# Updates and Corrections (as of April 19, 2018) <br> Data Analysis and Graphics Using R - An Example-Based Approach, $3^{\text {rd }}$ edn, 2013 reprint 

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Webpage: http://www.maths.anu.edu.au/~johnm/r-book/r-book.html.
Changes are of four types:

- Changes made necessary, or desirable, arising from changes in R functions;
- Corrections to errors;
- Code changes, designed to improve coding style;
- Removal of some obscurities.


## Note: Prospects for a 4th edition

An initial draft has been in place since late 2017. Cambridge University Press seems, at the moment, not to be interested in proceeding. New material will be posted on the web shortly.

## Graphs that need to be replaced

Figure 2.14, page 64, should be replaced if the figure now given by the existing code is preferred.

Figure 7.10, page 237 (the function used for the smooth is from a package that has been archived; hence use a different package)

Figure 12.2, page 381 (the graph should be printed with aspect="iso")
Figure 12.9, page 403 (the maximum number of features selected should be 23 , as in the code that is in the text; the attempt to select more than 23 generates a collinearity warning.)

## Chapter 1

page 27, Subsection 1.5.2

Replace the first paragraph, i.e.
The default palette, which can be changed, has eight colors including " white". by
The default color palette, shown in Plate 10, attaches the numbers $1, \ldots, 8$ to the eight colors that are included.

## page 27, bullet points in lines 3 to 18 [addition]

Insert as the second to last bullet point:

- Aspect ratio: asp=1 gives a plot with isometric scaling, i.e., $x$ and $y$ units occupy the same physical distances in both directions.


## page 32, following line -9 [additions]

Add, following the first bullet point, the further bullet points:

- Most common types of parameter settings can be handled by using simpleTheme() to create a theme, which the argument par. settings then supplies to the function that creates the graph. See the code for Figure 15.3 in Subsection 15.5.1.
- Use the argument aspect to control aspect. If a number, this specifies the ratio of graph height to width. For isometric scales, specify aspect="iso".


## page 33, final 2 lines

Edit the final sentence to read: Use either .RData or .rda as the extension for such files.

Omit footnote 9.
page 39 , line 2

Replace there is extreme value of area that by One extreme value for area

## Chapter 2

page 45, line 1 [better coding style]
Replace attach(fossum)
by ftotlngth <- fossum[, "totlngth"]
page 45 , lines $2,4,-17,-15,-12$ [better coding style]
Replace totlngth by ftotlngth

## page 45 , line-10, [better coding style]

```
Replace par(mfrow=c(1,1)); detach(fossum)
by par(mfrow=c(1,1))
```


## page 47 , line 7

Replace horiz by horizontal

## page 51, line 9

Figure 2.7B would be better plotted with asp=1, i.e., line 9 becomes:
plot(log(brain) ~ log(body), data=Animals, asp=1) \# Panel B
This ensures that equal changes on a logarithmic scale on the two axes (corresponding to equal relative changes) are the same physical distance apart. (The tick marks in the graph shown are a factor of 10 apart, on both axes.)

## page 64 , fnote 17 , line 3

Edit line 3 to read:
\#\# Panel function calls panel.dotplot(), then plots means

## page 64, fnote 17, lines 6-7

Replace

```
panel.average(x, y, type="p", col="black",
    pch=3, cex=1.25)},
```

by

```
av <- sapply(split(x,y),mean)
ypos <- unique(y)
lpoints(ypos~av, pch=3, cex=1.25, col="black")},
```

[The code in the text still works, but uses a line to join up the means. There is in this instance, where there is an interest in comparing the pattern across the three panels, in leaving the code as is and replacing the graph.]
[There are equivalent replacements on page 123 (footnote 13) and page 261 (lines -4 and -3).]

## page 75 , exercise 5 [better coding style]

Replace the first line of code (attach(cuckoohosts)) by
with(cuckoohosts, \{
Indent the next 5 lines by 5 spaces
Replace the final line of code (detach(cuckoohosts)) by
\})

## Chapter 3

page 105, line - 11
Replace Figure 4.1B by Figure 4.1

## Chapter 4

page 123, footnote 13 , lines 6-7
Replace

```
panel.average(x, y, type="p", col="black",
    pch=3, cex=1.25)},
```

by

```
av <- sapply(split(x,y),mean)
ypos <- factor(levels(y), levels=levels(y))
lpoints(ypos~av, pch=3, cex=1.25, col="black")},
```

[The code in the text still works, but uses a line to join up the means.]

## page 131, lines -15 to -12

Replace

```
median.fun <- function(data, indices){median(data[indices])}
## Call boot(), with statistic=median.fun, R = # of resamples
set.seed(23)
(wren.boot <- boot(data = wren, statistic = median.fun, R = 999))
```

by
\#\# Call the function boot.ci() , with boot.out=wren.boot
boot.ci(boot.out=wren.boot, type=c("perc"))

## page 140, Exercise 14, final 2 lines

Replace

```
# See help(bootci.object). The 4th and 5th elements of
# the percent list element hold the interval endpoints.
```

by

```
# The 4th and 5th elements of the list element percent
# hold the interval endpoints. See ?boot.ci
```


## Chapter 6

page 177 , line -5

Replace The models nihills.1mnihillsG.lm by The models nihills.lm and nihillsG.lm.

## page 186, lines of code preceding subsection 6.3.2

Replace lines 2 and 3 of the code with influencePlot(allbacks.lm)
[influencePlots() is not a direct replacement for leverage.plots(). It gives output that, while different from what is discussed in the text, is at the same more informative.]
page 203 , line prior to 6.3 Errors in $x$
Replace remedy by ) recourse
page 216 , exercise $14,1^{\text {st }}$ line of code
Replace \} by )

## Chapter 7

page 236 , Section 7.5.3, line 7
Replace All by Most

## page 237, final 13 lines

Replace with:
Monotone polynomials, as fitted using the function monpol () from the MonoPoly package, will often be satisfactory, as in Figure 7.10.

Code is:

```
library(MonoPoly)
u <- monpol(ohms~juice, data=fruitohms, degree=3)
plot(ohms ~ juice, data=fruitohms, xlab="Apparent juice content (%)",
    ylab="Resistance (ohms)", col="gray40")
ord <- with(fruitohms, order(juice))
lines(fitted(u)[ord] ~ juice[ord], data=fruitohms, col=2)
```

For fitting a monotonic spline curve, see help(mono.con, package="mgcv")
[Also, replace Figure 7.10 with a figure that has been generated using the above code. The monoProc package has been archived.]

## page 238, line - 2

Replace 1m by gam

## Chapter 8

Replace $15 / 60=0.25$ by $45 / 60=0.75$

## page 249 , footnote 4 , line 2

Add, at the end of the line asp=1, i.e., the line becomes

```
plot(northing ~ easting, data=frogs, pch=c(1,16)[frogs$pres.abs+1], asp=1,
```

page 260 , line - 17

Replace is ("suggesting that is should be") by it.
page 261, line -12
Replace The Number is the total number by Here, Number is the number

## page 261 , line -5 and -4

Replace

```
panel.average(x, y, pch=3, cex=1.25, type="p", col="gray45")
},
```

by

```
av <- sapply(split(x,y),mean)
ypos <- factor(levels(y), levels=levels(y))
lpoints(ypos~av, pch=3, cex=1.25, col="gray45")},
```

[The code in the text still works, but uses a line to join up the means.]
pages 262-266

Insert replacement pages, as supplied, in place of these.
page 282, exercise $7 \mathrm{c}, 3^{\text {rd }}$ line of code
Replace newdat by ndf

## page 285, Figures 9.2B \& 9.2C

Figure 1 does a better job than Figures 9.2B \& 9.2C.

## page 287

Replace the final 4 lines of Subsection 9.1.3 (While the plot . . . the needed additional flexibility.) by
Is an $A R(2)$ model adequate, or is something more needed? We now move to investigating use of an autoregressive moving average (ARMA) model, which adds one or more moving average terms, to examine whether this makes any worthwhile improvement.

## Chapter 9

page 291, line 1
Replace $m f r o w=c(3,2)$ by mfrow=c $(2,2)$

B: Autocorelation -- Data vs AR1 process


C: Partial autocorrelation -- Data vs AR1 process


Figure 1: Panel B compares the estimated autocorrelations with the autocorrelations for an $\operatorname{AR}(1)$ process, and for an $\operatorname{AR}(2)$ process. Panel C repeats the comparison, now for the partial autocorrelations. The dashed horizontal lines are approximate pointwise $5 \%$ critical values for the autocorrelations for a pure noise process, i.e., for independent normal data with mean 0 . Panel C shows the theoretical autocorrelations for an $\operatorname{AR}(1)$ process with $\alpha=0.8$. For an $\operatorname{AR}(1)$ process, partial autocorrelations after the first are all 0 - hence the single lightblue spike at lag 1 . For an $A R(2)$ process, partial autocorrelations beyond lag 2 are all 0 . - a positive spike at lag 1 is followed by a negative spike at lag 2 , with no further spikes.

## Chapter 10

The pdf for a rewrite of this chapter is available as a separate file.

## General

This chapter is strongly affected by changes in the lme4 package. The most important is that msmcsamp () is, because it did not work reliably, no longer available. The primary function for calculating confidence intervals is confint(), using method="profile") or method="boot" in preference to the less reliable method="Wald".

The following demonstrate the form of command that one might now expect to use, with objects returned by lmer():
library (lme4)
library (DAAG)
science1.lmer <- lmer(like ~ sex + PrivPub + (1 | school:class), data $=$ science, na.action=na.exclude)
\# science1.lmer is then a merMod object
\#\# Output from print()
print(science1.lmer, ranef.comp="Variance", digits=3)
\#\# Output from use of summary() with a merMod object
print(summary(science1.lmer), ranef.comp="Variance", digits=3)

```
## Variance components informatioc
print(VarCorr(science1.lmer), comp="Variance", digits=3)
```

See help(merMod) and help(VarCorr) for further details.

## More important Ch 10 changes, with explanatory comment

## page 310, following final paragraph

Insert
The object ant1111b.lmer has the class merMod. See help(merMod) for details of functions ("methods") that are available for use with this class.
page 312 , subsubsection in lines 3 to 26

Replace with:

## *Uncertainty in parameter estimates - profile likelihood \& alternatives

The limits of acceptance of a likelihood ratio test for the null hypothesis of no change in a parameter value can be used as approximate $95 \%$ confidence limits for that parameter. Where the likelihood is a function of more than one parameter, the profile likelihood may be used. For any parameter $\psi$, the profile likelihood is the function of $\psi$ that is obtained by maximizing the likelihood, for each value of $\psi$, over values of other parameters. ${ }^{1}$

The function confint() can be used to pull together the profile information, calculated using the profile method for merMod objects, to create approximate confidence intervals:

```
> prof.lmer <- profile(ant111b.lmer)
> CI95 <- confint(prof.lmer, level=0.95)
> rbind("sigmaL\^{}2"=CI95{[}1,{]}\^{}2, "sigma\^{}2"=CI95{[}2,{]}\^{}2)
    2.5 % 97.5 %
```

sigmaL^2 $0.796 \quad 6.94$
sigma^2 0.3441 .08

A $95 \%$ confidence interval for the intercept is:
> CI95[3,]
2.5 \% $97.5 \%$
(Intercept) 3.1285 .46

[^0]The function confint(), as used here, returned confidence intervals for $\sigma_{L}$ (row label .sig01, random), for $\sigma$ (row label .sigma, random), and for (Intercept) (fixed). The (Intercept) is the intercept in the fitted model, which estimates the overall mean.

The profile likelihoods, scaled so that the lower $2.5 \%$ limit transforms to -1.96 and the upper lower $97.5 \%$ limit, can be plotted thus:

```
library(lattice)
print(xyplot(prof.lmer, conf=c(50, 80, 95, 99)/100, between=list(x=0.35)))
```

For variances, the horizontal scales show Std.Dev.$=\sqrt{\text { Variance. }}$. On the vertical scale, the confidence interval limits are labeled according to the equivalent normal deviates. The $95 \%$ confidence interval limits are thus at -1.96 and 1.96 .

For details of this and other displays that can be used for the output from the profile() method for merMod objects, see help(xyplot.thpr).

See help(confint.merMod) for details of the confint method for merMod objects. Alternatives to method="profile" are method="Wald" or method="boot". The Wald method is fast, but based on approximations that can be highly inaccurate. The boot method uses repeated fits to suitably constructed bootstrap samples, and can be time consuming. The trustworthiness of results from this method may be questioned if more than an occasional fit fails. See help(bootMer) and help(simulate.merMod) for further details of method="boot", and for references.

## page 316 , line 7 :

Replace mcmcsamp() by confint()

## page 316, lines 9-18:

Replace by:

```
> ## Use profile likelihood
> pp <- profile(science1.lmer, which="theta_")
> # which="theta_": all random parameters
> # which="beta_": fixed effect parameters
> var95 <- confint(pp, level=0.95)^2
> # Square to get variances in place of SDs
> rownames(var95) <- c("sigma_Class^2", "sigma^2")
> signif(var95, 3)
    2.5 % 97.5 %
sigma_Class^2 0.178 0.511
sigma^2 2.830 3.300
```


## page 318, footnote 3

Replace with

```
print(VarCorr(science2.lmer), comp="Variance", digits=3)
## The component of variance that is labeled 'Residual' is
## the estimate of the within class variance.
```


## page 326 , lines -10 to -1

Replace line 10 by: The following agree with results from the preceding section:
Replace the code chunk in lines -9 to -2 by:

```
> print(kiwishade.lmer, ranef.comp="Variance", digits=3)
Random effects:
    Groups Name Variance
    block:plot (Intercept) 2.19
    block (Intercept) 4.08
    Residual 12.18
Number of obs: 48, groups: block:plot, 12; block, 3
```

Delete the final line.

## page 330 , line 4 :

Replace by:

```
library(R.utils)
levels(tinting$agegp) <- capitalize(levels(tinting$agegp))
```


## page 330, lines 7, 11 and 14:

Replace method="ML" with REML=FALSE
page 330 , line 15 :

Replace
method="ML"
with
REML=FALSE (use maximum likelihood in place of REML)
page 331, lines 8 and 13:
Replace method="REML" with REML=TRUE
pages 332-333:
Insert replacement pages, as supplied, in place of these.
page 340 , fnote 13 , line 4 :
Replace scale by scales

## page 349, exercise 1 :

Replace the final three lines of code, i.e.

```
vars <- c("(block:plot)^2"=as.vector(vcov[["block:plot"]]),
    "sigma`2"=as.vector(attributes(vcov, "sigmaREML")$sc^2))
print(vars)
by
```

```
print(vcov) # For variances, specify print(vcov^2)
```

```
print(vcov) # For variances, specify print(vcov^2)
```


## Chapter 12

p.381, line -7

Add at end of line aspect="iso", , i.e. the line becomes:

```
xyplot(possum.prc$scores[, 2] ~ possum.prc$scores[, 1], aspect="iso",
```

[Also, replace Figure 12.2 with a figure that has been printed with aspect="iso". (The vertical and horizontal scales are in the same units.)]

## p.383, line -6 ( $3^{\text {rd }}$ last line of code)

Replace value by values
page 388, function DAAG: :CVbinary ()
In line 6, replace leaf17.cv\$cv by leaf17.cv\$cvhat. (It is not good practice to abbreviate the names of list elements.)

Replace the third sentence in the 2nd paragraph (halfway down the page) by: The function CVbinary () returns the resubstitution estimate (other names are internal and training in the list element training, repeated in the list element internal.

## p.395, lines -8 to -4

Replace with:

```
chooseCols <- with(golubInfo, tissue.mf=="PB:f"& cancer=="allB")
df.PBf <- data.frame(t(Golub[ord15, chooseCols, drop=FALSE]))
scores.PBf <- predict(dfB15.lda, newdata=df.PBf, dimen=2)$x
```


## p.403, Figure 12.9

Replace with a figure in which the maximum number of features selected is 23, as in the text and in the code.

## p.409, lines 8 and 9

Replace

```
    # Alternatively, get primates.dna from the DAAGbio package
primates.dist <- dist.dna(primates.dna, model="F84")
with
```

```
# Alternatively, work with primateDNA from the DAAGbio package
```


# Alternatively, work with primateDNA from the DAAGbio package

library(DAAGbio)
library(DAAGbio)
primates.dna <- as.DNAbin(primateDNA)
primates.dna <- as.DNAbin(primateDNA)
primates.dist <- dist.dna(primates.dna, model="K80")

```
primates.dist <- dist.dna(primates.dna, model="K80")
```


## Chapter 13

p.413, footnote 2 , line 1

Replace componenet by component

## p.416, Figure 13.2

Code for Figure 13.2 is not given. See the separate file.

## p.421, code on lines 3 to 12

The code assumes an updated version of the function overlapDensity(). This will be included in the next version of $D A A G(>=1.21)$. It is provided as a separate file. Replace the current line 7 by the more explicit code:

```
overlapDensity(sc.rf[tnum==1], sc.rf[tnum==2], ratio=c(1/20, 50),
    ratio.number=TRUE, plotvalues="Density")
```

Replace the current lines 11 and 12 by the more explicit code:

```
overlapDensity(sc.lda[tnum==1], sc.lda[tnum==2], ratio=c(1/20, 50),
    ratio.number=TRUE, plotvalues="Density")
```


## p.423, line starting proba.rf

Replace probability by prob

## Chapter 14

p.429, line 2

Replace \} by )
page 442, line -6:
Replace scale by scales
page 448, Subsection 14.9.7, line 6:
Replace tsunits[,2] by jobts[,2] or jobts[,"Alberta"].
page 467 , line 10 :
Replace Packages can have their own namespaces, with
by
Each package has its own namespace. There can be

## Chapter 15

p.473, lines 6-8 (2nd para)

Omit lines 6-8.

Run together the two paragraphs in lines -9 to -4 . Omit for details at the end of the final sentence.

## p.473, line -6

Replace The function by The base graphics function.

## p.474, Color [addition]

Add
The function adjustcolor() can be used to set the opacity alpha for the color that is returned. Thus with alpha=0.4, $60 \%$ of the background shows through. Two overlapping points have a combined opacity of $80 \%$, so that $20 \%$ of the background shows through. Lattice and ggplot2 graphics functions accept the argument alpha directly.

## p.481, lines 1 to 7

Delete lines 4-7.
Modify (add a comment at the end) line 2, so that it reads:

```
trellis.device(color=TRUE) # Try also with color=TRUE
```


## p.481, following the current line 7 [addition]

## Insert

A theme settings1 that has been created earlier, e.g., by a call to simpleTheme(), can be modified thus:

```
settings1[["fontsize"]] <- list(text = 16, points = 8)
## Then do, e.g.
grogplot1 <- update(grogplot0, par.settings=settings1)
```


## p.491, Table 2

The current Table 15.2 has been overtaken by changes in ggplot2. Replace Table 15.2 by:

Table 1: Control of ggplot2 graphics features. Note that functions such as xlab() and scale_x_continuous() all have counterparts with y in place of x. .

|  | Argument to qplot() | ggplot() or qplot() ${ }^{1}$ |
| :---: | :---: | :---: |
| Title | main="mytitle" | + labs(title="mytitle") |
| Axes | see help(qplot) | + scale_x_continuous() ${ }^{2}$ [or scale_x_discrete() or scale_x_date()] |
| Axis labels | e.g., xlab="myxlab" | + xlab("myxlab") ${ }^{3}$ |
| $\log$ axes | log="x", (or "y", or "xy") | + scale_x_log10() ${ }^{4}$ |
| Facets ${ }^{5}$ | facets=sex ~ sport | + facet_grid(sex ${ }^{\sim}$ sport) |
| Aspect ratio | e.g., asp=1 | + coord_equal() ${ }^{6}$ |
| Theme | - |  |
| Graph title | e.g., main="maintitle" | + ggtitle("mytitle") |

[^1]
[^0]:    ${ }^{1}$ Note that convergence problems will sometimes occur in the calculation of the profile likelihood, generating warning messages.

[^1]:    ${ }^{1}$ Recall that quickplot() (or qplot()) returns a ggplot object. Functions such as xlab() or scale_x_continuous() can be used, just as for any other ggplot2 object, to update objects returned by quickplot().
    ${ }^{2}$ Available arguments include limits, breaks (locations for the ticks), labels (labels for the breaks), and trans (e.g., trans="log").
    ${ }^{3}$ This is an alternative to using name (e.g., name="myxlab") as an argument to scale_x_continuous() or scale_x_discrete().
    ${ }^{4}$ This is an alternative to using trans="log10" as an argument to scale_x_continuous() or scale_x_discrete(). Note also trans="log" and trans="log2".
    ${ }^{5}$ Facets give Lattice style conditioning.
    ${ }^{6}$ By default (ratio $=1$ ), a given distance, e.g., 1 cm , represents the same range along both $x-$ and $y$-axes.
    ${ }^{7}$ Themes control such graphical attributes as background color, gridlines, and size and color of fonts. See help (ggtheme) for details of other available themes.

