are measurements for six wines, some of which are red (*Merlot, Cabernet, Syrah*) and some of which are white (*Chardonnay, Riesling, Gewürztraminer*). We could compare *Red Wine* as a group to *White Wine* as a group. Or we could compare the treatments *within* the red wine group.

The packages *emmeans* and *multcomp* allow for unlimited tests of single-degree contrasts, with a p-value correction for multiple tests. They also allow for an F-test for multi-line contrasts, for example when testing within groups of multiple treatments. The *aov* function in the native *stats* package has more limited functionality.

See the chapters on *One-way Anova* and *Two-way Anova* for general considerations on conducting analysis of variance.

Packages used in this chapter

The following commands will install these packages if they are not already installed:

```
if(!require(car)){install.packages("car")}
if(!require(emmeans){install.packages("emmeans")}
if(!require(multcomp)){install.packages("multcomp")}
```

Example for single degree-of-freedom contrasts

This hypothetical example could represent an experiment with a factorial design two treatments (D and C) each at two levels (1 and 2), and a control treatment. The 2-by-2 factorial plus control is treated as a one-way anova with five treatments.

```
'D2:C1'
             1.4
 'D2:C1'
             1.6
 'D2:C1'
             1.7
 'D2:C2'
             2.5
 'D2:C2'
             2.6
 'D2:C2'
             2.8
 'Control'
             1.0
 'Control'
             0.9
 'Control'
            0.8
")
### Specify the order of factor levels. Otherwise R will alphabetize them.
Data$Treatment = factor(Data$Treatment,
                          levels=unique(Data$Treatment))
Data
boxplot(Response ~ Treatment,
        data = Data,
        ylab="Response",
        xlab="Treatment")
       2
       Si
   Response
       0
       ď
       5
       1.0
```

D1:C1

D1:C2

Example with emmeans

You need to look at order of factor levels to determine the contrasts

D2:C1

Treatment

D2:C2

Control

levels(Data\$Treatment) "D1:C2" "D2:C1" "D2:C2" [1] "D1:C1" "Control" library(emmeans) marginal = emmeans(model, ~ Treatment) Contrasts = list(D1vsD2)0), = c(1, 1, -1, -1,C1vsC2 = c(1, -1, 1, -1,0), = c(1, -1, -1,0), InteractionDC 1, 0, C1vsC2forD1only = c(1, -1,0. 0), C1vsC2forD2only = c(0, 0, 1, -1,0). TreatsvsControl = c(1, 1, 1, -4), = c(1, 0, 0, 0, -1),T1vsC = c(0, 1,T2vsC 0, 0, -1),T3vsC = c(0, 0, 1, 0, -1),T4vsC = c(0, 0, 0, 1, -1))### The column names match the order of levels of the treatment variable ### The coefficients of each row sum to 0 contrast(marginal, Contrasts, adjust="sidak") contrast estimate SE df t.ratio p.value D1vsD2 -0.83333333 0.1549193 10 -5.379 0.0031 C1vsC2 -2.10000000 0.1549193 10 -13.555 <.0001 InteractionDC 0.03333333 0.1549193 10 0.215 1.0000 ClvsC2forDlonly -1.03333333 0.1095445 10 -9.433 <.0001 ClvsC2forD2only -1.06666667 0.1095445 10 -9.737 <.0001 TreatsvsControl 3.96666667 0.3464102 10 11.451 <.0001 0.26666667 0.1095445 10 T1vsC 2.434 0.3011 T2vsC 1.30000000 0.1095445 10 11.867 <.0001 0.66666667 0.1095445 10 T3vsC 6.086 0.0012 1.73333333 0.1095445 10 15.823 <.0001 T4vsC ### Note that p-values are slightly different than those from multcomp ### due to different adjustment methods. If "none" is chosen as the adjustment method for both procedures. p-values and other statistics will be the same. ### ### With adjust="none", results will be the same as

Example with multcomp

###

the aov method.

You need to look at order of factor levels to determine the contrasts
levels(Data\$Treatment)

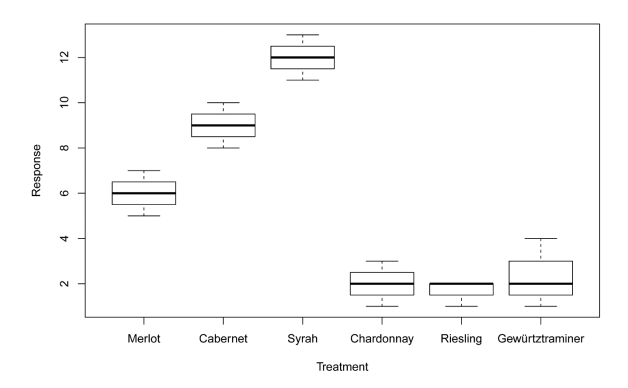
```
[1] "D1:C1"
                 "D1:C2"
                            "D2:C1"
                                      "D2:C2"
                                                 "Control"
Input = ("
                         D1C2 D2C1 D2C2
                   D1C2
                                          Control
Contrast.Name
 D1vsD2
                    1
                          1
                              -1
                                   -1
                                           0
                    1
                         -1
                               1
                                   -1
                                           0
 C1vsC2
                    1
                         -1
                              -1
                                    1
                                           0
 InteractionDC
 C1vsC2forD1only
                    1
                         -1
                               0
                                    0
                                           0
 C1vsC2forD2only
                    0
                          0
                               1
                                   -1
                                          0
 TreatsvsControl
                    1
                          1
                               1
                                    1
                                          -4
                    1
                          0
                               0
                                    0
                                          -1
 T<sub>1</sub>vsC
                    0
                          1
                               0
                                    0
                                          -1
 T2vsC
 T3vsC
                    0
                          0
                               1
                                    0
                                          -1
 T4vsC
                    0
                          0
                               0
                                    1
                                          -1
")
   ### The column names match the order of levels of the treatment variable
   ### The coefficients of each row sum to 0
Matriz = as.matrix(read.table(textConnection(Input),
                    header=TRUE,
                    row.names=1))
Matriz
library(multcomp)
G = glht(model,
         linfct = mcp(Treatment = Matriz))
G$1infct
summary(G,
        test=adjusted("single-step"))
   ### Adjustment options: "none", "single-step", "Shaffer",
                            "Westfall", "free", "holm", "hochberg",
   ###
                            "hommel", "bonferroni", "BH", "BY", "fdr"
   ###
                         Estimate Std. Error t value Pr(>|t|)
   D1vsD2 == 0
                         -0.83333
                                     0.15492
                                              -5.379 0.00218 **
   C1vsC2 == 0
                         -2.10000
                                     0.15492 -13.555
                                                      < 0.001 ***
   InteractionDC == 0
                                     0.15492
                                                0.215 0.99938
                          0.03333
                                              -9.433 < 0.001 ***
   C1vsC2forD1on1v == 0 -1.03333
                                     0.10954
                                                      < 0.001 ***
   C1vsC2forD2only == 0 -1.06667
                                              -9.737
                                     0.10954
   TreatsvsControl == 0 3.96667
                                     0.34641
                                              11.451 < 0.001 ***
   T1vsc == 0
                          0.26667
                                     0.10954
                                               2.434 0.17428
   T2vsC == 0
                                              11.867
                                                      < 0.001 ***
                          1.30000
                                     0.10954
                                                      < 0.001 ***
   T3vsC == 0
                                     0.10954
                                              6.086
                          0.66667
                                     0.10954 15.823 < 0.001 ***
   T4vsC == 0
                          1.73333
```

With test=adjusted("none"), results will be the same as aov method below.

Example for global F-test for a group of treatments

This example has treatments consisting of three red wines and three white wines. We will want to know if there is an effect of the larger wine group (red vs. white) on the response variable, while keeping the individual identities of the wines in the *Treatment* variable. This approach is advantageous because comparisons could still be made within the red wines, for example comparing Merlot to Cabernet.

```
Data = read.table(header=TRUE, stringsAsFactors=TRUE, text="
Treatment
                   Response
 Merlot
                     5
 Merlot
                    6
                    7
 Merlot
                    8
 Cabernet
                    9
 Cabernet
 Cabernet
                    10
                   11
 Syrah
 Syrah
                   12
                   13
 Syrah
 Chardonnay
                    1
                    2
 Chardonnay
 Chardonnay
                    3
 Riesling
                    1
 Riesling
                    2
                    2
 Riesling
 Gewürtztraminer
                    1
                    2
 Gewürtztraminer
 Gewürtztraminer
                    4
")
### Specify the order of factor levels. Otherwise R will alphabetize them.
Data$Treatment = factor(Data$Treatment,
                         levels=unique(Data$Treatment))
Data
boxplot(Response ~ Treatment,
        data = Data,
        ylab="Response",
        xlab="Treatment")
```



You need to look at order of factor levels to determine the contrasts
levels(Data\$Treatment)

Tests of contrasts with emmeans

Ouestion: Is there an effect within red wine?

```
### The column names match the order of levels of the treatment variable
     ### The coefficients of each row sum to 0
  Test = contrast(marginal, Contrasts)
   test(Test, joint=TRUE)
      df1 df2
                F p.value
       2 12 24.3 0.0001
      ### Note that two lines of contrasts resulted in one hypothesis test
            using 2 degrees of freedom. This investigated the effect within
            a group of 3 treatments.
      ###
      ### Results are essentially the same as those from multcomp.
Question: Is there an effect within white wine?
   library(emmeans)
  marginal = emmeans(model, ~ Treatment)
   Contrasts = list(White line1
                                 = c(0, 0, 0, 1, -1, 0),
                    White_line2
                                 = c(0, 0, 0, 0, 1, -1))
      ### The column names match the order of levels of the treatment variable
      ### The coefficients of each row sum to 0
  Test = contrast(marginal, Contrasts)
   test(Test, joint=TRUE)
      df1 df2 F p.value
       2 12 0.3 0.7462
      ### Note that two lines of contrasts resulted in one hypothesis test
            using 2 degrees of freedom. This investigated the effect within
            a group of 3 treatments.
      ### Results are the same as those from multcomp.
Ouestion: Is there a difference between red and white wines? And, mean separation for red wine
   library(emmeans)
  marginal = emmeans(model, ~ Treatment)
   Contrasts = list(Red_vs_white
                                   = c(1, 1, 1, -1, -1, -1),
                                    = c(1, -1, 0, 0, 0, 0),
                    Merlot_vs_Cab
                    Cab_vs_Syrah
                                    = c(0, 1, -1, 0, 0, 0),
```

The column names match the order of levels of the treatment variable

 $Syrah_vs_Merlot = c(-1, 0, 1, 0, 0, 0))$

```
### The coefficients of each row sum to 0
contrast(marginal, Contrasts, adjust="sidak")
                  estimate
                                 SE df t.ratio p.value
   contrast
   Red_vs_white
                         21 1.490712 12 14.087 <.0001
   Merlot_vs_Cab
                         -3 0.860663 12 -3.486 0.0179
                         -3 0.860663 12
   Cab_vs_Syrah
                                        -3.486 0.0179
   Syrah_vs_Merlot
                        6 0.860663 12
                                        6.971 0.0001
     ### Note that p-values are slightly different than those from multcomp
     ### due to different adjustment methods. If "none" is chosen as
     ### the adjustment method for both procedures,
          p-values and other statistics will be the same.
```

Tests of contrasts with multcomp

Question: Is there an effect within red wine?

```
Input = "
          Merlot Cabernet Syrah Chardonnay Riesling Gewürtztraminer
Contrast
 Red_line1 1
                           0
                  -1
Red_line2 0
                  1
                           -1
                                 0
                                            0
                                                     0
   ### Note: there are two lines of contrasts for a group of three treatments
   ### The column names match the order of levels of the treatment variable
   ### The coefficients of each row sum to 0
Matriz = as.matrix(read.table(textConnection(Input),
                   header=TRUE,
                   row.names=1))
Matriz
library(multcomp)
G = glht(model, linfct = mcp(Treatment = Matriz))
G$1infct
summary(G,
        test = Ftest())
   Global Test:
         F DF1 DF2
                         Pr(>F)
   1 24.3 2 12 6.029e-05
```

```
### Note that two lines of contrasts resulted in one hypothesis test
### using 2 degrees of freedom. This investigated the effect within
### a group of 3 treatments.
```

Question: Is there an effect within white wine?

```
Input = "
Contrast
           Merlot Cabernet Syrah Chardonnay
                                             Riesling Gewürtztraminer
White line1 0
                    0
                             0
                                    1
                                               -1
                                                          0
                    0
                             0
                                    0
                                                1
White_line2 0
                                                         -1
   ### Note: there are two lines of contrasts for a group of three treatments
   ### The column names match the order of levels of the treatment variable
   ### The coefficients of each row sum to 0
Matriz = as.matrix(read.table(textConnection(Input),
                   header=TRUE.
                   row.names=1))
Matriz
library(multcomp)
G = glht(model, linfct = mcp(Treatment = Matriz))
G$1infct
summary(G,
        test = Ftest())
   Global Test:
       F DF1 DF2 Pr(>F)
           2 12 0.7462
   ### Note that two lines of contrasts resulted in one hypothesis test
         using 2 degrees of freedom. This investigated the effect within
   ###
   ###
         a group of 3 treatments.
```

Ouestion: Is there a difference between red and white wines? And, mean separation for red wine

```
Input = "
Contrast
                  Merlot Cabernet Syrah
                                           Chardonnay Riesling
                                                                  Gewürtztraminer
Red_vs_white
                   1
                            1
                                      1
                                            -1
                                                         -1
                                                                   -1
                   1
                                      0
                                                          0
                                                                    0
Merlot_vs_Cab
                           -1
                                             0
Cab_vs_Syrah
                   0
                            1
                                     -1
                                             0
                                                          0
                                                                    0
                 -1
                            0
                                      1
                                             0
                                                          0
                                                                    0
Syrah_vs_Merlot
```

names match the order of levels of the treatment variable
The coefficients of each row sum to 0

```
Matriz = as.matrix(read.table(textConnection(Input),
                    header=TRUE,
                    row.names=1))
Matriz
library(multcomp)
G = glht(model,
         linfct = mcp(Treatment = Matriz))
G$1infct
summary(G.
        test=adjusted("single-step"))
   ### Adjustment options: "none", "single-step", "Shaffer".
                            "Westfall", "free", "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr"
   ###
   ###
   Linear Hypotheses:
                         Estimate Std. Error t value Pr(>|t|)
                                                         <0.001 ***
   Red_vs_white == 0
                          21.0000
                                       1.4907 14.087
   Merlot_vs_Cab == 0
                          -3.0000
                                       0.8607 -3.486
                                                         0.0157 *
                                                         0.0156 *
   Cab_vs_svrah == 0
                          -3.0000
                                       0.8607 -3.486
   Syrah_vs_Merlot == 0 6.0000
                                       0.8607
                                                         <0.001 ***
                                                6.971
   (Adjusted p values reported -- single-step method)
   ### With test=adjusted("none"), results will be the same as aov method
   below.
```

Tests of contrasts within aov

Another method to use single-degree-of-freedom contrasts within an anova is to use the *split* option within the *summary* function for an *aov* analysis. The number of degrees of freedom that a factor can be split into for contrast tests is limited.

```
Input =("
Treatment
             Response
 'D1:C1'
             1.0
 'D1:C1'
             1.2
 'D1:C1'
             1.3
 'D1:C2'
             2.1
 'D1:C2'
             2.2
 'D1:C2'
             2.3
 'D2:C1'
             1.4
 'D2:C1'
             1.6
 'D2:C1'
             1.7
 'D2:C2'
             2.5
 'D2:C2'
             2.6
 'D2:C2'
             2.8
```

```
AN R COMPANION FOR THE HANDBOOK OF BIOLOGICAL STATISTICS
 'Control'
            1.0
 'Control'
            0.9
 'Control'
            0.8
Data = read.table(textConnection(Input), header=TRUE)
Data$Treatment = factor(Data$Treatment, levels=unique(Data$Treatment))
   ### Specify the order of factor levels. Otherwise R will alphabetize them.
Data
boxplot(Response ~ Treatment,
        data = Data,
        ylab="Response",
        xlab="Treatment")
    2
    S
Response
    0
    Сi
    5
    1.0
                      D1:C2
            D1:C1
                                          D2:C2
                                D2:C1
                                                    Control
                              Treatment
```

levels(Data\$Treatment)

```
### You need to look at order of factor levels to determine the contrasts
[1] "D1:C1" "D1:C2"
                      "D2:C1" "D2:C2"
                                          "Control"
```

```
### Define contrasts
```

CList = list("D1vsD2" = 1,

```
D1vsD2 =
                 c(1, 1, -1, -1,
                 c(1, -1, 1, -1,
C1vsC2 =
                                  0)
                 c(1, -1, -1, 1,
InteractionDC =
TreatsvsControl = c(1, 1, 1, -4)
Matriz = cbind(D1vsD2, C1vsC2,
              InteractionDC, TreatsvsControl)
contrasts(Data$Treatment) = Matriz
```

```
"c1vsc2" = 2,
            "InteractionDC" = 3,
             "TreatsvsControl" = 4)
### Define model and display summary
model = aov(Response ~ Treatment, data = Data)
summary(model,
        split=list(Treatment=CList))
                               Df Sum Sq Mean Sq F value
                                                           Pr(>F)
                                           1.547 85.963 1.06e-07 ***
  Treatment
                                4 6.189
                                           0.521 28.935 0.00031 ***
    Treatment: D1vsD2
                                1 0.521
                                           3.307 183.750 9.21e-08 ***
    Treatment: C1vsC2
                                1 3.307
    Treatment: InteractionDC
                                1 0.001
                                           0.001
                                                   0.046 0.83396
     Treatment: TreatsvsControl 1 2.360
                                           2.360 131.120 4.53e-07 ***
   Residuals
                               10 0.180
                                           0.018
```

Cate-Nelson Analysis

Cate–Nelson analysis is used to divide bivariate data into two groups: one where a change in the *x* variable is likely to correspond to a change in the *y* variable, and the other group where a change in *x* is unlikely to correspond to a change *y*. Traditionally this method was used for soil test calibration in agronomy studies. For example, to determine if a certain level of soil test phosphorus would indicate that adding phosphorus to the soil would likely cause an increase in crop yield or not.

The method can be used for any case in which bivariate data can be separated into two groups, one with a large *x* variable is associated with a large *y*, and a small *x* associated with a small *y*. Or vice-versa.

For a fuller description of Cate–Nelson analysis and examples in soil-test and other applications, see <u>Mangiafico</u> (2013) and the references there.

Custom function to develop Cate-Nelson models

My *cateNelson* function follows the method of <u>Cate and Nelson</u> (1971). A critical *x* value is determined by iteratively breaking the data into two groups and comparing the explained sum of squares of the iterations. A critical *y* value is determined by using an iterative process which minimizes the number of data point which fall into Quadrant I and III for data with a positive trend.

Options in the *cateNelson* function:

- *plotit=TRUE* (the default) produces a plot of the data, a plot of the sum of squares of the iterations, a plot of the data points in error quadrants, and a final plot with critical x and critical y drawn as lines on the plot.
- *hollow=TRUE* (the default) for the final plot, points in the error quadrants as open circles
- *trend="negative"* (not the default) needs to be used if the trend of the data is negative.
- *xthreshold* and *ythreshold* determine how many options the function will return for critical x and critical y. A value of 1 would return all possibilities. A value of 0.10 returns values in the top 10% of the range of maximum sum of squares.
- *clx* and *cly* determine which of the listed critical x and critical y the function should use to build the final model. A value of 1 selects the first displayed value, and a value of 2 selects the second. This is useful when you have more than one critical x that maximizes or nearly maximizes the sum of squares, or if you want to force the critical y value to be close to some value such as 90% of maximum yield. Note that changing the clx value will also change the list of critical y values that is displayed. In the second example I set *clx=2* to select a critical x that more evenly divides the errors across the quadrants.

Example of Cate-Nelson analysis

```
_____
## Cate-Nelson analysis
## Data from Mangiafico, S.S., Newman, J.P., Mochizuki, M.J.,
    & Zurawski, D. (2008). Adoption of sustainable practices
    to protect and conserve water resources in container nurseries
##
    with greenhouse facilities. Acta horticulturae 797, 367-372.
##-----
size = c(68.55, 6.45, 6.98, 1.05, 4.44, 0.46, 4.02, 1.21, 4.03,
         6.05,48.39,9.88,3.63,38.31,22.98,5.24,2.82,1.61,
         76.61, 4.64, 0.28, 0.37, 0.81, 1.41, 0.81, 2.02, 20.16,
         4.04,8.47,8.06,20.97,11.69,16.13,6.85,4.84,80.65,1.61,0.10)
proportion = c(0.850, 0.729, 0.737, 0.752, 0.639, 0.579, 0.594, 0.534,
               0.541,0.759,0.677,0.820,0.534,0.684,0.504,0.662,
               0.624, 0.647, 0.609, 0.647, 0.632, 0.632, 0.459, 0.684,
               0.361, 0.556, 0.850, 0.729, 0.729, 0.669, 0.880, 0.774,
               0.729, 0.774, 0.662, 0.737, 0.586, 0.316
library(rcompanion)
cateNelson(x = size,
          y = proportion,
          plotit=TRUE,
          hollow=TRUE,
          xlab="Nursery size in hectares",
          ylab="Proportion of good practices adopted",
          trend="positive",
          c1x=1,
```

```
cly=1,
xthreshold=0.10,
ythreshold=0.15)
```

Critical x that maximize sum of squares:

```
Critical.x.value Sum.of.squares
1 4.035 0.2254775
2 4.740 0.2046979
```

Critical y that minimize errors:

```
Critical.y.value Q.i Q.ii Q.iii Q.iv Q.model Q.err Cramer.V
1
                                   2
             0.6355
                       3
                           20
                                       13
                                                33
                                                        5
                                                            0.7289
2
             0.6430
                       3
                           19
                                   3
                                       13
                                                32
                                                        6
                                                            0.6761
3
             0.6470
                       3
                           19
                                       13
                                                32
                                                        6
                                                            0.6761
4
             0.6545
                           18
                                       14
                                                32
                                   4
                                                        6
                                                            0.6854
5
             0.6620
                       2
                           18
                                   4
                                       14
                                                32
                                                        6
                                                            0.6854
6
                                       10
             0.6015
                           21
                                                31
                                                        7
                                                            0.6309
7
                           20
                                   2
                                                31
             0.6280
                       5
                                       11
                                                        7
                                                            0.6209
8
             0.6320
                       5
                           20
                                       11
                                                31
                                                            0.6209
```

```
n = Number of observations
CLX = Critical value of x
```

SS = Sum of squares for that critical value of x

CLy = Critical value of y

Q = Number of observations which fall into quadrants I, II, III, IV Q.Model = Total observations which fall into the quadrants predicted by the model

p.Model = Percent observations which fall into the quadrants predicted by the model

Q.Error = Observations which do not fall into the quadrants predicted by the model p.Error = Percent observations which do not fall into the quadrants predicted by the model

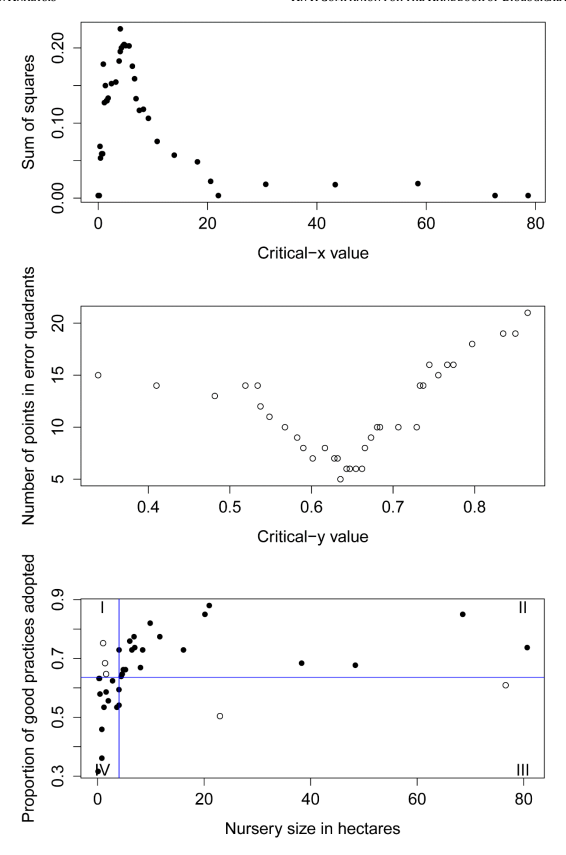
Fisher.p = p-value from Fisher exact test dividing data into these quadrants

Cramer.V = Cramer's V statistic from dividing data into these quadrants

Final model:

```
n CLx SS CLy Q.I Q.II Q.III Q.IV Q.Model p.Model Q.Error 1 38 4.035 0.2254775 0.6355 3 20 2 13 33 0.8684211 5
```

```
p.Error Fisher.p.value Cramer.V
0.1315789   8.532968e-06   0.7289
```



Plots showing the results of Cate—Nelson analysis. In the final plot, the critical x value is indicated with a vertical blue line, and the critical y value is indicated with a

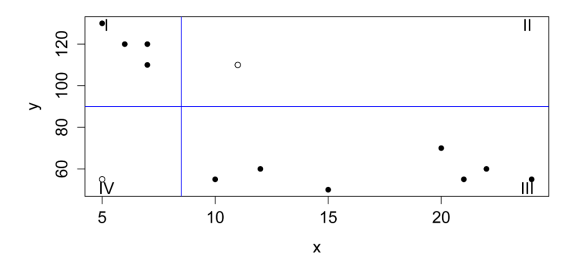
horizontal blue line. Points agreeing with the model are solid, while hollow points indicate data not agreeing with model. (Data from Mangiafico, S.S., Newman, J.P., Mochizuki, M.J., & Zurawski, D. (2008). Adoption of sustainable practices to protect and conserve water resources in container nurseries with greenhouse facilities. Acta horticulturae 797, 367–372.)

Example of Cate-Nelson analysis with negative trend data

```
## Cate-Nelson analysis
## Hypothetical data
Input =("
  Χ
  5
        55
  7
       110
  6
       120
  5
       130
  7
       120
 10
        55
 12
        60
 11
       110
 15
        50
 21
        55
 22
        60
 20
        70
 24
        55
")
Data = read.table(textConnection(Input),header=TRUE)
library(rcompanion)
cateNelson(x = Data$x,
           y = Data y,
           plotit=TRUE,
           hollow=TRUE,
           xlab="x",
           ylab="y"
           trend="negative",
                       # Normally leave as 1 unless you wish to
           c1x=2.
           cly=1,
                       # select a specific critical x value
           xthreshold=0.10,
           ythreshold=0.15)
  Critical x that maximize sum of squares:
     Critical.x.value Sum.of.squares
   1
                 11.5
                             5608.974
                  8.5
                             5590.433
   2
```

Critical y that minimize errors:

```
Critical.y.value O.i O.ii O.iii O.iv O.model O.err Cramer.V
1
                  90
                             1
                                    7
                                         1
                                                 11
                                                         2
                                                              0.6750
2
                             1
                                    7
                                         1
                                                 11
                                                         2
                 110
                       4
                                                              0.6750
3
                       3
                             0
                                         2
                 115
                                    8
                                                 11
                                                         2
                                                              0.6928
                                         2
4
                 120
                       3
                                                 11
                             0
                                    8
                                                              0.6928
          = Number of observations
          = Critical value of x
CLX
          = Sum of squares for that critical value of x
SS
          = Critical value of y
CLy
          = Number of observations which fall into quadrants I, II, III, IV
Q.Model
         = Total observations which fall into the quadrants predicted by the model
p.Model
        = Percent observations which fall into the quadrants predicted by the model
Q.Error = Observations which do not fall into the quadrants predicted by the model
p.Error = Percent observations which do not fall into the quadrants predicted by the model
Fisher.p = p-value from Fisher exact test dividing data into these quadrants
Cramer.V = Cramer's V statistic from dividing data into these quadrants
Final model:
                 SS CLy Q.I Q.II Q.III Q.IV Q.Model
                                                          p.Model Q.Error
   n CLX
1 13 8.5 5608.974 90
                                1
                                             1
                                                    11 0.8461538
                          4
                                       7
  p.Error Fisher.p.value Cramer.V
0.1538462
               0.03185703
                               0.675
```



Plot showing the final result of Cate—Nelson analysis, for data with a negative trend.

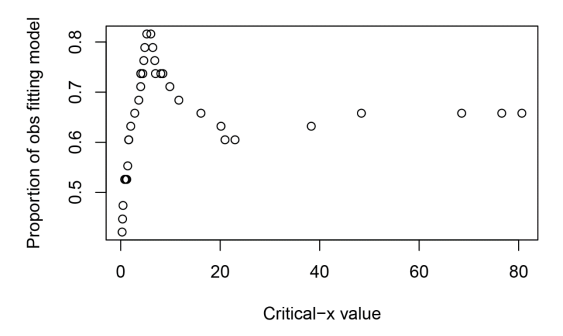
Example of Cate-Nelson analysis with a fixed critical y value

Often when using a Cate–Nelson analysis, we wish to set the critical *y* at some pre-determined value, and then find the critical *x* value that best divides the data. This is common, for example, in agromony studies where we might want to set the critical value at e.g. 90% of maximum potential yield for a crop.

The following example revisits the sustainable nursery practices data above but sets the critical y value at 0.70 (or, 70%).

Here, the first two critical *x* values in the results (5.24 and 6.05), both result in maximizing the count of observations fitting the model. It's possible to sort the resultant data frame by other statistics, like the Pearson *chi-square* value or the effect size statistics *phi*.

```
## Data from Mangiafico, S.S., Newman, J.P., Mochizuki, M.J.,
     & Zurawski, D. (2008). Adoption of sustainable practices
     to protect and conserve water resources in container nurseries
     with greenhouse facilities. Acta horticulturae 797, 367-372.
size = c(68.55, 6.45, 6.98, 1.05, 4.44, 0.46, 4.02, 1.21, 4.03,
          6.05,48.39,9.88,3.63,38.31,22.98,5.24,2.82,1.61,
          76.61,4.64,0.28,0.37,0.81,1.41,0.81,2.02,20.16,
          4.04,8.47,8.06,20.97,11.69,16.13,6.85,4.84,80.65,1.61,0.10)
proportion = c(0.850, 0.729, 0.737, 0.752, 0.639, 0.579, 0.594, 0.534,
                0.541, 0.759, 0.677, 0.820, 0.534, 0.684, 0.504, 0.662,
                0.624, 0.647, 0.609, 0.647, 0.632, 0.632, 0.459, 0.684,
                0.361, 0.556, 0.850, 0.729, 0.729, 0.669, 0.880, 0.774,
                0.729, 0.774, 0.662, 0.737, 0.586, 0.316
library(rcompanion)
cateNelsonFixedY (x
                             = size.
                             = proportion,
                   cly
                             = 0.70,
                   plotit = TRUE,
                   hollow
                             = TRUE,
                             = "Nursery size in hectares",
                   xlab
                             = "Proportion of good practices adopted",
                   ylab
                             = "positive",
                   trend
                   clx
                             = 1,
                   outlength = 5,
                   sortstat = "error")
      Critx Crity Q1 Q2 Q3 Q4 Model Error N
                                            pQ1
                                                  pQ2
                                                        pQ3
                                                                   pModel
                                                                          pError
                                      7 38 0.053 0.316 0.132 0.500
   1
      5.24
             0.7 2 12
                       5 19
                               31
                                                                   0.816
                                                                           0.184
                                      7 38 0.053 0.316 0.132 0.500
      6.05
             0.7
                 2 12
                       5 19
                               31
                                                                    0.816
                                                                           0.184
                 2 12 6 18
   3
      4.84
             0.7
                               30
                                      8 38 0.053 0.316 0.158 0.474
                                                                    0.789
                                                                           0.211
      6.45
             0.7 3 11 5 19
                               30
                                      8 38 0.079 0.289 0.132 0.500
                                                                    0.789
                                                                           0.211
             0.7 2 12 7 17
      4.64
                               29
                                      9 38 0.053 0.316 0.184 0.447
                                                                    0.763
                                                                           0.237
       Fisher.p Pearson.chisq
                                 Pearson.p
                                                phi
                                             -0.629
      0.0001517
                       12.5500
                                 0.0003972
      0.0001517
                       12.5500
                                 0.0003972
                                             -0.62
      0.0005311
                       10.7500
                                 0.0010420
                                             -0.587
      0.0007735
                       9.8400
                                 0.0017080
                                             -0.564
      0.0018910
                       9.1610
                                 0.0024730
                                            -0.546
```





Plots showing the final result of Cate-Nelson analysis, for an alysis with a fixed critical y value.

References

Mangiafico, S.S. 2013. Cate-Nelson Analysis for Bivariate Data Using R-project. *Journal of Extension* 51:5, 5TOT1. tigerprints.clemson.edu/cgi/viewcontent.cgi?article=2547&context=joe.

Cate, R. B., & Nelson, L.A. (1971). A simple statistical procedure for partitioning soil test correlation data into two classes. *Soil Science Society of America Proceedings* 35, 658–660.

Additional Helpful Tips

Reading SAS Datalines in R

Reading SAS datalines with *DescTools*

The *ParseSASDatalines* function in the *DescTools* package will read in data with simple SAS DATALINES code. More complex INPUT schemes may not work.

```
### Reading SAS datalines, DescTools::ParseSASDatalines example
Input = ("
DATA survey;
INPUT id sex $ age inc r1 r2 r3 @@;
DATALINES:
  F 35 17 7 2 2 17 M 50 14 5 5 3 33 F 45 6 7 2 7
49 M 24 14 7 5 7 65 F 52 9 4 7 7
                                    81 M 44 11 7 7 7
   F 34 17 6 5 3 18 M 40 14 7 5 2 34 F 47 6 6 5 6
50 M 35 17 5 7 5
;
")
library(DescTools)
Data = ParseSASDatalines(Input)
### You can omit the DATA statement, the @@, and the final semi-colon.
    The $ is required for factor variables.
Data
     id sex age inc r1 r2 r3
    1
        F 35 17 7
  2 17
         M
           50 14 5
  3 33
        F 45
               6 7
              14 7 5
  4 49
        м 24
  5 65
        F 52
  6 81 M 44 11 7 7
        F 34 17 6 5
  8 18
        M 40 14 7 5
  9 34
        F 47
               6 6 5
  10 50 M 35 17 5 7
```