R Code for 'A Practical Guide . . .'

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Preface

Code provided here is for the May 2024 text:

"A Practical Guide to Data Analysis Using R – An Example-Based Approach", by John H Maindonald, W John Braun, and Jeffrey L Andrews.

Code that is shown in the text is filled out to include all code for graphs. In chapter 2 and later, the text includes code only for those graphs that are specifically targeted at the methodology under discussion.

The text has been available in digital format since early May 2024. It has been available in print in the UK from May 30, is due to be available in print in the USA in July, and in Australia and New Zealand in August 2024.

For details for the UK, see: A Practical Guide to Data Analysis Using R – Cambridge University Press UK

1 Chapter 1: Learning from data

```
## xtras=TRUE  ## Set to TRUE to execute code 'extras'
xtras <- FALSE
library(knitr)
## opts_chunk[['set']](results="asis")
## opts_chunk[['set']](eval=FALSE)  ## Set to TRUE to execute main part of code
opts_chunk[['set']](eval=FALSE)</pre>
```

Packages required (plus any dependencies)

latticeExtra (lattice is a dependency); DAAG; car; MASS; AICcmodavg; BayesFactor; boot; MPV; ggplot2; tidyr

Additionally, knitr and Hmisc are required in order to process the Rmd source file. The prettydoc package is by default used to format the html output.

Chapter summary

A note on terminology — variables, factors, and more!

Section 1.1: Questions, and data that may point to answers

Subsection 1.1.1: A sample is a window into the wider population

```
## For the sequence below, precede with set.seed(3676)
set.seed(3696)
sample(1:9384, 12, replace=FALSE) # NB: `replace=FALSE` is the default

chosen1200 <- sample(1:19384, 1200, replace=FALSE)

## For the sequence below, precede with set.seed(366)
set.seed(366)
split(sample(seq(1:10)), rep(c("Control", "Treatment"), 5))
# sample(1:10) gives a random re-arrangement (permutation) of 1, 2, ..., 10</pre>
```

Cluster sampling

*A note on with-replacement samples

```
sample(1:10, replace=TRUE)
## sample(1:10, replace=FALSE) returns a random permutation of 1,2,...10
```

Subsection 1.1.2: Formulating the scientific question

Example: a question about cuckoo eggs

```
library(latticeExtra)  # Lattice package will be loaded and attached also cuckoos <- DAAG::cuckoos  ## Panel A: Dotplot without species means added dotplot(species ~ length, data=cuckoos)  ## `species ~ length` is a 'formula'  ## Panel B: Box and whisker plot bwplot(species ~ length, data=cuckoos)  ## The following shows Panel A, including species means & other tweaks av <- with(cuckoos, aggregate(length, list(species=species), FUN=mean))  dotplot(species ~ length, data=cuckoos, alpha=0.4, xlab="Length of egg (mm)") + as.layer(dotplot(species ~ x, pch=3, cex=1.4, col="black", data=av))  # Use `+` to indicate that more (another 'layer') is to be added.  # With `alpha=0.4`, 40% is the point color with 60% background color  # `pch=3`: Plot character 3 is '+'; `cex=1.4`: Default char size X 1.4
```

```
## Code
suppressPackageStartupMessages(library(latticeExtra, quietly=TRUE))
cuckoos <- DAAG::cuckoos
## For tidier labels replace ".", in several of the names, by a space
specnam <- with(cuckoos, sub(pattern=".", replacement=" ",</pre>
                             levels(species), fixed=TRUE))
# fixed=TRUE: "interpret "." as ".", not as a 'any single character'"
cuckoos <- within(cuckoos, levels(species) <- specnam)</pre>
## Panel A: Dotplot: data frame cuckoos (DAAG)
av <- with(cuckoos, aggregate(length, list(species=species), FUN=mean))
gphA <- dotplot(species ~ length, data=cuckoos, alpha=0.4) +</pre>
  as.layer(dotplot(species ~ x, pch=3, cex=1.4, col="black", data=av))
# alpha sets opacity. With alpha=0.4, 60% of the background shows through
# Enter `print(plt1)` or `plot(plt1)` or simply `plt1` to display the graph
## Panel B: Box plot
gphB <- bwplot(species ~ length, data=cuckoos)</pre>
update(c("A: Dotplot"=gphA, "B: Boxplot"=gphB), between=list(x=0.4),
       xlab="Length of egg (mm)")
## latticeExtra::c() joins compatible plots together.
## See `?latticeExtra::c`
```

Subsection 1.1.3: Planning for a statistical analysis

Understand the data

Causal inference

What was measured? Is it the relevant measure?

Use relevant prior information in the planning stages

Subject area knowledge and judgments

The importance of clear communication

Data-based selection of comparisons

Models must be fit for their intended use

Subsection 1.1.4: Results that withstand thorough and informed challenge

Subsection 1.1.5: Using graphs to make sense of data

Graphical comparisons

Subsection 1.1.6: Formal model-based comparison

```
options(width=70)
cuckoos <- DAAG::cuckoos
av <- with(cuckoos, aggregate(length, list(species=species), FUN=mean))
setNames(round(av[["x"]],2), abbreviate(av[["species"]],10))
with(cuckoos, scale(length[species=="wren"], scale=FALSE))[,1]</pre>
```

Section 1.2: Graphical tools for data exploration

Subsection 1.2.1: Displays of a single variable

```
library(latticeExtra, quietly=TRUE)
fossum <- subset(DAAG::possum, sex=="f")</pre>
femlen <- DAAG::bounce(fossum[["totlngth"]], d=0.1)</pre>
## Panel A
yaxpos <- c(0,5,10,15,20)/(5*nrow(fossum))
z <- boxplot(list(val = femlen), plot = FALSE)</pre>
gph1 <- bwplot(~femlen, ylim=c(0.55, 2.75), xlim=c(70, 100),
                scales=list(y=list(draw=FALSE)))+
        latticeExtra::layer(panel.rug(x,pch="|"))
legstat \leftarrow data.frame(x=c(z$out,z$stats), y=c(1.08, rep(1.3,5)),
  tx=c("Outlier?", "Smallest value", "lower quartile", "median",
       "upper quartile", "Largest value"),
  tx2= c("", "(outliers excepted)",rep("",3), "(no outliers)"))
gphA <- gph1+latticeExtra::layer(data=legstat,</pre>
  panel.text(x=x,y=y,labels=tx,adj=c(0,0.4),srt=90, cex=0.85),
  panel.text(x=x[c(2,6)]+0.75, y=c(1.125,1.38), labels=tx2[c(2,6)],
             adj=c(0,0.4),srt=90, cex=0.85)
```

```
fossum <- subset(DAAG::possum, sex=="f")
densityplot(~totlngth, plot.points=TRUE, pch="|", data=fossum) +
    layer_(panel.histogram(x, type="density", breaks=c(75,80,85,90,95,100)))</pre>
```

Comparing univariate displays across factor levels

```
library(latticeExtra, quietly=TRUE)
fossum <- subset(DAAG::possum, sex=="f")</pre>
femlen <- DAAG::bounce(fossum[["totlngth"]], d=0.1)</pre>
## Panel A
yaxpos <- c(0,5,10,15,20)/(5*nrow(fossum))
z <- boxplot(list(val = femlen), plot = FALSE)</pre>
gph1 \leftarrow bwplot(\sim femlen, ylim=c(0.55, 2.75), xlim=c(70, 100),
                scales=list(y=list(draw=FALSE)))+
        latticeExtra::layer(panel.rug(x,pch="|"))
legstat \leftarrow data.frame(x=c(z$out,z$stats), y=c(1.08, rep(1.3,5)),
  tx=c("Outlier?", "Smallest value", "lower quartile", "median",
       "upper quartile", "Largest value"),
  tx2= c("", "(outliers excepted)",rep("",3), "(no outliers)"))
gphA <- gph1+latticeExtra::layer(data=legstat,</pre>
  panel.text(x=x,y=y,labels=tx,adj=c(0,0.4),srt=90,cex=0.85),
  panel.text(x=x[c(2,6)]+0.75, y=c(1.125,1.38), labels=tx2[c(2,6)],
             adj=c(0,0.4),srt=90, cex=0.85)
## Panel B
gph2 <- densityplot(\negfemlen, ylim=c(0,0.108), xlim=c(70,100),
          plot.points=TRUE, pch="|",cex=1.75, ylab=c("","
                                                                   Density"))
```

```
## Create boxplot graph object --- Simplified code
gph <- bwplot(Pop~totlngth | sex, data=possum)
## plot graph, with dotplot distribution of points below boxplots
gph + latticeExtra::layer(panel.dotplot(x, unclass(y)-0.4))</pre>
```

Subsection 1.2.2: Patterns in univariate time series

```
layout(matrix(c(1,2)), heights=c(2.6,1.75))
measles <- DAAG::measles</pre>
## Panel A:
par(mgp=c(2.0,0.5,0))
plot(log10(measles), xlab="", ylim=log10 (c(1,5000*540)),
     ylab=" Deaths", yaxt="n", fg="gray", adj=0.16)
londonpop <-ts(c(1088, 1258, 1504, 1778, 2073, 2491, 2921, 3336, 3881,
  4266, 4563, 4541, 4498, 4408), start=1801, end=1931, deltat=10)
points(log10(londonpop*500), pch=16, cex=.5)
ytiks1 \leftarrow c(1, 10, 100, 1000)
axis(2, at=log10(ytiks1), labels=paste(ytiks1), lwd=0, lwd.ticks=1)
abline(h=log10(ytiks1), col = "lightgray", lwd=2)
par(mgp=c(-2,-0.5,0))
ytiks2 <- c(1000000, 5000000)  ## London population in thousands
abline(h=log10(ytiks2*0.5), col = "lightgray", lwd=1.5)
abline(v=seq(from=1650,to=1950,by=50), col = "lightgray", lwd = 1.5)
mtext(side=2, line=0.5, "Population", adj=1, cex=1.15, las=3)
axis(2, at=log10(ytiks2*0.6), labels=paste(ytiks2), tcl=0.3,
     hadj=0, lwd=0, lwd.ticks=1)
mtext(side=3, line=0.3, "A (1629-1939)", adj=0, cex=1.15)
##
```

Subsection 1.2.3: Visualizing relationships between pairs of variables

Subsection 1.2.4: Response lines (and/or curves)

```
par(pty="s")
plot(distance.traveled ~ starting.point, data=DAAG::modelcars, fg="gray",
xlim=c(0,12.5), xaxs="i", xlab = "Distance up ramp (cm)",
ylab="Distance traveled (cm)")
```

Subsection 1.2.5: Multiple variables and times

```
## Apply function range to columns of data frame jobs (DAAG)
sapply(DAAG::jobs, range) ## NB: `BC` = British Columbia
```

```
## Panel A: Basic plot; all series in a single panel; use log y-scale
formRegions <- Ontario+Quebec+BC+Alberta+Prairies+Atlantic ~ Date</pre>
basicGphA <-
  xyplot(formRegions, outer=FALSE, data=DAAG::jobs, type="1", xlab="",
         ylab="Number of workers", scales=list(y=list(log="e")),
         auto.key=list(space="right", lines=TRUE, points=FALSE))
  ## `outer=FALSE`: plot all columns in one panel
## Panel B: Separate panels (`outer=TRUE`); sliced log scale
basicGphB <-
  xyplot(formRegions, data=DAAG::jobs, outer=TRUE, type="l", layout=c(3,2),
         xlab="", ylab="Number of workers",
         scales=list(y=list(relation="sliced", log=TRUE)))
# Provinces are in order of number of workers in Dec96
## Create improved x- and y-axis tick labels; will update to use
datelabpos <- seq(from=95, by=0.5, length=5)</pre>
datelabs <- format(seq(from=as.Date("1Jan1995", format="%d%b%Y"),
                   by="6 month", length=5), "%b%y")
```

```
## Now create $y$-labels that have numbers, with log values underneath
ylabposA <- exp(pretty(log(unlist(DAAG::jobs[,-7])), 5))</pre>
ylabelsA <- paste(round(ylabposA),"\n(", log(ylabposA), ")", sep="")</pre>
## Repeat, now with 100 ticks, to cover all 6 slices of the scale
ylabposB <- exp(pretty(log(unlist(DAAG::jobs[,-7])), 100))</pre>
ylabelsB <- paste(round(ylabposB), "\n(", log(ylabposB), ")", sep="")</pre>
gphA <- update(basicGphA, scales=list(x=list(at=datelabpos, labels=datelabs),</pre>
              y=list(at=ylabposA, labels=ylabelsA)))
gphB <- update(basicGphB, xlab="", between=list(x=0.25, y=0.25),</pre>
                scales=list(x=list(at=datelabpos, labels=datelabs),
               y=list(at=ylabposB, labels=ylabelsB)))
layout.list <- list(layout.heights=list(top.padding=0,</pre>
                     bottom.padding=0, sub=0, xlab=0),
                     fontsize=list(text=8, points=5))
jobstheme <- modifyList(ggplot2like(pch=1, lty=c(4:6,1:3),</pre>
                                     col.line='black', cex=0.75),layout.list)
print(update(gphA, par.settings=jobstheme, axis=axis.grid,
      main=list("A: Same vertical log scale", y=0)),
      position=c(0.1,0.615,0.9,1), newpage=TRUE)
print(update(gphB, par.settings=jobstheme, axis=axis.grid,
      main=list("B: Sliced vertical log scale",y=0)),
      position=c(0,0,1,0.625), newpage=FALSE)
```

```
plot(c(1230,1860), c(0, 10.5), axes=FALSE, bty="n",
     xlab="", ylab="", type="n", log="x")
xpoints <- c(1366, 1436, 1752, 1840)
axis(1, at=xpoints, labels=FALSE, tck=0.01, lty=1, lwd=0, lwd.ticks=1)
for(i in 1:4){
  axis(1, at=xpoints[i],
       labels=substitute(italic(a), list(a=paste(xpoints[i]))),
  line=-2.25, lty=0, cex=0.8, lwd=0, lwd.ticks=1)
  lines(rep(xpoints[i],2), c(0, 0.15*par()[["cxy"]][2]), lty=1)
axpos <- 1250*cumprod(c(1, rep(1.2,2)))
axis(1, at=c(axpos,1840), labels=F, lwd.ticks=0)
lab <- round(axpos)</pre>
axis(1, at=axpos, labels=lab)
lab2 <- lapply(round(log2(xpoints),3), function(x)substitute(2^a, list(a=x)))</pre>
axis(1, at=xpoints, labels=as.expression(lab2), line=-3.5, lwd=0)
labe <- lapply(format(round(log(xpoints),3)), function(x)substitute(e^a, list(a=x)))</pre>
axis(1, at=xpoints, labels=as.expression(labe), line=-5, lwd=0)
lab10 <- lapply(round(log10(xpoints),3), function(x)substitute(10^a, list(a=x)))</pre>
```

```
axis(1, at=xpoints, labels=as.expression(lab10), line=-6.5, lwd=0)
par(family="mono", xpd=TRUE)
axis(1, at=1220, labels="log=2", line=-3.5, hadj=0, lwd=0)
axis(1, at=1220, labels='log="e"', line=-5, hadj=0, lwd=0)
axis(1, at=1220, labels="log=10", line=-6.5, hadj=0, lwd=0)
wid2 <- strwidth("log=2")
par(family="sans")</pre>
```

Subsection 1.2.6: *Labeling technicalities

Subsection 1.2.7: Graphical displays for categorical data

Subsection 1.2.8: What to look for in plots

Outliers

Asymmetry of the distribution

Changes in variability

Clustering

Nonlinearity

Time trends in the data

Section 1.3: Data Summary

Subsection 1.3.1: Counts

```
## Table of counts example: data frame nswpsid1 (DAAG)
## Specify `useNA="ifany"` to ensure that any NAs are tabulated
tab <- with(DAAG::nswpsid1, table(trt, nodeg, useNA="ifany"))
dimnames(tab) <- list(trt=c("none", "training"), educ = c("completed", "dropout"))
tab</pre>
```

Tabulation that accounts for frequencies or weights - the xtabs() function

```
gph <- lattice::bwplot(log(nassCDS$weight+1), xlab="Inverse sampling weights",
  scales=list(x=list(at=c(0,log(c(10^(0:5)+1))), labels=c(0,10^(0:5)))))
update(gph, par.settings=DAAG::DAAGtheme(color=F, col.points='gray50'))
sampNum <- table(nassCDS$dead)</pre>
popNum <- as.vector(xtabs(weight ~ dead, data=nassCDS))</pre>
rbind(Sample=sampNum, "Total number"=round(popNum,1))
nassCDS <- DAAG::nassCDS</pre>
Atab <- xtabs(weight ~ airbag + dead, data=nassCDS)/1000
## Define a function that calculates Deaths per 1000
DeadPer1000 <- function(x)1000*x[2]/sum(x)
Atabm <- ftable(addmargins(Atab, margin=2, FUN=DeadPer1000))</pre>
print(Atabm, digits=2, method="compact", big.mark=",")
SAtab <- xtabs(weight ~ seatbelt + airbag + dead, data=nassCDS)
## SAtab <- addmarqins(SAtab, marqin=3, FUN=list(Total=sum)) ## Gdet Totals
SAtabf <- ftable(addmargins(SAtab, margin=3, FUN=DeadPer1000), col.vars=3)
print(SAtabf, digits=2, method="compact", big.mark=",")
```

```
FSAtab <- xtabs(weight ~ dvcat + seatbelt + airbag + dead, data=nassCDS)
FSAtabf <- ftable(addmargins(FSAtab, margin=4, FUN=DeadPer1000), col.vars=3:4)
print(FSAtabf, digits=1)
```

Subsection 1.3.2: Summaries of information from data frames

```
## Individual vine yields, with means by block and treatment overlaid
kiwishade <- DAAG::kiwishade
kiwishade$block <- factor(kiwishade$block, levels=c("west","north","east"))
keyset <- list(space="top", columns=2,</pre>
```

The benefits of data summary – dengue status example

Subsection 1.3.3: Measures of variation

Cuckoo eggs example

Degrees of freedom

Subsection 1.3.4: Inter-quartile range (IQR) and median absolute deviation (MAD)

Subsection 1.3.5: A pooled standard deviation estimate

Elastic bands example

```
<>= sapply(DAAG::two65, function(x) c(Mean=mean(x), sd=sd(x))) |> round(2) @
```

Subsection 1.3.6: Effect size

```
setNames(diff(c(ambient=244.1, heated=253.5))/c(sd=10.91), "Effect size")
```

Data are available in the data frame DAAG::two65.

```
vignette('effectsize', package='effectsize')
```

Subsection 1.3.7: Correlation

```
set.seed(17)
x1 \leftarrow x2 \leftarrow x3 \leftarrow (11:30)/5
y1 <- x1 + rnorm(20, sd=0.5)
y2 \leftarrow 2 - 0.05 * x1 + 0.1 * ((x1 - 1.75))^4 + rnorm(20, sd=1.5)
y3 \leftarrow (x1 - 3.85)^2 + 0.015 + rnorm(20)
theta \leftarrow ((2 * pi) * (1:20))/20
x4 < -10 + 4 * cos(theta)
y4 < -10 + 4 * sin(theta) + rnorm(20, sd=0.6)
xy \leftarrow data.frame(x = c(rep(x1, 3), x4), y = c(y1, y2, y3, y4),
                 gp = factor(rep(1:4, rep(20, 4))))
xysplit <- split(xy, xy$gp)</pre>
rho <- sapply(xysplit, function(z)with(z,cor(x,y, method=c("pearson"))))</pre>
rhoS <- sapply(xysplit, function(z)with(z,cor(x,y, method=c("spearman"))))</pre>
rnam <- as.list(setNames(round(c(rho,rhoS),2), paste0("r",1:8)))</pre>
striplabs <- bquote(expression(paste(r==.(r1), " ",r[s]==.(r5)),
                                 paste(r==.(r2), " ",r[s]==.(r6)),
                                 paste(r==.(r3), " ",r[s]==.(r7)),
                                 paste(r==.(r4), " ",r[s]==.(r8))), rnam)
xyplot(y ~ x | gp, data=xy, layout=c(4,1), xlab="", ylab="",
  strip=strip.custom(factor.levels=striplabs), aspect=1,
  scales=list(relation='free', draw=FALSE), between=list(x=0.5,y=0)
```

Section 1.4: Distributions: quantifying uncertainty

Subsection 1.4.1: Discrete distributions

```
## dbinom(0:10, size=10, prob=0.15)
setNames(round(dbinom(0:10, size=10, prob=0.15), 3), 0:10)
pbinom(q=4, size=10, prob=0.15)
```

```
qbinom(p = 0.70, size = 10, prob = 0.15)

## Check that this lies between the two cumulative probabilities:

## pbinom(q = 1:2, size=10, prob=0.15)
```

```
rbinom(15, size=4, p=0.5)
```

```
## dpois(x = 0:8, lambda = 3)
setNames(round(dpois(x = 0:8, lambda = 3),4), 0:8)
## Probability of > 8 \ raisins
## 1-ppois(q = 8, lambda = 3) ## 0r, ppois(q=8, lambda=3, lower.tail=FALSE)
```

```
1 - ppois(q = 8, lambda = 3)
ppois(q=8, lambda=3, lower.tail=FALSE) ## Alternative
1-sum(dpois(x = 0:8, lambda = 3)) ## Another alternative
```

```
raisins <- rpois(20, 3)
raisins</pre>
```

Initializing the random number generator

```
set.seed(23286) # Use to reproduce the sample below
rbinom(15, size=1, p=0.5)
```

Means and variances

Subsection 1.4.2: Continuous distributions

```
polygon(c(z[z \le 1.0], 1.0), c(dnorm(z[z \le 1.0]), dnorm(-3)), col="grey")
chh <- par()$cxy[2]</pre>
arrows(-1.8, 0.32, -0.25, 0.2, length=0.07, xpd=T)
cump <- round(pnorm(1), 3)</pre>
text(-1.8, 0.32+0.75*chh, paste("pnorm(1)\n", "=", cump), xpd=T, cex=0.8)
pnormExs <- c('pnorm(0)', 'pnorm(1)', 'pnorm(-1.96)', 'pnorm(1.96)',</pre>
'pnorm(1.96, mean=2)', 'pnorm(1.96, sd=2)')
Prob <- sapply(pnormExs, function(x)eval(parse(text=x)))</pre>
df <- as.data.frame(Prob)</pre>
df$Prob <- round(df$Prob,3)</pre>
print(df)
## Plot the normal density, in the range -3 to 3
z \leftarrow pretty(c(-3,3), 30) # Find ~30 equally spaced points
                           # Equivalent to dnorm(z, mean=0, sd=1)
ht <- dnorm(z)</pre>
plot(z, ht, type="l", xlab="Normal variate", ylab="Density", yaxs="i")
# yaxs="i" locates the axes at the limits of the data
qnorm(.9)
                    # 90th percentile; mean=0 and SD=1
## Additional examples:
setNames(qnorm(c(.5,.841,.975)), nm=c(.5,.841,.975))
qnorm(c(.1,.2,.3)) # -1.282 -0.842 -0.524 (10th, 20th and 30th percentiles)
```

Different ways to represent distributions

Generating simulated samples from the normal and other continuous distributions

qnorm(.1, mean=100, sd=10) # 87.2 (10th percentile, mean=100, SD=10)

```
options(digits=2)  # Suggest number of digits to display
rnorm(10)  # 10 random values from the normal distribution
```

```
mu <- 10
sigma <- 1
n <- 1
m <- 50
four <- 4
nrep <- 5
```

```
seed <- 21
totrows <- 1
if(is.null(totrows))
totrows <- floor(sqrt(nrep))</pre>
totcols <- ceiling(nrep/totrows)</pre>
z \leftarrow range(pretty(mu + (c(-3.4, 3.4) * sigma), 50))
xy <- data.frame(x=rep(0,nrep),y=rep(0,nrep),n=rep(n,nrep),</pre>
                  mm=rep(m,nrep),four=rep(four,nrep))
fac <- factor(paste("Simulation", 1:nrep),</pre>
               lev <- paste("Simulation", 1:nrep))</pre>
xlim<-z
## ylim<-c(0, dnorm(0)*sqrt(n))
ylim \leftarrow c(0,1)
xy <- split(xy,fac)</pre>
xy<-lapply(1:length(xy),function(i){c(as.list(xy[[i]]), list(xlim=xlim,</pre>
            ylim=ylim))})
panel.mean <- function(data, mu = 10, sigma = 1, n2 = 1,
                        mm = 100, nrows, ncols, ...)
{
  vline <- function(x, y, lty = 1, col = 1)
  lines(c(x, x), c(0, y), lty = lty, col = col)
  n2<-data$n[1]
  mm<-data$mm[1]
  our \leftarrow data four [1] ## Four characters in each unit interval of x
  nmid <- round(four * 4)</pre>
  nn \leftarrow array(0, 2 * nmid + 1)
  z <- mu+seq(from=-3.4*sigma, to=3.4*sigma, length=mm)
  atx<-pretty(z)
  qz <- pnorm((z - mu)/sigma)
  dz <- dnorm((z - mu)/sigma)</pre>
  chw <- sigma/four</pre>
  chh <- strheight("0")*0.75
  htfac <- (mm * chh)/four
  if(nrows==1&&ncols==1)
  lines(z, dz * htfac)
  if(nrows==1)axis(1,at=atx, lwd=0, lwd.ticks=1)
  y <- rnorm(mm, mu, sigma/sqrt(n2))
  pos <- round((y - mu)/sigma * four)</pre>
  for(i in 1:mm) {
    nn[nmid + pos[i]] \leftarrow nn[nmid + pos[i]] + 1
    xpos <- chw * pos[i]</pre>
```

```
text(mu + xpos, nn[nmid + pos[i]] * chh - chh/4, "x")
}

DAAG::panelplot(xy,panel=panel.mean,totrows=totrows,totcols=totcols,
    oma=c(1.5, 0, rep(0.5,2)), fg='gray')

## The following gives a conventional histogram representations:
set.seed (21)  # Use to reproduce the data in the figure
df <- data.frame(x=rnorm(250), gp=rep(1:5, rep(50,5)))
lattice::histogram(~x|gp, data=df, layout=c(5,1))

runif(n = 20, min=0, max=1) # 20 numbers, uniform distn on (0, 1)
rexp(n=10, rate=3) # 10 numbers, exponential, mean 1/3.</pre>
```

Subsection 1.4.3: Graphical checks for normality

```
tab <- t(as.matrix(DAAG::pair65))
rbind(tab, "heated-ambient"=tab[1,]-tab[2,])

## Normal quantile-quantile plot for heated-ambient differences,
## compared with plots for random normal samples of the same size
plt <- with(DAAG::pair65, DAAG::qreference(heated-ambient, nrep=10, nrows=2))
update(plt, scales=list(tck=0.4), xlab="")</pre>
```

Subsection 1.4.4: Population parameters and sample statistics

The sampling distribution of the mean

The standard error of the median

Simulation in learning and research

Subsection 1.4.5: The *t*-distribution

```
x \leftarrow seq(from=-4.2, to = 4.2, length.out = 50)
ylim \leftarrow c(0, dnorm(0))
ylim[2] <- ylim[2]+0.1*diff(ylim)</pre>
h1 \leftarrow dnorm(x)
h3 < -dt(x, 3)
h8 \leftarrow dt(x,8)
plot(x, h1, type="l", xlab = "", xaxs="i", ylab = "", yaxs="i",
bty="L", ylim=ylim, fg="gray")
mtext(side=3,line=0.5, "A: Normal (t8 overlaid)", adj=-0.2)
lines(x, h8, col="grey60")
mtext(side=1, line=1.75, "No. of SEMs from mean")
mtext(side=2, line=2.0, "Probability density")
chh <- par()$cxy[2]
topleft <- par()$usr[c(1,4)] + c(0, 0.6*chh)
legend(topleft[1], topleft[2], col=c("black", "grey60"),
lty=c(1,1), legend=c("Normal","t (8 d.f.)"), bty="n", cex=0.8)
plot(x, h1, type="l", xlab = "", xaxs="i",
ylab = "", yaxs="i", bty="L", ylim=ylim, fg="gray")
mtext(side=3,line=0.5, "B: Normal (t3 overlaid)", adj=-0.2)
lines(x, h3, col="grey60")
mtext(side=1, line=1.75, "No. of SEMs from mean")
## mtext(side=2, line=2.0, "Probability density")
legend(topleft[1], topleft[2], col=c("black", "grey60"),
lty=c(1,1), legend=c("Normal","t (3 d.f.)"), bty="n", cex=0.8)
## Panels C and D
cump <- 0.975
```

```
x \leftarrow seq(from=-3.9, to = 3.9, length.out = 50)
ylim \leftarrow c(0, dnorm(0))
plotfun <- function(cump, dfun = dnorm, qfun=qnorm,</pre>
ytxt = "Probability density",
txt1="qnorm", txt2="", ...)
h \leftarrow dfun(x)
plot(x, h, type="l", xlab = "", xaxs="i", xaxt="n",
ylab = ytxt, yaxs="i", bty="L", ylim=ylim, fg="gray",
axis(1, at=c(-2, 0), cex=0.8, lwd=0, lwd.ticks=1)
axis(1, at=c((-3):3), labels=F, lwd=0, lwd.ticks=1)
tailp <- 1-cump
z <- qfun(cump)</pre>
ztail <- pretty(c(z,4),20)
htail <- dfun(ztail)</pre>
polygon(c(z,z,ztail,max(ztail)), c(0,dfun(z),htail,0), col="gray")
text(0, 0.5*dfun(z)+0.08*dfun(0),
paste(round(tailp, 3), " + ", round(1-2*tailp,3),
"\n= ", round(cump, 3), sep=""), cex=0.8)
lines(rep(z, \frac{2}{2}), c(\frac{0}{2}, dfun(z)))
lines(rep(-z, 2), c(0, dfun(z)), col="gray60")
chh \leftarrow par()$cxy[2]
arrows(z, -1.5*chh, z, -0.1*chh, length=0.1, xpd=T)
text(z, -2.5*chh, paste(txt1, "(", cump, txt2, ")", "\n= ",
round(z,2), sep=""), xpd=T)
x1 < -z + .3
y1 <- dfun(x1)*0.35
y0 < -dfun(0)*0.2
arrows(-2.75, y0, -x1, y1, length=0.1, col="gray60")
arrows(2.75, y0, x1, y1, length=0.1)
text(-2.75, y0+0.5*chh, tailp, col="gray60")
text(2.75, y0+0.5*chh, tailp)
}
## ytxt <- "t probability density (8 d.f.)"
plotfun(cump=cump, cex.lab=1.05)
mtext(side=3, line=1.25, "C: Normal distribution", adj=-0.2)
ytxt <- "t probability density (8 d.f.)"</pre>
plotfun(cump=cump, dfun=function(x)dt(x, 8),
        qfun=function(x)qt(x, 8),
        ytxt="", txt1="qt", txt2=", 8", cex.lab=1.05)
mtext(side=3, line=1.25, "D: t distribution (8 df)", adj=-0.2)
```

```
qnorm(c(0.975,0.995), mean=0) # normal distribution qt(c(0.975, 0.995), df=8) # t-distribution with 8 d.f.
```

Subsection 1.4.6: The likelihood, and maximum likelihood estimation

Section 1.5: Simple forms of regression model

Subsection 1.5.1: Line or curve?

```
roller <- DAAG::roller
t(cbind(roller, "depression/weight ratio"=round(roller[,2]/roller[,1],2)))</pre>
```

Using models to predict

```
y <- DAAG::roller$depression
x <- DAAG::roller$weight
pretext <- c(reg = "A", lo = "B")</pre>
for(curve in c("reg", "lo")) {
  plot(x, y, xlab = "Roller weight (t)", xlim=c(0,12.75), fg="gray",
       ylab = "Depression in lawn (mm)", type="n")
  points(x, y, cex=0.8, pch = 4)
  mtext(side = 3, line = 0.25, pretext[curve], adj = 0)
  topleft <- par()$usr[c(1, 4)]</pre>
  chw <- strwidth("0"); chh <- strheight("0")</pre>
  points(topleft[1]+rep(0.75,2)*chw,topleft[2]-c(0.75,1.8)*chh,
         pch=c(4,1), col=c("black", "gray40"), cex=0.8)
  text(topleft[1]+rep(1.2,2)*chw, topleft[2]-c(0.75,1.8)*chh,
       c("Data values", "Fitted values"),adj=0, cex=0.8)
  if(curve=="lo")
    text(topleft[1]+1.2*chw, topleft[2]-2.85*chh, "(smooth)", adj=0, cex=0.8)
  if(curve[1] == "reg") {
    u < -1m(y \sim -1 + x)
  abline(0, u$coef[1])
  yhat <- predict(u)</pre>
else {
  lawn.lm < -lm(y \sim x + I(x^2))
  yhat<-predict(lawn.lm)</pre>
  xnew < -pretty(x, 20)
```

```
b<-lawn.lm$coef
  ynew < -b[1] + b[2] * xnew + b[3] * xnew^2
  lines(xnew, ynew)
}
here <- y < yhat
yyhat <- as.vector(rbind(y[here], yhat[here], rep(NA, sum(here))))
xx <- as.vector(rbind(x[here], x[here], rep(NA, sum(here))))</pre>
lines(xx, yyhat, lty = 2, col="gray")
here <- y > yhat
yyhat <- as.vector(rbind(y[here], yhat[here], rep(NA, sum(here))))
xx <- as.vector(rbind(x[here], x[here], rep(NA, sum(here))))</pre>
lines(xx, yyhat, lty = 1, col="gray")
n <- length(y)</pre>
ns \leftarrow min((1:n)[y - yhat >= 0.75*max(y - yhat)])
ypos \leftarrow 0.5 * (y[ns] + yhat[ns])
chw <- par()$cxy[1]</pre>
text(x[ns] - 0.25*chw, ypos, "+ve residual", adj = 1,cex=0.75, col="gray30")
points(x, yhat, pch = 1, col="gray40")
ns \leftarrow (1:n)[y - yhat == min(y - yhat)][1]
ypos \leftarrow 0.5 * (y[ns] + yhat[ns])
text(x[ns] + 0.4*chw, ypos, "-ve residual", adj = 0,cex=0.75,col="gray30")
```

Which model is best — line or curve?

Subsection 1.5.2: Fitting models – the model formula

```
## Fit line - by default, this fits intercept & slope.
roller.lm <- lm(depression ~ weight, data=DAAG::roller)
## Compare with the code used to plot the data
plot(depression ~ weight, data=DAAG::roller)
## Add the fitted line to the plot
abline(roller.lm)</pre>
```

```
## For a model that omits the intercept term, specify
lm(depression ~ 0 + weight, data=roller) # Or, if preferred, replace `0` by `-1`
```

Model objects

```
roller.lm <- lm(depression ~ weight, data=DAAG::roller)
names(roller.lm)  # Get names of list elements</pre>
```

```
roller.lm$coef # An alternative is roller.lm[["coef"]]
```

```
print(summary(roller.lm), digits=3)
```

Residual plots

```
## Normal quantile-quantile plot, plus 7 reference plots
DAAG::qreference(residuals(roller.lm), nrep=8, nrows=2, xlab="")
```

Simulation of regression data

```
roller.lm <- lm(depression ~ weight, data=DAAG::roller)
roller.sim <- simulate(roller.lm, nsim=20) # 20 simulations
with(DAAG::roller, matplot(weight, roller.sim, pch=1, ylim=range(depression)))
points(DAAG::roller, pch=16)</pre>
```

Subsection 1.5.3: The model matrix in regression

```
model.matrix(roller.lm)
## Specify coef(roller.lm) to obtain the column multipliers.
```

From straight line regression to multiple regression

```
mouse.lm <- lm(brainwt ~ lsize+bodywt, data=DAAG::litters)
coef(summary(mouse.lm))</pre>
```

Section 1.6: Data-based judgments – frequentist, in a Bayesian world

Subsection 1.6.1: Inference with known prior probabilities

Relating 'incriminating' evidence to the probability of guilt

Subsection 1.6.2: Treatment differences that are on a continuous scale

```
## Sum of tail probabilities
2*pt(1.580/0.389, 9, lower.tail=FALSE)
```

```
## 95% CI for mean of heated-ambient: data frame DAAG::pair65
t.test(sleep, conf.level=0.95)
```

An hypothesis test

```
pt(4.06, 9, lower.tail=F)
```

The p-value probability relates to a decision process

Subsection 1.6.3: Use of simulation with p-values

eff2stat <- function(eff=c(.2,.4,.8,1.2), n=c(10,40), numreps=100,

```
FUN=function(x,N)pt(sqrt(N)*mean(x)/sd(x), df=N-1,
                                           lower.tail=FALSE)){
  simStat \leftarrow function(eff=c(.2,.4,.8,1.2), N=10, nrep=100, FUN){
    num <- N*nrep*length(eff)</pre>
    array(rnorm(num, mean=eff), dim=c(length(eff),nrep,N)) |>
      apply(2:1, FUN, N=N)
  mat <- matrix(nrow=numreps*length(eff),ncol=length(n))</pre>
  for(j in 1:length(n)) mat[,j] <-</pre>
    as.vector(simStat(eff, N=n[j], numreps, FUN=FUN)) ## length(eff)*numep
  data.frame(effsize=rep(rep(eff, each=numreps), length(n)),
             N=rep(n, each=numreps*length(eff)), stat=as.vector(mat))
set.seed(31)
df200 <- eff2stat(eff=c(.2,.4,.8,1.2), n=c(10, 40), numreps=200)
labx \leftarrow c(0.001, 0.01, 0.05, 0.2, 0.4, 0.8)
gph <- bwplot(factor(effsize) ~ I(stat^0.25) | factor(N), data=df200,</pre>
               layout=c(2,1), xlab="P-value", ylab="Effect size",
               scales=list(x=list(at=labx^0.25, labels =labx)))
update(gph+latticeExtra::layer(panel.abline(v=labx[1:3]^0.25, col='lightgray')),
```

```
eff10 <- with(subset(df200, N==10&effsize==0.2), c(gt5pc=sum(stat>0.05), lohi=fivenum(stat)[eff40 <- with(subset(df200, N==40&effsize==0.2), c(gt5pc=sum(stat>0.05), c(gt5pc=sum(stat)[eff40 <- with(subset(df200, N==40&effsize==0.2), c(gt5pc=sum(stat>0.05), c(gt5pc=sum(stat)[eff50 <- with(subset(df200, N==40&effsize==0.2), c(gt5pc=sum(stat)[eff50 <- with(subset(df200, N==40&effsize==0.2),
```

Subsection 1.6.4: Power — minimizing the chance of false positives

strip=strip.custom(factor.levels=paste0("n=",c(10,40))),
par.settings=DAAG::DAAGtheme(color=F, col.points="gray50"))

```
tf1 <- rbind('R=0.2'=c(0.8*50, 0.05*250),
'R=1'=c(0.8*150, 0.05*150),
'R=5'=c(0.8*200, 0.05*50))
tf2 <- rbind(c('0.8 x50', '0.05x250'),
```

```
c('0.8x150', '0.05x150'),
c('0.8x250', '0.05 x50'))
tf <- cbind("True positives"=paste(tf2[,2],tf1[,2],sep="="),
"False positives"=paste(tf2[,1],tf1[,1],sep="="))
rownames(tf) <- rownames(tf1)
print(tf, quote=FALSE)</pre>
```

Power calculations – examples

```
power.t.test(d=0.5, sig.level=0.05, type="one.sample", power=0.8)
pwr1 <- power.t.test(d=0.5, sig.level=0.005, type="one.sample", power=0.8)
pwr2 <- power.t.test(d=0.5, sig.level=0.005, type="two.sample", power=0.8)
## d=0.5, sig.level=0.005, One- and two-sample numbers
c("One sample"=pwr1$n, "Two sample"=pwr2$n)</pre>
```

```
print(tab[,1:3], quote=F)
```

```
print(tab[,4:6], quote=F)
```

Positive Predictive Values

Subsection 1.6.5: The future for p-values

How small a p-value is needed?

Subsection 1.6.6: Reporting results

Is there an alternative that is more likely?

Section 1.7: Information statistics and Bayesian methods with Bayes Factors

Subsection 1.7.1: Information statistics – using likelihoods for model choice

```
## Calculations using mouse brain weight data
mouse.lm <- lm(brainwt ~ lsize+bodywt, data=DAAG::litters)
n <- nrow(DAAG::litters)
RSSlogLik <- with(mouse.lm, n*(log(sum(residuals^2)/n)+1+log(2*pi)))
p <- length(coef(mouse.lm))+1  # NB: p=4 (3 coefficients + 1 scale parameter)</pre>
```

```
k <- 2*n/(n-p-1)
c("AICc" = AICcmodavg::AICc(mouse.lm), fromlogL=k*p-2*logLik(mouse.lm)[1],
fromFit=k*p + RSSlogLik) |> print(digits=4)
```

The sampling properties of the difference in AIC statistics

```
simOvs1 <- function(mu=0, n=15, ntimes=200){</pre>
a0 <- a1 <- numeric(ntimes)
for(i in 1:ntimes){
  y <- rnorm(n, mean=mu, sd=1)
  m0 \leftarrow lm(y \sim 0); m1 \leftarrow lm(y \sim 1)
  a0[i] <- AIC(m0); a1[i] <- AIC(m1)
data.frame(a0=a0, a1=a1, diff01=a0-a1, mu=rep(paste0("mu=",mu)))
library(latticeExtra)
sim0 <- sim0vs1(mu=0)</pre>
sim0.5 <- sim0vs1(mu=0.5)
simboth <- rbind(sim0, sim0.5)</pre>
cdiff <- with(list(n=15, p=2), 2*(p+1)*p/(n-(p+1)-1))
xyplot(diff01 ~ a0 | mu, data=simboth, xlab="AIC(m0)", ylab="AIC(m0) - AIC(m1)") +
  latticeExtra::layer({panel.abline(h=0, col='red');
         panel.abline(h=cdiff, lwd=1.5, lty=3, col='red', alpha=0.5);
         panel.abline(h=-2, lty=2, col='red')})
tab <- rbind(c(with(sim0, sum(diff01>0))/200, with(sim0.5, sum(diff01>0))/200),
  c(with(sim0,sum(diff01>-cdiff))/200, with(sim0.5, sum(diff01>-cdiff))/200))
dimnames(tab) <- list(c("AIC: Proportion choosing m1",</pre>
                         "AICc: Proportion choosing m1"),
                       c("True model is m0", "True model is m1"))
tab
```

Subsection 1.7.2: Bayesian methods with Bayes Factors

The Cauchy prior with different choices of scale parameter

```
x \leftarrow seq(from=-4.5, to=4.5, by=0.1)
densMed <- dcauchy(x,scale=sqrt(2)/2)</pre>
densUltra <- dcauchy(x, scale=sqrt(2))</pre>
denn <- dnorm(x, sd=1)</pre>
plot(x,densMed, type='l', mgp=c(2,0.5,0), xlab="",
     ylab="Prior density", col="red", fg='gray')
mtext(side=1, line=2, expression("Effect size "==phantom(0)*delta), cex=1.1)
lines(x, denn, col="blue", lty=2)
lines(x, densUltra,col=2, lty=2)
legend("topleft", title="Normal prior",
       y.intersp=0.8, lty=2, col="blue", bty='n', cex=0.8,
       legend=expression(bold('sd=1')))
legend("topright", title="Cauchy priors", y.intersp=0.8,
       col=c('red', 'red'),lty=c(1,2), cex=0.8,
         legend=c(expression(bold('medium')),
         expression(bold('ultrawide'))),bty="n")
mtext(side=3, line=0.25, adj=0, cex=1.15,
      expression("A: Alternative priors for "*delta==frac(mu,sigma)))
## Panel B
pairedDiffs <- with(datasets::sleep, extra[group==2] - extra[group==1])</pre>
ttBF0 <- BayesFactor::ttestBF(pairedDiffs)</pre>
simpost <- BayesFactor::posterior(ttBF0, iterations=10000)</pre>
plot(density(simpost[,'mu']), main="", xlab="", col="red",
     mgp=c(2,0.5,0), ylab="Posterior density", fg='gray')
mtext(side=1, line=2, expression(mu), cex=1.1)
abline(v=mean(pairedDiffs), col="gray")
mtext(side=3, line=0.5, expression("B: Posterior density for "*mu), adj=0, cex=1.15)
## Calculate and plot density for default prior - Selected lines of code
x \leftarrow seq(from=-4.5, to=4.5, by=0.1)
densMed <- dcauchy(x, scale=sqrt(2)/2)</pre>
plot(x, densMed, type='1')
## Panel B
pairedDiffs <- with(datasets::sleep, extra[group==2] - extra[group==1])</pre>
ttBF0 <- BayesFactor::ttestBF(pairedDiffs)</pre>
## Sample from posterior, and show density plot for mu
simpost <- BayesFactor::posterior(ttBF0, iterations=10000)</pre>
plot(density(simpost[,'mu']))
```

A thought experiment

```
tval <- setNames(qt(1-c(.05,.01,.005)/2, df=19), paste(c(.05,.01,.005)))
bf01 <- setNames(numeric(3), paste(c(.05,.01,.005)))
for(i in 1:3)bf01[i] <- BayesFactor::ttest.tstat(tval[i],n1=20, simple=T)

pairedDiffs <- with(datasets::sleep, extra[group==2] - extra[group==1])
ttBF0 <- BayesFactor::ttestBF(pairedDiffs)</pre>
```

```
pval <- t.test(pairedDiffs)[['p.value']]
1/(-exp(1)*pval*log(pval))</pre>
```

A null interval may make better sense

```
min45 <- round(0.75/sd(pairedDiffs),2) ## Use standardized units
ttBFint <- BayesFactor::ttestBF(pairedDiffs, nullInterval=c(-min45,min45))
round(as.data.frame(ttBFint)['bf'],3)</pre>
```

```
bf01 <- as.data.frame(ttBFint)[['bf']]</pre>
```

The effect of changing sample size

```
plabpos <- with(subset(bfDF, n==max(bfDF$n)), log((bf+bfInterval)/2))</pre>
gphA1 <- lattice::xyplot(log(bf)~log(n), groups=factor(p), data=bfDF,</pre>
                         panel=function(x,y,...){
                         lattice::panel.xyplot(x,y,type='b',...)})
ylabA <-10^{((-3):6/2)}
scalesA <- list(x=list(at=log(nval), labels=nval),</pre>
                y=list(at=log(ylabA), labels=signif(ylabA,2)))
keyA \leftarrow list(corner=c(0.99,0.98), lines=list(col=c(1,1), lty=1:2),
             text=list(c('Point null at 0', "null=(-0.1,0.1)")))
ylim2 \leftarrow log(c(min(bfDF[['bfInterval']])-0.05,150))
gphA2 <- lattice::xyplot(log(bfInterval)~log(n), groups=factor(p), lty=2,</pre>
  xlim=c(log(3.5),log(max(nval)*3.25)), ylim=ylim2, data=bfDF,
  panel=function(x,y,...){
    panel.xyplot(x,y,type='b',...)
    panel.grid(h=-1,v=-1)
    panel.text(rep(log(max(nval*0.975)),3), plabpos,
      labels=c('p=0.05','0.01','0.001'), pos=4)
  },
  par.settings=DAAG::DAAGtheme(color=T),
  main="A: Bayes factor vs sample size",
  xlab="Sample size", ylab="Bayes factor", scales=scalesA, key=keyA)
## Panel B
bfDF[['eff']] = bfDF[["t"]]/sqrt(bfDF[['n']])
ylabB < 10^{((-3):2/3)}
scalesB= list(x=list(at=log(nval), labels=nval),
              y=list(at=log(ylabB), labels=signif(ylabB,2)))
keyB \leftarrow list(corner=c(0.98,0.975), lines=list(lty=1:3),
             points=list(pch=1:3), text=list(c('p=0.001','p=0.01','p=0.05')))
gphB <- xyplot(log(eff)~log(n), groups=log(p), data=bfDF, pch=1:3, lty=1:3,</pre>
               type='b', xlab="Sample size", ylab="Effect size",
               par.settings=DAAG::DAAGtheme(color=T),
  main="B: Effect size vs sample size", key=keyB, scales=scalesB) +
  latticeExtra::layer(panel.grid(h=-1,v=-1))
```

```
plot(gphA2+latticeExtra::as.layer(gphA1), position=c(0, 0, 0.525, 1), more=T)
plot(gphB, position=c(0.52, 0, 1, 1), par.settings=DAAG::DAAGtheme(color=T))
```

Different statistics give different perspectives

```
n1 <- BayesFactor::ttest.tstat(qt(0.00001, df=40), n1=40, simple=T)
n2 <- BayesFactor::ttest.tstat(qt(0.000001, df=40), n1=40, simple=T)

bf1 <- BayesFactor::ttest.tstat(qt(0.00001, df=40), n1=40, simple=T)
bf2 <- BayesFactor::ttest.tstat(qt(0.000001, df=40), n1=40, simple=T)
rbind("Bayes Factors"=setNames(c(bf1,bf2), c("p=0.00001", "p=0.000001")),
    "t-statistics"=c(qt(0.00001, df=40), qt(0.000001, df=40)))

knitr::kable(matrix(c("A bare mention", "Positive", "Strong", "Very strong"), nrow=1),
    col.names=c("1 -- 3", "3 -- 20", "20 -- 150", ">150"), align='c',
    midrule='', vline="")
```

Section 1.8: Resampling methods for SEs, tests and confidence intervals

Subsection 1.8.1: The one-sample permutation test

```
tab <- t(as.matrix(DAAG::pair65))
rbind(tab,"heated-ambient"=tab[1,]-tab[2,])</pre>
```

Subsection 1.8.2: The two-sample permutation test

```
## First of 3 curves; permutation distribution of difference in means
two65 <- DAAG::two65
set.seed(47)  # Repeat curves shown here
nsim <- 2000; dsims <- numeric(nsim)
x <- with(two65, c(ambient, heated))
n <- length(x); n2 <- length(two65$heated)
dbar <- with(two65, mean(heated)-mean(ambient))
for(i in 1:nsim){</pre>
```

^{*}Technical details of the family of priors used in the BayesFactor package

```
mn <- sample(n,n2,replace=FALSE); dsims[i] <- mean(x[mn]) - mean(x[-mn]) }</pre>
plot(density(dsims), xlab="", main="", lwd=0.5, yaxs="i", ylim=c(0,0.08), bty="n")
abline(v=c(dbar, -dbar), lty=3)
pval1 <- (sum(dsims >= abs(dbar)) + sum (dsims <= -abs(dbar)))/nsim</pre>
mtext(side=3,line=0.25,
  text=expression(bar(italic(x))[2]-bar(italic(x))[1]), at=dbar)
mtext(side=3,line=0.25,
  text=expression(-(bar(italic(x))[2] - bar(italic(x))[1])), at=-dbar)
## Second permutation density
for(i in 1:nsim){
mn <- sample(n,n2,replace=FALSE)</pre>
dsims[i] \leftarrow mean(x[mn]) - mean(x[-mn])
pval2 <- (sum(dsims >= abs(dbar)) + sum (dsims <= -abs(dbar)))/nsim</pre>
lines(density(dsims), lty=2, lwd=1)
## Third permutation density
for(i in 1:nsim){
mn <- sample(n,n2,replace=FALSE)</pre>
dsims[i] \leftarrow mean(x[mn]) - mean(x[-mn])
}
pval3 <- (sum(dsims >= abs(dbar)) + sum (dsims <= -abs(dbar)))/nsim</pre>
lines(density(dsims),lty=3,lwd=1.25)
box(col="gray")
leg3 <- paste(c(pval1,pval2,pval3))</pre>
legend(x=20, y=0.078, title="P-values are", cex=1, xpd=TRUE,
  bty="n", lty=c(1,2,3), lwd=c(1,1,1,1.25), legend=leg3, y.intersp=0.8)
```

Subsection 1.8.3: Estimating the standard error of the median: bootstrapping

```
## Bootstrap estimate of median of wren length: data frame cuckoos
wren <- subset(DAAG::cuckoos, species=="wren")[, "length"]
library(boot)
## First define median.fun(), with two required arguments:
## data specifies the data vector,
## indices selects vector elements for each resample
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 = 4999))</pre>
```

Subsection 1.8.4: Bootstrap estimates of confidence intervals

Bootstrap 95% confidence intervals for the median

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

The correlation coefficient

```
## Bootstrap estimate of 95% CI for `cor(chest, belly)`: `DAAG::possum`
corr.fun <- function(data, indices)
  with(data, cor(belly[indices], chest[indices]))
set.seed(29)
corr.boot <- boot(DAAG::possum, corr.fun, R=9999)</pre>
```

```
library(boot)
boot.ci(boot.out = corr.boot, type = c("perc", "bca"))
```

The bootstrap – parting comments

Section 1.9: Organizing and managing work, and tools that can assist

The RStudio Integrated Development Environment

Subsection 1.9.1: Reproducible reporting — the knitr package

Section 1.10: The changing environment for data analysis

Subsection 1.10.1: Models and machine learning

The limits of current machine learning systems

Traps in big data analysis

Of mice and machine learning — missing data values

Humans are not good intuitive statisticians

Subsection 1.10.2: Replicability is the definitive check

To what extent is published work replicable? What is the evidence?

Some major replication studies

Replicability in pre-clinical cancer research

Studies where there may be strong social influences

The scientific study of scientific processes

Would lowering the p-value threshold help?

Peer review at the study planning stage

Section 1.11: Further, or supplementary, reading

Exercises (1_12)

1.4

```
Animals <- MASS::Animals
manyMals <- rbind(Animals, sqrt(Animals), Animals^0.1, log(Animals))
manyMals$transgp <- rep(c("Untransformed", "Square root transform",
    "Power transform, lambda=0.1", "log transform"),
rep(nrow(Animals),4))
manyMals$transgp <- with(manyMals, factor(transgp, levels=unique(transgp)))
lattice::xyplot(brain~body|transgp, data=manyMals,
    scales=list(relation='free'), layout=c(2,2))</pre>
```

1.5

```
with(Animals, c(cor(brain,body), cor(brain,body, method="spearman")))
with(Animals, c(cor(log(brain),log(body)),
    cor(log(brain),log(body), method="spearman")))
```

```
usableDF <- DAAG::cuckoohosts[c(1:6,8),]
nr <- nrow(usableDF)</pre>
with(usableDF, {
  plot(c(clength, hlength), c(cbreadth, hbreadth), col=rep(1:2,c(nr,nr)))
  for(i in 1:nr)lines(c(clength[i], hlength[i]), c(cbreadth[i], hbreadth[i]))
  text(hlength, hbreadth, abbreviate(rownames(usableDF),8), pos=c(2,4,2,1,2,4,2))
})
1.10
## Take a random sample of 100 values from the normal distribution
x <- rnorm(100, mean=3, sd=5)
(xbar \leftarrow mean(x))
## Plot, against `xbar`, the sum of squared deviations from `xbar`
lsfun <- function(xbar) apply(outer(x, xbar, "-")^2, 2, sum)</pre>
curve(lsfun, from=xbar-0.01, to=xbar+0.01)
boxplot(avs, meds, horizontal=T)
1.15
x \leftarrow rpois(7, 78.3)
mean(x); var(x)
1.16
nvals100 <- rnorm(100)</pre>
heavytail \leftarrow rt(100, df = 4)
veryheavytail \leftarrow rt(100, df = 2)
boxplot(nvals100, heavytail, veryheavytail, horizontal=TRUE)
1.19
boxdists <- function(n=1000, times=10){</pre>
  df <- data.frame(normal=rnorm(n*times), t=rt(n*times, 7),</pre>
  sampnum <- rep(1:times, rep(n,times)))</pre>
  lattice::bwplot(sampnum ~ normal+t, data=df, outer=TRUE, xlab="",
                   horizontal=T)
}
```

```
a <- 1
form <- \mbox{rchisq}(1000,1)^a + \mbox{rchisq}(1000,25)^a + \mbox{rchisq}(1000,500)^a
lattice::qqmath(form, scales=list(relation="free"), outer=TRUE)
1.21
y \leftarrow rnorm(51)
ydep <- y1[-1] + y1[-51]
           # acf plots `autocorrelation function'(see Chapter 6)
acf(y)
acf(ydep)
1.24
ptFun <- function(x,N)pt(sqrt(N)*mean(x)/sd(x), df=N-1, lower.tail=FALSE)
simStat <- function(eff=.4, N=10, nrep=200, FUN)</pre>
    array(rnorm(n=N*nrep*length(eff), mean=eff), dim=c(length(eff),nrep,N)) |>
      apply(2:1, FUN, N=N)
pval <- simStat(eff=.4, N=10, nrep=200, FUN=ptFun)</pre>
# Suggest a power transform that makes the distribution more symmetric
car::powerTransform(pval) # See Subsection 2.5.6
labx \leftarrow c(0.0001, 0.001, 0.005, 0.01, 0.05, 0.1, 0.25)
bwplot(~I(pval^0.2), scales=list(x=list(at=labx^0.2, labels=paste(labx))),
       xlab=expression("P-value (scale is "*p^{0.2}*")") )
1.24a
pvalDF <- subset(df200, effsize==0.4 & N==10)$stat</pre>
plot(sort(pval^0.2), sort(pvalDF^0.2))
abline(0,1)
1.24c
## Estimated effect sizes: Set `FUN=effFun` in the call to `eff2stat()`
effFun <- function(x,N)mean(x)/sd(x)</pre>
  # Try: ' = \log(labx)' ; 'v = \log(labx)'
## NB also, Bayes Factors: Set `FUN=BFfun` in the call to `eff2stat()`
BFfun <- function(x,N)BayesFactor::ttest.tstat(sqrt(N)*mean(x)/sd(x),
```

A few very large Bayes Factors are likely to dominate the plots

n1=N, simple=T)

```
(\deg C \leftarrow \operatorname{setNames}(c(21,30,38,46),\operatorname{paste}(\operatorname{rep},1:4)))
1.27a
radonC <- tidyr::pivot_longer(MPV::radon, names_to='key',</pre>
                                 cols=names(degC), values_to='percent')
radonC$temp <- degC[radonC$key]</pre>
lattice::xyplot(percent ~ temp|factor(diameter), data = radonC)
matplot(scale(t(MPV::radon[,-1])), type="l", ylab="scaled residuals")
1.27d
radon.res <- aggregate(percent ~ diameter, data = radonC, FUN = scale,
   scale = FALSE)
1.30
diamonds <- ggplot2::diamonds
with(diamonds, plot(carat, price, pch=16, cex=0.25))
with(diamonds, smoothScatter(carat, price))
t2bfInterval \leftarrow function(t, n=10, rscale="medium", mu=c(-.1,.1)){}
     null0 <- BayesFactor::ttest.tstat(t=t, n1=n, nullInterval=mu,</pre>
                                           rscale=rscale,simple=TRUE)
alt0 <- BayesFactor::ttest.tstat(t=t, n1=n, nullInterval=mu, rscale=rscale,
                                    complement=TRUE, simple=TRUE)
alt0/null0
pval \leftarrow c(0.05, 0.01, 0.001); nval \leftarrow c(10, 40, 160)
bfDF <- expand.grid(p=pval, n=nval)</pre>
pcol <- 1; ncol <- 2; tcol <- 3
bfDF[,'t'] <- apply(bfDF,1,function(x){qt(x[pcol]/2, df=x[ncol]-1,
other <- apply(bfDF,1,function(x)</pre>
    c(BayesFactor::ttest.tstat(t=x[tcol], n1=x[ncol], rscale="medium",
                                  simple=TRUE),
      BayesFactor::ttest.tstat(t=x[tcol], n1=x[ncol], rscale="wide",
```

```
simple=TRUE),
## Now specify a null interval
    t2bfInterval(t=x[tcol], n=x[ncol], mu=c(-0.1,0.1),rscale="medium"),
   t2bfInterval(t=x[tcol], n=x[ncol], mu=c(-0.1,0.1),rscale="wide")
  ))
bfDF <- setNames(cbind(bfDF, t(other)),</pre>
    c('p','n','t','bf','bfInterval'))
df <- data.frame(d = with(datasets::sleep, extra[group==2] - extra[group==1]))</pre>
library(statsr)
BayesFactor::ttestBF(df$d, rscale=1/sqrt(2)) # Or, `rscale="medium"`
  # `rscale="medium"` is the default
bayes_inference(d, type='ht', data=df, statistic='mean', method='t', rscale=1/sqrt(2),
                alternative='twosided', null=0, prior family = "JZS")
  # Set `rscale=1/sqrt(2)` (default is 1.0)
 # as for BayesFactor; gives same BF
# Compare with `prior_family = "JUI"` (`"JZS"` is the default),
# with (if not supplied) default settings
bayes_inference(d, type='ht', data=df, statistic='mean', method='t',
                alternative='twosided', null=0, prior_family = "JUI")
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()</pre>
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))</pre>
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch1.R")
```

2 Chapter 2: Generalizing from models

Packages required (plus any dependencies)

DAAG MASS qra investr HistData BHH2 xtable BayesFactor boot zoo boot MCMCpack, Additionally, knitr and Hmisc are required in order to process the Rmd source file.

Section 2.1 Model assumptions

Subsection 2.1.1: Inferences are never assumption free

Subsection 2.1.2: Has account been taken of all relevant effects?

```
## Tabulate by Admit and Gender
byGender <- 100*prop.table(margin.table(UCBAdmissions, margin=1:2), margin=2)
round(byGender,1)
## Admission rates, by department
pcAdmit <- 100*prop.table(UCBAdmissions, margin=2:3)["Admitted", , ]</pre>
round(pcAdmit,1)
applied <- margin.table(UCBAdmissions, margin=2:3)</pre>
pcAdmit <- 100*prop.table(UCBAdmissions, margin=2:3)["Admitted", , ]</pre>
      byGender <- 100*prop.table(margin.table(UCBAdmissions,
      margin=1:2), margin=2)
dimnam <- dimnames(UCBAdmissions)</pre>
mfStats <- data.frame(Admit=c(pcAdmit[1,],pcAdmit[2,]),</pre>
  Applicants=c(applied[1,], applied[2,]),
  mf=factor(rep(dimnam[['Gender']],c(6,6)),
  levels=dimnam[['Gender']]), Department=rep(dimnam[["Dept"]],2))
xlim <- c(0, max(mfStats$Admit)*1.025)</pre>
vlim <- c(0, max(mfStats$Applicants)*1.075)</pre>
plot(Applicants~Admit, data=mfStats, type="h",lwd=2, xlim=xlim, ylim=ylim,
     fg="gray", cex.lab=1.2, col=rep(c("blue", "red"), rep(6,2)),
```

```
xlab="UCB Admission rates (%), 1973", ylab="Number of applicants")
pcA <- rbind(pcAdmit[1,], apply(pcAdmit,2, mean)+2, pcAdmit[2,], rep(NA,6))
pcA[2,3] \leftarrow pcA[2,3]+1
appA <- rbind(applied[1,], apply(applied,2, mean)+80,
              applied[2,], rep(NA,6))
deptNam <- dimnam[[3]]</pre>
for(j in 1: ncol(appA)) lines(pcA[,j], appA[,j], col="gray", lwd=0.8)
points(pcA[2,],appA[2,], pch=16, cex=1.1, col="white")
text(pcA[2,],appA[2,],deptNam, cex=0.85)
par(xpd=TRUE)
text(byGender[1,1:2], rep(par()$usr[4],2)+0.5*strheight("^"),
     labels=c("^","^"), col=c("blue","red"),cex=1.2,srt=180)
text(byGender[1,], par()$usr[4]+1.4*strheight("A"),
     labels=paste(round(byGender[1,],1)),cex=0.85)
text(byGender[1,1:2]+c(-3.5,3.5), rep(par() usr[4],2)+2.65*strheight("A"),
     labels=c("All males", "All females"), pos=c(4,2), cex=1.2)
par(xpd=FALSE)
abline(h=200*(0:4),col="lightgray",lty="dotted")
abline(v=20*(0:4),col="lightgray",lty="dotted")
legend("topleft", col=c('blue','red'),lty=c(1,1), lwd=0.75, cex=0.9,
       y.intersp=0.65, legend=c("Males", "Females"), bty="n")
```

```
## Calculate totals, by department, of males & females applying margin.table(UCBAdmissions, margin=2:3)
```

Subsection 2.1.3: The limitations of models

Subsection 2.1.4: Use the methodology that best suits the task in hand?

Section 2.2: t-statistics, binomial proportions, and correlations

Subsection 2.2.1: One- and two-sample t-tests

Subsection 2.2.2: A two-sample comparison

```
c('av1','sd1','n1','av2','sd2','n2','pooledsd'))
print(stats2, digits=4)
with(DAAG::two65, t.test(heated, ambient, var.equal=TRUE))
```

When is pairing helpful?

What if the standard deviations are unequal?

Subsection 2.2.3: The normal approximation to the binomial

Subsection 2.2.4: The Pearson or product-moment correlation

Section 2.3 Extra-binomial and extra-Poisson variation

```
maleDF <- data.frame(number=0:12, freq=unname(qra::malesINfirst12[["freq"]]))</pre>
N <- sum(maleDF$freq)</pre>
pihat <- with(maleDF, weighted.mean(number, freq))/12</pre>
probBin <- dbinom(0:12, size=12, prob=pihat)</pre>
rbind(Frequency=setNames(maleDF$freq, nm=0:12),
      binomialFit=setNames(probBin*N, nm=0:12),
      rawResiduals = maleDF$freq-probBin*N,
      SDbinomial=sqrt(probBin*(1-probBin)*N)) |>
  formatC(digits=2, format="fg") |> print(digits=2, quote=F, right=T)
set.seed(29)
rgres.plot(doBI, plot.type='all', type="QQ", main=""); box(col='white')
mtext(side=3, line=0.5, "A: Binomial model, Q-Q", adj=0, cex=1.25)
rqres.plot(doBI, plot.type='all', type="wp", main=""); box(col='white')
## Plots C, D, E, F: Set object name; set type="wp" (C, E, F), or "QQ" (D)
mtext(side=3, line=0.5, "B: Binomial, worm plot 1", adj=-0.05, cex=1.25)
rqres.plot(doBI, plot.type='all', type="wp", main=""); box(col='white')
mtext(side=3, line=0.5, "C: Binomial, worm plot 2", adj=-0.05, cex=1.25)
rqres.plot(doBB, plot.type='all', type="QQ", main="", ylab=''); box(col='white')
mtext(side=3, line=0.5, "D: BB model, Q-Q", adj=0, cex=1.25)
rqres.plot(doBB, plot.type='all', type="wp", main="", ylab=''); box(col='white')
mtext(side=3, line=0.5, "E: BB, worm plot 1", adj=0, cex=1.25)
rqres.plot(doBB, plot.type='all', type="wp", main="", ylab=''); box(col='white')
mtext(side=3, line=0.5, "F: BB, worm plot 2", adj=0, cex=1.25)
aicStat <- AIC(doBI, doBB)</pre>
rownames(aicStat) <-
  c(doBI="Binomial", doBB="Betabinomial")[rownames(aicStat)]
aicStat$dAIC <- with(aicStat, round(AIC-AIC[1],1))</pre>
aicStat
## Numbers of accidents in three months, with Poisson fit
machinists <- data.frame(number=0:8, freq=c(296, 74, 26, 8, 4, 4, 1, 0, 1))
N <- sum(machinists[['freq']])</pre>
lambda <- with(machinists, weighted.mean(number, freq))</pre>
fitPoisson <- dpois(0:8, lambda)*sum(machinists[['freq']])</pre>
rbind(Frequency=with(machinists, setNames(freq, number)),
```

formatC(digits=2, format="fg") |> print(quote=F, digits=2, right=T)

poissonFit=fitPoisson) |>

```
set.seed(23)
rqres.plot(doPO, plot.type='all', type="QQ", main=""); box(col='white')
## Repeat, changing the argument, for remaining plots
mtext(side=3, line=0.5, "A: Poisson, Q-Q plot", adj=0, cex=1.25)
rqres.plot(doPO, plot.type='all', type="wp", main="", ylab=''); box(col='white')
mtext(side=3, line=0.5, "B: Poisson, worm plot", adj=0, cex=1.25)
rqres.plot(doNBI, plot.type='all', type="wp", main="", ylab='')
mtext(side=3, line=0.5, "C: NBI, worm plot", adj=0, cex=1.25); box(col='white')
```

Subsection 2.3.2: *Technical details – extra-binomial or extra-Poisson variation

```
sigma <- exp(coef(doBB, "sigma"))
cat("Phi =", (1+12*sigma)/(1+sigma))

mu <- exp(coef(doNBI, "mu"))
sigma <- exp(coef(doNBI, "sigma"))
cat("Phi =", (1+sigma*mu))</pre>
```

Section 2.4 Contingency tables

```
# To agree with hand calculation below, specify correct=FALSE
chisq.test(degTAB, correct=FALSE)
```

The mechanics of the chi-squared test

An example where a chi-squared test may not be valid

```
## Engine man data
engineman <- matrix(c(5,3,17,85), 2,2)
chisq.test(engineman)</pre>
```

Rare and endangered plant species

```
fisher.test(engineman)

## Enter the data thus:
rareplants <- matrix(c(37,190,94,23, 59,23,10,141, 28,15,58,16), ncol=3,
    byrow=TRUE, dimnames=list(c("CC","CR","RC","RR"), c("D","W","WD")))

(x2 <- chisq.test(rareplants))</pre>
```

Examination of departures from a consistent overall row pattern

```
## Expected values
x2$expected

options(digits=2)
## Standardized residuals
residuals(x2)
```

Interpretation issues

Section 2.5 Issues for Regression with a single explanatory variable

Subsection 2.5.1: Iron slag example — check residuals with care!

```
leg <- c("A: Fitted line", "B: Residuals from line", "C: Variance check")
ord <- order(DAAG::ironslag[["magnetic"]])
ironslag <- DAAG::ironslag[ord,]
slagAlpha.lm <- lm(chemical~magnetic, data=ironslag)
resval <- residuals(slagAlpha.lm)
fitchem <- fitted(slagAlpha.lm)
sqrtabs2 <- sqrt(abs(resval))
plot(chemical~magnetic, xlab = "Magnetic", ylab = "Chemical",</pre>
```

```
pch = 1, data=ironslag, fg="gray")
lines(fitchem~ironslag[["magnetic"]])
mtext(side = 3, line = 0.25, leg[1], adj=-0.1, cex=0.925)
scatter.smooth(resval~ironslag[["magnetic"]], lpars=list(col="red"), span=0.8,
               xlab = "Magnetic", ylab = "Residual", fg="gray")
mtext(side = 3, line = 0.25, leg[2], adj = -0.1, cex=0.925)
scatter.smooth(sqrtabs2 ~ fitchem, lpars=list(col="red"), span=0.8,
xlab = "Predicted chemical", fg="gray",
ylab = expression(sqrt(abs(residual))))
mtext(side = 3, line = 0.25, leg[3], adj = -0.1, cex=0.8)
## Diagnostics from fit using loess()
leg2 <- c("D: Smooth, using loess()",</pre>
          "E: Residuals from smooth",
          "F: Variance check")
slag.loess <- loess(chemical~magnetic, data=ironslag, span=0.8)</pre>
resval2 <- slag.loess[["residuals"]]</pre>
fitchem2 <- slag.loess[["fitted"]]</pre>
sqrtabs2 <- sqrt(abs(resval2))</pre>
plot(chemical~magnetic, xlab = "Magnetic", ylab = "Chemical",
pch = 1, data=ironslag, fg="gray")
lines(fitchem2 ~ ironslag[["magnetic"]], col="red")
mtext(side = 3, line = 0.25, leg2[1], adj=-0.1, cex=0.925)
scatter.smooth(resval2~ironslag[["magnetic"]], span=0.8,
lpars=list(col="red"),
xlab = "Magnetic", ylab = "Residual", fg="gray")
mtext(side = 3, line = 0.25, leg2[2], adj = -0.1, cex=0.925)
scatter.smooth(sqrtabs2 ~ fitchem2, lpars=list(col="red"),
span=0.8, xlab = "Predicted chemical", fg="gray",
ylab = expression(sqrt(abs(residual))))
mtext(side = 3, line = 0.25, leg2[3], adj = -0.1, cex=0.925)
```

Subsection 2.5.2: The analysis of variance table

```
roller.lm <- lm(depression ~ weight, data=DAAG::roller)
anova(roller.lm)</pre>
```

Subsection 2.5.3: Outliers, influence, and robust regression

```
softbacks <- DAAG::softbacks
x <- softbacks[,"volume"]</pre>
y <- softbacks[,"weight"]</pre>
u \leftarrow lm(y \sim x)
yhat <- predict(u)</pre>
res <- resid(u)
r <- with(softbacks, cor(x, y))</pre>
xlim <- with(softbacks, range(volume))</pre>
xlim[2] \leftarrow xlim[2]+diff(xlim)*0.08
plot(y ~ x, xlab = "Volume (cc)", xlim=xlim,
data=softbacks, ylab = "Weight (g)", pch = 4,
ylim = range(c(y, yhat)), cex.lab=0.9, fg="gray")
abline(u\$coef[1], u\$coef[2], lty = 1)
bottomright <- par()$usr[c(2, 3)]</pre>
chw <- par()$cxy[1]</pre>
chh <- par()$cxy[2]</pre>
z <- summary(u)$coef</pre>
btxt <- c(paste("a =", format(round(z[1, 1], 1)),
" SE =", format(round(z[1, 2], 1))),
paste("b =", format(round(z[2, 1], 2)),
   SE =", format(round(z[2, 2], 2))))
legend(bottomright[1], bottomright[2],
legend=btxt, xjust=1, yjust=0, cex=0.8, bty="n")
softbacks.lm <- lm(weight ~ volume, data=DAAG::softbacks)</pre>
print(coef(summary(softbacks.lm)), digits=3)
plot(softbacks.lm, fg="gray",
caption = c("A: Residuals vs Fitted", "B: Normal Q-Q",
"C: Scale-Location", "", "D: Resids vs Leverage"))
```

Robust regression

Subsection 2.5.4: Standard errors and confidence intervals

Confidence intervals and tests for the slope

```
SEb <- coef(summary(roller.lm))[2, 2]
coef(roller.lm)[2] + qt(c(0.025,.975), 8)*SEb</pre>
```

SEs and confidence intervals for predicted values

```
## Code to obtain fitted values and standard errors (SE, then SE.OBS)
fit.with.se <- predict(roller.lm, se.fit=TRUE)</pre>
fit.with.se$se.fit
                                                                # SE
sqrt(fit.with.se[["se.fit"]]^2+fit.with.se$residual.scale^2) # SE.OBS
predict(roller.lm, interval="confidence", level=0.95)
predict(roller.lm, interval="prediction", level=0.95) # CI for a new observation
## Depression vs weight, with 95\% pointwise bounds for both
## the fitted line and predicted values
investr::plotFit(roller.lm, interval="both", col.conf="red", fg="gray")
mtext(side=3,line=0.75, "A: Lawn roller data", cex=1.2, adj=-0.25)
## Male child vs father height, Galton's data
galtonMales <- subset(HistData::GaltonFamilies, gender=="male")</pre>
galton.lm <- lm(childHeight~father, data=galtonMales)</pre>
investr::plotFit(galton.lm, interval="both", col.conf="red", hide=FALSE,
                 col=adjustcolor('black',alpha=0.5), fg="gray")
```

mtext(side=3,line=0.75, "B: Son vs father heights", cex=1.2, adj=-0.25)

Implications for design

```
panelci<-function(data,...)
{
    nrows<-list(...)$nrows
    ncols<-list(...)$ncols
    if(ncols==1)axis(2, lwd=0, lwd.ticks=1)
    if(ncols==1)axis(1, lwd=0, lwd.ticks=1) else
    axis(3, lwd=0, lwd.ticks=1)
    x<-data$stretch; y<-data$distance
    u <- lm(y ~ x)
    upred <- predict(u, interval="confidence")
    ci <- data.frame(fit=upred[,"fit"],lower=upred[,"lwr"], upper=upred[,"upr"])
    ord<-order(x)
    lines(x[ord], ci[["fit"]][ord], lty=1, lwd=2)</pre>
```

```
lines(lowess(x[ord], ci[["upper"]][ord]), lty=2, lwd=2, col="grey")
lines(lowess(x[ord], ci[["lower"]][ord]), lty=2, lwd=2, col="grey")
elastic1 <- DAAG::elastic1</pre>
elastic2 <- DAAG::elastic2</pre>
xy<-rbind(elastic2,elastic1)</pre>
nam <- c("Range of stretch 30-65 mm", "Range of stretch 42-54 mm")</pre>
trial<-rep(nam, c(dim(elastic2)[1],dim(elastic1)[1]))</pre>
xlim<-range(elastic2$stretch)</pre>
ylim<-range(elastic2$distance)</pre>
xy<-split(xy,trial)</pre>
xy<-lapply(1:length(xy),function(i){c(as.list(xy[[i]]), list(xlim=xlim,</pre>
ylim=ylim))})
names(xy) <- nam</pre>
DAAG::panelplot(xy,panel=panelci,totrows=1,totcols=2,
                 par.strip.text=list(cex=.9), oma=c(4,4,2.5,2), fg='gray')
mtext(side = 2, line = 3.35, "Distance moved (cm)", cex=1.1, las=0)
mtext(side=1,line=3,"Amount of stretch (mm)", cex=1.1)
```

Subsection 2.5.5: There are two regression lines!

```
## There are two regression lines!
pair65 <- DAAG::pair65
bothregs <- function(x=pair65[, "ambient"], y=pair65[, "heated"],
  xlab="Stretch (band at ambient)", ylab = "Stretch (heated band)", pch=16){
    plot(y ~ x, xlab = xlab, ylab = ylab, pch = pch, fg="gray")
    topleft \leftarrow par() susr[c(1, 4)] + c(0.5, -0.5) * par() scxy
    text(topleft[1], topleft[2], paste("r = ", round(cor(x, y), 2)), adj = 0)
    u1 \leftarrow lm(y \sim x)
    abline(u1$coef[1], u1$coef[2])
    u2 \leftarrow lm(x \sim y)
    abline( - coef(u2)[1]/coef(u2)[2], 1/coef(u2)[2], 1ty = 2)
}
bothregs()
mtext(side = 3, line = 0.5, "A", adj = 0)
bothregs(x=trees[, "Girth"], y=trees[, "Height"],
         xlab="Girth (in)", ylab <- "Height (ft)", pch=16)</pre>
mtext(side = 3, line = 0.5, "B", adj = 0)
```

An alternative to a regression line

Subsection 2.5.6: Logarithmic and Power Transformations

```
## Logarithmic and Power Transformations
DAAG::powerplot(expr="sqrt(x)", xlab="")
DAAG::powerplot(expr="x^0.25", xlab="", ylab="")
DAAG::powerplot(expr="log(x)", xlab="", ylab="")
DAAG::powerplot(expr="x^2")
DAAG::powerplot(expr="x^4", ylab="")
DAAG::powerplot(expr="exp(x)", ylab="")
```

General power transformations — Box-Cox and Yeo-Johnson

Subsection 2.5.7: General forms of nonlinear response

Subsection 2.5.8: Size and shape data – allometric growth

```
## Heart weight versus body weight, for 30 Cape fur seals.
g2.12 <- function()</pre>
cfseal <- DAAG::cfseal
x <- log(cfseal[,"weight"])</pre>
y <- log(cfseal[, "heart"])
ylim <- log(c(82.5,1100))
xlim < -log(c(17,180))
ylab <- "Heart weight (g, log scale)"</pre>
xlab <- "Body weight (kg, log scale)"</pre>
xtik < -c(20,40,80,160)
ytik <- c(100,200,400,800)
plot(x, y, xlab = xlab, ylab = ylab, axes = F, xlim =
xlim, ylim = ylim, pch = 16, cex=0.85, fg="gray", cex.lab=1.1)
axis(1, at = log(xtik), labels = paste(xtik), lwd=0, lwd.ticks=1)
axis(2, at = log(ytik), labels = paste(ytik), lwd=0, lwd.ticks=1)
box(col="gray")
form1 <- formula(y ~ x)</pre>
u <- lm(form1, data = cfseal)</pre>
abline(u$coef[1], u$coef[2])
usum <- summary(u)$coef</pre>
options(digits=3)
print(usum)
```

```
cwh <- par()$cxy
eqn <- paste("log y =", round(usum[1, 1], 2), " [",
round(usum[1, 2], 2), "] +", round(usum[2, 1], 3),
" [", round(usum[2, 2], 3), "] log x")
mtext(side=3, line=1.15, eqn, adj = 0.4, cex = 0.8)
mtext(side=3, line=0.25, "(Values in square brackets are SEs)", adj = 0.4, cex = 0.8)
}
g2.12()</pre>
```

The allometric growth equation

```
options(scipen=4)
cfseal.lm <- lm(log(heart) ~ log(weight), data=DAAG::cfseal)
print(coef(summary(cfseal.lm)), digits=4)</pre>
```

Section 2.6 Empirical assessment of predictive accuracy

Subsection 2.6.1: The training/test approach, and cross-validation

Cross-validation - a tutorial example

```
houseprices <- DAAG::houseprices
df <- DAAG::CVlm(houseprices, form.lm = formula(sale.price ~ area), m=3, printit=F, plotit=FALS
panelfun <- function(x,y,subscripts,groups, ...){</pre>
  lattice::panel.superpose(x,y,subscripts,groups, ...)
  lattice::panel.superpose(x,df[["cvpred"]],subscripts,groups,type="b", cex=0.5, ...)
gph <- lattice::xyplot(sale.price ~ area, groups=fold, data=df, pch=1:3, panel=panelfun)</pre>
parset <- DAAG::DAAGtheme(color=T, lty=1:3, pch=1:3, lwd=2)</pre>
keylist <- list(lines=TRUE, columns=3, between.columns=1.5, between=1, cex=0.85)
update(gph, par.settings=parset, auto.key=keylist)
set.seed(29)
                    # Generate results shown
rand <- sample(rep(1:3, length=15))</pre>
## sample() randomly permutes the vector of values 1:3
for(i in 1:3) cat(paste0(i,":"), (1:15)[rand == i],"\n")
houseprices <- DAAG::houseprices
row.names(houseprices) <- (1:nrow(houseprices))</pre>
DAAG::CVlm(houseprices, form.lm = formula(sale.price ~ area), plotit=FALSE)
```

```
## Estimate of sigma^2 from regression output
houseprices <- DAAG::houseprices
houseprices.lm <- lm(sale.price ~ area, houseprices)
summary(houseprices.lm)[["sigma"]]^2</pre>
```

Subsection 2.6.2: Bootstrapping in regression

```
houseprices <- DAAG::houseprices
houseprices.lm <- lm(sale.price ~ area, houseprices)
print(coef(summary(houseprices.lm)),digits=2)
houseprices.fn <-
  function (houseprices, index,
            statfun=function(obj)coef(obj)[2]){
            house.resample <- houseprices[index, ]</pre>
            house.lm <- lm(sale.price ~ area, data=house.resample)</pre>
            statfun(house.lm) # slope estimate for resampled data
            }
set.seed(1028)
                    # use to replicate the exact results below
library(boot)
                   # ensure that the boot package is loaded
## requires the data frame houseprices (DAAG)
(houseprices.boot <- boot(houseprices, R=999, statistic=houseprices.fn))
statfun1200 <- function(obj)predict(obj, newdata=data.frame(area=1200))</pre>
price1200.boot <- boot(houseprices, R=999, statistic=houseprices.fn,</pre>
statfun=statfun1200)
boot.ci(price1200.boot, type="perc") # "basic" is an alternative to "perc"
set.seed(1111)
library(boot)
par(las=0)
houseprices2.fn<-function (houseprices,index){</pre>
house.resample<-houseprices[index,]
house.lm<-lm(sale.price~area,data=house.resample)</pre>
houseprices$sale.price-predict(house.lm,houseprices)
# resampled prediction errors
houseprices <- DAAG::houseprices
```

Commentary

Section 2.7 One- and two-way comparisons

Subsection 2.7.1: One-way comparisons

```
BHH2::anovaPlot(tomato.aov)
```

The analysis of variance table

```
## Do analysis of variance calculations
anova(tomato.aov)
```

Other multiple comparison tests

Subsection 2.7.2: Regression versus qualitative comparisons – issues of power

```
gph <- DAAG::simulateLinear(alpha=0.6, seed=17, aspect='iso')
update(gph, par.settings=DAAG::DAAGtheme(color=FALSE, alpha=0.4))</pre>
```

Subsection 2.7.3: *Severe multiplicity — the false discovery rate

*Microarrays and alternatives — technical note

The false discovery rate (FDR)

```
coralPval <- DAAG::coralPval
pcrit <- c(0.05, 0.02, 0.01, 0.001)
under <- sapply(pcrit, function(x)sum(coralPval<=x))</pre>
```

```
expected <- pcrit*length(coralPval)</pre>
```

```
fdrtab <- data.frame(Threshold=pcrit, Expected=expected,
Discoveries=under, FDR=round(expected/under, 4))
print(xtable::xtable(fdrtab), include.rownames=FALSE, hline.after=FALSE)</pre>
```

```
fdr <- p.adjust(coralPval, method="BH")</pre>
```

```
fdrcrit <- c(0.05, 0.04, 0.02, 0.01)
under <- sapply(fdrcrit, function(x)sum(coralPval<=x))
setNames(under, paste(fdrcrit))</pre>
```

Subsection 2.7.4: Data with a two-way structure, i.e., two factors

```
par(fig=c(0.525,1,0,1), mgp=c(1.5,0.4,0))
lev <- c("F10", "NH4C1", "NH4NO3", "F10 +ANU843",</pre>
         "NH4Cl +ANU843", "NH4NO3 +ANU843")
rice <- within(DAAG::rice, trt <- factor(trt, levels=lev))</pre>
with(rice, interaction.plot(fert, variety, ShootDryMass, fg="gray",
     legend = FALSE, xlab="Fertiliser", cex.lab=0.95, mex=0.65))
xleg <- par()$usr[2]</pre>
yleg \leftarrow par() usr[4] - 0.72 * diff(par() usr[3:4])
leginfo <- legend(xleg, yleg, bty = "n", legend = levels(rice$variety),</pre>
                   col = 1, lty = 2:1, lwd=1, xjust = 1, cex = 0.8,
                   y.intersp=0.8)$rect
text(leginfo$left + 0.5 * leginfo$w, leginfo$top, " variety",
      adj = 0.5, cex = 0.8
mtext(side=3, line=0.65, cex=0.9, adj=-0.15, "B")
gph <- dotplot(trt ~ ShootDryMass, pch=1, cex=0.9, las=2,</pre>
               xlab="Shoot dry mass (g)", data=rice,
               panel=function(x,y,...){panel.dotplot(x,y,...)
                  av <- sapply(split(x,y),mean);</pre>
                  ypos <- unique(y)</pre>
                  lpoints(ypos~av, pch=3, col="gray40", cex=1.25)},
               main=list("A", cex=0.88, just="left", x=0.1, y=-0.7, font=1))
pars <- DAAG::DAAGtheme(fontsize=list(text=9, points=6), color=FALSE)</pre>
print(update(gph, scales=list(tck=0.5), par.settings=pars, aspect=0.9),
      position=c(-0.065,0.0,0.6,1), newpage=FALSE)
```

Subsection 2.7.5: Presentation issues

Section 2.8 Data with a nested variation structure

Subsection 2.8.1: Degrees of freedom considerations

Subsection 2.8.2: General multi-way analysis of variance designs

Section 2.9 Bayesian estimation – further commentary and approaches

Subsection 2.9.1: Bayesian estimation with normal priors and likelihood

Subsection 2.9.2: Further comments on Bayes Factors

The Sellke calibration upper limit

A note on the Bayesian Information Criterion

```
pval \leftarrow c(.05,.01,.001); np \leftarrow length(pval)
Nval \leftarrow c(4,6,10,20,40,80,160); nlen \leftarrow length(Nval)
## Difference in BIC statistics, interpreted as Bayes factor
t2BFbic <- function(p,N){t <- qt(p/2, df=N-1, lower.tail=FALSE)
                           \exp((N*\log(1+t^2/(N-1))-\log(N))/2))
bicVal <- outer(pval, Nval, t2BFbic)</pre>
## Bayes factor, calculated using BayesFactor::ttest.tstat()
t2BF \leftarrow function(p, N)\{t \leftarrow qt(p/2, df=N-1, lower.tail=FALSE)\}
           BayesFactor::ttest.tstat(t=t, n1=N, simple=TRUE, rscale = "medium")}
BFval <- matrix(nrow=np, ncol=nlen)</pre>
for(i in 1:np)for(j in 1:nlen) BFval[i,j] <- t2BF(pval[i], Nval[j])</pre>
cfVal <- rbind(BFval, bicVal)[c(1,4,2,5,3,6),]
dimnames(cfVal) <- list(</pre>
  paste(rep(pval,rep(2,np)), rep(c("- from ttest.tstat", "- from BIC"),np)),
         paste0(c("n=",rep("",nlen-1)),Nval))
round(cfVal,1)
```

Subsection 2.9.3: Bayesian regression estimation using the MCMCpack package

```
suppressPackageStartupMessages(library(MCMCpack))
roller.mcmc <- MCMCregress(depression ~ weight, data=DAAG::roller)
summary(roller.mcmc)</pre>
```

```
mat <- matrix(c(1:6), byrow=TRUE, ncol=2)
layout(mat, widths=rep(c(2,1.1),3), heights=rep(0.9,8))
    # NB: widths & heights are relative
plot(roller.mcmc, auto.layout=FALSE, ask=FALSE, col="gray", fg="gray")</pre>
```

Section 2.10: Recap

Regress y on x, or x on $\sim y$?

Section 2.11: Further reading

Exercises (2.12)

2.2

```
## UCBAdmissions is in the datasets package
## For each combination of margins 1 and 2, calculate the sum
UCBtotal <- apply(UCBAdmissions, c(1,2), sum)</pre>
```

2.2b

```
apply(UCBAdmissions, 3, function(x)(x[1,1]*x[2,2])/(x[1,2]*x[2,1]))
```

2.3

```
tabA <- array(c(30,30,10,10,15,5,30,10), dim=c(2,2,2))
tabB <- array(c(30,30,20,10,10,5,20,25), dim=c(2,2,2))
```

2.5

```
z.transform <- function(r) .5*log((1+r)/(1-r))
z.inverse <- function(z) (exp(2*z)-1)/(exp(2*z)+1)
possum.fun <- function(data, indices) {
   chest <- data$chest[indices]
   belly <- data$belly[indices]
   z.transform(cor(belly, chest))}
possum.boot <- boot::boot(DAAG::possum, possum.fun, R=999)
z.inverse(boot.ci(possum.boot, type="perc")$percent[4:5])
# The 4th and 5th elements of the percent list element
# hold the interval endpoints. See ?boot.ci</pre>
```

```
with(pressure, MASS::boxcox(pressure ~ I(1/(temperature+273))))
```

2.14

```
"funRel" <-
function(x=leafshape$logpet, y=leafshape$loglen, scale=c(1,1)){
    ## Find principal components rotation; see Subsection 9.1.2
    ## Here (unlike 9.1.2) the interest is in the final component
    xy.prc <- prcomp(cbind(x,y), scale=scale)
    b <- xy.prc$rotation[,2]/scale
    c(bxy = -b[1]/b[2])  # slope of functional equation line
}
## Try the following:
leafshape <- DAAG::leafshape
funRel(scale=c(1,1))  # Take x and y errors as equally important
    # Note that all lines pass through (mean(x), mean(y))</pre>
```

2.15

```
P <- rbind(
    c(1,0,0,0,0,0),
    c(.5,0,.5,0,0,0),
    c(0,.5,0,.5,0,0),
    c(0,0,.5,0,.5,0),
    c(0,0,0,0,.5,0,.5),
    c(0,0,0,0,0,0))
dimnames(P) <- list(0:5,0:5)
P</pre>
```

```
Markov <- function(N=15, initial.value=1, transition=P, stopval=NULL)
    {X <- numeric(N)
        X[1] <- initial.value + 1  # States 0:(n-1); subscripts 1:n
        n <- nrow(transition)
        for (i in 2:N){
            X[i] <- sample(1:n, size=1, prob=transition[X[i-1], ])
            if(length(stopval)>0)if(X[i] %in% (stopval+1)){X <- X[1:i]; break}}
        X - 1
}
# Set `stopval=c(0,5)` to stop when the player's fortune is $0 or $5</pre>
```

```
Pb <- rbind(
   Sun = c(Sun=0.6, Cloud=0.2, Rain=0.2),
   Cloud= c(0.2, 0.4, 0.4),
   Rain= c(0.4, 0.3, 0.3))
Pb</pre>
```

2.16b

```
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch2.R")
}</pre>
```

3 Chapter 3: Multiple linear regression

Packages required (plus any dependencies)

DAAG car MASS AICcmodavg leaps BayesFactor splines

Additionally, Hmisc and knitr are required in order to process the Rmd source file.

Section 3.1 Basic ideas: the allbacks book weight data

```
allbacks <- DAAG::allbacks # Place the data in the workspace
allbacks.lm <- lm(weight ~ volume+area, data=allbacks)
print(coef(summary(allbacks.lm)), digits=2)</pre>
```

```
xlim <- range(allbacks$volume)
xlim <- xlim+c(-.075,.075)*diff(xlim)
## Plot of weight vs volume: data frame allbacks (DAAG)
plot(weight ~ volume, data=allbacks, pch=c(16,1)[unclass(cover)],
lwd=1.25, xlim=xlim, fg="gray")
## unclass(cover) gives the integer codes that identify levels
## As text() does not accept the parameter data, use with()
## to specify the data frame.
with(allbacks, text(weight ~ volume, labels=paste(1:15), cex=0.75, offset=0.35,
pos=c(2,4)[unclass(cover)]))
legend(x='topleft', pch=c(16,1), legend=c("hardback ","softback"),
horiz=T, bty="n", xjust=0.5, x.intersp=0.75, )</pre>
```

```
## Correlations between estimates -- model with intercept
round(summary(allbacks.lm, corr=TRUE)$correlation, 3)
```

```
out <- capture.output(summary(allbacks.lm,digits=2))
cat(out[15:17], sep='\n')</pre>
```

```
## 5% critical value; t-statistic with 12 d.f.
qt(0.975, 12)
cat(out[5:7], sep='\n')
```

Subsection 3.1.1: A sequential analysis of variance table

```
anova(allbacks.lm)
```

Omission of the intercept term

```
## Show rows 1, 7, 8 and 15 only
model.matrix(allbacks.lm)[c(1,7,8,15),]
## NB, also, code that returns the data frame used
model.frame(allbacks.lm)

allbacks.lm0 <- lm(weight ~ -1+volume+area, data=allbacks)
print(coef(summary(allbacks.lm0)), digits=2)

## Correlations between estimates -- no intercept
print(round(summary(allbacks.lm0, corr=TRUE)$correlation, 3))</pre>
```

Subsection 3.1.2: Diagnostic plots

```
allbacks.lm13 <- lm(weight ~ -1+volume+area, data=allbacks[-13, ])
print(coef(summary(allbacks.lm13)), digits=2)</pre>
```

Section 3.2 The interpretation of model coefficients

Subsection 3.2.1: Times for Northern Irish hill races

```
oldpar <- par(fg='gray20',col.axis='gray20',lwd=0.5,col.lab='gray20')
nihr <- within(DAAG::nihills, {mph <- dist/time; gradient <- climb/dist})</pre>
nihr <- nihr[, c("time", "dist", "climb", "gradient", "mph")]</pre>
varLabs <- c("\ntime\n(hours)","\ndist\n(miles)","\nclimb\n(feet)",</pre>
             "\ngradient\n(ft/mi)", "\nmph\n(mph)")
smoothPars <- list(col.smooth='red', lty.smooth=2, lwd.smooth=0.5, spread=0)</pre>
car::spm(nihr, cex.labels=1.2, regLine=FALSE, col='blue',
         oma=c(1.95,3,4, 3), gap=.25, var.labels=varLabs, smooth=smoothPars)
title(main="A: Untransformed scales:", outer=TRUE,
adj=0, line=-1.0, cex.main=1, font.main=1)
## Panel B: Repeat with log(nihills) in place of nihills,
## and with variable labels suitably modified.
varLabs <- c("\ntime\n(log h)","\ndist\n(log miles)", "\nclimb\n(log feet)",</pre>
             "\ngradient\n(log ft/mi)", "\nmph\n(log mph)")
car::spm(log(nihr), regLine=FALSE, col="blue", oma=c(1.95,2.5,4, 2.5),
         gap=.25, var.labels=varLabs, smooth=smoothPars)
title("B: Logarithmic scales", outer=TRUE,
      adj=0, line=-1.0, cex.main=1, font.main=1)
par(oldpar)
```

What is special about logarithmic transformations?

Subsection 3.2.2: An equation that predicts dist/time

```
## Hold climb constant at mean on logarithmic scale
mphClimb.lm <- lm(mph ~ log(dist)+log(climb), data = nihr)
## Hold `gradient=climb/dist` constant at mean on logarithmic scale
mphGradient.lm <- lm(mph ~ log(dist)+log(gradient), data = nihr)
avRate <- mean(nihr$mph)
bClimb <- coef(mphClimb.lm)</pre>
```

```
constCl <- c(bClimb[1]+bClimb[3]*mean(log(nihr$climb)), bClimb[2])</pre>
bGradient <- coef(mphGradient.lm)</pre>
constS1 <- c(bGradient[1]+bGradient[3]*mean((log(nihr$climb/nihr$dist))),</pre>
             bGradient[2])
# Use `dist` and `climb` as explanatory variables
coef(mphClimb.lm)
# Use `dist` and `gradient` as explanatory variables
coef(mphGradient.lm)
opar \leftarrow par(mfrow=c(1,2), mgp=c(2.25,0.5,0), mar=c(3.6,4.1,2.1,1.6))
lineCols <- c("red", adjustcolor("magenta",0.4))</pre>
yaxlab<-substitute(paste("Minutes per mile (Add ", ym, ")"), list(ym=round(avRate,2)))</pre>
car::crPlots(mphClimb.lm, terms = . ~ log(dist), xaxt='n',
             xlab="Distance", col.lines=lineCols, ylab=yaxlab)
axis(2, at=4:7, labels=paste(4:7))
labx \leftarrow c(4,8,16,32)
axis(1, at=log(2^{(2:5)}), labels=paste(2^{(2:5)}))
box(col="white")
mtext("A: Hold climb constant at mean value", adj=0,
      line=0.8, at=0.6, cex=1.15)
car::crPlots(mphGradient.lm, terms = . ~log(dist), xaxt='n',
             xlab="Distance", col.lines=lineCols, ylab=yaxlab)
axis(1, at=log(2^{(2:5)}), labels=paste(2^{(2:5)}))
axis(2, at=4:7, labels=paste(4:7))
box(col="white")
mtext("B: Hold log(gradient) constant at mean", adj=0, line=0.8, at=0.6, cex=1.15)
par(opar)
summary(mphClimb.lm, corr=T)$correlation["log(dist)", "log(climb)"]
summary(mphGradient.lm, corr=T)$correlation["log(dist)", "log(gradient)"]
## Show the plots, with default captions
plot(mphClimb.lm, fg='gray')
plot(mphGradient.lm, caption=c('A: Resids vs Fitted', 'B: Normal Q-Q',
'C: Scale-Location', '', 'D: Resids vs Leverage'),
cex.caption=0.85, fg='gray')
```

Subsection 3.2.3: Equations that predict log(time)

```
lognihr <- setNames(log(nihr), paste0("log", names(nihr)))
timeClimb.lm <- lm(logtime ~ logdist + logclimb, data = lognihr)

print(coef(summary(timeClimb.lm)), digits=2)

timeGradient.lm <- lm(logtime ~ logdist + loggradient, data=lognihr)
print(coef(summary(timeGradient.lm)), digits=3)</pre>
```

Subsection 3.2.4: Book dimensions — the oddbooks dataset

```
oldpar <- par(fg='gray40',col.axis='gray20',lwd=0.5,col.lab='gray20')
## Code for Panel A
oddbooks <- DAAG::oddbooks
pairs(log(oddbooks), lower.panel=panel.smooth, upper.panel=panel.smooth,
      labels=c("log(thick)", "log(breadth)", "log(height)", "log(weight)"),
      gap=0.25, oma=c(1.95,1.95,4, 1.95), col='blue'
title(main="A: Columns from log(oddbooks)",
      outer=TRUE, adj=0, line=-1.0, cex.main=1.1, font.main=1)
## Panel B
oddothers <-
  with(oddbooks, data.frame(density = weight/(breadth*height*thick),
area = breadth*height, thick=thick, weight=weight))
pairs(log(oddothers), lower.panel=panel.smooth, upper.panel=panel.smooth,
labels=c("log(density)", "log(area)", "log(thick)", "log(weight)"),
gap=0.5, oma=c(1.95,1.95,4, 1.95), col='blue')
title("B: Add density & area; omit breadth & height",
outer=TRUE, adj=0, line=-1.0, cex.main=1.1, font.main=1)
par(oldpar)
lob3.lm <- lm(log(weight) ~ log(thick)+log(breadth)+log(height),</pre>
```

```
lob2.lm <- lm(log(weight) ~ log(thick)+log(breadth), data=oddbooks)
coef(summary(lob2.lm))</pre>
```

Subsection 3.2.5: Mouse brain weight example

```
## Regression of brainwt on lsize
summary(lm(brainwt ~ lsize, data = litters), digits=3)$coef
## Regression of brainwt on lsize and bodywt
summary(lm(brainwt ~ lsize + bodywt, data = litters), digits=3)$coef
```

Subsection 3.2.6: Issues for causal interpretation

Effects of lifestyle on health

The studies mostly agree. But what do they say?

Adjusting for confounders

Section 3.3 Choosing the model, and checking it out

Effects of lifestyle on health

The studies mostly agree. But what do they say?

Adjusting for confounders

```
oddbooks.lm <- lm((weight) ~ log(thick)+log(height)+log(breadth),
data=DAAG::oddbooks)
yterms <- predict(oddbooks.lm, type="terms")</pre>
```

Subsection 3.3.3: A more formal approach to the choice of transformation

```
## Use car::powerTransform
nihr <- within(DAAG::nihills, {mph <- dist/time; gradient <- climb/dist})
summary(car::powerTransform(nihr[, c("dist", "gradient")]), digits=3)

form <- mph ~ log(dist) + log(gradient)
summary(car::powerTransform(form, data=nihr))</pre>
```

The use of transformations — further comments

Subsection 3.3.4: Accuracy estimates, fitted values and new observations

```
## First 4 rows
print(ci_then_pi[1:4,], digits=2)
```

```
timeClimb2.lm <- update(timeClimb.lm, formula = . ~ . + I(logdist^2))
g3.10 <-
function(model1=timeClimb.lm, model2=timeClimb2.lm)
## Panel A
citimes <- predict(model1, interval="confidence")</pre>
ord <- order(citimes[,"fit"])</pre>
citimes <- citimes[ord,]</pre>
hat <- citimes[,"fit"]</pre>
pitimes <- predict(model1, newdata=lognihr, interval="prediction")[ord,]</pre>
logobs <- log(nihr[ord,"time"])</pre>
xtiks <- pretty(exp(hat))</pre>
ylim <- range(c(pitimes[,"lwr"], pitimes[,"upr"], logobs)-rep(hat,3))</pre>
logytiks <- pretty(ylim,5)</pre>
ytiks <- round(exp(logytiks),2)</pre>
xlim <- range(hat)</pre>
plot(hat, citimes[,"lwr"]-hat, type="n", xlab = "Time (fitted)",
ylab = "Difference from fit",
xlim=xlim, ylim = ylim, xaxt="n", yaxt="n", fg="gray")
mtext(side=3, line=0.75, adj=0, at=-2.0, "A: CIs and PIs: Mean, prediction")
mtext(side=4, line=1.25, "exp(Difference from fit)", las=0)
axis(1, at=log(xtiks), labels=paste(xtiks), lwd=0, lwd.ticks=1)
axis(2, at=logytiks, las=1, lwd=0, lwd.ticks=1)
axis(4, at=logytiks, labels=paste(ytiks), las=0, lwd=0, lwd.ticks=1)
points(hat, logobs-hat, pch=16, cex=0.65)
lines(hat, citimes[,"lwr"]-hat, col = "red")
lines(hat, citimes[,"upr"]-hat, col = "red")
lines(hat, pitimes[,"lwr"]-hat, col = "black")
lines(hat, pitimes[,"upr"]-hat, col = "black")
## Panel B
citimes2 <- predict(model2, interval="confidence")[ord,]</pre>
plot(hat, citimes[,"lwr"]-hat, type="n", xlab = "Time (fitted)",
ylab = "Difference from fit",
xlim=xlim, ylim = ylim, xaxt="n", yaxt="n", fg="gray")
mtext(side=3, line=0.75, adj=0, at=-2.0,
"B: CIs for fit, compare two models")
mtext(side=4, line=1.25, "exp(Difference from fit)", las=0)
axis(1, at=log(xtiks), labels=paste(xtiks), lwd=0, lwd.ticks=1)
axis(2, at=logytiks,las=1, lwd=0, lwd.ticks=1)
```

```
axis(4, at=logytiks, labels=paste(ytiks), las=0,, lwd=0, lwd.ticks=1)
points(hat, logobs-hat, pch=16, cex=0.65)
lines(hat, citimes[,"lwr"]-hat, col = "red")
lines(hat, citimes[,"upr"]-hat, col = "red")
hat2 <- citimes2[,"fit"]
lines(hat, citimes2[,"lwr"]-hat2, col = "blue", lty=2, lwd=1.5)
lines(hat, citimes2[,"upr"]-hat2, col = "blue", lty=2, lwd=1.5)
}</pre>
```

```
timeClimb2.lm <- update(timeClimb.lm, formula = . ~ . + I(logdist^2))</pre>
```

Subsection 3.3.5: Choosing the model — deaths from Atlantic hurricanes

```
oldpar <- par(fg='gray20',col.axis='gray20',lwd=0.5,col.lab='gray20')
hurric <- DAAG::hurricNamed[,c("LF.PressureMB", "BaseDam2014", "deaths")]</pre>
thurric <- car::powerTransform(hurric, family="yjPower")</pre>
transY <- car::yjPower(hurric, coef(thurric, round=TRUE))</pre>
smoothPars <- list(col.smooth='red', lty.smooth=2, lwd.smooth=1, spread=0)</pre>
car::spm(transY, lwd=0.5, regLine=FALSE, oma=rep(2.5,4), gap=0.5,
         col="blue", smooth=smoothPars, cex.labels=1)
par(oldpar)
modelform <- deaths ~ log(BaseDam2014) + LF.PressureMB</pre>
powerT <- car::powerTransform(modelform, data=as.data.frame(hurric),</pre>
                                family="yjPower")
summary(powerT, digits=3)
deathP <- with(hurric, car::yjPower(deaths, lambda=-0.2))</pre>
power.lm <- MASS::rlm(deathP ~ log(BaseDam2014) + LF.PressureMB, data=hurric)</pre>
print(coef(summary(power.lm)),digits=2)
## Use (deaths+1)^{(-0.2)} as outcome variable
```

Subsection 3.3.6: Strategies for fitting models — suggested steps

Diagnostic checks

Section 3.4 Robust regression, outliers, and influence

Subsection 3.4.1: Making outliers obvious — robust regression

```
hills2000 <- DAAG::hills2000[,c("dist", "climb", "time")]
varLabels <- c("\ndist\n(log miles)", "\nclimb\n(log feet)", "\ntime\n(log hours)")</pre>
smoothPars <- list(col.smooth='red', lty.smooth=2, lwd.smooth=1, spread=0)</pre>
hills2000 <- DAAG::hills2000[,c("dist", "climb", "time")]
varLabels <- c("\ndist\n(log miles)", "\nclimb\n(log feet)", "\ntime\n(log hours)")</pre>
car::spm(log(hills2000), smooth=smoothPars, regLine=FALSE, cex.labels=1.5,
var.labels = varLabels, lwd=0.5, gap=0.5, oma=c(1.95,1.95,1.95,1.95))
## Panel A
lhills2k.lm <- lm(log(time) ~ log(climb) + log(dist), data = hills2000)</pre>
plot(lhills2k.lm, caption="", which=1, fg="gray", col=adjustcolor("black", alpha=0.8))
mtext(side=3, line=0.75, "A: Least squares (lm) fit", adj=0, cex=1.1)
## Panel B
lhills2k.lqs <- MASS::lqs(log(time) ~ log(climb) + log(dist), data = hills2000)</pre>
reres <- residuals(lhills2k.lqs)</pre>
refit <- fitted(lhills2k.lqs)</pre>
big3 <- which(abs(reres) >= sort(abs(reres), decreasing=TRUE)[3])
plot(reres ~ refit, xlab="Fitted values (resistant fit)",
ylab="Residuals (resistant fit)", col=adjustcolor("black", alpha=0.8), fg="gray")
lines(lowess(reres ~ refit), col=2)
text(reres[big3] ~ refit[big3], labels=rownames(hills2000)[big3],
pos=4-2*(refit[big3] > mean(refit)), cex=0.8)
mtext(side=3, line=0.75, "B: Resistant (lqs) fit", adj=0, cex=1.1)
## Show only the 2nd diagnostic plot, i.e., a normal Q-Q plot
## plot(lhills2k.lm, which=2)
```

Outliers, influential or not, should be taken seriously

Subsection 3.4.2: Leverage, influence, and Cook's distance

* Leverage and the hat matrix — technical details

```
round(unname(hatvalues(timeClimb.lm)),2)
```

Influential points and Cook's distance

Dynamic graphics

```
## Residuals versus leverages
nihills <- DAAG::nihills
timeClimb.lm <- lm(log(time) ~ log(dist) + log(climb), data = nihills)
plot(timeClimb.lm, which=5, add.smooth=FALSE, ps=9, sub.caption="",
     cex.caption=1.1, fg="gray")
  ## The points can alternatively be plotted using
 ## plot(hatvalues(model.matrix(timeClimb.lm)), residuals(timeClimb.lm))
## Residuals versus leverages
plot(timeClimb.lm, which=5, add.smooth=FALSE)
## The points can alternatively be plotted using
## plot(hatvalues(model.matrix(timeClimb.lm)), residuals(timeClimb.lm))
## This code is designed to be evaluated separately from other chunks
with(nihills, scatter3d(x=log(dist), y=log(climb), z=log(time), grid=FALSE,
                        point.col="black", surface.col="gray60",
                        surface.alpha=0.2, axis.scales=FALSE))
with(nihills, Identify3d(x=log(dist), y=log(climb), z=log(time),
                labels=row.names(DAAG::nihills), minlength=8), offset=0.05)
## To rotate display, hold down the left mouse button and move the mouse.
## To put labels on points, right-click and drag a box around them, perhaps
## repeatedly. Create an empty box to exit from point identification mode.
```

Influence on the regression coefficients

*Additional diagnostic plots

```
## As an indication of what is available, try
car::influencePlot(allbacks.lm)
```

Section 3.5 Assessment and comparison of regression models

Subsection 3.5.1: *AIC, AICc, BIC, and Bayes Factors for normal theory regression models

The functions drop1() and add1()

```
## Obtain AIC or BIC using `drop1()` or `add1()`
n <- nrow(DAAG::litters)
drop1(mouse.lm, scope=~lsize)  # AIC, with/without `lsize`
drop1(mouse.lm, scope=~lsize, k=log(n))  # BIC, w/wo `lsize`
add1(mouse0.lm, scope=~bodywt+lsize)  # AIC, w/wo `lsize`, alternative</pre>
```

The use of Bayesfactor::ImBF to compare the two models

```
suppressPackageStartupMessages(library(BayesFactor))
bf1 <- lmBF(brainwt ~ bodywt, data=DAAG::litters)
bf2 <- lmBF(brainwt ~ bodywt+lsize, data=DAAG::litters)
bf2/bf1

## Relative support statistics
setNames(exp(-apply(infstats[,-1],2,diff)/2), c("AIC","AICc","BIC"))</pre>
```

Subsection 3.5.2: Using anova() to compare models — the ihills data

```
lognihr <- log(DAAG::nihills)
lognihr <- setNames(log(nihr), pasteO("log", names(nihr)))
timeClimb.lm <- lm(logtime ~ logdist + logclimb, data = lognihr)
timeClimb2.lm <- update(timeClimb.lm, formula = . ~ . + I(logdist^2))
print(anova(timeClimb.lm, timeClimb2.lm, test="F"), digits=4)

print(anova(timeClimb.lm, timeClimb2.lm, test="Cp"), digits=3)
## Compare with the AICc difference
sapply(list(timeClimb.lm, timeClimb2.lm), AICcmodavg::AICc)

form1 <- update(formula(timeClimb.lm), ~ . + I(logdist^2) + logdist:logclimb)
addcheck <- add1(timeClimb.lm, scope=form1, test="F")
print(addcheck, digits=4)</pre>
```

Subsection 3.5.3: Training/test approaches, and cross-validation

```
pf(mspe/trainVar, mspeDF, trainDF, lower.tail=FALSE)

scot.lm <- lm(logtime ~ logdist+logclimb, data=logscot)
signif(summary(scot.lm)[['sigma']]^2, 4)</pre>
```

Subsection 3.5.4: Further points and issues

Patterns in the diagnostic plots – are they more than hints?

```
{r 3_18, eval=F|
```

What is the scatter about the fitted response

Model selection and tuning risks

Generalization to new contexts requires a random sample of contexts

What happens if we do not transform the hillrace data?

Are "errors in x" an issue?

Section 3.6 Problems with many explanatory variables

Subsection 3.6.1: Variable selection issues

Variable selection - a simulation with random data

```
y <- rnorm(100)
## Generate a 100 by 40 matrix of random normal data
xx <- matrix(rnorm(4000), ncol = 40)
dimnames(xx)<- list(NULL, paste("X",1:40, sep=""))

## ## Find the best fitting model. (The 'leaps' package must be installed.)
xx.subsets <- leaps::regsubsets(xx, y, method = "exhaustive", nvmax = 3, nbest = 1)
subvar <- summary(xx.subsets)$which[3,-1]
best3.lm <- lm(y ~ -1+xx[, subvar])
print(summary(best3.lm, corr = FALSE))</pre>
```

```
## DAAG::bestsetNoise(m=100, n=40)
best3 <- capture.output(DAAG::bestsetNoise(m=100, n=40))
cat(best3[9:14], sep='\n')</pre>
```

The extent of selection effects – a detailed simulation:

```
oldpar <- par(fg='gray20',col.axis='gray20',lwd=0.5,col.lab='gray20')
set.seed(41)
library(splines)
DAAG::bsnVaryNvar(nvmax=3, nvar = 3:35, xlab="")
mtext(side=1, line=1.75, "Number selected from")</pre>
```

Cross-validation that accounts for the variable selection process

*Regularization approaches

par(oldpar)

Subsection 3.6.2: Multicollinearity

An example – compositional data

```
coxiteAll.lm <- lm(porosity ~ A+B+C+D+E+depth, data=coxite)</pre>
print(coef(summary(coxiteAll.lm)), digits=2)
coxiteAll.lm <- lm(porosity ~ A+B+C+D+E+depth, data=coxite)</pre>
coxite.hat <- predict(coxiteAll.lm, interval="confidence")</pre>
hat <- coxite.hat[,"fit"]</pre>
plot(porosity ~ hat, data=coxite, fg="gray", type="n", xlab="Fitted values",
vlab="Fitted values, with 95% CIs\n(Points are observed porosities)",
tc1=-0.35)
with(coxite, points(porosity ~ hat, cex=0.75, col="gray45"))
lines(hat, hat, lwd=0.75)
ord <- order(hat)</pre>
sebar <- function(x, y1, y2, eps=0.15, lwd=0.75){
lines(rep(x,2), c(y1,y2), lwd=lwd)
lines(c(x-eps,x+eps), rep(y1,2), lwd=lwd)
lines(c(x-eps,x+eps), rep(y2,2), lwd=lwd)
}
q <- ord[round(quantile(1:length(hat), (1:9)/10))]
for(i in q)sebar(hat[i], coxite.hat[i,"lwr"], coxite.hat[i,"upr"])
coxiteAll.lm <- lm(porosity ~ A+B+C+D+E+depth, data=coxite)</pre>
coxite.hat <- predict(coxiteAll.lm, interval="confidence")</pre>
hat <- coxite.hat[,"fit"]</pre>
## Pointwise confidence bounds can be obtained thus:
hat <- predict(coxiteAll.lm, interval="confidence", level=0.95)
```

Subsection 3.6.3: The variance inflation factor (VIF)

```
print(DAAG::vif(lm(porosity ~ A+B+C+D+depth, data=coxite)), digits=2)

b <- leaps::regsubsets(porosity ~ ., data=coxite, nvmax=4, method='exhaustive')
## The calculation fails for nvmax=5
inOut <- summary(b)[["which"]]
## Extract and print the coefficents for the four regressions
dimnam <- list(rep("",4),c("Intercept", colnames(coxite)[-7]))
cmat <- matrix(nrow=4, ncol=7, dimnames=dimnam)
for(i in 1:4)cmat[i,inOut[i,]] <- signif(coef(b,id=1:4)[[i]],3)
outMat <- cbind(cmat," "=rep(NA,4),
as.matrix(as.data.frame(summary(b)[c("adjr2", "cp", "bic")])))
print(signif(outMat,3),na.print="")</pre>
```

```
BC.lm <- lm(porosity ~ B+C, data=coxite)
print(signif(coef(summary(BC.lm)), digits=3))
car::vif(BC.lm)</pre>
```

```
## Diagnostic plots can be checked thus:
plot(BC.lm, eval=xtras)
```

Numbers that do not quite add up

```
coxiteR <- coxite
coxiteR[, 1:5] <- round(coxiteR[, 1:5])
coxiteR.lm <- lm(porosity ~ ., data=coxiteR)
print(coef(summary(coxiteR.lm)), digits=2)
print(DAAG::vif(lm(porosity ~ .-E, data=coxiteR)), digits=2)</pre>
```

Remedies for multicollinearity

Section 3.7 Errors in x

Measurement of dietary intake

Simulations of the effect of measurement error

Two explanatory variables, one measured without error – a simulation

^{*}Two explanatory variables

An arbitrary number of variables

*The classical error model versus the Berkson error model

Using missing value approaches to address measurement error

Section 3.8 Multiple regression models – additional points

```
coef(lm(area ~ volume + weight, data=allbacks))
b <- as.vector(coef(lm(weight ~ volume + area, data=allbacks)))
c("_Intercept_"=-b[1]/b[3], volume=-b[2]/b[3], weight=1/b[3])</pre>
```

Unintended correlations

Subsection 3.8.2: Missing explanatory variables

```
gaba <- DAAG::gaba
gabalong <- stack(gaba["30", -match('min', colnames(gaba))])
gabalong$sex <- factor(rep(c("male", "female", "all"), rep(2,3)),
levels=c("female", "male", "all"))
gabalong$treatment <- factor(rep(c("Baclofen", "No baclofen"), 3),
levels=c("No baclofen", "Baclofen"))
gph <- lattice::stripplot(sex~values, groups=treatment, data=gabalong,
panel=function(x,y,...){
lattice::panel.stripplot(x,y,...)
lattice::ltext(x,y,paste(c(3,9,15,7,22,12)), pos=1, cex=0.8))
}, auto.key=list(space="right", points=TRUE, cex=0.8))
bw9 <- list(fontsize=list(text=9, points=5),
cex=c(1.5,1.5), pch=c(1,16))</pre>
```

```
update(gph, par.settings=parset,
xlab=list("Average reduction: 30 min vs 0 min", cex=1.0),
scales=list(cex=1.0, tck=0.35))
```

Strategies

Subsection 3.8.3: Added variable plots

```
yONx.lm <- lm(logtime ~ logclimb, data=lognihr)</pre>
e_y0Nx <- resid(y0Nx.lm)</pre>
print(coef(yONx.lm), digits=4)
zONx.lm <- lm(logdist ~ logclimb, data=lognihr)</pre>
e_z0Nx <- resid(z0Nx.lm)</pre>
print(coef(yONx.lm), digits=4)
ey_x0Nez_x.lm <- lm(e_y0Nx ~ 0+e_z0Nx)
e_y0Nxz <- resid(ey_x0Nez_x.lm)</pre>
print(coef(ey_x0Nez_x.lm), digits=4)
oldpar <- par(fg='gray')</pre>
## Code for added variable plots
logtime.lm <- lm(logtime ~ logclimb+logdist, data=lognihr)</pre>
car::avPlots(logtime.lm, lwd=1, terms="logdist", fg="gray")
mtext(side=3, line=0.5, "A: Added var: 'logdist'", col="black", adj=0, cex=1.15)
car::avPlots(logtime.lm, lwd=1, terms="logclimb", fg="gray")
mtext(side=3, line=0.5, "B: Added var: 'logclimb'", col="black", adj=0, cex=1.15)
par(oldpar)
## One call to show both plots
car::avPlots(timeClimb.lm, terms=~.)
## Alternative code for first plot
plot(e_y0Nx ~ e_z0Nx)
```

```
plot(y0Nx.lm, which=1, caption="", fg="gray")
mtext(side=3, line=0.5, "A: From 'logtime' on 'logclimb'", adj=0, cex=0.85)
plot(z0Nx.lm, which=1, caption="", fg="gray")
mtext(side=3, line=0.5, "B: From 'logdist' on 'logclimb'", adj=0, cex=0.85)
plot(ey_x0Nez_x.lm, which=1, caption="", fg="gray")
mtext(side=3, line=0.5, "C: From AVP", adj=-0, cex=0.85)
```

Alternatives to Added Variable Plots

*Algebraic details

```
ab1 <- coef(y0Nx.lm)

ab2 <- coef(z0Nx.lm)

b2 <- coef(ey_x0Nez_x.lm)

b1 <- ab1[2] - b2*ab2[2]

a <- ab1[1] - b2*ab2[1]
```

```
coef(lm(logtime ~ logclimb + logdist, data=lognihr))
```

Subsection 3.8.4: Nonlinear methods – an alternative to transformation?

```
signif(coef(summary(nihr.nls0)),3)
```

```
nihr.nls <- nls(time ~ gamma + delta1*dist^alpha + delta2*climb.mi^beta,
start=c(gamma = .045, delta1 = .09, alpha = 1,
delta2=.9, beta = 1.65), data=nihr)
## plot(residuals(nihr.nls) ~ log(predict(nihr.nls)))</pre>
```

```
signif(coef(summary(nihr.nls)),3)
```

Section 3.9: Recap

Section 3.10: Further reading

Exercises (3.11)

3.1

```
cities.lm1 <- lm(POP1996 ~ have+POP1992, data=cities)
cities.lm2 <- lm(log(POP1996) ~ have+log(POP1992), data=cities)</pre>
```

3.8a

```
nihills.lm <- lm(time ~ dist+climb, data=DAAG::nihills)
nihillsX.lm <- lm(time ~ dist+climb+dist:climb, data=DAAG::nihills)
anova(nihills.lm, nihillsX.lm) # Use `anova()` to make the comparison
coef(summary(nihillsX.lm)) # Check coefficient for interaction term
drop1(nihillsX.lm)</pre>
```

3.11

```
log(time) ~ log(dist) + log(climb) ## lm model
time ~ alpha*dist + beta*I(climb^2) ## nls model
```

3.13

```
x1 <- runif(10)  # predictor which will be missing
x2 <- rbinom(10, 1, 1-x1)
  ## observed predictor, depends on missing predictor
y <- 5*x1 + x2 + rnorm(10,sd=.1)  # simulated model; coef of x2 is positive
y.lm <- lm(y ~ factor(x2))  # model fitted to observed data
coef(y.lm)
y.lm2 <- lm(y ~ x1 + factor(x2))  # correct model
coef(y.lm2)</pre>
```

3.16

```
bomData <- DAAG::bomregions2021
nraw.lqs <- MASS::lqs(northRain ~ SOI + CO2, data=bomData)
north.lqs <- MASS::lqs(I(northRain^(1/3)) ~ SOI + CO2, data=bomData)
plot(residuals(nraw.lqs) ~ Year, data=bomData)
plot(residuals(north.lqs) ~ Year, data=bomData)</pre>
```

3.17f

```
socpsych <- subset(DAAG::repPsych, Discipline=='Social')
with(socpsych, scatter.smooth(T_r.R~T_r.0))
abline(v=.5)</pre>
```

```
soc.rlm <- MASS::rlm(T_r.R~T_r.0, data=subset(socpsych, T_r.0<=0.5))
## Look at summary statistics
termplot(soc.rlm, partial.resid=T, se=T)</pre>
```

```
plot(soc.rlm)
```

```
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch3.R")
}</pre>
```

4 Chapter 4: Exploiting the linear model framework

Packages required (with dependencies)

 ${\rm DAAG}$ effects m
gcv splines scam MASS lattice Extra car WDI AICcmodav
g ggplot2 kable Extra qgam patchwork

Additionally, Hmisc and knitr are required in order to process the Rmd source file.

Note the use of the 'patchwork' package to make it easy to place two ggplot2 plots side by side.

Section 4.1 Levels of a factor – using indicator variables

Subsection 4.1.1: Example – sugar weight

```
sugar <- DAAG::sugar # Copy dataset 'sugar' into the workspace
## Ensure that "Control" is the first level
sugar[["trt"]] <- relevel(sugar[["trt"]], ref="Control")
options()[["contrasts"]] # Check the default factor contrasts
## If your output does not agree with the above, then enter
## options(contrasts=c("contr.treatment", "contr.poly"))</pre>
```

```
sugar.aov <- aov(weight ~ trt, data=sugar)
## To display the model matrix, enter: model.matrix(sugar.aov)
## Note the use of summary.lm(), not summary() or summary.aov()
round(signif(coef(summary.lm(sugar.aov)), 3), 4)

sem <- summary.lm(sugar.aov)$sigma/sqrt(3) # 3 results/trt
# Alternatively, sem <- 6.33/sqrt(2)
qtukey(p=.95, nmeans=4, df=8) * sem</pre>
```

Subsection 4.1.2: Different choices for the model matrix when there are factors

```
contrasts(sugar$trt) <- 'contr.sum'
sugarSum.aov <- aov(weight ~ trt, data = sugar)
round(signif(coef(summary.lm(sugarSum.aov)), 3),4)
dummy.coef(sugarSum.aov)</pre>
```

Factor contrasts - further details

```
contrasts(sugar$trt) <- "contr.sum"

fish <- factor(1:3, labels=c("Trout","Cod","Perch"))

contr.treatment(fish)
# Base is "Trout"

contr.SAS(fish)
# Base is "Perch"

contr.sum(fish)
# Base is mean of levels</pre>
```

^{*}Tests for main effects in the presence of interactions?

Section 4.2 Block designs and balanced incomplete block designs

Subsection 4.2.1: Analysis of the rice data, allowing for block effects

```
rice <- DAAG::rice
ricebl.aov <- aov(ShootDryMass ~ Block + variety * fert, data=rice)
print(summary(ricebl.aov), digits=3)

round(signif(coef(summary.lm(ricebl.aov)), 3), 5)
with(summary.lm(ricebl.aov),
cat("Residual standard error: ", sigma, "on", df[2], "degrees of freedom"))

## AOV calculations, ignoring block effects
rice.aov <- aov(ShootDryMass ~ variety * fert, data=rice)
summary.lm(rice.aov)$sigma

ricebl.aov <- aov(ShootDryMass ~ factor(Block) + variety * fert, data=rice)

model.tables(ricebl.aov, type="means", se=TRUE, cterms="variety:fert")</pre>
```

Subsection 4.2.2: A balanced incomplete block design

```
appletaste <- DAAG::appletaste
with(appletaste, table(product, panelist))

sapply(appletaste, is.factor) # panelist & product are factors
appletaste.aov <- aov(aftertaste ~ product + panelist, data=appletaste)
summary(appletaste.aov)

as.data.frame(effects::Effect("product", appletaste.aov, confidence.level=0.95))

## NB that 'product' was first term in the model formula
## Thus, the 1st 4 coefficients have the information required
coef(summary.lm(appletaste.aov))[1:4,]</pre>
```

Section 4.3 Fitting multiple lines

Section 4.4 Methods for fitting smooth curves

print(coef(summary(leaf.lm3)), digits=2)

Subsection 4.4.1: Polynomial Regression

```
seedrates <- DAAG::seedrates
form2 <- grain ~ rate + I(rate^2)
# Without the wrapper function I(), rate^2 would be interpreted
# as the model formula term rate:rate, and hence as rate.
quad.lm2 <- lm(form2, data = seedrates)
## Alternative, using gam()
## quad.gam <- mgcv::gam(form2, data = seedrates)</pre>
```

suppressPackageStartupMessages(library(ggplot2))

*An alternative formulation using orthogonal polynomials

ggln <- ggplot_build(gph+geom_smooth(method=lm,</pre>

```
seedratesP.lm2 <- lm(grain ~ poly(rate,2), data = seedrates)
print(coef(summary(seedratesP.lm2)), digits=2)

## Alternative, using mgcv::gam()
seedratesP.gam <- mgcv::gam(grain ~ poly(rate,2), data = seedrates)

logseed.lm <- lm(log(grain) ~ log(rate), data=DAAG::seedrates)
coef(summary(logseed.lm))

## Use ggplot2 functions to plot points, line, curve, & 95% CIs
## library(ggplot2)
gph <- ggplot(DAAG::seedrates, aes(rate,grain)) +
    geom_point(size=3, color="magenta")+xlim(c(25,185))
colors <- c("Loglinear"="gray40", "Quadratic"="red")</pre>
```

fullrange=TRUE))\$data[[2]]

ggdat <- ggplot_build(gph+geom_smooth(method=lm, formula=y~poly(x,2),</pre>

```
linewidth = 0.75) +
geom_ribbon(data=ggln, aes(x=x,y=exp(y), ymin=exp(ymin), ymax=exp(ymax),
            color="Loglinear"), fill=NA, linewidth=0.75, linetype=3,
            outline.type='both', show.legend=FALSE)+
  scale_color_manual(values=colors, aesthetics = "color")+
  coord_cartesian(expand=FALSE) +
  xlab("Seeding rate (kg/ha)") + ylab("Grains per head") +
  labs(title="A: Loglinear fit vs quadratic fit", color="Model") +
  guides(size='none',
         color = guide_legend(override.aes = list(fill="transparent") ) ) +
  theme(legend.position=c(.8,.78))
df <- data.frame(rate=rep(DAAG::seedrates$rate,2), res=c(resid(logseed.lm),</pre>
  log(DAAG::seedrates$grain)-log(fitted(quad.lm2))),
  Model=rep(c("Loglinear","Quadratic"),rep(nrow(DAAG::seedrates),2)))
## Assign to gphB rather than (as in text) plotting at this point
gphB <- ggplot(df, aes(x=rate, y=res, shape=Model,color=Model))+</pre>
geom_point(size=2.5) + scale_color_manual(values=colors) +
xlab("Seeding rate (kg/ha)") + ylab("Residuals on log scale") +
labs(title="B: Residuals") +
  guides(size='none',
         color = guide_legend(override.aes = list(fill="transparent") ) ) +
  theme(legend.position=c(.8,.78))
## Now take advantage of the magic of the 'patchwork' package
library(patchwork)
gphA+gphB
## detach("package:ggplot2")
aic <- AIC(quad.lm2, logseed.lm)
aic["logseed.lm",2] <- aic["logseed.lm",2] + sum(2*log(seedrates$grain))
round(aic,1)
seedrates<-DAAG::seedrates
quad.lm2 <- lm(grain ~ poly(rate, degree=2), data=seedrates)</pre>
ns.lm2 <- lm(grain ~ splines::ns(rate,df=2), data=seedrates)</pre>
tps.gam2 <- mgcv::gam(grain ~ s(rate, k=3, fx=T), data=seedrates)
mflist <- lapply(list(quad=quad.lm2, nsplines=ns.lm2, tps=tps.gam2), model.matrix)</pre>
mftab <- with(mflist, cbind(quad, nsplines, tps))</pre>
colnames(mftab) <- c("(Int)", "poly2.1", "poly2.2", "(Int)", "ns2.1", "ns2.2", "(Int)", "s3.
library(kableExtra)
```

 $geom_line(data = ggln, aes(x = x, y = exp(y), color="Loglinear"),$

GAM models versus models fitted using Im()

Alternative fits – what is the best choice?

```
## Load required packages
suppressPackageStartupMessages(library(splines))
suppressPackageStartupMessages(library(mgcv))

ohms.tp <- gam(kohms~s(juice, bs="tp"), data=fruitohms)
ohms.cs <- gam(kohms~s(juice, bs="cs"), data=fruitohms)
range(fitted(ohms.tp)-fitted(ohms.cs))

summary(ohms.tp)

summary(ohms.tpBIC)</pre>
```

Subsection 4.4.3: The contributions of basis curves to the fit

Subsection 4.4.4: Checks on the fitted model

```
## Printed output from `gam.check(ohms.tpBIC)`
cat(out, sep="\n")
```

Subsection 4.3.5: Monotone curves

```
ohms.scam <- scam::scam(kohms ~ s(juice,bs="mpd"), data=fruitohms)
summary(ohms.scam)</pre>
```

```
AIC(ohms.scam, ohms.tp)
BIC(ohms.scam, ohms.tp)
```

Subsection 4.4.6: Different smooths for different levels of a factor

```
whiteside <- MASS::whiteside
gas.gam <- gam(Gas ~ Insul+s(Temp, by=Insul), data=whiteside)

summary(gas.gam)

Box.test(resid(gas.gam)[whiteside$Insul=='Before'], lag=1)
Box.test(resid(gas.gam)[whiteside$Insul=='After'], lag=1)</pre>
```

Subsection 4.4.7: The remarkable reach of mgcv and related packages

Departures from independence assumptions

Subsection 4.4.8: Multiple spline smoothing terms — dewpoint data

```
## GAM model -- `dewpoint` data
dewpoint <- DAAG::dewpoint
ds.gam <- gam(dewpt ~ s(mintemp) + s(maxtemp), data=dewpoint)
plot(ds.gam, resid=TRUE, pch=".", se=2, cex=2, fg="gray")</pre>
```

Using residuals as a check for non-additive effects

*A smooth surface

```
## Fit surface
ds.tp <- gam(dewpt ~ s(mintemp, maxtemp), data=DAAG::dewpoint)
vis.gam(ds.tp, plot.type="contour")  # gives a contour plot of the
# fitted regression surface
vis.gam(ds.gam, plot.type="contour")  # cf, model with 2 smooth terms</pre>
```

Subsection 4.4.9: Atlantic hurricanes that made landfall in the US

```
hurricNamed <- DAAG::hurricNamed
hurricS.gam <- gam(car::yjPower(deaths, lambda=-0.2) ~
  s(log(BaseDam2014)) + s(LF.PressureMB),
  data=hurricNamed, method="ML")
anova(hurricS.gam)</pre>
```

```
plot(hurricS.gam, resid=TRUE, pch=16, cex=0.5, select=1, fg="gray")
mtext(side=3, line=1, "A: Term in log(BaseDam2014)", cex=1.0, adj=0, at=-3.75)
plot(hurricS.gam, resid=TRUE, pch=16, cex=0.5, select=2, fg="gray")
mtext(side=3, line=1, "B: Term in LF.PressureMB", cex=1.0, adj=0, at=878)
qqnorm(resid(hurricS.gam), main="", fg="gray")
mtext(side=3, line=1, "C: Q-Q plot of residuals", cex=1.0, adj=0, at=-4.25)
```

An explanatory variable with an overly long-tailed distribution

```
hurricSlog1.gam <- gam(log(deaths+1) ~ s(log(BaseDam2014)), data=hurricNamed)
hurricSlog2.gam <- gam(log(deaths+1) ~ s(BaseDam2014), data=hurricNamed)</pre>
```

```
plot(hurricSlog1.gam, resid=TRUE, pch=16, cex=0.5, adj=0, fg="gray")
mtext(side=3, "A: Use log(BaseDam2014)", cex=1.4, adj=0, line=1, at=-3.15)
plot(hurricSlog2.gam, resid=TRUE, pch=16, cex=0.5, fg="gray")
mtext(side=3, "B: Use BaseDam2014", cex=1.4, adj=0, line=1, at=-28500)
```

Subsection 4.4.10: Other smoothing methods

Section 4.5 Quantile regression

2020 World Bank data on fertility and life expectancy

Section 4.6: Further reading and remarks

Exercises (4.7)

4.2

```
roller.lm <- lm(depression~weight, data=DAAG::roller)
roller.lm2 <- lm(depression~weight+I(weight^2), data=DAAG::roller)</pre>
```

4.4

4.4a

```
parLines.lm <- lm(distance ~ 0+factor(car)+angle, data=toycars)</pre>
sepLines.lm <- lm(distance ~ factor(car)/angle, data=toycars)</pre>
4.4b
sepPol3.lm <- lm(distance ~ factor(car)/angle+poly(angle,3)[,2:3], data=toycars)</pre>
4.4c
sapply(list(parLines.lm, sepLines.lm, sepPol3.lm), AICcmodavg::AICc)
4.4e
setNames(sapply(list(parLines.lm, sepLines.lm, sepPol3.lm),
  function(x)summary(x)$adj.r.squared), c("parLines", "sepLines", "sepPol3"))
4,7
seedrates.lm <- lm(grain ~ rate + I(rate^2), data=seedrates)</pre>
seedrates.pol <- lm(grain ~ poly(rate,2), data=seedrates)</pre>
4.10a
geo.gam <- gam(thickness ~ s(distance), data=DAAG::geophones)</pre>
4.11
plot(DAAG::geophones$distance, acf(resid(geo.gam), lag.max=55)$acf)
Box.test(resid(geo.gam), lag=10)
Box.test(resid(geo.gam), lag=20)
Box.test(resid(geo.gam), lag=20, type="Ljung")
4.15
library(mgcv)
xy \leftarrow data.frame(x=1:200, y=arima.sim(list(ar=0.75), n=200))
df.gam \leftarrow gam(y \sim s(x), data=xy)
plot(df.gam, residuals=TRUE)
```

4.16

```
library(mgcViz)
ohms.tpBIC <- gam(kohms ~ s(juice, bs="tp"), data=fruitohms,
                  gamma=log(nrow(fruitohms))/2, method="REML")
ohms.gamViz <- mgcViz::getViz(ohms.tpBIC) # Convert to a `gamViz` object</pre>
g1 <- plot(sm(ohms.gamViz, 1)) # Graphics object for term 1 (of 1)</pre>
g1 + l_fitLine(colour = "red") + l_rug(mapping = aes(x=x, y=y), alpha = 0.4) +
     l_ciLine(mul = 2, colour = "blue", linetype = 2) + # Multiply SE by `mul`
     l_points(shape = 19, size = 1, alpha = 0.5)
4.16a
plot(sm(ohms.gamViz, 1), nsim = 20) + l_ciLine() + l_fitLine() + l_simLine()
4.16b
gam(Gas ~ Insul+s(Temp, by=Insul), data=whiteside) |>
   getViz() -> gas.gamViz
plot(sm(gas.gamViz,1), nsim = 20) + l_ciLine() + l_fitLine() + l_simLine()
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()</pre>
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))</pre>
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch4.R")
```

5 Chapter 5: Generalized linear models and survival analysis

Packages required (with dependencies)

DAAG car mgcv colorspace HistData gamlss dplyr tidyr MASS ggplot2 latticeExtra qgam VGAM survival HistData

Additionally, knitr is required in order to process the Rmd source file.

Section 5.1 Generalized linear models

Subsection 5.1.1: Linking the expected value to the covariate

```
## Simplified plot showing the logit link function
p <- (1:39)/40
logitp <- log(p/(1 - p))
plot(p, logitp, xlab = "Proportion", ylab = "logit(p)", type = "l", pch = 1)</pre>
```

```
par(las=0)
p <- seq(from=1, to=99, by=1)/100; n<- 150; eps=0.001
gitp \langle -\log(p/(1-p))\rangle
plot(p, gitp, xlab = "", ylab = "", type = "l", pch = 1,
las=1, xlim=0:1, xaxs="i", fg="gray")
mtext(side = 1, line = 1.75, expression("Proportion "*pi))
mtext(side = 2, line = 1.75,
expression("logit("*pi*") = log(Odds)"))
mtext(side = 3, line = 0.5, "A: Logit link", adj=0, cex=1.0)
pval \leftarrow c(0.001, 0.01, 0.1, 0.5, 0.9, 0.99, 0.999)
par(mgp = c(2.5, 0.5, 0))
## axis(1, at=c(0,1), lwd=0, labels=c(0,1), xpd=TRUE)
axis(4, adj=0.075, at = log(pval/(1 - pval)), las=1,
col="gray", labels = paste(pval), lwd=0, lwd.ticks=1)
seP \leftarrow sqrt(p*(1-p)/100)
plot(p, seP, xlab = "", ylab = "", type = "l", pch = 1,
```

```
las=1, xlim=0:1, xaxs="i", fg="gray")
## axis(1, at=c(0,1), lwd=0, labels=c(0,1), xpd=TRUE)
mtext(side = 1, line = 1.75, expression("Proportion "*pi))
mtext(side = 2, line = 2.25, expression("SD["*p*"], "*n*"=100"))
mtext(side = 3, line = 0.5,
expression("B: SD["*p*"], "*n*"=100"), adj=0, cex=1.0)
seLP <- (p*(1-p)/n)*((p+eps)*(1-p+eps))^-2
plot(p, seLP, xlab = "", ylab = "", type = "l", pch = 1,
las=1, xlim=0:1, xaxs="i", fg="gray")
## axis(1, at=c(0,1), lwd=0, labels=c(0,1), xpd=TRUE)
mtext(side = 1, line = 1.75, expression("Proportion "*pi))
mtext(side = 2, line = 1.75, expression("SD[logit("*p*")], "*n*"=100"))
mtext(side = 3, line = 0.5, "C: SD[logit(p)]", adj=0, cex=1.0)</pre>
```

Subsection 5.1.2: Noise terms need not be normal

Subsection 5.1.3: Variation that is greater than binomial or Poisson

Least squares versus logistic regression

Subsection 5.1.4: Log odds in contingency tables

Subsection 5.1.5: Logistic regression with a continuous explanatory variable

```
anestot <- aggregate(DAAG::anesthetic[, c("move","nomove")],
by=list(conc=DAAG::anesthetic$conc), FUN=sum)
## The column 'conc', because from the 'by' list, is then a factor.
## The next line recovers the numeric values
anestot$conc <- as.numeric(as.character(anestot[["conc"]]))
anestot$total <- apply(anestot[, c("move","nomove")], 1 , sum)
anestot$prop <- anestot$nomove/anestot$total</pre>
```

```
par(mgp=c(2.5,.5,0))
anesthetic <- DAAG::anesthetic
z <- table(anesthetic$nomove, anesthetic$conc)
tot <- apply(z, 2, sum)
prop <- z[2, ]/(tot)
oprop <- sum(z[2, ])/sum(tot)
conc <- as.numeric(dimnames(z)[[2]])</pre>
```

```
DAAG::sumry(anes.glm, digits=2)
```

A note on model output

```
## Tp get coefficients, SEs, and associated statistics, specify:
print(coef(summary(anes.glm)), digits=2)
## Get full default output
summary(anes.glm, digits=2)
```

Section 5.2 Logistic multiple regression

```
frogs <- DAAG::frogs

## Presence/absence information: data frame frogs (DAAGS)
suppressMessages(library(ggplot2))
p <- ggplot(frogs, aes(easting, northing)) +
  geom_point(size=3, alpha=0.25) + coord_fixed() +
  xlab("Meters east of reference point")+ylab("Meters north") +</pre>
```

Implications for the Variance Inflation Factor

Subsection 5.2.1: Choose explanatory terms, and fit model

```
## Find power transformations
useCols <- c('distance','NoOfPools','NoOfSites','avrain','maxAddmin','maxSubmin')</pre>
tfrogs <- car::powerTransform(frogs[,useCols], family="yjPower")</pre>
## Create, for later use, a matrix with variables transformed as suggested
transY <- car::yjPower(frogs[,useCols], coef(tfrogs, round=TRUE))</pre>
summary(tfrogs, digits=2)
frogs0.glm <- glm(formula = pres.abs ~ log(distance) + log(NoOfPools)+</pre>
                  sqrt(NoOfSites) + avrain + maxAddmin + maxSubmin,
                  family = binomial, data = frogs)
DAAG::sumry(frogs0.glm, digits=1)
## Check effect of omitting sqrt(NoOfSites) and avrain from the model
## ~ . takes the existing formula. Precede terms to be
## omitted by '-'. (For additions, precede with '+')
frogs.glm <- update(frogs0.glm, ~ . -sqrt(NoOfSites)-avrain)</pre>
frogsAlt.glm <- update(frogs.glm, ~ . -maxAddmin+altitude)</pre>
AIC(frogs0.glm, frogs.glm,frogsAlt.glm)
'frogs0.glm'=coef(frogs0.glm)[c('log(distance)','log(NoOfPools)','maxAddmin','maxSubmin')],
'frogs.glm'=coef(frogs.glm)[c('log(distance)','log(NoOfPools)','maxAddmin','maxSubmin')]
coef(frogsAlt.glm)[c('log(distance)','log(NoOfPools)','altitude','maxSubmin')]
```

```
coef(summary(frogs.glm))
```

Subsection 5.2.2: Fitted values

```
## Use of `predict()` and `fitted()` --- examples
fitted(frogs.glm) # Fitted values' scale of response
predict(frogs.glm, type="response") # Same as fitted(frogs.glm)
                                  # Scale of linear predictor
predict(frogs.glm, type="link")
## For approximate SEs, specify
predict(frogs.glm, type="link", se.fit=TRUE)
library(ggplot2)
frogs$Prob. <- fitted(frogs.glm)</pre>
frogs$presAbs <- factor(frogs$pres.abs)</pre>
p <- ggplot(frogs, aes(easting, northing, color=Prob.)) +</pre>
  geom_point(size=2, alpha=0.5) + coord_fixed() +
  xlab("Meters east of reference point")+ylab("Meters north") +
   theme(axis.title=element_text(size=9), axis.text=element_text(size=6))
p2 <- p+scale_color_gradientn(colours=colorspace::heat_hcl(10,h=c(0,-100),
                              1=c(75,40), c=c(40,80), power=1)) +
  guides(fill=guide_legend(title=NULL))
p2 + geom_point(data=subset(frogs, presAbs==1),
```

Subsection 5.2.3: Plots that show the contributions of explanatory variables

aes(easting, northing), alpha=1, shape=3, col="white", size=1)

Subsection 5.2.4: Cross-validation estimates of predictive accuracy

Subsection 5.2.5: Cholera deaths in London — 1849 to 1855

By air, or by water — the 1849 epidemic

```
ylabs=rep("Partial residual",3), terms='poly(poor_rate, 2)', fg="gray")
par(opar)
```

The 1854 epidemic — a natural experiment

Section 5.3 Logistic models for categorical data – an example

```
## Create data frame from multi-way table UCBAdmissions (datasets)
## dimnames(UCBAdmissions) # Check levels of table margins
UCB <- as.data.frame.table(UCBAdmissions["Admitted", , ], responseName="admit")
UCB$reject <- as.data.frame.table(UCBAdmissions["Rejected", , ])$Freq
UCB$Gender <- relevel(UCB$Gender, ref="Male")
## Add further columns total and p (proportion admitted)
UCB$total <- UCB$admit + UCB$reject
UCB$pAdmit <- UCB$admit/UCB$total</pre>
UCB.glm <- glm(pAdmit ~ Dept*Gender, family=binomial, data=UCB, weights=total)
## Abbreviated `anova() ` output:
anova(UCB.glm, test="Chisq") |>
capture.output() |> tail(8) |> (\(x)x[-c(2,3)])() |> cat(sep='\n')

round(signif(coef(summary(UCB.glm)),4), 3)
```

Section 5.4 Models for counts — poisson, quasipoisson, and negative binomial

Subsection 5.4.1: Data on aberrant crypt foci

```
sum(resid(ACF.glm, type="pearson")^2)/19
ACFq.glm <- glm(formula = count ~ endtime + I(endtime^2),
family = quasipoisson, data = DAAG::ACF1)
print(coef(summary(ACFq.glm)), digits=2)
sapply(split(residuals(ACFq.glm), DAAG::ACF1$endtime), var)
fligner.test(resid(ACFq.glm) ~ factor(DAAG::ACF1$endtime))
Subsection 5.4.2: Moth habitat example
## Number of moths by habitat: data frame DAAG::moths
moths <- DAAG::moths
tab <- rbind(Number=table(moths[, 4]),</pre>
             sapply(split(moths[, -4], moths$habitat), apply, 2, sum))
## Number of zero counts, by habitats
with(droplevels(subset(moths, A==0)), table(habitat))
library(lattice)
gph <- dotplot(habitat ~ A+P, data=DAAG::moths, xlab="Number of moths", outer=TRUE,</pre>
               strip=strip.custom(factor.levels=paste("Number of species",c("A","B"))),
               panel=function(x, y, ...){
panel.dotplot(x,y, pch=1, ...)
av <- sapply(split(x,y),mean)</pre>
ypos <- factor(names(av), levels=names(av))</pre>
lpoints(ypos~av, pch=3, col="gray45", cex=1.25)
},
key=list(text=list(c("Individual transects", "Mean")),
points=list(pch=c(1,3), cex=c(1,1.25), col=c("black","gray45")),
columns=2), scales=list(tck=0.5, alternating=1))
bw9 <- list(fontsize=list(text=9, points=5))</pre>
update(gph, par.settings=bw9)
Astats <- with(DAAG::moths, sapply(split(A, habitat),
function(x)c(Amean=mean(x),Avar=var(x))))
```

avlength <- with(DAAG::moths, sapply(split(meters, habitat), mean))</pre>

round(rbind(Astats, avlen=avlength),1)

A quasipoisson model

```
A.glm <- glm(A ~ habitat + log(meters), family=quasipoisson,
data=DAAG::moths)
DAAG::sumry(A.glm, digits=1)

subset(DAAG::moths, habitat=="Bank")

## Analysis with tighter convergence criterion
A.glm <- update(A.glm, epsilon=1e-10)
print(coef(summary(A.glm)), digits=2)

AfitSE <- predict(A.glm, se=TRUE)$se.fit
cfSE <- with(DAAG::moths, c(AfitSE[habitat=="Bank"],
range(AfitSE[habitat!="Bank"])))
round(setNames(cfSE, c("SEbank", "SEotherMIN", "SEotherMAX")), digits=2)</pre>
```

A more satisfactory choice of reference level

Subsection 5.4.3: Models with negative binomial errors

```
dframe <- data.frame(sigma1A =(Astats[2,]-Astats[1,])/Astats[1,]^2,
    sigma2A =(Astats[2,]-Astats[1,])/Astats[1,]^1,
    mu = Astats[1,],    habitat=colnames(Astats))
bw9 <- list(fontsize=list(text=9, points=5), pch=1:7)
xyplot(sigma1A+sigma2A ~ mu, groups=habitat, outer=TRUE,
    data=subset(dframe,habitat!="Bank"),
    par.settings=bw9, auto.key=list(columns=4),
    strip=strip.custom(factor.levels=paste("Model",c("NBI","NBII"))),
    xlab="Mean number of species A moths",
    ylab=expression("Estimate of "*sigma))</pre>
```

```
LR.test(mothsCon.lss, mothsVary.lss)
```

```
## mothsCon.lss <- gamlss(A ~ log(meters)+habitat,family=NBI(),data=noBank)
## summary(mothsCon.lss, type="qr") ## Main part of output</pre>
```

Diagnostic plots

```
plot(mothsCon.lss, panel=panel.smooth)
```

Use of the square root link function

```
Asqrt.lss <- gamlss(A ~ habitat + sqrt(meters), trace=FALSE,
family = NBI(mu.link='sqrt'), data = moths)
```

```
## Asqrt.lss <- gamlss(A ~ habitat + sqrt(meters),
## family = NBI(mu.link='sqrt'), data = moths)
## summary(Asqrt.lss, type="qr") ## Main part of output
out <- capture.output(summary(Asqrt.lss, digits=1))[-(3:10)]
cat(out, sep="\n")</pre>
```

Subsection 5.4.4: Negative binomial versus alternatives — hurricane deaths

Aside – a quasibinomial binomial fit

Negative binomial versus power transformed scale

Fit a negative binomial (NBI) model

Fit linear model to power transformed response

```
hurr.lm <- lm(car::yjPower(deaths,-0.2) ~ log(BaseDam2014), data=hurric[-56,])
## Use the following function to transform from power scale to log scale
powerTOlog <- function(z, lambda)log(lambda*z+1)/lambda
## Calculate fitted values, and transform to log(deaths+1) scale
hatPower <- powerTOlog(predict(hurr.lm), lambda=-0.2)
resPower <- log(hurric[-56,"deaths"]+1) - hatPower</pre>
```

```
table(sign(resPower))
```

Compare NBI and power transform fits with smoothed quantiles

```
xvar <- log(hurric$BaseDam2014)[-56]
plot(log(deaths+1) ~ log(BaseDam2014), data=hurric, xaxt="n", yaxt="n",
    cex=4, pch=".", fg="gray", col=adjustcolor("black",alpha.f=0.65),
    xlab="Damage, millions of US$ in 2014", ylab="Deaths")
axis(1, at=log(c(1,10,1000, 100000)),
    labels=paste(c(1,10,1000, 100000)), lwd=0, lwd.ticks=1)
axis(2, at=log(c(0,10,100,1000)+1),
    labels=paste(c(0,10,100,1000)), lwd=0, lwd.ticks=1)</pre>
```

```
## Negative binomial regression fitted values
hatNB <- fitted(hurrNB.gamlss)</pre>
lines(xvar, log(hatNB+1), col="blue", lty=2)
with(hurric, text(log(BaseDam2014)[56], log(deaths+1)[56], "Audrey", pos=3),
     cex=0.72)
## Show fit from power transform model
lines(xvar, hatPower, col="blue", lty=1)
## Show 68.8\% and 40.1\% fits from regression smooths
lines(hat68.8 ~ xvar, lty=2, col='red')
lines(hat40.9 ~ xvar, lty=1, col='red')
legend("topleft", col=rep(c('blue','red'),c(2,2)), lty=rep(2:1,2), cex=0.8,
       y.intersp=0.75, legend=c("Negative binomial fit", "Power transform fit",
                                 "68.8% quantile", "40.9% quantile"), bty="n")
mtext(side=3, "A: Deaths vs damage", line=0.5, cex=1.15, adj=0)
## Quantile-quantile plot -- negative binomial model
qqnorm(zres, main="", fg="gray", cex=0.5,
  col=adjustcolor("black",alpha.f=0.65)); qqline(zres, col=2)
mtext(side=3, "B: Q-Q plot", line=0.5, cex=1.15, adj=0)
## a) Fitted and empirical centiles from hurrNB.gamlss
pc <- t(centiles.split(hurrNB.gamlss, xvar=log(hurric$BaseDam2014)[-56],</pre>
   cent=c(5,10,25,50,75,90,95), xcut.points=log(c(150, 1500)),
   plot=FALSE))
rownames(pc) <- c("up to 150M", "150M to 1500M", "above 1500M")
round(pc,2)
hurrP.gamlss <- gamlss(car::yjPower(deaths, -0.2) ~ log(BaseDam2014), data=hurric)
## Fitted and empirical centiles from hurrP.gamlss
pc <- t(centiles.split(hurrP.gamlss, xvar=log(hurric$BaseDam2014),</pre>
cent=c(5,10,25,50,75,90,95),
xcut.points=log(c(150, 1500)), plot=FALSE))
rownames(pc) <- c("up to 150M", "150M to 1500M", "above 1500M")
round(pc,2)
```

Section 5.5 Fitting smooths

Subsection 5.5.1: Handedness of first-class cricketers in the UK

```
tab <- with(DAAG::cricketer, table(left,dead))
colnames(tab) <- c('live','dead')
tab <- cbind(addmargins(tab, margin=2), prop.table(tab, margin=1))
tab</pre>
```

```
library(mgcv)
library(latticeExtra)
DAAG::cricketer |> dplyr::count(year, left, name="Freq") -> handYear
names(handYear)[2] <- "hand"</pre>
byYear <- tidyr::pivot_wider(handYear, names_from='hand', values_from="Freq")
hand.gam <- gam(cbind(left,right) ~ s(year), data=byYear, family=binomial)
const <- attr(predict(hand.gam, type='terms'), "constant")</pre>
  ## `const` is the mean on the scale of the linear predictor
plot(hand.gam, shift=const, trans=function(x)exp(x)/(1+exp(x)), ylim=c(.05,.4),
     xlab="", ylab="Proportion lefhanded", rug=FALSE, fg="gray",
     main=list("Proportion lefthanded, with 2SE limits",font=1,cex=1.2))
  ## Add `const`, then apply inverse link function.
  ## Plots estimated proportions (i.e., on the scale of the response)
with(byYear, points(year, I(left/(left+right)), cex=0.8, col="gray50"))
leftrt.gam <- gam(Freq ~ hand + s(year, by=factor(hand)), data=handYear,</pre>
                  family=poisson)
leftrt.pred <- predict(leftrt.gam, se=T, type='response')</pre>
handYear <- cbind(handYear, as.data.frame(leftrt.pred))</pre>
col2 <- DAAG::DAAGtheme(color=T)$superpose.symbol$col[c(2,2,1)]
gph.key <- list(space="top", columns=3, lines=list(lty=c(1,2,1), lwd=2, col=col2),
                text=list(c("left",expression(4.4%*%"left"),"right")), cex=1.2)
gph <- xyplot(leftrt.pred$fit ~ year, groups=hand, ylab=list("Number born", cex=1.2),</pre>
              type="1", xlab="", data=handYear, key=gph.key, col=col2[c(3,1)], lwd=2)
gph1 <- xyplot(Freq~year, groups=hand, data=handYear, col=col2[c(3,1)])</pre>
gph2 <- xyplot(I(4.4*fit) ~ year, data=subset(handYear, hand=="left"),</pre>
               type="1", lty=2, lwd=2, col=col2[2])
update(gph+as.layer(gph1)+as.layer(gph2), par.settings=DAAG::DAAGtheme(color=TRUE),
       scales=list(cex=1.2))
```

Section 5.6 Additional notes on generalized linear models

Subsection 5.6.1: Residuals, and estimating the dispersion

Other choices of link function for binomial models

Quasi models — estimating the dispersion

Subsection 5.6.2: Standard errors and z- or t-statistics for binomial models

```
fac <- factor(LETTERS[1:4])
p <- c(73, 30, 11, 2)/500
n <- rep(500,4)
round(signif(coef(summary(glm(p ~ fac, family=binomial, weights=n))), 6), 6)</pre>
```

```
p \leftarrow c(0.001, 0.002, (1:99)/100, 0.998, 0.999)
for(i in 1:3){
link <- c("logit", "probit", "cloglog")[i]</pre>
fun <- make.link(link)$linkfun</pre>
x \leftarrow fun(p)
u <- glm(p ~ x, family=binomial(link=link), weights=rep(1000,103))
if (i==1)
plot(x, hatvalues(u), type="l", ylab="Leverage", xaxt="n", fg='gray',
yaxt="n",
ylim=c(0, 0.0425), yaxs="i", xlab="Fitted proportion") else {
phat <- predict(u, type="response")</pre>
lines(log(phat/(1-phat)), hatvalues(u), type="l",
col=c("black","black","gray")[i], lwd=0.75,
lty=c(1,2,1)[i])
}
}
pos=c(0.001,0.002, 0.005, 0.01,0.02,0.05,0.1,0.25,0.5,0.75,0.9,0.95,0.98,0.99, 0.995, 0.998,
sub1 \leftarrow seq(from=1, to=17, by=2)
sub3 \leftarrow seq(from=2, to=16, by=2)
axis(1, at=log(pos/(1-pos))[sub1], labels=paste(pos)[sub1],
cex.axis=0.7, lwd=0, lwd.ticks=1)
axis(3, at=log(pos/(1-pos))[sub3], labels=paste(pos)[sub3],
cex.axis=0.7, lwd=0, lwd.ticks=1)
axis(2, at=c(0,.01,.02,.03), cex.axis=.7, lwd=0, lwd.ticks=1)
legend("topleft", lty=c(1,2,1),
legend=c("logit link", "probit link", "cloglog link"),
col=c("black","black","gray"), bty="n", cex=0.8)
```

Section 5.7 Models with an ordered categorical or categorical response

```
library(VGAM)
inhaler \leftarrow data.frame(freq=c(99,76,41,55,2,13),
  choice=rep(c("inh1", "inh2"), 3),
  ease=ordered(rep(c("easy", "re-read", "unclear"), rep(2,3))))
inhaler1.vglm <- vglm(ease ~ 1, weights=freq, data=inhaler,</pre>
  cumulative(link="logitlink"), subset=inhaler$choice=="inh1")
inhaler2.vglm <- vglm(ease ~ 1, weights=freq, data=inhaler,</pre>
  cumulative(link="logitlink"), subset=inhaler$choice=="inh2")
## Inhaler 1
round(coef(summary(inhaler1.vglm)),3)
## Inhaler 2
round(coef(summary(inhaler2.vglm)),3)
inhaler.vglm <- vglm(ease ~ choice, weights=freq, data=inhaler,
cumulative(link="logitlink", parallel=FALSE))
round(coef(summary(inhaler.vglm)),3)
inhalerP.vglm <- vglm(ease ~ choice, weights=freq, data=inhaler,</pre>
cumulative(link="logitlink", parallel=TRUE))
round(coef(summary(inhalerP.vglm)),3)
pred <- predict(inhalerP.vglm, se.fit=TRUE, newdata=inhaler[1:2,])</pre>
colnames(pred$se.fit) <- paste("SE", colnames(pred$se.fit))</pre>
fitvals <- with(pred, cbind(fitted.values, se.fit))</pre>
colnames(fitvals) <- gsub('link', '', colnames(fitvals))</pre>
round(fitvals, 2)
d <- deviance(inhalerP.vglm) - deviance(inhaler.vglm)</pre>
## Refer to chi-squared distribution with 1 degree of freedom
c(Difference=d, "p-Value"=pchisq(3.416, df=1, lower.tail=FALSE))
```

Subsection 5.7.2: Loglinear Models

Section 5.8 Survival analysis

```
df \leftarrow data.frame(x0 = c(1, 5, 1, 2, 14, 10, 12, 19)*30,
x1 = c(46, 58, 85, 67, 17, 85, 18, 42)*30,
fail = c(1, 0, 0, 1, 1, 0, 0, 1)
plot(c(0, 2610), c(0.65, 8.15), type = "n",
xlab = "Days from beginning of study",
ylab = "Subject number", axes = F)
## mtext(side = 1, line = 2.5, "Days from beginning of study", adj = 0.5)
m <- dim(df)[1]
par(las=2)
axis(2, at = (1:m), labels = paste((m:1)), lwd=0, lwd.ticks=1)
par(las=1)
abline(v = 600, lty = 4, col="gray40")
abline(v = 2550)
mtext(side = 3, line = 0.5, at = c(600, 2550),
text = c("\nEnd of recruitment",
"\nEnd of study"), cex = 0.9)
lines(rep((0:8) * 300, rep(3, 9)), rep(c(-0.4, -0.2, NA), 9),
xpd = T)
mtext(side = 1, line = 1.0, at = (0:8) * 300,
text = paste((0:8) * 300), adj = 0.5)
chw <- par()$cxy[1]</pre>
xx \leftarrow as.vector(t(cbind(df[, 1], df[, 2] - 0.25 * chw,
rep(NA, m))))
yy <- as.vector(t(cbind(matrix(rep(m:1, 2), ncol = 2),</pre>
rep(NA, m))))
lines(as.numeric(xx), as.numeric(yy))
points(df[, 1], m:1, pch = 16)
text(df[, 1]-0.25*chw, m:1, paste(df[,1]), pos=1, cex=0.75)
fail <- as.logical(df$fail)</pre>
points(df[fail, 2], (m:1)[fail], pch = 15)
points(df[!fail, 2], (m:1)[!fail], pch = 0)
text(df[, 2]+0.25*chw, m:1, paste(df[,2]), pos=1, cex=0.75)
par(xpd=TRUE)
legend(0, 11.5, pch = 16, legend = "Entry", y.intersp=0.15)
legend(1230, 11.5, pch = c(15, 0),
legend = c("Dead", "Censored"), ncol=2, y.intersp=0.15)
```

Subsection 5.8.1: Analysis of the Aids2 data

```
str(MASS::Aids2, vec.len=2)
bloodAids <- subset(MASS::Aids2, T.categ=="blood")</pre>
bloodAids$days <- bloodAids$death-bloodAids$diag
bloodAids$dead <- as.integer(bloodAids$status=="D")</pre>
bloodAids <- subset(MASS::Aids2, T.categ=="blood")</pre>
bloodAids$days <- bloodAids$death-bloodAids$diag</pre>
bloodAids$dead <- as.integer(bloodAids$status=="D")</pre>
plot(survfit(Surv(days, dead) ~ sex, data=bloodAids),
     col=c(2,4), conf.int=TRUE, lty=1, fg="gray",
     xlab="Days from diagnosis", ylab="Survival probability")
legend("top", legend=levels(bloodAids$sex), lty=c(1,1),
       col=c(2,4), horiz=TRUE, bty="n")
## Pattern of censoring for male homosexuals
hsaids <- subset(MASS::Aids2, sex=="M" & T.categ=="hs")
hsaids$days <- hsaids$death-hsaids$diag
hsaids$dead <- as.integer(hsaids$status=="D")</pre>
table(hsaids$status,hsaids$death==11504)
hsaids <- subset(MASS::Aids2, sex=="M" & T.categ=="hs")
hsaids$days <- hsaids$death-hsaids$diag
hsaids$dead <- as.integer(hsaids$status=="D")</pre>
hsaids.surv <- survfit(Surv(days, dead) ~ 1, data=hsaids)
plot(hsaids.surv, col="gray", conf.int=F, tcl=-0.4, fg="gray")
par(new=TRUE)
plot(hsaids.surv,col=1, conf.int=F,mark.time=F, fg="gray",
xlab="Days from diagnosis", ylab="Estimated survival probabality")
chw <- par()$cxy[1]
chh <- par()$cxy[2]
surv <- hsaids.surv$surv</pre>
xval <- c(200,700,1400,1900)
hat <- approx(hsaids.surv$time, surv, xout=xval)$y
for(i in 1:2) arrows(xval[i], hat[i], 0, hat[i],
length=0.05, col="gray")
lines(rep(xval[1],2), hat[1:2], col="gray")
      lines(rep(xval[3],2), hat[3:4], col="gray")
```

```
## Offset triangle 1
chw \leftarrow par()$cxy[1]
lines(xval[c(1,2,1,1)]+650, hat[c(2,2,1,2)]+0.2,col="gray40")
xy1 \leftarrow c(mean(xval[c(1,1,2)]), mean(hat[c(1,2,2)]))
arrows(xy1[1], xy1[2], xy1[1]+650, xy1[2]+0.2, col="gray40", length=0.1)
text(xval[1]-0.1*chw+650, hat[1]+0.2,
paste(round(hat[1],2)), col="gray20",cex=0.75, adj=1)
text(xval[1]+650-0.1*chw, hat[2]+0.2,
paste(round(hat[2],2)), col="gray20",cex=0.75, adj=1)
text(mean(xval[1:2])+650, hat[2]+0.2-0.5*chh,
paste(round(diff(xval[1:2]))), col="gray20", cex=0.75)
text(xval[1]+650-0.5*chw, mean(hat[1:2]+0.2), paste(round(hat[1]-hat[2],3)),
srt=90, adj=0.5, col="gray20", cex=0.75)
```

Subsection 5.8.4: Hazard rates

Subsection 5.8.5: The Cox proportional hazards model

```
bloodAids.coxph <- coxph(Surv(days, dead) ~ sex, data=bloodAids)</pre>
print(summary(bloodAids.coxph), digits=6)
## Add `age` as explanatory variable
bloodAids.coxph1 <- coxph(Surv(days, dead) ~ sex+age, data=bloodAids)
bloodAids <- subset(MASS::Aids2,T.categ=="blood")</pre>
bloodAids <- within(bloodAids, {days <- death-diag
dead <- as.integer(status=="D")})</pre>
bloodAids.coxph <- coxph(Surv(days, dead) ~ sex, data = bloodAids)
plot(cox.zph(bloodAids.coxph), cex=0.75, bty="n")
box(col="gray")
cox.zph(bloodAids.coxph)
```

```
cricketer <- DAAG::cricketer</pre>
kia4.coxph <- coxph(Surv(life, kia) ~ left/poly(year,4),</pre>
                     data = cricketer, model=T)
kia6.coxph <- update(kia4.coxph, . ~ left/poly(year,6),</pre>
                      data = cricketer, model=T)
# Type `plot(cox.zph(kia6.coxph)` to plot the two graphs
```

```
# Perhaps check also `AIC(kia4.coxph, kia6.coxph)`
cox.zph(kia6.coxph)
```

```
plot(cox.zph(kia6.coxph), cex=0.75, bty="n")
box(col="gray")
```

Section 5.9: Transformations for proportions and counts

Section 5.10: Further reading

Exercises (5.11)

5.1

```
inhibition <- rbind(
conc =c(0.1,0.5, 1,10,20,30,50,70,80,100,150),
no = c(7, 1, 10, 9, 2, 9, 13, 1, 1, 4, 3),
yes = c(0, 0, 3, 4, 0, 6, 7, 0, 0, 1, 7)
)
colnames(inhibition) <- rep("", ncol(inhibition))
inhibition</pre>
```

```
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch5.R")
}</pre>
```

6 Chapter 6: Time series models

Packages required (plus any dependencies)

DAAG ggsci latticeExtra ggplot2 mice car forecast mgcv tseries

Additionally, knitr and Hmisc are required in order to process the Rmd source file.

Section 6.1: Time series – some basic ideas

Subsection 6.1.1: Time series objects

```
class("lakeHuron")

[1] "character"

## Use `time()` to extract the `time` attribute
range(time(LakeHuron))

[1] 1875 1972

## Use `window()` to subset a time series
LHto1925 <- window(LakeHuron, from=1875, to=1925)

jobs <- DAAG::jobs
names(jobs)

[1] "BC" "Alberta" "Prairies" "Ontario" "Quebec" "Atlantic" "Date"

allRegions <- ts(jobs[, -7]) # Create multivariate time series
time(allRegions) # Times run from 1 to 24</pre>
```

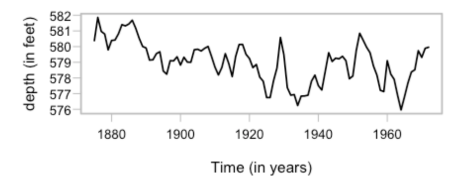
```
Time Series:
Start = 1
End = 24
Frequency = 1
[1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24
```

```
allRegions <- ts(jobs[, -7], start=c(1995,1), frequency=12)
allRegions[,"BC"] # Extract jobs data for British Columbia
```

Jan Feb Mar Apr May Jun Jul Aug Sep Oct Nov Dec 1995 1752 1737 1765 1762 1754 1759 1766 1775 1777 1771 1757 1766 1996 1786 1784 1791 1800 1800 1798 1814 1803 1796 1818 1829 1840

```
jobsBC <- ts(jobs[, "BC"], start=c(1995,1), frequency=12)
# Obtain equivalent of `allRegions[,"BC"]` directly from `jobs` dataset</pre>
```

Subsection 6.1.2: Preliminary graphical exploration



```
## Plot depth measurements: ts object LakeHuron (datasets)
plot(LakeHuron, ylab="depth (in feet)", xlab = "Time (in years)", fg="gray")
```

```
lag.plot(LakeHuron, lags=3, do.lines=FALSE)
```

Subsection 6.1.3: The autocorrelation and partial autocorrelation function

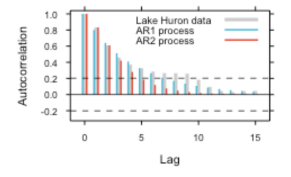
578

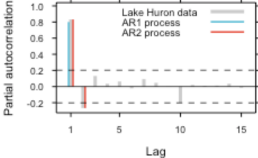
A: Lag plots 576 578 580 582 576 578 580 582 581 580 .akeHuron akeHuror 579 🚽 578 577 576 lag 2 lag 1 lag 3 lag 4

```
par(oma=c(0,0,1.5,0))
par(pty="s")
lag.plot(LakeHuron, set.lags=1:4,do.lines=F, oma=c(0,1.5,1.5,1.5),
fg="gray", layout=c(1,4), cex.lab=1.15, asp=1)
mtext(side=3, line=0.5, "A: Lag plots", adj=0, cex=0.85, outer=TRUE)
```

B: Autocorelation -- Data vs AR processes

C: Partial autocorrelation -- Data vs AR processes





578

580

582

```
library(lattice)
col3 <- c("gray80",rev(ggsci::pal_npg()(2)))
lag.max <- 15
offset <- 0.18</pre>
```

```
ci95 <- 2/sqrt(length(LakeHuron))</pre>
ar2 <- ar(LakeHuron)
gph.key < -list(x=0.975, y=0.965, corner=c(1,1), columns=1, cex=0.85,
                  text=list(c("Lake Huron data", "AR1 process", "AR2 process")),
                  lines=list(lwd=c(3,1.5,1.5), col=col3, lend=2),
                  padding.text=1)
parsetBC <- list(fontsize=list(text=8, points=5),</pre>
                  superpose.line=list(col=col3, lty=rep(1,3),
                  lwd=c(3,1.5,1.5))
parsetBC <- modifyList(parsetBC,list(grid.pars = list(lineend = "butt")))</pre>
lev3 <- factor(c("acfData","acfAR1","acfAR2"),</pre>
                levels=c("acfData","acfAR1","acfAR2"))
acfData <- acf(LakeHuron, main="", plot=FALSE, lag.max=lag.max)$acf
pacfData <- pacf(LakeHuron, main="", plot=FALSE, lag.max=lag.max)$acf</pre>
acfAR1 <- ARMAacf(ar=0.8, lag.max=lag.max)</pre>
acfAR2 <- ARMAacf(ar=ar2$ar, ma=0, lag.max=lag.max)</pre>
pacfAR1 <- ARMAacf(ar=0.8, lag.max=lag.max, pacf=TRUE)</pre>
pacfAR2 <- ARMAacf(ar=ar2$ar, ma=0, lag.max=lag.max, pacf=TRUE)</pre>
xy <- data.frame(acf=c(acfData,acfAR1,acfAR2),</pre>
Lag=c(rep(0:lag.max,3))+rep(c(0,-offset,offset),
rep(lag.max+1,3)),
gp=rep(lev3, rep(lag.max+1,3)))
gphB <- xyplot(acf ~ Lag, data = xy, groups=gp, type=c("h"),</pre>
               par.strip.text = list(cex = 0.85), lend=2, origin=0,
               ylim=c(-0.325, 1.04), key=gph.key, par.settings=parsetBC,
          panel=function(x,y,...){
            panel.xyplot(x,y,...)
            panel.abline(h=0, lwd=0.8)
            panel.abline(h=ci95, lwd=0.8, lty=2)
            panel.abline(h=-ci95, lwd=0.8, lty=2) } )
xyp <- data.frame(pacf=c(pacfData,pacfAR1,pacfAR2),</pre>
                   Lag=c(rep(1:lag.max,3))+c(rep(c(0,-offset,offset),
                   rep(lag.max,3))), gp=rep(lev3, rep(lag.max,3)))
gphC <- xyplot(pacf ~ Lag, data = xyp, groups=gp, type=c("h"),</pre>
                par.strip.text = list(cex = 0.85), lend=2,
                ylab = "Partial correlation", origin=0, ylim=c(-0.325, 1.04),
               key=gph.key, par.settings=parsetBC,
               panel=function(x,y,...){
                  panel.xyplot(x,y,...)
                  panel.abline(h=0, lwd=0.8)
                  panel.abline(h=ci95, lwd=0.8, lty=2)
                  panel.abline(h=-ci95, lwd=0.8, lty=2) } )
```

```
acf(LakeHuron)
## pacf(LakeHuron) gives the plot of partial autocorrelations
```

Subsection 6.1.4: Autoregressive (AR) models

The AR(1) model

```
## Yule-Walker autocorrelation estimate
LH.yw <- ar(LakeHuron, order.max = 1, method = "yw") # autocorrelation estimate
# order.max = 1 for AR(1) model
LH.yw$ar # autocorrelation estimate of alpha</pre>
```

[1] 0.8319112

[1] 0.837546

```
LH.mle$x.mean # estimated series mean
```

[1] 579.1141

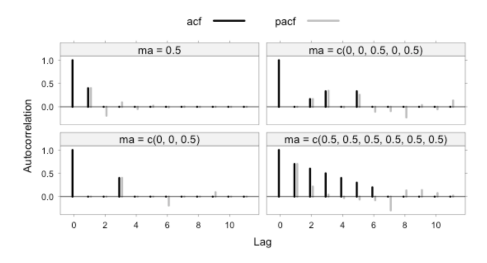
[1] 0.5092867

The general AR(p) model

```
ar(LakeHuron, method="mle")
```


Order selected 2 sigma^2 estimated as 0.4788

~Moving average (MA) processes



Subsection 6.1.5: ~Autoregressive moving average (ARMA) models – theory

Subsection 6.1.6: Automatic model selection?

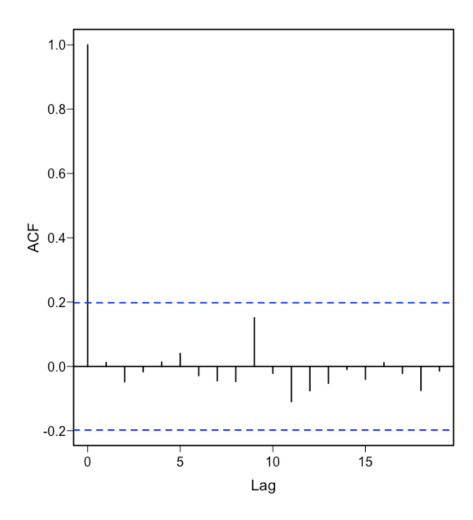
```
library(forecast, quietly=TRUE)
(aaLH <- auto.arima(LakeHuron, approximation=F, stepwise=F))</pre>
```

Series: LakeHuron
ARIMA(2,1,1)

Coefficients:

ar1 ar2 ma1 0.9712 -0.2924 -0.9108 s.e. 0.1137 0.1030 0.0712

Check that model removes most of the correlation structure
acf(resid(aaLH, type="innovation")) # `type="innovation"` is the default



auto.arima(LakeHuron)

Series: LakeHuron ARIMA(0,1,0)

 $sigma^2 = 0.5588$: log likelihood = -109.11

AIC=220.22 AICc=220.26 BIC=222.79

(aaLHO <- auto.arima(LakeHuron, d=0, approximation=F, stepwise=F))</pre>

Series: LakeHuron

ARIMA(1,0,1) with non-zero mean

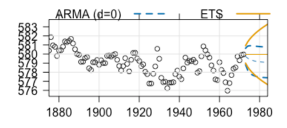
Coefficients:

ar1 ma1 mean 0.7449 0.3206 579.0555 s.e. 0.0777 0.1135 0.3501

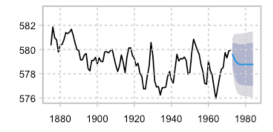
 $sigma^2 = 0.4899$: log likelihood = -103.25

AIC=214.49 AICc=214.92 BIC=224.83

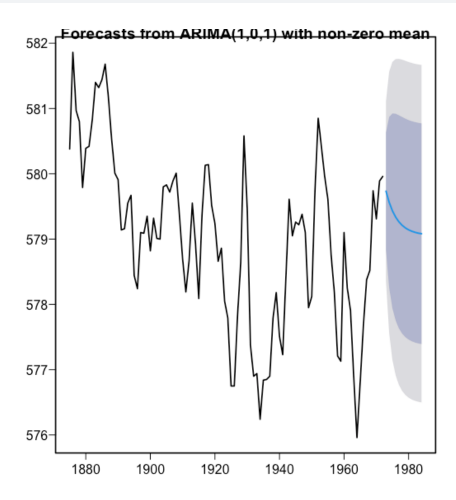
A: ARMA and ETS forecasts with 80% limits

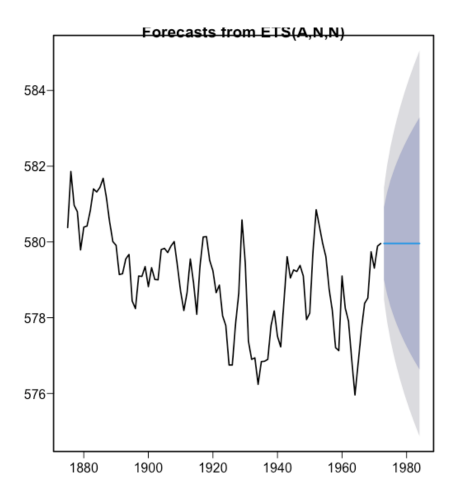


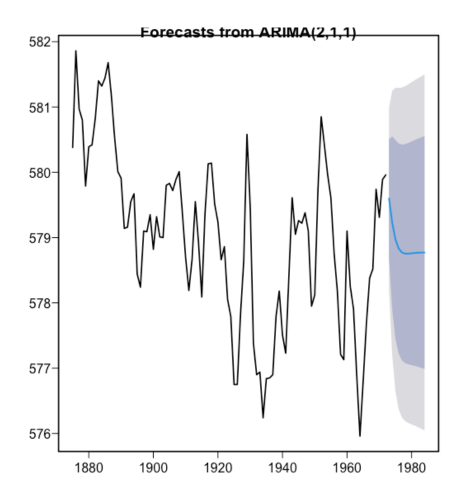
B: ARIMA forecast with 80% and 95% limits



```
plot(forecast(aaLH0, h=12)) ## `level=c(80,95)` is the default
fcETS <- forecast(LakeHuron, h=12)
plot(fcETS)
plot(forecast(aaLH, h=12, level=c(80,95))) # Panel B; ARIMA(2,1,1)</pre>
```







auto.arima(LakeHuron, d=0, max.Q=0, approximation=F, stepwise=F)

Series: LakeHuron

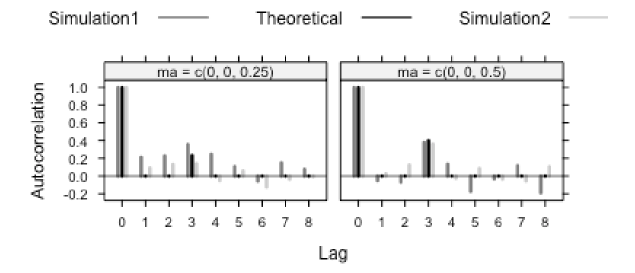
ARIMA(1,0,1) with non-zero mean

Coefficients:

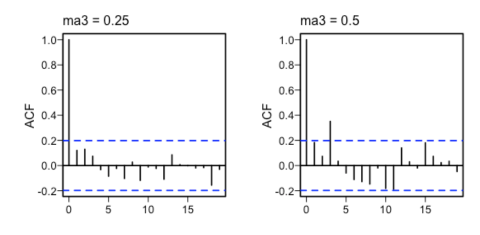
ar1 ma1 mean 0.7449 0.3206 579.0555 s.e. 0.0777 0.1135 0.3501

sigma^2 = 0.4899: log likelihood = -103.25
AIC=214.49 AICc=214.92 BIC=224.83

Use of simulation as a check



```
oldpar <- par(mfrow=c(2,2), mar=c(3.1,4.6,2.6, 1.1))
for(i in 1:2){
    simts <- arima.sim(model=list(order=c(0,0,3), ma=c(0,0,0.25*i)), n=98)
    acf(simts, main="", xlab="")
    mtext(side=3, line=0.5, paste("ma3 =", 0.25*i), adj=0)
}
par(oldpar)</pre>
```

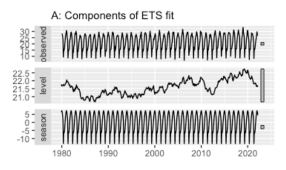


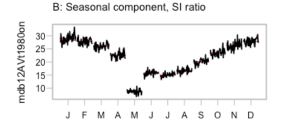
```
set.seed(29)  # Ensure that results are reproducible
estMAord <- matrix(0, nrow=4, ncol=20)
for(i in 1:4){
  for(j in 1:20){
    simts <- arima.sim(n=98, model=list(ma=c(0,0,0.125*i)))
    estMAord[i,j] <- auto.arima(simts, start.q=3)$arma[2] }
}
detectedAs <- table(row(estMAord), estMAord)
dimnames(detectedAs) <- list(ma3=paste(0.125*(1:4)),
Order=paste(0:(dim(detectedAs)[2]-1)))</pre>
```

print(detectedAs)

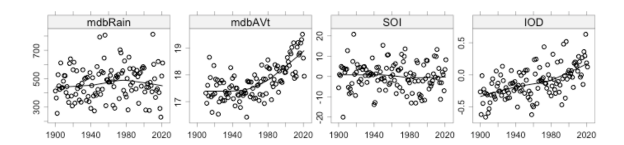
Order

Subsection 6.1.7: Seasonal effects





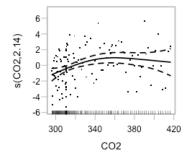
```
suppressPackageStartupMessages(library(ggplot2))
mdb12AVt1980on <- window(DAAG::mdbAVtJtoD, c(1980,1))
AVt.ets <- ets(mdb12AVt1980on)
autoplot(AVt.ets, main="", fg="gray") +
    ggplot2::ggtitle("A: Components of ETS fit") +
    theme(plot.title = element_text(hjust=0, vjust=0.5, size=11))
monthplot(mdb12AVt1980on, col.base=2, fg="gray")
title("B: Seasonal component, SI ratio",
    font.main=1, line=1, adj=0, cex=1.25)</pre>
```

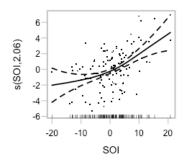


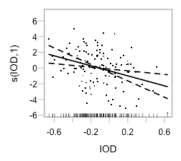
```
suppressPackageStartupMessages(library(mgcv))
bomreg <- within(DAAG::bomregions2021, mdbrtRain <- mdbRain^0.5)
## Check first for a sequential correlation structure, after
## fitting smooth terms s(CO2), s(SOI), and s(IOD)
library(mgcv)
mdbrtRain.gam <- gam(mdbrtRain~s(CO2) + s(SOI) + s(IOD), data=bomreg)
auto.arima(resid(mdbrtRain.gam))</pre>
```

Series: resid(mdbrtRain.gam)
ARIMA(0,0,0) with zero mean

sigma^2 = 4.424: log likelihood = -263.82 AIC=529.64 AICc=529.67 BIC=532.44







```
plot(mdbrtRain.gam, residuals=T, cex=2, fg="gray")
## Do also `gam.check(mdbrtRain.gam)` (Output looks fine)
```

anova(mdbrtRain.gam)

Family: gaussian

Link function: identity

Formula:

 $mdbrtRain \sim s(CO2) + s(SOI) + s(IOD)$

Approximate significance of smooth terms:

edf Ref.df F p-value

s(CO2) 2.144 2.673 4.345 0.01090

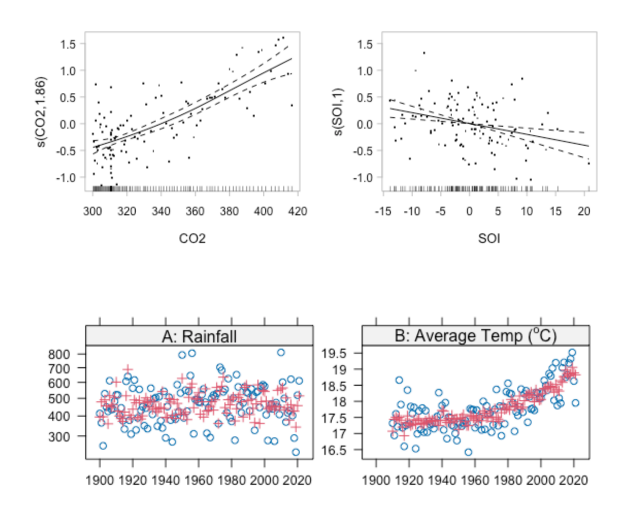
s(SOI) 2.058 2.637 12.581 2.72e-06

s(IOD) 1.000 1.000 9.696 0.00233

```
Box.test(resid(mdbrtRain.gam), lag=10, type="Ljung")
   Box-Ljung test
data: resid(mdbrtRain.gam)
X-squared = 11.935, df = 10, p-value = 0.2894
## Examine normality of estimates of "residuals"
qqnorm(resid(mdbrtRain.gam))
The mdbAVt series
mdbAVt.gam <- gam(mdbAVt ~ s(CO2)+s(SOI)+s(IOD), data=bomreg)</pre>
auto.arima(resid(mdbAVt.gam))
Series: resid(mdbAVt.gam)
ARIMA(0,0,0) with zero mean
sigma^2 = 0.1689: log likelihood = -59.33
AIC=120.67 AICc=120.7 BIC=123.39
anova(mdbAVt.gam)
Family: gaussian
Link function: identity
Formula:
mdbAVt \sim s(CO2) + s(SOI) + s(IOD)
Approximate significance of smooth terms:
         edf Ref.df
                     F p-value
s(CO2) 1.855 2.312 41.842 < 2e-16
s(SOI) 1.000 1.000 10.688 0.00145
s(IOD) 1.000 1.000 0.005 0.94403
```

```
mdbAVt1.gam <- gam(mdbAVt ~ s(CO2)+s(SOI), data=bomreg)</pre>
```

plot(mdbAVt1.gam, residuals=TRUE)



```
labels=list((3:8)*100, (33:39)/2)), x=list(alternating=rep(1,2))),
 strip=strip.custom(factor.levels=faclevs))
gph + latticeExtra::as.layer(xyplot(fitrain+fitAVt~Year, outer=T,
                                     scales=list(y=list(relation='free')),
                                     data=bomreg, pch=3, col=2))
## Use `auto.arima()` to choose the ARIMA order:
aaFitCO2 <- with(bomreg[-(1:10),], auto.arima(mdbAVt, xreg=cbind(CO2,SOI)))</pre>
## Try including a degree 2 polynomial term
aaFitpol2CO2 \leftarrow with(bomreg[-(1:10),],
                     auto.arima(mdbAVt, xreg=cbind(poly(CO2,2),SOI)))
cbind(AIC(aaFitCO2, aaFitpol2CO2), BIC=BIC(aaFitCO2, aaFitpol2CO2))
                     AIC BIC.df BIC.BIC
             df
aaFitCO2
              7 125.3765
                             7 144.4060
```

Subsection 6.1.8: The gamm() function, with a correlated errors model

5 142.6033

aaFitpol2CO2 5 129.0109

```
SOI.gam <- gam(SOI~s(Year), data=bomreg)</pre>
auto.arima(resid(SOI.gam))
                                        \# sigma^2 = 43.4
Series: resid(SOI.gam)
ARIMA(0,0,2) with zero mean
Coefficients:
         ma1
      0.2070 - 0.1557
s.e. 0.0943 0.1008
sigma^2 = 43.43: log likelihood = -402.2
AIC=810.41 AICc=810.61
                          BIC=818.82
## The following breaks the model into two parts -- gam and lme
SOI.gamm <- gamm(SOI~s(Year), data=bomreg)</pre>
res <- resid(SOI.gamm$lme, type="normalized")</pre>
auto.arima(res)
                                        \# sigma^2 = 0.945
```

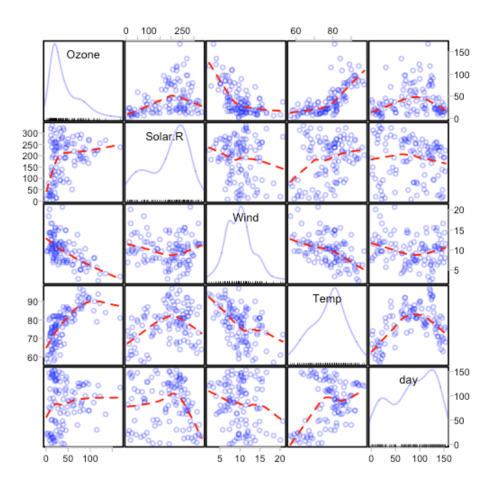
```
Series: res
ARIMA(0,0,2) with zero mean
Coefficients:
        ma1
                  ma2
      0.2070 -0.1557
s.e. 0.0943 0.1008
sigma^2 = 0.9446: log likelihood = -168.68
AIC=343.37 AICc=343.57
                          BIC=351.78
## Extract scale estimate for `gam` component of SOI.gamm
summary(SOI.gamm$gam)[['scale']]
[1] 45.98091
# Note that 45.98 x .945 ~= 43.4
SOIma2.gamm <- gamm(SOI~s(Year), data=bomreg, correlation=corARMA(q=2))
coef(SOIma2.gamm$lme$modelStruct$corStruct, unconstrained = FALSE) # MA2 ests
    Theta1
               Theta2
 0.2070115 -0.1557351
SOIar2.gamm <- gamm(SOI~s(Year), data=bomreg, correlation=corARMA(p=2))
coef(SOIar2.gamm$lme$modelStruct$corStruct, unconstrained = FALSE) # AR2 ests
      Phi1
                 Phi2
 0.2257966 -0.2097561
cbind(AIC(SOI.gam, SOIma2.gamm$lme, SOIar2.gamm$lme),
      BIC=BIC(SOI.gam, SOIma2.gamm$lme, SOIar2.gamm$lme)[,2])
                df
                        AIC
                                 BIC
                 3 819.2646 827.6767
SOI.gam
```

The dataset airquality (153 days, New York, 1972)

SOIma2.gamm\$lme 6 816.4090 833.2331 SOIar2.gamm\$lme 6 815.5117 832.3358

```
## Add time in days from May 1 to data.
airq <- cbind(airquality[, 1:4], day=1:nrow(airquality))
  # Column 5 ('day' starting May 1) replaces columns 'Month' & 'Day')
## Check numbers of missing values # Solar.R:7; Ozone:37
mice::md.pattern(airq, plot=FALSE) # Final row has totals missing.</pre>
```

Wind Temp day Solar.R Ozone 1 0 0 1 1 1 1 1 1 1 0 2 37 44



car::powerTransform(gam(Ozone ~ s(Solar.R)+s(Wind)+s(Temp)+s(day), data=airq))

Estimated transformation parameter Y1

0.2306506

airq\$rt40zone <- airq\$0zone^0.25</pre>

Ozone.gam <- gam(rt4Ozone ~ s(Solar.R)+s(Wind)+s(Temp)+s(day), data=airq)
auto.arima(resid(Ozone.gam)) # Independent errors model appears OK

Series: resid(Ozone.gam)
ARIMA(0,0,0) with zero mean

```
sigma^2 = 0.05897: log likelihood = -0.4
AIC=2.79
          AICc=2.83 BIC=5.5
## Check model terms
anova(Ozone.gam) # For GAM models, this leaves out terms one at a time
Family: gaussian
Link function: identity
Formula:
rt40zone ~ s(Solar.R) + s(Wind) + s(Temp) + s(day)
Approximate significance of smooth terms:
             edf Ref.df
                             F p-value
s(Solar.R) 2.284 2.871 7.701 0.000159
s(Wind)
          2.448 3.104 9.497 1.29e-05
s(Temp)
          4.275 5.287 13.305 < 2e-16
s(day)
          1.000 1.000 0.495 0.483507
## The term in `day` has no explanatory, and will be removed
Ozone1.gam <- update(Ozone.gam, formula=rt40zone ~ s(Solar.R)+s(Wind)+s(Temp))
Subsection 6.1.9: A calibration problem with time series errors
flakes <- DAAG::frostedflakes</pre>
calib.arima <- with(flakes, auto.arima(IA400, xreg=Lab))</pre>
calib.arima
Series: IA400
Regression with ARIMA(0,0,1) errors
Coefficients:
         ma1 intercept
                           xreg
      0.3876 6.9235 0.8323
s.e. 0.0859
                2.5634 0.0679
sigma^2 = 3.278: log likelihood = -199.81
```

AIC=407.62 AICc=408.04 BIC=418.04

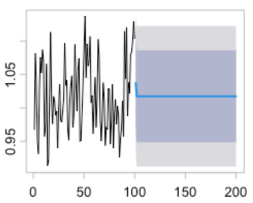
```
with(flakes, coef(auto.arima(IA400/Lab, approximation=F, stepwise=F)))
```

```
ma1 intercept 0.3718151 1.0173303
```

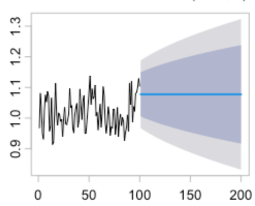
```
with(flakes, coef(auto.arima(IA400-Lab, approximation=F, stepwise=F)))
```

ma1 intercept 0.3838733 0.6201945





B: Forecast from ETS(M,N,N)



Section 6.2: Nonlinear time series

```
x <- numeric(999) # x will contain the ARCH(1) errors
x0 <- rnorm(1)
for (i in 1:999){
x0 <- rnorm(1, sd=sqrt(.25 + .95*x0^2))
x[i] <- x0
}</pre>
```

```
suppressPackageStartupMessages(library(tseries))
garch(x, order = c(0, 1), trace=FALSE)
```

Section 6.3: Further reading

Spatial statistics

Other time series models and packages

Section 6.4: Exercises

6.4

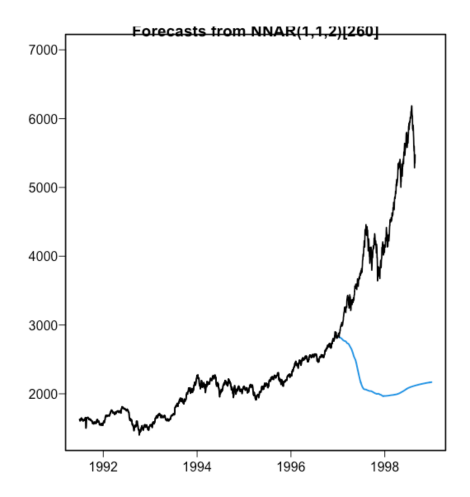
```
xx <- matrix(x, ncol=1000)
```

6.7

```
library(tseries)
data(ice.river)
river1 <- diff(log(ice.river[, 1]))</pre>
```

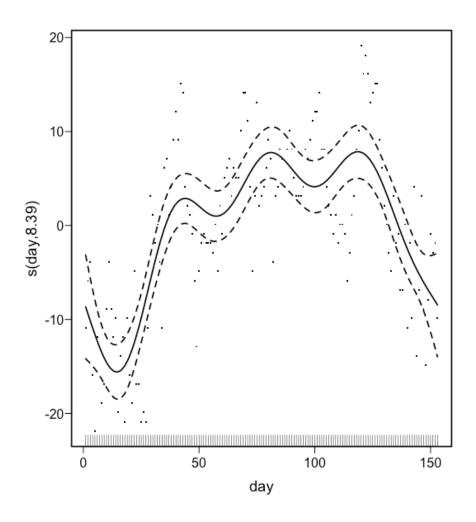
6.9

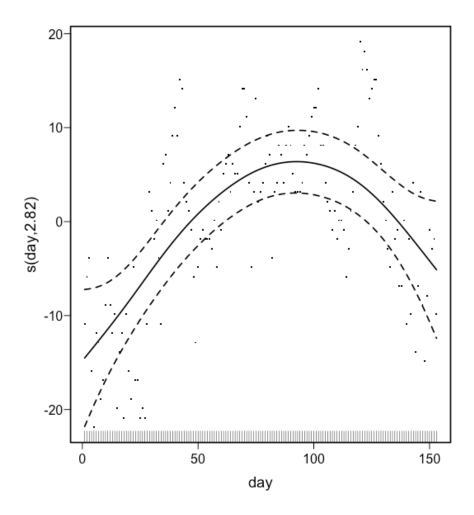
```
library(forecast)
Eu1 <- window(EuStockMarkets[,1], end = c(1996, 260))
Eu1nn <- nnetar(Eu1)
Eu1f <- forecast(Eu1nn, end=end(EuStockMarkets[,1]))
plot(Eu1f, ylim=c(1400, 7000))
lines(EuStockMarkets[,1])</pre>
```



6.10a

```
airq <- cbind(airquality[, 1:4], day=1:nrow(airquality))
  # Column 5 ('day' starting May 1) replaces columns 'Month' & 'Day')
library(mgcv)
temp.gam <- gam(Temp~s(day), data=airq)
tempAR1.gamm <- gamm(Temp~s(day), data=airq, correlation=corAR1())
plot(temp.gam, res=T, cex=2)
plot(tempAR1.gamm$gam, res=T, cex=2)</pre>
```





6.10b

```
(Phi <- coef(tempAR1.gamm$lme$modelStruct$corStruct, unconstrained = FALSE) )
Sigma <- sqrt(tempAR1.gamm$gam$sig2)
## Simulate an AR1 process with this parameter
AR1.sim <- arima.sim(model=list(ar=Phi), n=nrow(airq), sd=Sigma)
simSeries <- AR1.sim+fitted(tempAR1.gamm$gam)
plot(I(1:nrow(airq)), simSeries)
## Compare with initial series
plot(I(1:nrow(airq)), airq$Temp)</pre>
```

```
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))</pre>
```

```
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch6.R")
}
```

7 Chapter 7: Multilevel models, and repeated measures

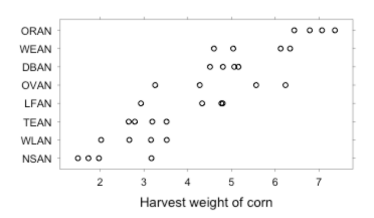
Packages required (plus any dependencies)

DAAG lme4 afex MASS utils devtools qra glmmTMB DHARMa MEMSS forecast splines gamlss plotrix nlme

Additionally, knitr and Hmisc are required in order to process the Rmd source file.

Section 7.1 Corn yield data — analysis using aov()

Corn yield measurements example



```
ant111b <- DAAG::ant111b
ant111b.aov <- aov(harvwt ~ 1 + Error(site), data=ant111b)</pre>
```

```
summary(ant111b.aov)
```

Error: site

Df Sum Sq Mean Sq F value Pr(>F)

Residuals 7 70.3 10.1

Error: Within

Df Sum Sq Mean Sq F value Pr(>F)

Residuals 24 13.9 0.578

Interpreting the mean squares

Details of the calculations

Practical use of the analysis of variance results

Random effects vs. fixed effects

Nested factors - a variety of applications

Subsection 7.1.1: A More Formal Approach

Relations between variance components and mean squares

Interpretation of variance components

Intra-class correlation

Section 7.2 Analysis using lme4::lmer()

```
library(lme4)
ant111b.lmer <- lmer(harvwt ~ 1 + (1 | site), data=ant111b)
## Note that there is no degrees of freedom information.
print(ant111b.lmer, ranef.comp="Variance")
Linear mixed model fit by REML ['lmerMod']
Formula: harvwt ~ 1 + (1 | site)
   Data: ant111b
REML criterion at convergence: 94.42
Random effects:
 Groups
          Name
                      Variance
          (Intercept) 2.368
 site
 Residual
                      0.578
Number of obs: 32, groups: site, 8
Fixed Effects:
(Intercept)
       4.29
```

The processing of output from Imer()

Fitted values and residuals in Imer()

```
s2W <- 0.578; s2L <- 2.37; n <- 4
sitemeans <- with(ant111b, sapply(split(harvwt, site), mean))
grandmean <- mean(sitemeans)
shrinkage <- (n*s2L)/(n*s2L+s2W)
## Check that fitted values equal BLUPs, and compare with site means
BLUP <- grandmean + shrinkage*(sitemeans - grandmean)
BLUP <- fitted(ant111b.lmer)[match(names(sitemeans), ant111b$site)]
BLUP <- grandmean + ranef(ant111b.lmer)$site[[1]]</pre>
```

rbind(BLUP=BLUP, sitemeans=sitemeans)

```
DBAN LFAN NSAN ORAN OVAN TEAN WEAN WLAN
BLUP 4.851 4.212 2.217 6.764 4.801 3.108 5.455 2.925
sitemeans 4.885 4.207 2.090 6.915 4.833 3.036 5.526 2.841
```

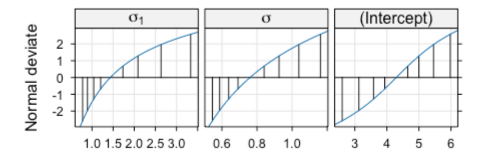
*Uncertainty in the parameter estimates — profile likelihood and alternatives

```
prof.lmer <- profile(ant111b.lmer)
CI95 <- confint(prof.lmer, level=0.95)
rbind("sigmaL^2"=CI95[1,]^2, "sigma^2"=CI95[2,]^2)</pre>
```

2.5 % 97.5 % sigmaL^2 0.7965 6.936 sigma^2 0.3444 1.079

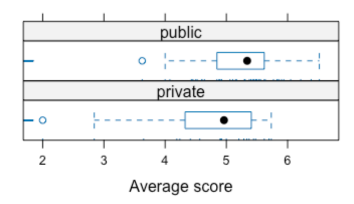
CI95[3,]

2.5 % 97.5 % 3.128 5.456



Modeling more than two levels of random variation

Section 7.3 Survey data, with clustering



Subsection 7.3.1: Alternative models

```
print(VarCorr(science.lmer), comp="Variance", digits=2)
```

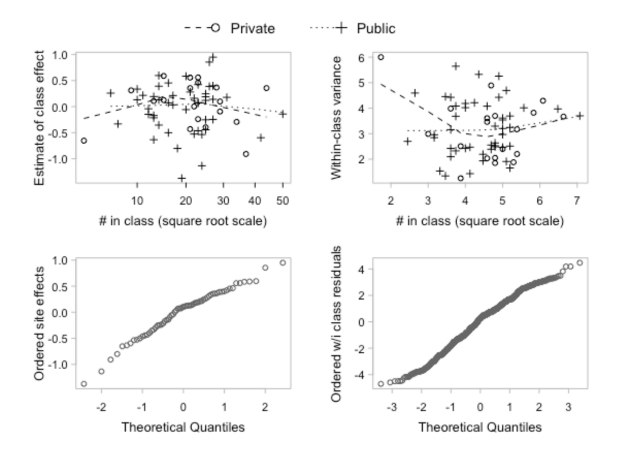
```
Groups
              Name
                           Variance
 school:class (Intercept) 0.32
 school
              (Intercept) 0.00
 Residual
                           3.05
print(coef(summary(science.lmer)), digits=2)
              Estimate Std. Error t value
(Intercept)
                  4.72
                             0.162
                                      29.1
                  0.18
                                       1.9
                             0.098
sexm
                  0.41
                                       2.2
PrivPubpublic
                             0.186
summary(science.lmer)$ngrps
school:class
                   school
          66
                       41
science1.lmer <- lmer(like ~ sex + PrivPub + (1 | school:class),</pre>
                      data = DAAG::science, na.action=na.exclude)
print(VarCorr(science1.lmer), comp="Variance", digits=3)
 Groups
              Name
                           Variance
 school:class (Intercept) 0.321
 Residual
                           3.052
print(coef(summary(science1.lmer)), digits=2)
              Estimate Std. Error t value
(Intercept)
                  4.72
                            0.162
                                      29.1
                             0.098
                                       1.9
                  0.18
sexm
                  0.41
                             0.186
                                       2.2
PrivPubpublic
opt <- options(contrasts=c("contr.sum","contr.poly"))</pre>
  # Change is otherwise made as and if required for individual factors
  # prior to fitting model, and a warning message is generated.
afex::mixed(like ~ sex + PrivPub + (1 | school:class), method="KR", type=2,
            data = na.omit(science), sig_symbols=rep("",4), progress=FALSE)
```

More detailed examination of the output

```
## 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)</pre>
```

```
2.5 % 97.5 % sigma_Class^2 0.178 0.511 sigma^2 2.830 3.300
```

```
## Fit model and generate quantities that will be plotted
science1.lmer <- lmer(like ~ sex + PrivPub + (1 | school:class),
data = science, na.action=na.omit)
## Panel A: random site effects vs number in class
ranf <- ranef(obj = science1.lmer, drop=TRUE)[["school:class"]]
flist <- science1.lmer@flist[["school:class"]]
privpub <- science[match(names(ranf), flist), "PrivPub"]
num <- unclass(table(flist)); numlabs <- pretty(num)
## Panel B: Within class variance estimates vs numbers
res <- residuals(science1.lmer)
vars <- tapply(res, INDEX=list(flist), FUN=var)*(num-1)/(num-2)
## Panel C: Normal probability of random site effects (`ranf`)
## Panel D: Normal probability of residuals (`res`)</pre>
```



```
opar \leftarrow par(oma=c(0,0,1.5,0))
## Panel A: Plot effect estimates vs number
xlab12 <- "# in class (square root scale)"</pre>
plot(sqrt(num), ranf, xaxt="n", pch=c(1,3)[as.numeric(privpub)], cex=0.8,
     xlab=xlab12, ylab="Estimate of class effect", fg="gray")
lines(lowess(sqrt(num[privpub=="private"]),
ranf[privpub=="private"], f=1.1), lty=2)
lines(lowess(sqrt(num[privpub=="public"]),
ranf[privpub=="public"], f=1.1), lty=3)
axis(1, at=sqrt(numlabs), labels=paste(numlabs), lwd=0, lwd.ticks=1)
## Panel B: Within class variance estimates vs numbers
plot(sqrt(num), vars, pch=c(1,3)[unclass(privpub)], cex=0.8,
     xlab=xlab12, ylab="Within-class variance", fg="gray")
lines(lowess(sqrt(num[privpub=="private"]),
      as.vector(vars)[privpub=="private"], f=1.1), lty=2)
lines(lowess(sqrt(num[privpub=="public"]),
as.vector(vars)[privpub=="public"], f=1.1), lty=3)
```

Subsection 7.3.2: Instructive, though faulty, analyses

Ignoring class as the random effect

```
science2.lmer <- lmer(like ~ sex + PrivPub + (1 | school),
data = science, na.action=na.exclude)
print(coef(summary(science2.lmer)), digits=3)</pre>
```

```
Estimate Std. Error t value (Intercept) 4.738 0.163 29.00 sexm 0.197 0.101 1.96 PrivPubpublic 0.417 0.185 2.25
```

```
## NB: Output is misleading
print(VarCorr(science2.lmer), comp="Variance", digits=3)
```

```
Groups Name Variance school (Intercept) 0.166
Residual 3.219
```

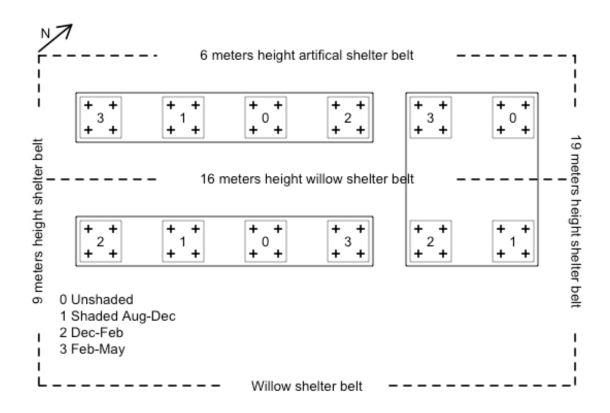
Ignoring the random structure in the data

```
## Faulty analysis, using lm
science.lm <- lm(like ~ sex + PrivPub, data=science)
round(coef(summary(science.lm)), digits=4)</pre>
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 4.7402 0.0996 47.616 0.0000
sexm 0.1509 0.0986 1.531 0.1261
PrivPubpublic 0.3951 0.1051 3.759 0.0002
```

Subsection 7.3.3: Predictive accuracy

Section 7.4 A multilevel experimental design



```
 \begin{aligned} & \text{par}(\text{mar} = \text{rep}(0.25, 4)) \\ & \text{MASS}: = \text{eqscplot}(c(0, 13), c(4.0, 13), \text{type} = \text{"n", xlab} = \text{"", ylab} = \text{"", asp} = 1, axes} = F) \\ & \text{eps} < -0.1 \\ & \text{suby} < -12 \\ & \text{vines} < -\text{function}(x = 1, y = 1, \text{subp} = 0, \text{suby} = 12) \{ \\ & \text{lines}(c(y, y, y + 1, y + 1, y), \text{suby} - c(x, x + 1, x + 1, x, x), \text{lwd} = 0.5) \\ & \text{points}(c(y + .2, y + .2, y + .8, y + .8), \text{suby} - c(x + .2, x + .8, x + .8, x + .2), \text{pch} = 3, \text{cex} = 0.65) \\ & \text{text}(y + .5, \text{suby} - (x + .5), \text{paste}(\text{subp})) \} \end{aligned}
```

```
k<-0
for(i in c(1,3,5,7)){k<-k+1; vines(1,i,c(3,1,0,2)[k])}
k<-0
for(i in c(1,3,5,7)){k<-k+1; vines(4,i,c(2,1,0,3)[k])}
k < -0
for(i in c(1,4,4,1)){k<-k+1
j < -c(9,9,11,11)[k]
vines(i,j,c(3,2,1,0)[k])
lines(c(2*eps, 2.85, NA, 10.15, 13-2*eps), suby-c(3,3,NA,3,3), lty=2)
lines(c(0,2.85,NA,10.15,13), suby-c(0,0,NA,0,0), lty=2)
lines(c(0,4.5,NA,8.5,13), suby-c(8,8,NA,8,8), lty=2)
lines(rep(0,5), suby-c(0,1.25, NA, 6.75, 8), lty=2)
lines(rep(13,5), suby-c(0,1.25, NA, 6.75, 8), lty=2)
lines(c(9,9,12,12,9)+c(-eps,-eps,eps,-eps),
      suby-(c(1,5,5,1,1)+c(-eps,eps,-eps,-eps,-eps)), lwd=1)
lines(c(1,1,8,8,1)+c(-eps,-eps,eps,-eps),
      suby-c(c(1,2,2,1,1)+c(-eps,eps,-eps,-eps,-eps)), lwd=1)
lines(c(1,1,8,8,1)+c(-eps,-eps,eps,-eps),
      suby-c(c(1,2,2,1,1)+3+c(-eps,eps,-eps,-eps,-eps)), lwd=1)
text(6.5, suby, "6 meters height artifical shelter belt")
text(0,suby-4,"9 meters height shelter belt", srt=90)
text(13, suby-4, "19 meters height shelter belt", srt=-90)
text(6.5, suby-8, "Willow shelter belt")
text(0.5, suby-6.5, "0 Unshaded \n1 Shaded Aug-Dec \n2 Dec-Feb \n3 Feb-May", adj=0)
text(6.5, suby-3, "16 meters height willow shelter belt")
offset <-c(4.75, 4.75-sqrt(3)*0.5)/6
arrows(x0=0,y0=12.1, x1=0+offset[1], y1=12.1+offset[2], length=0.15)
text(0.17, 12.55, "N")
```

Subsection 7.4.1: The analysis of variance (anova) table

```
## Analysis of variance: data frame kiwishade (DAAG)
kiwishade <- DAAG::kiwishade
kiwishade.aov <- aov(yield ~ shade + Error(block/shade),
data=kiwishade)
summary(kiwishade.aov)</pre>
```

Error: block

```
Df Sum Sq Mean Sq F value Pr(>F)
Residuals 2
                172
                       86.2
Error: block:shade
          Df Sum Sq Mean Sq F value Pr(>F)
              1395
                        465
                               22.2 0.0012
shade
Residuals 6
               126
                         21
Error: Within
          Df Sum Sq Mean Sq F value Pr(>F)
Residuals 36
               439
                       12.2
```

Subsection 7.4.2: Expected values of mean squares

```
model.tables(kiwishade.aov, type="means", cterms="shade")

Tables of means
Grand mean

96.53

shade
shade
none Aug2Dec Dec2Feb Feb2May
100.20 103.23 89.92 92.77

## Calculate treatment means
with(kiwishade, sapply(split(yield, shade), mean))

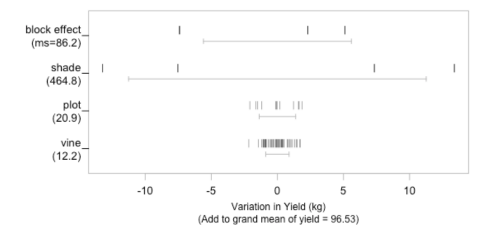
none Aug2Dec Dec2Feb Feb2May
100.20 103.23 89.92 92.77
```

Subsection 7.4.3:* The analysis of variance sums of squares breakdown

```
## For each plot, calculate mean, and differences from the mean
vine <- paste("vine", rep(1:4, 12), sep="")
vine1rows <- seq(from=1, to=45, by=4)
kiwivines <- unstack(kiwishade, yield ~ vine)</pre>
```

```
kiwimeans <- apply(kiwivines, 1, mean)
kiwivines <- cbind(kiwishade[vine1rows, c("block", "shade")],
Mean=kiwimeans, kiwivines-kiwimeans)
kiwivines <- with(kiwivines, kiwivines[order(block, shade), ])
mean(kiwimeans)  # the grand mean</pre>
```

[1] 96.53



```
kiwishade <- DAAG::kiwishade
kiwimeans <- with(kiwishade, aggregate(yield,by=list(block,shade),mean))</pre>
names(kiwimeans) <- c("block", "shade", "meanyield")</pre>
plotmeans.lm <- lm(meanyield~block+shade, data=kiwimeans)</pre>
effects <- predict(plotmeans.lm, type="terms")</pre>
kiwishade.lm <- lm(yield ~ block*shade, data=kiwishade)</pre>
y <- c(effects[,"block"]/sqrt(2) * sqrt(16),
effects[,"shade"]/sqrt(3) * sqrt(12),
residuals(plotmeans.lm)/sqrt(6) * sqrt(4),
residuals(kiwishade.lm)/sqrt(12))
n \leftarrow rep(4:1, c(12, 12, 12, 48))
gps <- rep(c("block effect\n(ms=86.2)", "shade\n(464.8)",
"plot\n(20.9)", "vine\n(12.2)"), c(12, 12, 12, 48))
gps <- factor(gps, levels = rev(c("block effect\n(ms=86.2)",</pre>
"shade\n(464.8)", "plot\n(20.9)", "vine\n(12.2)")))
gps.sd <- sapply(split(y,gps), sd)</pre>
gps.av <- sapply(split(y,gps), mean)</pre>
plot(range(y), range(n)+c(-0.5, 0.5), xlab="", ylab="", yaxt="n", type="n", fg="gray")
text(y, n+0.15, "|", cex=0.8, col=adjustcolor("black", alpha.f=0.5))
un <- 1:4
```

```
sapply(un, function(j){lines(gps.av[j]+c(-gps.sd[j], gps.sd[j]),
    rep(j-0.15,2), col="gray")
lines(rep(gps.av[j],2)-gps.sd[j],
    j-0.15+c(-.06, .06), col="gray")
lines(rep(gps.av[j],2)+gps.sd[j],
    j-0.15+c(-.06, .06), col="gray")
})
mtext(side=1,line=2.25, cex=0.9,
text="Variation in Yield (kg)\n(Add to grand mean of yield = 96.53)")
par(las=2)
axis(2, at=1:4, labels=levels(gps), lwd=0, lwd.ticks=1)
```

Subsection 7.4.4: The variance components

Subsection 7.4.5: The mixed model analysis

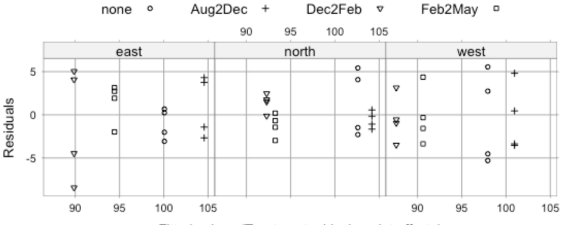
```
kiwishade.lmer <- lmer(yield ~ shade + (1|block) + (1|block:plot),
data=kiwishade)
# block:shade is an alternative to block:plot</pre>
```

```
print(kiwishade.lmer, ranef.comp="Variance", digits=3)
```

```
Linear mixed model fit by REML ['lmerMod']
Formula: yield ~ shade + (1 | block) + (1 | block:plot)
   Data: kiwishade
REML criterion at convergence: 252
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
Fixed Effects:
 (Intercept) shadeAug2Dec shadeDec2Feb shadeFeb2May
      100.20
                     3.03
                                 -10.28
                                                -7.43
```

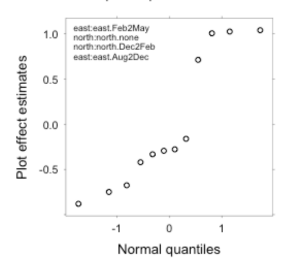
Residuals and estimated effects

A: Standardized residuals after fitting block and plot effects

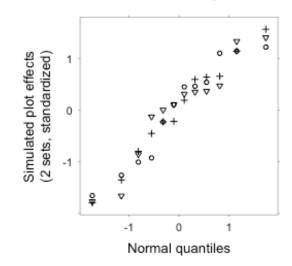


Fitted values (Treatment + block + plot effects)

B: Normal Q-Q plot of plot effects



C: 3 simulations -- normal Q-Q plots



```
xlab="Fitted values (Treatment + block + plot effects)",
              ylab="Residuals",
  main=list(label="A: Standardized residuals after fitting block and plot effects",
          cex=1.05, x=0.01, y=0, font=1, just="left"),
             auto.key=list(space='top', points=TRUE, columns=4))
print(update(pk1, scales=list(x=list(alternating=TRUE), tck=0.35),
             par.settings=parsetA), position=c(0,.52,1,1))
## Panel B
ploteff <- ranef(kiwishade.lmer, drop=TRUE)[[1]]</pre>
nam <- names(sort(ploteff, decreasing=TRUE)[1:4])</pre>
parsetB <- modifyList(DAAG::DAAGtheme(color=FALSE),</pre>
                       list(layout.heights=list(axis.top=0.85)))
pk2 <- qqmath(ploteff, ylab="Plot effect estimates",</pre>
              key=list(x=0, y=0.98, corner=c(0,1),
              text=list(labels=nam, cex=0.65)),
              main=list(label="B: Normal Q-Q plot of plot effects",
                         cex=1.05, x=0.01, y=0.25, font=1, just="left"),
              xlab="Normal quantiles")
print(update(pk2, scales=list(tck=0.35), par.settings=parsetB),
      newpage=FALSE, position=c(0,0,.5,.5))
## Panel C
simvals <- simulate(kiwishade.lmer, nsim=3, seed=23)</pre>
simranef <- function(y)ranef(lme4::refit(kiwishade.lmer, y))[[1]]</pre>
simeff <- apply(simvals, 2, function(y) scale(simranef(y)))</pre>
simeff <- as.data.frame(simeff)</pre>
pk3 <- qqmath(~ sim_1+sim_2+sim_3, data=simeff,</pre>
              ylab="Simulated plot effects\n(2 sets, standardized)",
              Qs=list(tck=0.35), aspect=1,
               main=list(label="C: 3 simulations -- normal Q-Q plots",
                          x=0.01, y=0.25, cex=1.05, font=1, just="left"),
               xlab="Normal quantiles")
print(update(pk3, scales=list(tck=0.35), par.settings=parsetB),
      newpage=FALSE, position=c(0.48,0,1,.5))
```

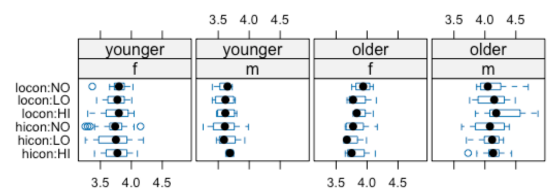
```
with(kiwishade, sapply(split(yield, shade), mean))
```

```
none Aug2Dec Dec2Feb Feb2May 100.20 103.23 89.92 92.77
```

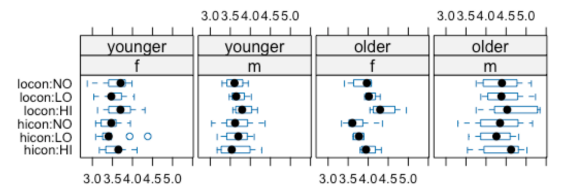
Subsection 7.4.6: Predictive accuracy

Section 7.5 Within and between subject effects

A: Boxplots for `log(csoa)`, by combinations of `target` and `tint`



B: Boxplots for `log(it)`, by combinations of `target` and `tint`



Model fitting criteria

Subsection 7.5.1: Model selection

```
## Capitalize tinting$agegp
levels(tinting$agegp) <- R.utils::capitalize(levels(tinting$agegp))</pre>
## Fit all interactions: data frame tinting (DAAG)
it3.lmer <- lmer(log(it) ~ tint*target*agegp*sex + (1 | id),
                 data=DAAG::tinting, REML=FALSE)
it2.lmer <- lmer(log(it) ~ (tint+target+agegp+sex)^2 + (1 | id),
                 data=DAAG::tinting, REML=FALSE)
it1.lmer <- lmer(log(it)~(tint+target+agegp+sex) + (1 | id),
                 data=DAAG::tinting, REML=FALSE)
anova(it1.lmer, it2.lmer, it3.lmer)
Data: DAAG::tinting
Models:
it1.lmer: log(it) ~ (tint + target + agegp + sex) + (1 | id)
it2.lmer: log(it) \sim (tint + target + agegp + sex)^2 + (1 | id)
it3.lmer: log(it) ~ tint * target * agegp * sex + (1 | id)
        npar AIC BIC logLik deviance Chisq Df Pr(>Chisq)
it1.lmer
          8 1.14 26.8 7.43
                                  -14.9
it2.lmer 17 -3.74 50.7 18.87 -37.7 22.88 9
                                                     0.0065
it3.lmer 26 8.15 91.5 21.93
                                  -43.9 6.11 9
                                                     0.7288
## Code that gives the first four such plots, for the observed data
interaction.plot(tinting$tint, tinting$agegp, log(tinting$it))
interaction.plot(tinting$target, tinting$sex, log(tinting$it))
interaction.plot(tinting$tint, tinting$target, log(tinting$it))
interaction.plot(tinting$tint, tinting$sex, log(tinting$it))
```

Subsection 7.5.2: Estimates of model parameters

```
it2.reml <- update(it2.lmer, REML=TRUE)
print(coef(summary(it2.reml)), digits=2)</pre>
```

```
Estimate Std. Error t value
(Intercept)
                         3.6191
                                     0.130
                                             27.82
tint.L
                         0.1609
                                     0.044
                                              3.64
tint.Q
                         0.0210
                                     0.045
                                              0.46
                                     0.042
targethicon
                        -0.1181
                                            -2.79
                                     0.233
                                              2.02
agegpolder
                         0.4712
                                              0.35
sexm
                         0.0821
                                     0.233
tint.L:targethicon
                        -0.0919
                                     0.046
                                             -2.00
                                            -0.15
tint.Q:targethicon
                        -0.0072
                                     0.048
tint.L:agegpolder
                        0.1308
                                     0.049
                                              2.66
tint.Q:agegpolder
                                              1.34
                         0.0697
                                     0.052
                                             -1.99
tint.L:sexm
                                     0.049
                        -0.0979
tint.Q:sexm
                         0.0054
                                     0.052
                                              0.10
targethicon:agegpolder -0.1389
                                             -2.38
                                     0.058
targethicon:sexm
                         0.0779
                                     0.058
                                              1.33
agegpolder:sexm
                         0.3316
                                     0.326
                                              1.02
```

```
# NB: The final column, giving degrees of freedom, is not in the # summary output. It is our addition.
```

```
subs <- with(tinting, match(unique(id), id))
with(tinting, table(sex[subs], agegp[subs]))</pre>
```

```
Younger Older f 9 4 m 4 9
```

Section 7.6 A mixed model with a betabinomial error

Subsection 7.6.1: The betabinomial distribution

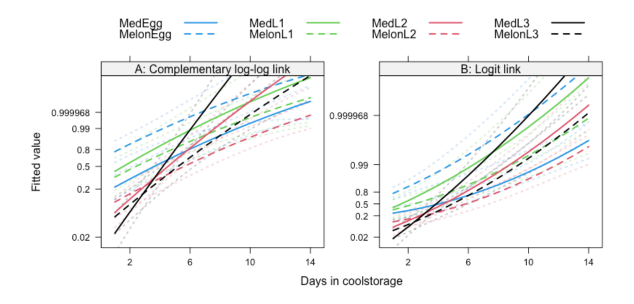
```
## devtools::install_github("jhmaindonald/qra") # Use if not found on CRAN
```

Notation

Source of data

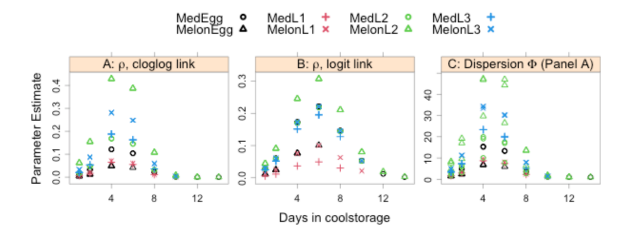
```
HawCon <- within(as.data.frame(qra::HawCon), {</pre>
  trtGp <- gsub("Fly", "", paste0(CN,LifestageTrt))</pre>
  trtGp <- factor(trtGp, levels=sort(unique(trtGp))[c(1,5,2,6,3,7,4,8)])
  trtGpRep <- paste0(CN,LifestageTrt,":",RepNumber)</pre>
  scTime <- scale(TrtTime) # Centering and scaling can help model fit</pre>
  })
## Load packages that will be used
suppressMessages(library(lme4)); suppressMessages(library(glmmTMB))
library(splines)
form <- cbind(Dead,Live)~0+trtGp/TrtTime+(1|trtGpRep)</pre>
## Add the quadratic term from a degree 2 orthogonal polynomial
form2s <- update(form, . ~ . + scale(scTime^2))</pre>
  ## Scale "corrections" to reduce risk of potential numerical problems
HCbb.cll <- glmmTMB(form, dispformula=~trtGp+ns(scTime,2),</pre>
                    family=glmmTMB::betabinomial(link="cloglog"), data=HawCon)
HCbb2s.cll <- update(HCbb.cll, formula=form2s)</pre>
HCbb.logit <- glmmTMB(form, dispformula=~trtGp+ns(scTime,2),</pre>
                      family=glmmTMB::betabinomial(link="logit"), data=HawCon)
HCbb2s.logit <- update(HCbb.logit, formula=form2s)</pre>
                                                               ## CLL
summary(HCbb2s.cll)$coefficients$cond["scale(scTime^2)",]
  Estimate Std. Error z value Pr(>|z|)
  -0.09130 0.06143 -1.48627
                                     0.13721
summary(HCbb2s.logit)$coefficients$cond["scale(scTime^2)",] ## Logit
  Estimate Std. Error z value Pr(>|z|)
   0.43219 0.17766 2.43268 0.01499
AICtab <- AIC(BB=HCbb.cll, HCbb2s.cll, HCbb.logit, HCbb2s.logit)
AICtab[['Details']] <- c(paste0('BB: Compl. log-log', c('', ', added curve')),
                           paste0('BB: Logit', c('', ', added curve')))
AICtab[order(AICtab[["AIC"]]), ]
```

```
dfAICDetailsHCbb2s.cll28 709.9BB: Compl. log-logHCbb.cll27 710.0 BB: Compl. log-log, added curveHCbb2s.logit28 717.2BB: Logit, added curveHCbb.logit27 721.7BB: Logit
```



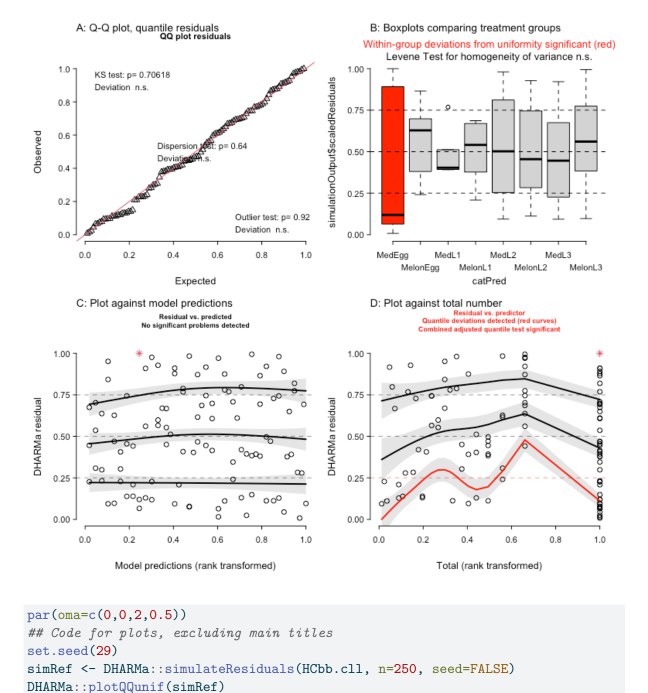
```
dat <- expand.grid(trtGp=factor(levels(HawCon$trtGp), levels=levels(HawCon$trtGp)),</pre>
TrtTime=pretty(range(HawCon$TrtTime),15), trtGpRep=NA)
ab <- qra::getScaleCoef(HawCon$scTime)</pre>
dat$scTime <- with(dat,(TrtTime-ab[1])/ab[2])</pre>
hatClog <- predict(HCbb2s.cll, newdata=dat)</pre>
hatClog <- predict(HCbb2s.cll, se=T, newdata=dat)</pre>
hatLgt <- predict(HCbb2s.logit, newdata=dat)</pre>
hatLgt <- predict(HCbb2s.logit, se=T, newdata=dat)</pre>
dat2 <- cbind(rbind(dat,dat), link=rep(c('clog','logit'), rep(nrow(dat),2)))</pre>
dat2 <- within(dat2, {fit<-c(hatClog$fit,hatLgt$fit);</pre>
se.fit<-c(hatClog$se.fit,hatLgt$se.fit);</pre>
link=rep(c('clog','logit'), rep(nrow(dat),2))})
dat2 <- within(dat2, {lwr<-fit-2*se.fit; upr<-fit+2*se.fit})</pre>
library(lattice)
my.panel.bands <- function(x, y, upper, lower, fill, col,</pre>
subscripts, ..., font, fontface)
upper <- upper[subscripts]</pre>
```

```
lower <- lower[subscripts]</pre>
panel.lines(x,lower, ...)
panel.lines(x,upper, ...)
panel2 <- function(x, y, ...){
panel.superpose(x, y, panel.groups = my.panel.bands, type='l', lty=3, alpha=0.25,...)
panel.xyplot(x, y, type='l', lwd=2, cex=0.6, ...)
parset <- simpleTheme(col=rep(4:1,rep(2,4)), lty=rep(1:2, 4), lwd=2)</pre>
## c('solid','1141')
p \leftarrow c(.02, .2, .5, .8, .99, .999968)
cloglog <- make.link('cloglog')$linkfun</pre>
logit <- make.link('logit')$linkfun</pre>
fitpos <- list(cloglog(p), logit(p))</pre>
lab <- paste(p)</pre>
\lim \leftarrow \text{list}(\text{cloglog}(c(0.02, 0.99998)), \log it(c(0.02, 0.99998)))
\lim \leftarrow \text{lapply}(\lim, \text{function}(x)c(x[1],x[1]+\text{diff}(x)*1.2))
gph <- xyplot(fit~TrtTime|link, outer=TRUE, data=dat2, groups=trtGp,</pre>
               upper = dat2$upr, lower = dat2$lwr, panel = panel2,
               xlab="Days in coolstorage", ylab="Fitted value",
               auto.key=list(text=levels(HawCon$trtGp), columns=4,
                               points=FALSE, lines=TRUE),
               par.settings=parset, layout=c(2,1),
               scales=list(x=list(at=c(2,6,10,14)),
               y=list(relation='free',
               at=fitpos, labels=lab, limits=lim), alternating=c(1,1)))
update(gph, strip=strip.custom(factor.levels=c("A: Complementary log-log link",
"B: Logit link")))
```



```
parset <- DAAG::DAAGtheme(color=TRUE, col.points=rep(1:4,rep(2,4)),</pre>
                           pch=rep(1:4,2), lwd=2)
HawCon$rho2clog <- qra::getRho(HCbb.cll)</pre>
HawCon$dispClog <- with(HawCon, 1+(Total-1)*rho2clog)</pre>
par(oma=c(0,0,2,0))
titles=c(expression("A: "*rho*", cloglog link"), expression("B: "*rho*", logit link"),
expression("C: Dispersion "*Phi*" (Panel A)"))
library(lattice)
HawCon$rho2logit <- qra::getRho(HCbb.logit)</pre>
xyplot(rho2clog+rho2logit+dispClog ~ TrtTime, groups=trtGp, data=HawCon,
       outer=TRUE, between=list(x=0.25), par.settings=parset,
   scales=list(x=list(alternating=FALSE,at=c(4,8,12), labels=paste(c(4,8,12))),
               y=list(relation='free',tick.number=4)),
   auto.key=list(columns=4, between.columns=2, between=1),
   xlab="Days in coolstorage", ylab="Parameter Estimate",
   strip=strip.custom(factor.levels=titles))
```

Subsection 7.6.2: Diagnostic checks



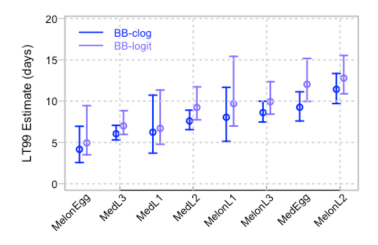
title(main="A: Q-Q plot, quantile residuals", adj=0, line=2.5,

Subsection 7.6.3: Lethal time estimates and confidence intervals

```
shorten <- function(nam, leaveout=c('trtGp','Fly',':')){
for(txt in leaveout){
  nam <- gsub(txt,'', nam, fixed=TRUE)
}
nam
}</pre>
```

```
LTbb.cll <- qra::extractLT(p=0.99, obj=HCbb.cll, link="cloglog", a=1:8, b=9:16, eps=0, df.t=NULL)[,-2] rownames(LTbb.cll) <- gsub("trtGp|Fly|:", '', rownames(LTbb.cll), perl=T)
```

```
LTbb.logit <- qra::extractLT(p=0.99, obj=HCbb.logit, link="logit", a=1:8, b=9:16, eps=0, offset=0, df.t=NULL)[,-2]
rownames(LTbb.logit) <- shorten(rownames(LTbb.logit))
```



```
library(plotrix)
gpNam <- rownames(LTbb.cll)</pre>
ordEst <- order(LTbb.cll[,1])</pre>
col5 <- c("blue","lightslateblue","blueviolet",'gray50','gray80')</pre>
plotCI(1:8-0.1, y=LTbb.cll[ordEst,1], ui=LTbb.cll[ordEst,3],
  li=LTbb.cll[ordEst,2], lwd=2, col=col5[1], xaxt="n",
  fg="gray", xlab="", ylab="LT99 Estimate (days)",
  xlim=c(0.8,8.2), ylim=c(0,19.5))
plotCI(1:8+0.1, y=LTbb.logit[ordEst,1], ui=LTbb.logit[ordEst,3],
  li=LTbb.logit[ordEst,2], lwd=2, col=col5[2], xaxt="n", add=TRUE)
axis(1, labels=FALSE, tck=0.02, col.ticks="gray40")
text(x = 1:length(gpNam)+0.1,
  ## Move labels to just below bottom of chart.
  y = par("usr")[3] - 0.8,
  labels = gpNam[ordEst], ## Use names from the data list.
    xpd = NA,
                    ## Change the clipping region
  srt = 45,
                    ## Rotate the labels by 45 degrees.
  adj = 0.95)
                    ## Adjust the labels to almost 100% right-justified
grid()
legend("topleft", legend=c("BB-clog", "BB-logit"),
       inset=c(0.01,0.01), lty=c(1,1), col=col5[1:2],
text.col=col5[1:2], bty="n",y.intersp=0.85)
```

```
LTbin.cll <- qra::extractLT(p=0.99, obj=HCbin.cll,
a=1:8, b=9:16, eps=0, df.t=NULL)[,-2]
rownames(LTbin.cll) <- shorten(rownames(LTbin.cll))
```

A warning against over-interpretation

Section 7.7 Observation level random effects — the moths dataset

```
moths <- DAAG::moths
moths$transect <- 1:41 # Each row is from a different transect
moths$habitat <- relevel(moths$habitat, ref="Lowerside")</pre>
A.glmer <- glmer(A~habitat+sqrt(meters)+(1|transect),
family=poisson(link=sqrt), data=moths)
print(summary(A.glmer), show.resid=FALSE, correlation=FALSE, digits=3)
Generalized linear mixed model fit by maximum likelihood (Laplace
  Approximation) [glmerMod]
 Family: poisson (sqrt)
Formula: A ~ habitat + sqrt(meters) + (1 | transect)
   Data: moths
     ATC
             BIC
                   logLik deviance df.resid
   212.6
           229.7
                   -96.3
                             192.6
                                         31
Random effects:
                     Variance Std.Dev.
 Groups
         Name
 transect (Intercept) 0.319
                              0.564
Number of obs: 41, groups: transect, 41
Fixed effects:
                Estimate Std. Error z value
                                              Pr(>|z|)
                                       4.93 0.00000082
(Intercept)
                  1.7322
                             0.3513
habitatBank
                 -2.0415
                             0.9377
                                      -2.18
                                                 0.029
habitatDisturbed -1.0359
                             0.4071 - 2.54
                                                 0.011
habitatNEsoak
                 -0.7319
                             0.4323
                                      -1.69
                                                 0.090
habitatNWsoak
                 2.6787
                             0.5101 5.25 0.00000015
habitatSEsoak
                 0.1178
                             0.3923
                                     0.30
                                                 0.764
habitatSWsoak
                  0.3900
                             0.5260
                                    0.74
                                                 0.458
habitatUpperside -0.3135
                             0.7549 - 0.42
                                                 0.678
sqrt(meters)
                  0.0675
                             0.0631
                                      1.07
                                                 0.285
```

```
suppressPackageStartupMessages(library(gamlss))
A1quasi.glm <- glm(A~habitat, data=moths, family=quasipoisson(link=sqrt))
Asqrt.lss <- gamlss(A ~ habitat + sqrt(meters), trace=FALSE,
                    family = NBI(mu.link='sqrt'), data = moths)
A1.glmer <- glmer(A~habitat+(1|transect), data=moths, family=poisson(link=sqrt))
Cglm <- coef(summary(A1quasi.glm))</pre>
Cglmer <- coef(summary(A1.glmer))</pre>
fitAll <- cbind("quasi-Coef"=Cglm[-1,1], "quasi-SE"=Cglm[-1,2],</pre>
  "NBI-Coef"=coef(Asqrt.lss)[2:8], "NBI-SE"=c(0.94,.41,.43,.51,.39,.53,.75),
  "glmer-Coef"=Cglmer[-1,1], "glmer-SE"=Cglmer[-1,2])
rownames(fitAll) <- substring(rownames(fitAll),8)</pre>
round(fitAll, 2) # NB, all SEs are for the difference from 'Lowerside'
          quasi-Coef quasi-SE NBI-Coef NBI-SE glmer-Coef glmer-SE
              -2.13
                         0.86 -2.19 0.94
                                                  -1.99
                                                             0.95
Bank
                                                   -1.13
              -1.07
Disturbed
                         0.41
                                -1.00 0.41
                                                             0.40
NEsoak
              -0.61
                         0.43 -0.80 0.43
                                                   -0.57
                                                             0.41
```

<pre>detach("package:glmmTMB",</pre>	<pre>character.only=TRUE)</pre>	

-0.41 0.75

2.67 0.51

0.28 0.53

0.09 0.39

2.73

0.19

0.54

0.36

0.51

0.39 0.51

0.43

Section 7.8 Repeated measures in time

2.73

0.16

0.45

0.23

0.54

0.54

0.45

0.41

The theory of repeated measures modeling

NWsoak

SEsoak

SWsoak

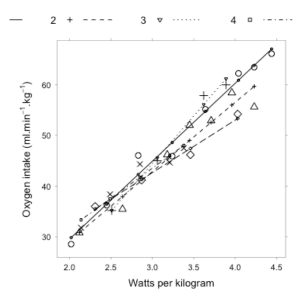
Upperside

Different approaches to repeated measures analysis

Subsection 7.8.1: Example – random variation between profiles

```
humanpower1 <- DAAG::humanpower1
```

^{*}Correlation structure



```
## Plot points and fitted lines
xyplot(o2 ~ wattsPerKg, groups=id, data=humanpower1,
    par.settings=DAAG::DAAGtheme(color=F), scales=list(tck=0.5),
    panel=function(x,y,subscripts,groups,...){
        yhat <- fitted(lm(y ~ groups*x))
        panel.superpose(x, y, subscripts, groups, pch=1:5, cex=1.2)
        panel.superpose(x, yhat, subscripts, groups, type="b", cex=0.5)
        },
    auto.key=list(columns=5, lines=T),
    xlab="Watts per kilogram",
    ylab=expression("Oxygen intake ("*ml.min^{-1}*.kg^{-1}*")"))</pre>
```

Separate lines for different athletes

```
## Calculate intercepts and slopes; plot Slopes vs Intercepts
## Uses the function lmList() from the lme4 package
humanpower1 <- DAAG::humanpower1
hp.lmList <- lmList(o2 ~ wattsPerKg | id, data=humanpower1)
coefs <- coef(hp.lmList)
round(coefs,3)</pre>
```

```
(Intercept) wattsPerKg
1 -1.155 15.35
```

```
2
        1.916
                   13.65
3
      -12.008
                   18.81
                   11.83
4
        8.029
5
       11.553
                   10.36
c("Correlation between intercept and slope"=cor(coefs[,1],coefs[,2]))
Correlation between intercept and slope
                                -0.9975
A random coefficients model
hp.lmer <- lmer(o2 ~ wattsPerKg + (wattsPerKg | id), data=humanpower1)</pre>
hp.lmer <- lmer(o2 ~ wattsPerKg + (0+wattsPerKg | id), data=humanpower1)</pre>
print(summary(hp.lmer), digits=3)
Linear mixed model fit by REML ['lmerMod']
Formula: o2 ~ wattsPerKg + (0 + wattsPerKg | id)
   Data: humanpower1
REML criterion at convergence: 129.7
Scaled residuals:
    Min
             1Q Median
                                    Max
                             3Q
-1.9117 -0.8978 0.0598 0.7854 1.5382
Random effects:
 Groups
          Name
                     Variance Std.Dev.
 id
          wattsPerKg 0.211
                              0.46
                     5.776
                              2.40
 Residual
```

Fixed effects:

Estimate Std. Error t value (Intercept) 1.299 2.220 0.59 wattsPerKg 14.204 0.715 19.88

Number of obs: 28, groups: id, 5

Correlation of Fixed Effects:

(Intr)

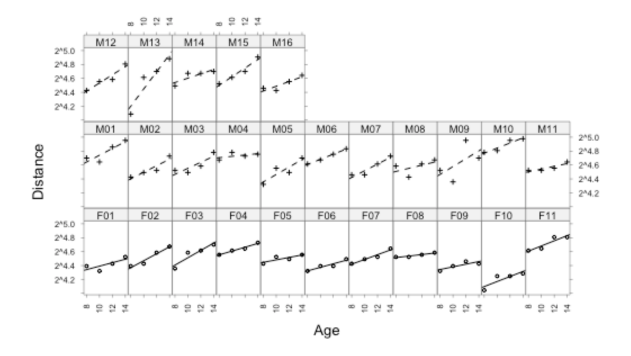
wattsPerKg -0.936

```
sort(coef(lmList(o2 ~ wattsPerKg | id, data=humanpower1))[,1])
```

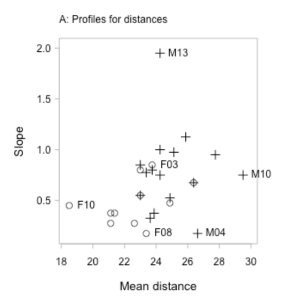
```
[1] -12.008 -1.155 1.916 8.029 11.553
```

Subsection 7.8.2: Orthodontic measurements on children

```
Orthodont <- MEMSS::Orthodont
Orthodont$logdist <- log(Orthodont$distance)
```



Preliminary data exploration



B: Logarithms of distances + M130.08 0.06 Slope 0.04 o F10 + M10 0.02 + M04 2.9 3.0 3.1 3.2 3.3 3.4 Mean log distance

```
## For Panel B, repeat with logdist replacing distance
plot(blog ~ ylogbar, col=c(F="gray40", M="black")[sex], data=ab, fg="gray",
     pch=c(F=1, M=3)[sex], xlim=lims[,"ylogbar"], ylim=lims[,"blog"],
     xlab="Mean log distance", ylab="Slope")
with(subset(ab, extremes),
     text(blog ~ ylogbar, labels=rownames(ab)[extremes], pos=4, xpd=TRUE))
mtext(side=3, line=0.75, "B: Logarithms of distances", adj=0)
## Compare male slopes with female slopes
extreme.males <- match(c("M04","M13"), rownames(ab))</pre>
with(ab[-extreme.males,],
t.test(blog[sex=="F"], blog[sex=="M"], var.equal=TRUE))
    Two Sample t-test
data: blog[sex == "F"] and blog[sex == "M"]
t = -2.3, df = 23, p-value = 0.03
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
 -0.0160529 -0.0009191
sample estimates:
mean of x mean of y
  0.02115
           0.02963
# Specify var.equal=TRUE, to allow comparison with anova output
```

A random coefficients model

```
rePCA(orthdiffx.lmer)

$Subject
Standard deviations (1, ..., p=2):
[1] 1.632 0.000
```

```
Rotation (n \times k) = (2 \times 2):
                 [,2]
         [,1]
[1,] -0.99993 -0.01153
[2,] -0.01153 0.99993
attr(,"class")
[1] "prcomplist"
orthdiff.lmer <- lmer(logdist ~ Sex * scAge + (1 | Subject),
                      data=Orthodont, subset=keep)
summary(orthdiff.lmer)
Linear mixed model fit by REML ['lmerMod']
Formula: logdist ~ Sex * scAge + (1 | Subject)
   Data: Orthodont
 Subset: keep
REML criterion at convergence: -232
Scaled residuals:
          1Q Median
                         3Q
-3.312 -0.510 0.014 0.543 3.945
Random effects:
 Groups
         Name
                      Variance Std.Dev.
 Subject (Intercept) 0.00633 0.0796
                      0.00238 0.0488
Number of obs: 100, groups: Subject, 25
Fixed effects:
              Estimate Std. Error t value
(Intercept)
              3.11451
                         0.02510 124.11
SexMale
              0.09443
                         0.03354
                                    2.82
                       0.00329
                                    6.42
scAge
               0.02115
SexMale:scAge 0.00849
                         0.00440
                                    1.93
Correlation of Fixed Effects:
            (Intr) SexMal scAge
SexMale
           -0.748
             0.000 0.000
scAge
SexMal:scAg 0.000 0.000 -0.748
```

```
Orthodont2 <- droplevels(subset(Orthodont, keep))</pre>
opt <- options(contrasts=c("contr.sum", "contr.poly"))</pre>
orthdiff.mixed <- afex::mixed(logdist ~ Sex * scAge + (1 | Subject), type=2,
                              method='S', data=Orthodont2)
  ## NB `type` refers to type of test, NOT `error` type.
             # Reset to previous contrasts setting
options(opt)
orthdiff.mixed
Mixed Model Anova Table (Type 2 tests, S-method)
Model: logdist ~ Sex * scAge + (1 | Subject)
Data: Orthodont2
     Effect
                  df
                              F p.value
        Sex 1, 23.00 7.93 **
1
                                   .010
      scAge 1, 73.00 140.72 ***
                                  <.001
3 Sex:scAge 1, 73.00 3.72 +
                                  .058
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '+' 0.1 ' ' 1
contrasts(Orthodont2[['Subject']]) <- 'contr.sum'</pre>
contrasts(Orthodont2[['Sex']]) <- 'contr.sum'</pre>
orthdiffa.lmer <- update(orthdiff.lmer, formula=. ~ . -Sex:scAge)
AIC(orthdiffa.lmer,orthdiff.lmer)
                     AIC
               df
orthdiffa.lmer 5 -227.4
orthdiff.lmer
                6 - 220.1
Correlation between successive times
```

```
F08 M06
[1,] 1 1
[2,] 0 0
[3,] 1 1
```

Fitting a sequential correlation structure

```
Correlation structure of class corCAR1 representing Phi 0.000000003311
```

Section 7.9: Further notes on multilevel models

Subsection 7.9.1: Sources of variation – complication or focus of interest?

Subsection 7.9.2: Predictions from models with a complex error structure

Consequences from assuming an overly simplistic error structure

Subsection 7.9.3: An historical perspective on multilevel models

Subsection 7.9.4: Meta-analysis

Subsection 7.9.5: Functional data analysis

Subsection 7.9.6: Error structure in explanatory variables

Exercises (7.12)

7.1

^{*}The variance for the difference in slopes

```
n.omit <- 2
take <- rep(TRUE, 48)
take[sample(1:48,2)] <- FALSE</pre>
kiwishade.lmer <- lmer(yield ~ shade + (1|block) + (1|block:plot),</pre>
                        data = kiwishade,subset=take)
vcov <- VarCorr(kiwishade.lmer)</pre>
print(vcov, comp="Variance")
 Groups
            Name
                         Variance
 block:plot (Intercept)
                          2.35
 block
             (Intercept) 3.28
 Residual
                         12.45
7.6
cult.lmer <- lmer(ct ~ Cultivar + Dose + factor(year) +</pre>
                        (-1 + Dose | gp), data = DAAG::sorption, REML=TRUE)
cultdose.lmer <- lmer(ct ~ Cultivar/Dose + factor(year) +</pre>
                             (-1 + Dose | gp), data = DAAG::sorption, REML=TRUE)
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()</pre>
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))</pre>
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch7.R")
}
```

8 Chapter 8: Tree-based Classification and Regression

Packages required (plus any dependencies)

DAAG latticeExtra plot rpart rpart.plot MASS ggplot2 car randomForest Additionally, knitr and Hmisc are required in order to process the Rmd source file.

suppressPackageStartupMessages(library(latticeExtra))

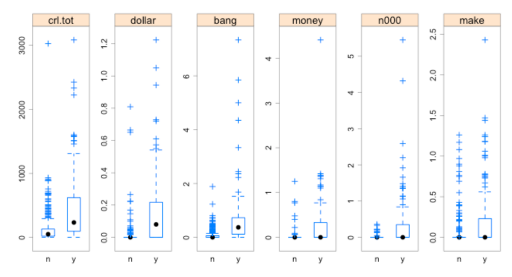
When are tree-based methods appropriate?

Section 8.1: Tree-based methods — uses and basic notions

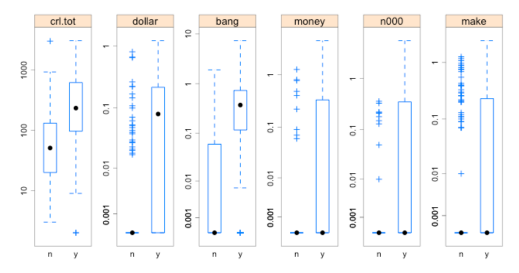
Examples that will be used to demonstrate the methodology

Subsection 8.1.1: Detecting email spam~- an initial look

A: Raw data values

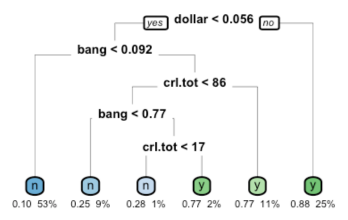


B: Boxplots, using log(x+1) scale



```
nam <- c("crl.tot", "dollar", "bang", "money", "n000", "make")
nr <- sample(1:dim(DAAG::spam7)[1],500)
yesno<-DAAG::spam7$yesno[nr]
spam7a <- DAAG::spam7[nr,c(nam,"yesno")]
formab <- as.formula(paste(paste(nam, collapse='+'), '~ yesno'))</pre>
```

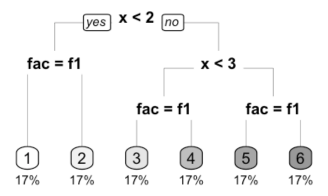
```
## Obtain 500-row sample; repeat the first plot (of crl.tot)
spam.sample <- spam7[sample(seq(1,4601), 500, replace=FALSE), ]
boxplot(split(spam.sample$crl.tot, spam.sample$yesno))</pre>
```



```
Classification tree:
rpart(formula = yesno ~ crl.tot + dollar + bang + money + n000 +
    make, data = DAAG::spam7, method = "class", model = TRUE)
Variables actually used in tree construction:
[1] bang
            crl.tot dollar
Root node error: 1813/4601 = 0.394
n = 4601
      CP nsplit rel error xerror
                                   xstd
1 0.4766
              0
                    1.000 1.000 0.0183
2 0.0756
                    0.523 0.547 0.0154
3 0.0116
                    0.372 0.388 0.0135
              3
4 0.0105
              4
                    0.361 0.384 0.0134
5 0.0100
              5
                    0.350 0.382 0.0134
```

Subsection 8.1.2: Choosing the number of splits

Section 8.2: Splitting criteria, with illustrative examples

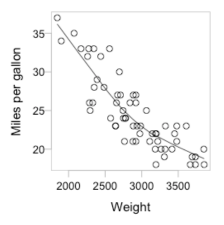


Choosing the split~- regression trees

Subsection 8.2.1: Within and between sums of squares

Subsection 8.2.2: Choosing the split~- classification trees

Subsection 8.2.3: Tree-based regression versus loess regression smoothing



```
## loess fit to Mileage vs Weight: data frame car.test.frame (rpart)
with(rpart::car.test.frame, scatter.smooth(Mileage ~ Weight))
```



```
dat <- data.frame(Weight=seq(from=min(car.test.frame$Weight),
    to=max(car.test.frame$Weight)))
pred <- predict(car.tree, newdata=dat)
pred2 <- predict(car2.tree, newdata=dat)
lwr <- dat$Weight[c(1,diff(pred)) != 0]
upr <- dat$Weight[c(diff(pred),1) != 0]
xy2 <- with(car.test.frame, loess.smooth(Weight, Mileage, evaluation=2011))
lwrL0 <- xy2$y[c(1,diff(pred)) != 0]</pre>
```

```
uprL0 <- xy2$y[c(diff(pred),1) != 0]
round(rbind(lwr,upr,lwrL0,uprL0,
pred[c(diff(pred),1)!=0],pred2[c(diff(pred),1)!=0]),1)</pre>
```

```
723
                903
                       1243
                              2011
      1845.0 2568.0 2748.0 3088.0
lwr
      2567.0 2747.0 3087.0 3855.0
upr
lwrL0
        36.2
               27.1
                       25.1
                              22.3
uprL0
        27.1
               25.1
                       22.3
                              18.8
               25.6
        30.9
                       23.8
                              20.4
        28.9
               25.6
                       24.1
                              18.7
```

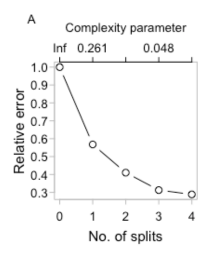
Subsection 8.2.4: Predictive accuracy, and the cost-complexity tradeoff

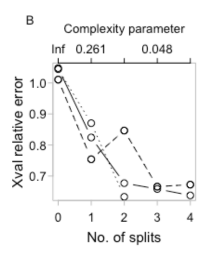
Subsection 8.2.5: Cross-validation

Subsection 8.2.6: The cost-complexity parameter

```
vignette("longintro", package="rpart")
```

Subsection 8.2.7: Prediction error versus tree size





Section 8.3: The practicalities of tree construction – two examples

Subsection 8.3.1: Data for female heart attack patients

```
mifem <- DAAG::mifem
summary(mifem) # data frame mifem (DAAG)
outcome
               age
                           yronset
                                      premi
                                               smstat
                                                       diabetes
         Min.
                 :35.0 Min.
live:974
                               :85.0
                                      y:311
                                               c :390
                                                       y:248
dead:321 1st Qu.:57.0 1st Qu.:87.0 n :928
                                               x :280
                                                       n:978
           Median:63.0 Median:89.0 nk: 56
                                                       nk: 69
                                               n:522
           Mean :60.9 Mean :88.8
                                               nk:103
           3rd Qu.:66.0 3rd Qu.:91.0
                               :93.0
           Max. :69.0 Max.
highbp
       hichol angina stroke
y:813 y:452 y:472 y:153
n:406 n:655 n:724 n:1063
nk: 76 nk:188 nk: 99 nk: 79
set.seed(29)
                # Make results reproducible
mifem.rpart <- rpart(outcome ~ ., method="class", data = mifem, cp = 0.0025)</pre>
## Tabular equivalent of Panel A from `plotcp(mifem.rpart)`
printcp(mifem.rpart, digits=3)
Classification tree:
rpart(formula = outcome ~ ., data = mifem, method = "class",
   cp = 0.0025)
Variables actually used in tree construction:
[1] age
            angina
                    diabetes hichol
                                    premi
                                             smstat
                                                     stroke
[8] yronset
Root node error: 321/1295 = 0.248
n= 1295
```

```
CP nsplit rel error xerror
                                   xstd
1 0.20249
              0
                    1.000 1.000 0.0484
2 0.00561
              1
                    0.798 0.829 0.0453
3 0.00467
                           0.875 0.0462
             13
                    0.717
4 0.00312
             17
                    0.698 0.860 0.0459
5 0.00250
                    0.695 0.863 0.0460
             18
```

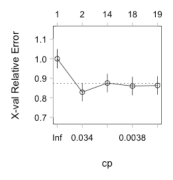
```
cat(c("...", capture.output(printcp(mifem.rpart, digits=3))[-(1:9)]),
    sep="\n")
```

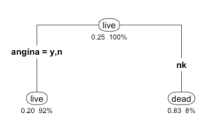
. . .

Root node error: 321/1295 = 0.248

n= 1295

	CP	nsplit	rel	error	xerror	xstd
1	0.20249	0		1.000	1.000	0.0484
2	0.00561	1		0.798	0.829	0.0453
3	0.00467	13		0.717	0.875	0.0462
4	0.00312	17		0.698	0.860	0.0459
5	0.00250	18		0.695	0.863	0.0460





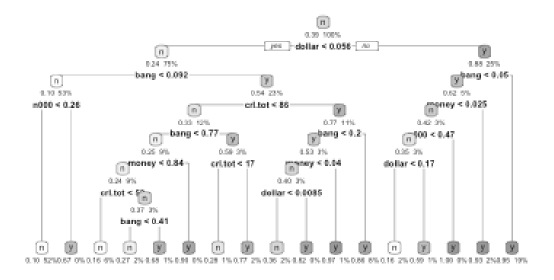
Subsection 8.3.2: The one-standard-deviation rule

Subsection 8.3.3: Printed Information on Each Split

```
print(mifemb.rpart)
n = 1295
node), split, n, loss, yval, (yprob)
      * denotes terminal node
1) root 1295 321 live (0.7521 0.2479)
  2) angina=y,n 1196 239 live (0.8002 0.1998) *
  3) angina=nk 99 17 dead (0.1717 \ 0.8283) *
set.seed(59)
spam7a.rpart <- rpart(formula = yesno ~ crl.tot + dollar + bang +</pre>
                     money + n000 + make, method="class", cp = 0.002,
                     model=TRUE, data = DAAG::spam7)
printcp(spam7a.rpart, digits=3)
Classification tree:
rpart(formula = yesno ~ crl.tot + dollar + bang + money + n000 +
    make, data = DAAG::spam7, method = "class", model = TRUE,
    cp = 0.002)
Variables actually used in tree construction:
[1] bang
         crl.tot dollar money
                                   n000
Root node error: 1813/4601 = 0.394
n = 4601
        CP nsplit rel error xerror
                                    xstd
1 0.47656
               0
                     1.000 1.000 0.0183
2 0.07557
               1
                     0.523 0.550 0.0154
               3
3 0.01158
                     0.372 0.390 0.0135
           4 0.361 0.386 0.0134
4 0.01048
```

```
5 0.00634
               5
                      0.350 0.374 0.0133
6 0.00552
               10
                      0.317 0.360 0.0130
  0.00441
7
               11
                      0.311 0.357 0.0130
  0.00386
               12
                      0.307 0.352 0.0129
8
  0.00276
                      0.291
                            0.339 0.0127
               16
10 0.00221
               17
                      0.288
                            0.339 0.0127
11 0.00200
               18
                      0.286 0.336 0.0127
```

```
cpdf <- signif(as.data.frame(spam7a.rpart$cptable),3)
minRow <- which.min(cpdf[,"xerror"])
upr <- sum(cpdf[minRow, c("xerror","xstd")])
takeRow <- min((1:minRow)[cpdf[1:minRow,"xerror"]<upr])
newNsplit <- cpdf[takeRow, 'nsplit']
cpval <- mean(cpdf[c(takeRow-1,takeRow),"CP"])</pre>
```



```
spam7b.rpart <- prune(spam7a.rpart, cp=cpval)
rpart.plot::rpart.plot(spam7b.rpart, under=TRUE, box.palette="Grays", tweak=1.65)</pre>
```

How does the one standard error rule affect accuracy of estimates?

```
requireNamespace('randomForest', quietly=TRUE)
DAAG::compareTreecalcs(data=DAAG::spam7, fun="rpart")
 rpSEcvI
           rpcvI rpSEtest rptest nSErule nREmin
  0.1396
           0.1387
                    0.1269 0.1217 7.0000
                                                8.0000
acctree.mat <- matrix(0, nrow=100, ncol=6)</pre>
spam7 <- DAAG::spam7</pre>
for(i in 1:100)
acctree.mat[i,] <- DAAG::compareTreecalcs(data=spam7, fun="rpart")</pre>
How is the standard error calculated?
When are tree-based methods appropriate?
Section 8.4: From one tree to a forest – a more global optimality
suppressPackageStartupMessages(library(randomForest))
spam7.rf <- randomForest(yesno ~ ., data=spam7, importance=TRUE)</pre>
spam7.rf
Call:
 randomForest(formula = yesno ~ ., data = spam7, importance = TRUE)
               Type of random forest: classification
                     Number of trees: 500
No. of variables tried at each split: 2
        OOB estimate of error rate: 11.56%
Confusion matrix:
          y class.error
n 2651 137
              0.04914
```

y 395 1418

0.21787

z <- tuneRF(x=spam7[, -7], y=spam7\$yesno, plot=FALSE)

```
mtry = 2 00B error = 11.8%
Searching left ...
mtry = 1 00B error = 12.67%
-0.07366 0.05
Searching right ...
mtry = 4 00B error = 11.76%
0.003683 0.05

zdash <- t(z[,2,drop=F])
colnames(zdash) <- paste0(c("mtry=",rep("",2)), z[,1])
round(zdash,3)</pre>
mtry=1 2 4
```

00BError 0.127 0.118 0.118

importance(spam7.rf)

n	у	${\tt MeanDecreaseAccuracy}$	${\tt MeanDecreaseGini}$
46.73	54.19	70.57	248.10
56.21	55.35	76.13	431.75
91.66	100.46	115.95	588.53
33.09	51.87	53.49	206.51
58.25	15.74	62.29	115.46
13.67	21.76	26.72	41.13
	46.73 56.21 91.66 33.09 58.25	n y 46.73 54.19 56.21 55.35 91.66 100.46 33.09 51.87 58.25 15.74 13.67 21.76	46.73 54.19 70.57 56.21 55.35 76.13 91.66 100.46 115.95 33.09 51.87 53.49 58.25 15.74 62.29

Subsection 8.4.1: Prior probabilities

```
Pima.tr <- MASS::Pima.tr
table(Pima.tr$type)</pre>
```

No Yes 132 68

```
set.seed(41)  # This seed should reproduce the result given here
Pima.rf <- randomForest(type ~ ., data=Pima.tr)
## The above is equivalent to:
## Pima.rf <- randomForest(type ~ ., data=Pima.tr, sampsize=200)
round(Pima.rf$confusion,3)</pre>
```

```
No Yes class.error
No 110 22 0.167
Yes 32 36 0.471

tab <- prop.table(table(Pima.tr$type))

Pima.rf <- randomForest(type ~ ., data=Pima.tr, sampsize=c(68,68))
```

Pima.rf <- randomForest(type ~ ., data=Pima.tr, sampsize=c(132,68))

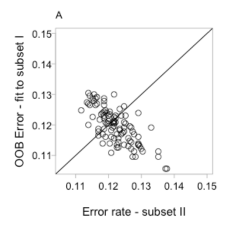
Subsection 8.4.2: A low-dimensional representation of observations

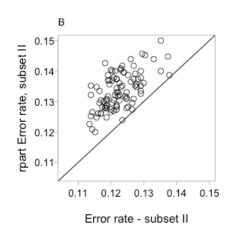
Subsection 8.4.3: Models with a complex error structure

Section 8.5: Additional notes – one tree, or many trees?

Subsection 8.5.1: Differences between rpart() and randomForest()

Error rates - rpart() versus randomForest()





Times required for computation

Subsection 8.5.2: Tree-based methods, versus other approaches

Tree-based methods may usefully complement other approaches

Subsection 8.5.3: Further notes

Pruning as variable selection

Section 8.6: Further reading and extensions

Exercises (8.7)

8.5

```
sapply(MASS::biopsy, function(x)sum(is.na(x))) ## Will omit rows with NAs
                      VЗ
                                               ۷7
   ID
         V1
                ۷2
                            ۷4
                                   ۷5
                                         ۷6
                                                      8V
                                                            V9 class
    0
          0
                 0
                       0
                             0
                                    0
                                         16
                                                0
                                                       0
                                                             0
biops <- na.omit(MASS::biopsy)[,-1]</pre>
                                                     ## Omit also column 1 (IDs)
## Examine list element names in randomForest object
names(randomForest(class ~ ., data=biops))
 [1] "call"
                        "type"
                                           "predicted"
 [4] "err.rate"
                                           "votes"
                        "confusion"
 [7] "oob.times"
                        "classes"
                                           "importance"
[10] "importanceSD"
                        "localImportance" "proximity"
[13] "ntree"
                        "mtry"
                                           "forest"
[16] "y"
                        "test"
                                           "inbag"
[19] "terms"
8.5a
## Repeated runs, note variation in OOB accuracy.
for(i in 1:10) {
  biops.rf <- randomForest(class ~ ., data=biops)</pre>
  OOBerr <- mean(biops.rf$err.rate[,"OOB"])</pre>
  print(paste(i, ": ", round(OOBerr, 4), sep=""))
  print(round(biops.rf$confusion,4))
[1] "1: 0.0288"
          benign malignant class.error
benign
              432
                         12
                                  0.0270
malignant
                        232
                                  0.0293
[1] "2: 0.0344"
          benign malignant class.error
             431
                                  0.0293
benign
                         13
malignant
                        230
                                  0.0377
[1] "3: 0.0308"
          benign malignant class.error
             433
                         11
                                 0.0248
benign
                        230
                9
                                  0.0377
malignant
[1] "4: 0.0307"
```

```
benign malignant class.error
             431
                                  0.0293
benign
                         13
                        233
malignant
               6
                                  0.0251
[1] "5: 0.0311"
          benign malignant class.error
             433
                         11
                                  0.0248
benign
malignant
               8
                        231
                                  0.0335
[1] "6: 0.0301"
          benign malignant class.error
             431
                         13
benign
                                  0.0293
                        233
                                 0.0251
malignant
               6
[1] "7: 0.0312"
          benign malignant class.error
             433
                         11
                                  0.0248
benign
                        232
malignant
                                  0.0293
[1] "8: 0.0301"
          benign malignant class.error
             433
                         11
                                  0.0248
benign
malignant
               8
                        231
                                 0.0335
[1] "9: 0.0285"
          benign malignant class.error
             433
benign
                         11
                                  0.0248
malignant
               6
                        233
                                  0.0251
[1] "10: 0.0301"
          benign malignant class.error
             432
                         12
                                  0.0270
benign
               7
                        232
                                  0.0293
malignant
8.5b
## Repeated train/test splits: OOB accuracy vs test set accuracy.
for(i in 1:10){
  trRows <- sample(1:dim(biops)[1], size=round(dim(biops)[1]/2))</pre>
  biops.rf <- randomForest(class ~ ., data=biops[trRows, ],</pre>
    xtest=biops[-trRows,-10], ytest=biops[-trRows,10])
  oobErr <- mean(biops.rf$err.rate[,"00B"])</pre>
  testErr <- mean(biops.rf$test$err.rate[,"Test"])</pre>
```

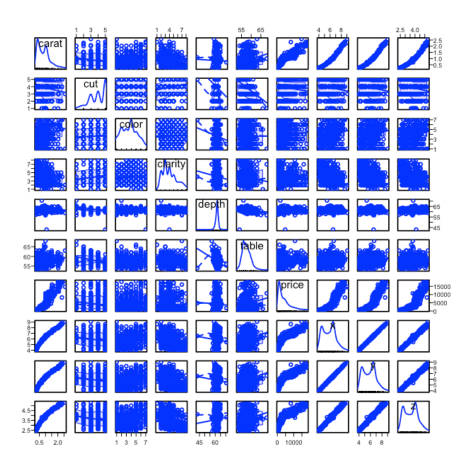
[1] 0.0282 0.0340

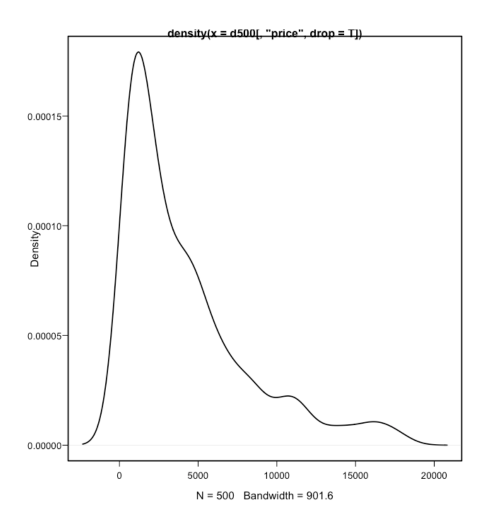
print(round(c(oobErr,testErr),4))

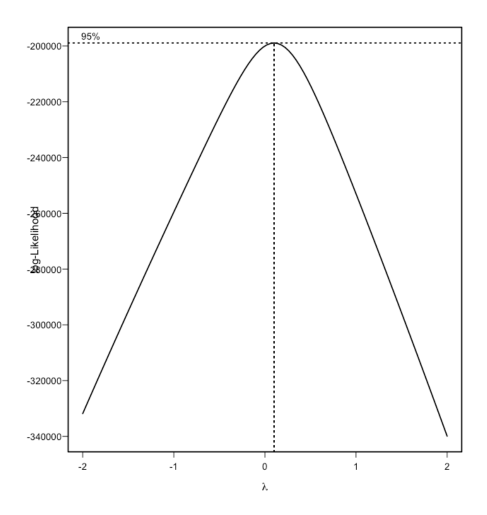
[1] 0.0366 0.0327

```
[1] 0.0360 0.0226
[1] 0.0330 0.0311
[1] 0.0360 0.0384
[1] 0.0357 0.0353
[1] 0.0248 0.0399
[1] 0.0442 0.0270
[1] 0.0407 0.0268
[1] 0.0347 0.0242
8.5c
randomForest(class ~ ., data=biops, xtest=biops[,-10], ytest=biops[,10])
Call:
 randomForest(formula = class ~ ., data = biops, xtest = biops[, -10], ytest = biops[,
               Type of random forest: classification
                     Number of trees: 500
No. of variables tried at each split: 3
        OOB estimate of error rate: 2.93%
Confusion matrix:
          benign malignant class.error
             433
                     11
                               0.02477
benign
                       230
malignant
                               0.03766
                Test set error rate: 0%
Confusion matrix:
          benign malignant class.error
             444
                      0
benign
malignant
              0
                       239
                                     0
8.7
## Run model on total data
randomForest(as.factor(type) ~ ., data=Pima.tr)
Call:
 randomForest(formula = as.factor(type) ~ ., data = Pima.tr)
               Type of random forest: classification
                     Number of trees: 500
```

```
No. of variables tried at each split: 2
        OOB estimate of error rate: 28%
Confusion matrix:
     No Yes class.error
No 109 23
                0.1742
Yes 33 35
                0.4853
rowsamp <- sample(dim(Pima.tr)[1], replace=TRUE)</pre>
randomForest(as.factor(type) ~ ., data=Pima.tr[rowsamp, ])
Call:
 randomForest(formula = as.factor(type) ~ ., data = Pima.tr[rowsamp,
                                                                         ])
              Type of random forest: classification
                    Number of trees: 500
No. of variables tried at each split: 2
        OOB estimate of error rate: 8.5%
Confusion matrix:
    No Yes class.error
No 129
        3
               0.02273
Yes 14 54
               0.20588
8.8a
d500 <- ggplot2::diamonds[sample(1:nrow(ggplot2::diamonds), 500),]
unlist(sapply(d500, class)) # Check the class of the 10 columns
              cut1
                        cut2
                                color1
                                          color2 clarity1 clarity2
    carat
"numeric" "ordered" "factor" "ordered" "factor" "ordered" "factor"
             table
                       price
                                     X
"numeric" "numeric" "numeric" "numeric" "numeric"
                    # If screen space is limited do two plots, thus:
car::spm(d500)
  # 1) variables 1 to 5 and 7 (`price`); 2) variables 6 to 10
plot(density(d500[, "price", drop = T]))
                                                # Distribution is highly skew
MASS::boxcox(price~., data=ggplot2::diamonds) # Suggests log transformation
```







8.8b

```
diamonds <- ggplot2::diamonds; Y <- diamonds[,"price", drop=T]
library(rpart)
d7.rpart <- rpart(log(Y) ~ ., data=diamonds[,-7], cp=5e-7) # Complex tree
d.rpart <- prune(d7.rpart, cp=0.0025)
printcp(d.rpart) # Relative to `d7.rpart`, simpler and less accurate</pre>
```

```
Regression tree:
rpart(formula = log(Y) ~ ., data = diamonds[, -7], cp = 0.0000005)
Variables actually used in tree construction:
[1] carat clarity color x y
```

```
Root node error: 55531/53940 = 1
n = 53940
      CP nsplit rel error xerror
                                    xstd
              0
                    1.000 1.000 0.00409
2 0.0885
              1
                    0.276 0.276 0.00141
3 0.0661
              2
                   0.187 0.188 0.00104
4 0.0290
                    0.121 0.121 0.00075
              3
5 0.0105
              4
                   0.092 0.092 0.00059
6 0.0072
              5
                  0.082 0.082 0.00054
7 0.0071
              6
                   0.074 0.072 0.00049
              7
8 0.0030
                   0.067 0.068 0.00047
9 0.0026
              8 0.064 0.065 0.00046
10 0.0026
              9
                   0.062 0.061 0.00044
11 0.0025
             10
                    0.059 0.059 0.00044
nmin <- which.min(d7.rpart$cptable[,'xerror'])</pre>
dOpt.rpart <- prune(d7.rpart, cp=d7.rpart$cptable[nmin,'CP'])</pre>
print(dOpt.rpart$cptable[nmin])
[1] 0.0000008743
(xerror12 <- dOpt.rpart$cptable[c(nrow(d.rpart$cptable),nmin), "xerror"])</pre>
           1931
     11
0.05934 0.01074
## Subtract from 1.0 to obtain R-squared statistics
rbind("d.rpart"=d.rpart[['variable.importance']],
      "dOpt.rpart"=dOpt.rpart[['variable.importance']]) |>
  (\x)100*apply(x,1,function(x)x/sum(x)))() > round(1) > t()
                  x carat
                             z clarity table color depth cut
d.rpart
           23.8 23.1 22.9 22.4
                                   4.8
                                         2.8
                                               0.2
                                                     0.1 0.1
dOpt.rpart 23.5 22.9 22.7 22.2
                                   5.2
                                         2.7
                                               0.5
                                                     0.2 0.1
```

```
Y <- ggplot2::diamonds[,"price", drop=T]
samp5K <- sample(1:nrow(diamonds), size=5000)</pre>
(diamond5K.rf <- randomForest(x=diamonds[samp5K,-7], y=log(Y[samp5K]),</pre>
                   xtest=diamonds[-samp5K,-7], ytest=log(Y[-samp5K])))
Call:
 randomForest(x = diamonds[samp5K, -7], y = log(Y[samp5K]), xtest = diamonds[-samp5K,
               Type of random forest: regression
                     Number of trees: 500
No. of variables tried at each split: 3
          Mean of squared residuals: 0.01434
                    % Var explained: 98.59
                       Test set MSE: 0.01
                    % Var explained: 98.65
## Omit arguments `xtest` and `ytest` if calculations take too long
sort(importance(diamond5K.rf)[,1], decreasing=T) |>
  ((x)100*x/sum(x))() > round(1) > t()
        y carat x
                      z clarity color depth table cut
[1,] 33.5 23.6 21 17.2
                            2.6
                                  1.2 0.4
                                              0.3 0.2
8.9a
(diamond5KU.rf <- randomForest(x=diamonds[samp5K,-7], y=Y[samp5K],</pre>
                   xtest=diamonds[-samp5K,-7], ytest=Y[-samp5K]))
Call:
 randomForest(x = diamonds[samp5K, -7], y = Y[samp5K], xtest = diamonds[-samp5K,
                                                                                        -7], y
               Type of random forest: regression
                     Number of trees: 500
No. of variables tried at each split: 3
          Mean of squared residuals: 450742
```

% Var explained: 97.11 Test set MSE: 493916 % Var explained: 96.9

```
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch8.R")
}</pre>
```

9 Chapter 9: Multivariate data exploration and discrimination

Packages required (plus any dependencies)

Packages used are: DAAG MASS RColorBrewer teigen BiocManager DAAGbio hddplot lmtest splines cobalt mice datasets car micemd oz randomForest ggplot2 latticeExtra mvtnorm teigen limma hddplot mgcv MatchIt sandwich gridExtra DAAGbio mlbench (exercise).

Additionally, knitr and Hmisc are required in order to process the Rmd source file.

Section 9.1: Multivariate exploratory data analysis

```
## Make the lattice package and the possum dataset available
library(latticeExtra)
possum <- DAAG::possum</pre>
```

Subsection 9.1.1: Scatterplot matrices

```
xlab="", varnames=vnames, key=key, axis.line.tck=0.6))
gphB <- with(possum, cloud(earconch~taill+footlgth, data=possum,
    col=colr[sex], key=key, pch = (0:6)[site],
    zlab=list("earconch", rot=90), zoom=0.925))
update(c("A: Scatterplot matrix"=gphA, "B: Cloud plot"=gphB),
    between=list(x=1))</pre>
```

Subsection 9.1.2: Principal components analysis

Preliminary data scrutiny

```
## Ratios of largest to smallest values: possum[, 6:14] (DAAG)
possum <- DAAG::possum</pre>
sapply(na.omit(possum[, 6:14]), function(x)round(max(x)/min(x),2))
## Principal components calculations: possum[, 6:14] (DAAG)
here <- complete.cases(possum[, 6:14])
possum.prc <- prcomp(log(possum[here, 6:14]))</pre>
scores <- cbind(predict(possum.prc), possum[here, c('sex', 'site')])</pre>
## For parset, key and colr; see code for Fig 9.1
pchr \leftarrow c(3,4,0,8,2,10,1)
parset <- list(fontsize=list(text=10, points=6), cex=0.75, pch=pchr, alpha=0.8)</pre>
key <- modifyList(key, list(columns=1, space="right"))</pre>
gph <- with(scores, xyplot(PC2 ~ PC1, aspect="iso", key = key,</pre>
            col = colr[sex], pch = (0:6)[site])
update(gph, scales=list(tck=0.5), par.settings=parset,
       xlab="1st Principal Component", ylab="2nd Principal Component")
print(summary(possum.prc),digits=2)
cat("\nRotations (otherwise called Loadings)\n")
print(possum.prc$rotation, digits=2)
## By default, blanks are shown for loadings < 0.1 in magnitude
```

The stability of the principal components plot

```
suppressPackageStartupMessages(library(ggplot2))
theme_set(theme_gray(base_size=8))
## Bootstrap principal components calculations: possum (DAAG)
```

```
## Sample from rows where there are no missing values
rowsfrom <- (1:nrow(possum))[complete.cases(possum[, 6:14])]</pre>
logpossum6to14 <- log(possum[rowsfrom, 6:14])</pre>
sexPop <- possum[rowsfrom, c("sex", "Pop")]</pre>
n <- length(rowsfrom); ntimes <- 3</pre>
bootscores <- data.frame(scores1=numeric(ntimes*n), scores2=numeric(ntimes*n))</pre>
for (i in 1:ntimes){
  samprows <- sample(1:n, n, replace=TRUE)</pre>
  bootscores[n*(i-1)+(1:n), 1:2] <-
  prcomp(logpossum6to14[samprows, ])$x[, 1:2]
bootscores[, c("sex","Pop")] <- sexPop[samprows, ]</pre>
bootscores$sampleID <- factor(rep(1:ntimes, rep(n,ntimes)))</pre>
gph <- quickplot(x=scores1, y=scores2, colour=sex, size=I(1.0),</pre>
  asp=1, shape=Pop, facets=.~sampleID, data=bootscores) +
  scale_shape_discrete(solid=F)
gph + scale_colour_manual(values=c("m"="blue", "f"="red")) +
  xlab("First Principal Component") + ylab("Second Principal Component")
```

Subsection 9.1.3: Multi-dimensional scaling

Distance measures

Ordination

Binary data

Section 9.2: Principal component scores in regression

```
## Principal components: data frame socsupport (DAAG)
socsupport <- DAAG::socsupport
ss.pr1 <- prcomp(as.matrix(na.omit(socsupport[, 9:19])), retx=TRUE, scale=TRUE)</pre>
oldpar <- par(fg='gray40',col.axis='gray20',lwd=0.5,col.lab='gray20')
pairs(ss.pr1$x[, 1:3], col='gray40', gap=0.2)
par(oldpar)
summary(sort(ss.pr1$rotation[,1]))
## Note the very large maximum value
which.max(ss.pr1$x[,1])
## Try also boxplot(ss.pr1$x[,1])
## ss.pr1$x["36",1] ## Check that this returns 42
use <- complete.cases(socsupport[, 9:19])</pre>
use[36] <- FALSE
ss.pr <- prcomp(as.matrix(socsupport[use, 9:19]))</pre>
## Output from summary()
print(summary(ss.pr), digits=1) # Compare contributions
comp <- as.data.frame(ss.pr$x[,1:6])</pre>
ss.lm <- lm(socsupport[use, "BDI"] ~ ., data=comp)
signif(round(coef(summary(ss.lm)),5), digits=3)
print(ss.pr$rotation[, 1], digits=2)
## Plot BDI against first principal component score
gph <- xyplot(BDI ~ ss.pr$x[ ,1], groups=gender, data=socsupport[use,],</pre>
par.settings=simpleTheme(pch=1:2), auto.key=list(columns=2))
bw9 <- list(pch=c(1,3), list(text=9, points=5))</pre>
update(gph, scales=list(tck=0.5), par.settings=bw9,
xlab ="1st principal component")
```

Section 9.3: Cluster analysis

Subsection 9.3.1: Hierarchical Clustering

font=1, line=1, cex=1.15)

```
library(mvtnorm)
makeClust \leftarrow function(n=6, d1=4, d2=4, sigs=c(1, 1, 1, 1), seed=NULL){
  if(!is.null(seed))set.seed(seed)
  g1 \leftarrow rmvnorm(n, mean = c(-d1, d2), sigma=sigs[1]*diag(2))
  g2 \leftarrow rmvnorm(n, mean = c(d1,d2), sigma=sigs[2]*diag(2))
  g3 \leftarrow rmvnorm(n, mean = c(-d1, -d2), sigma=sigs[3]*diag(2))
  g4 \leftarrow rmvnorm(n, mean = c(d1, -d2), sigma=sigs[4]*diag(2))
  rbind(g1,g2,g3,g4)
}
## Code for the plots
datA <- makeClust(seed=35151)</pre>
datB <- makeClust(d2=16, seed=35151)
datC <- makeClust(d1=2,d2=2, seed=35151)</pre>
plot(datA, xlab="X1", ylab="X2", fg="gray")
title(main="A: 4blobsA", adj=0, line=0.5, font.main=1)
## Repeat previous two lines for datB and datC
plot(datB, xlab="X1", ylab="X2", fg="gray")
title(main="B: 4blobsB", adj=0, line=0.5, font.main=1)
plot(datC, xlab="X1", ylab="X2", fg="gray")
title(main="C: 4blobsC", adj=0, line=0.5, font.main=1)
## Possible alternative
config <- c('Equidistant blobs', 'Pulled vertically', 'Closer centers')</pre>
dat123 <- cbind(as.data.frame(rbind(datA, datB, datC)),</pre>
                                gp=factor(rep(1:3, rep(6*4,3)), labels=config))
xyplot(V2 ~ V1 | gp, data=dat123, scales=list(relation='free'),
       strip=strip.custom(factor.levels=config), between=list(x=0.5),
       par.settings=DAAG::DAAGtheme(color=F))
## Code for single linkage plots: `?plot.hclust` gives help for the plot method
clusres_sing <- hclust(dist(datA), method="single")</pre>
par(fig=c(0,0.75,0,1))
plot(clusres_sing, sub="", xlab="", ylab="Distance joined", adj=0.5,
     main="", fg="gray")
mtext('A: Single linkage cluster dendrogram, for 4blobsA layout', side=3, adj=0,
```

```
par(fig=c(0.72,1,0,1), new=TRUE)
membs <- cutree(clusres_sing, 4)</pre>
col4= RColorBrewer::brewer.pal(4,'Set1')
plot(datA, xlab="X1", ylab="X2", col=col4[membs], fg='gray', pch=membs+1)
mtext('B: 4blobsA, by color', side=3, adj=1.0, font=1, cex=1.15, line=1)
## To see plots from 'average' and 'complete' linkage methods, do:
# plot(hclust(dist(datB), method="average"))
# plot(hclust(dist(datC), method="complete"))
## Dendrograms from data where blobs were pulled vertically
## Follow each use of `hclust()` with a `plot()` command
sclusres_sing <- hclust(dist(datB), method="single")</pre>
plot(sclusres_sing, sub="", xlab="", ylab="Distance Joined", main="")
title(main='A: Single linkage, blobs pulled vertically (4blobsB)',
      adj=0, font.main=1)
sclusres_sing_s <- hclust(dist(scale(datA)), method="single")</pre>
plot(sclusres_sing_s, sub="", xlab="", ylab="Distance Joined", main="")
title(main='B: Single linkage, (4blobsB, rescaled to variance 1)',
      adj=0, font.main=1)
# sclusres_avg_s <- hclust(dist(scale(datB)), method="average")</pre>
# #plot(sclusres_avg_s, sub="", xlab="", ylab="")
# sclusres_comp_s <- hclust(dist(scale(datB)), method="complete")</pre>
# #plot(sclusres_comp_s, sub="", xlab="", ylab="")
## Code. Follow each use of `hclust()` with a `plot()` command
clusres_sing2 <- hclust(dist(datC), method="single")</pre>
plot(clusres_sing2, sub="", xlab="", ylab="", cex=1.25, cex.main=1.65,
     main="A: Single linkage, closer clusters (4blobsC)", adj=0, font.main=1)
clusres_avg2 <- hclust(dist(datC), method="average")</pre>
plot(clusres_avg2, sub="", xlab="", ylab="", cex=1.25, cex.main=1.65,
```

Subsection 9.3.2: *k*-Means Clustering

clusres comp2 <- hclust(dist(datC), method="complete")</pre>

```
set.seed(35151)
kdat <- makeClust(n=100, d1=5, d2=5, sigs=c(.5, .5, 6, 6))</pre>
```

plot(clusres_comp2, sub="", xlab="", ylab="", cex=1.25, cex.main=1.65,

main="B: Average linkage, closer clusters (4blobsC)", adj=0, font.main=1)

main="C: Complete linkage, closer clusters (4blobsC)", adj=0, font.main=1)

Comments on k-means and hierarchical clustering

Subsection 9.3.3: Mixture model-based clustering

```
## Code
plotMix2 < -function(taus=c(.5, .5), means=c(10,15), sds=c(3,1), xlims=c(0,20)){
  curve(taus[1]*dnorm(x, mean=means[1], sd=sds[1]) +
          taus[2]*dnorm(x, mean=means[2], sd=sds[2]),
        from=xlims[1], to=xlims[2], ylab="Density", fg="gray")
  curve(taus[1]*dnorm(x, mean=means[1], sd=sds[1]),
        from=xlims[1], to=xlims[2], col="red", lty=2, add=TRUE, fg="gray")
  curve(taus[2]*dnorm(x, mean=means[2], sd=sds[2]),
        from=xlims[1], to=xlims[2], col="blue", lty=3, add=TRUE, fg="gray")
plotMix2(taus=c(.2, .8))
plotMix2(taus=c(.5, .5))
plotMix2(taus=c(.9, .1))
library(teigen)
possml \leftarrow na.omit(DAAG::possum[,c(3,9:11)])
set.seed(513451)
gaus_fit <- teigen(possml[,2:4], models="UUUU", gauss=TRUE, verbose=FALSE,</pre>
                    scale=FALSE)
## BIC values are plotted against number of groups
gaus_fit$allbic
plot(gaus_fit$allbic, type="b", ylab="", xlab="Number of Groups", fg="gray")
mtext(side=2, line=3.5, "BIC", las=0)
axis(1, at=1:9, fg="gray")
```

```
table(possml$Pop, gaus_fit$classification)
```

```
par(fig=c(0, 0.5, 0.5, 1))
plot(gaus_fit, what="contour", xmarg=1, ymarg=2, draw.legend=FALSE, fg="gray")
## See ?teigen::plot.teigen for details of the plot command used here.
par(fig=c(0, 0.5, 0, 0.5), new=TRUE)
plot(gaus_fit, what="contour", xmarg=1, ymarg=3, draw.legend=FALSE, fg="gray")
par(fig=c(0.5, 1, 0, 0.5), new=TRUE)
plot(gaus_fit, what="contour", xmarg=2, ymarg=3, draw.legend=FALSE, fg="gray")
par(fig=c(0,1,0,1))
```

Issues of high parametrization and scaling

Subsection 9.3.4: Relationship between k-means and mixture models

Section 9.4: Discriminant analysis

Subsection 9.4.1: Example – plant architecture

```
leafshape17 <- DAAG::leafshape17
plot(bladelen ~ bladewid, data=leafshape17, pch=c(1,3)[arch+1])
## For panel B, specify log="xy" in the call to plot()</pre>
```

Logistic regression, versus linear discriminant analysis

Subsection 9.4.2: Logistic regression

```
## Fit logistic regression model
leafshape17 <- DAAG::leafshape17
leaf17.glm <- glm(arch ~ logwid + loglen, family=binomial(link=logit),
data=leafshape17)
print(DAAG::sumry(leaf17.glm)$coef, digits=2)</pre>
```

Predictive accuracy

```
set.seed(29)
leaf17.cv <- DAAG::CVbinary(leaf17.glm)
tCV <- table(DAAG::leafshape17[["arch"]], round(leaf17.cv$cvhat))
rownames(tCV) <- colnames(tCV) <- c("0=Plagiotropic","1=Orthotropic")
cbind(tCV, "Proportion correct"=c(tCV[1,1], tCV[2,2])/(tCV[,1]+tCV[,2]))</pre>
```

```
round(unlist(leaf17.cv[c("acc.training","acc.cv")]),3)
```

Subsection 9.4.3: Linear discriminant analysis

```
suppressPackageStartupMessages(library(MASS))
## Discriminant analysis; data frame leafshape17 (DAAG)
leaf17.lda <- lda(arch ~ logwid+loglen, data=DAAG::leafshape17)
print(leaf17.lda)</pre>
```

Assessments of predictive accuracy

```
set.seed(29)
leaf17cv.lda <- lda(arch ~ logwid+loglen, data=leafshape17, CV=TRUE)
## the list element 'class' gives the predicted class
## The list element 'posterior' holds posterior probabilities
tab <- table(leafshape17$arch, leaf17cv.lda$class)
rownames(tab) <- colnames(tab) <- c("0=Plagiotropic","1=Orthotropic")
cbind(tab, "Proportion correct"=c(tCV[1,1], tCV[2,2])/(tCV[,1]+tCV[,2]))
cbind(tab, c(tab[1,1], class.acc=tab[2,2])/(tab[,1]+tab[,2]))
cat("Overall proportion correct =", sum(tab[row(tab)==col(tab)])/sum(tab), "\n")</pre>
```

The function qda(), and other alternatives to lda()

Subsection 9.4.4: An example with more than two groups

```
plot(possum.lda, dimen=3, col=1:9)
# Scatterplot matrix - scores on 1st 3 canonical variates
# See `?plot.lda` for details of the generic lda plot function
```

```
## Linear discriminant calculations for possum data
print(possum.lda, digits=3)
```

Section 9.5: *High-dimensional data — RNA-Seq gene expression

Setup for installing and using Bioconductor packages

```
## For latest details, see: https://www.bioconductor.org/install/
if (!require("BiocManager", quietly = TRUE))
    install.packages("BiocManager")
BiocManager::install()
BiocManager::install('limma', 'multtest')
```

*Brief note on mRNA technical issues

Subsection 9.5.1: Data and design matrix setup

```
counts <- DAAGbio::plantStressCounts
colSums(counts)

## Require at least 3 counts per million that are > 1
keep <- rowSums(counts)>=3
counts <- counts[keep,]

treatment <- factor(rep(c("CTL", "L", "D"), rep(3,3)))
design <- model.matrix(~0+treatment)
colnames(design) <- levels(treatment)</pre>
```

A two-dimensional representation

```
library(limma)
v <- voom(counts, design, plot=TRUE)</pre>
```

```
par(oma=c(0,0,1,0))
library(limma)
v <- voom(counts, design, plot=TRUE)
firstchar <- substring(colnames(counts),1,1)
plotMDS(counts, labels=paste0(firstchar, rep(1:3,3)), cex=0.8)
box(col="gray")
mtext(side=3, line=0.4, adj=0, "MDS summary plot")
mtext(side=3, line=-0.25, adj=0.105, "A", outer=TRUE)
mtext(side=3, line=-0.25, adj=0.605, "B", outer=TRUE)</pre>
```

Fitting the model

```
fit <- lmFit(v, design)

contrs <- c("D-CTL", "L-CTL", "L-D")
contr.matrix <- makeContrasts(contrasts=contrs,
levels=levels(treatment))
fit2 <- contrasts.fit(fit, contr.matrix)
efit2 <- eBayes(fit2)</pre>
```

Subsection 9.5.2: From p-values to false discovery rate (FDR)

```
## First contrast only; Drought-CTL
print(round(topTable(efit2, coef=1, number=4),15), digits=3)

round(sort(p.adjust(p=efit2$p.value[,1], method="BH"))[1:4], 15) # Not run

round(topTable(efit2, number=4), 15)

head(decideTests(fit2),5)

summary(decideTests(fit2))
## Try also
## summary(decideTests(fit2, p.value=0.001))
```

Section 9.6: High dimensional data from expression arrays

Subsection 9.6.1: Molecular classification of cancer — an older technology

Breakdown of ALL B-type data, with one observation excluded

```
library(hddplot)
data(golubInfo)
with(golubInfo, table(cancer, tissue.mf))
```

```
## Identify allB samples that are BM:f or BM:m or PB:m
subsetB <- with(golubInfo,
cancer=="allB" & tissue.mf%in%c("BM:f","BM:m","PB:m"))
## Separate off the relevant columns of the matrix Golub
## NB: variables (rows) by cases (columns)
GolubB <- with(golubInfo, Golub[, subsetB])
## Form vector that identifies these as BM:f or BM:m or PB:m
tissue.mfB <- with(golubInfo, tissue.mf[subsetB, drop=TRUE])
## Change the level names to leave out the colons
levels(tissue.mfB) <- list("b_f"="BM:f", "b_m"="BM:m", "PBm"="PB:m")</pre>
```

Subsection 9.6.2: Classifications and associated graphs

Preliminary data manipulation

```
## Display distributions for the first 20 observations
boxplot(data.frame(GolubB[, 1:20])) # First 20 columns (observations)
## Random selection of 20 rows (features)
boxplot(data.frame(GolubB[sample(1:7129, 20), ]))
```

Flawed graphs

```
colr <- c("red","blue","gray40", "magenta")
tissue.mf <- golubInfo[, "tissue.mf"]
cancer <- golubInfo[, "cancer"]
G.PBf <- Golub[, tissue.mf=="PB:f" & cancer=="allB", drop=FALSE]
set.seed(41)
rGolubB <- matrix(rnorm(prod(dim(GolubB))), nrow=dim(GolubB)[1])
rownames(rGolubB) <- rownames(Golub)
rG.PBf <- matrix(rnorm(prod(dim(G.PBf))), nrow=dim(G.PBf)[1])</pre>
```

```
plot2 <- function(x = GolubB, cl=tissue.mfB, x.omit=Golub.PBf, cl.omit="PBf",</pre>
                   ncol = length(cl), nfeatures=12, device = "", seed = 37,
                   pretext="", colr=1:3, levnames = NULL,
                   ylab="2nd discriminant function"){
  cl <- factor(cl)</pre>
  if(!is.null(levnames))levels(cl) <- levnames</pre>
  ord15 <- orderFeatures(x, cl=cl)[1:nfeatures]</pre>
  dfB \leftarrow t(x[ord15, ])
  dfB.lda <- lda(dfB, grouping=cl)</pre>
  scores <- predict(dfB.lda, dimen=2)$x</pre>
  df.PBf <- data.frame(t(x.omit[ord15, drop=FALSE]))</pre>
  colnames(df.PBf) <- colnames(dfB)</pre>
  scores.other <- predict(dfB.lda, newdata=df.PBf)$x</pre>
  scoreplot(list(scores=scores, cl=cl, other=scores.other, cl.other=cl.omit,
                  fg="gray", params=list(other=list(pch=4, cex=1.5)),
            xlab="1st discriminant function", ylab=ylab)
plot2(x = GolubB, cl = tissue.mfB, x.omit=G.PBf, cl.omit="PBf",
      nfeatures=15, device = "", seed = 37, ylab="2nd discriminant function",
      colr=colr, pretext="A: ALL B-cell:")
plot2(x = rGolubB, cl = tissue.mfB, x.omit=rG.PBf, cl.omit="Gp 4",
     device = "", seed = 37, colr=colr, levnames = c("Gp 1", "Gp 2", "Gp 3"),
     pretext="B: Random data:", ylab="")
## Uses orderFeatures() (hddplot); see below
ord15 <- orderFeatures(GolubB, cl=tissue.mfB)[1:15]</pre>
## Panel A: Take 1st 15 features & transpose to observations by features
dfB15 <- data.frame(t(GolubB[ord15, ]))</pre>
dfB15.lda <- MASS::lda(dfB15, grouping=tissue.mfB)
```

```
## Panel B: Repeat plot, now with random normal data
simscores <- simulateScores(nrow=7129, cl=rep(1:3, c(19,10,2)),
cl.other=4, nfeatures=15, seed=41)
# Returns list elements: scores, cl, scores.other & cl.other
scoreplot(simscores)</pre>
```

Subsection 9.6.3: The mean-variance relationship

```
par(oma=c(0,0,1,0))
designG <- model.matrix(~0+tissue.mfB)
colnames(designG) <- levels(tissue.mfB)
vG <- vooma(GolubB, designG, plot=TRUE)  # Panel A
plotMDS(vG, pch=unclass(tissue.mfB), cex=0.8)  # Panel B
leglabs <- c("BM:female", "BM:male", "PB:female")
legend(x="bottomright", bty="n", legend=leglabs, pch=1:3)
mtext(side=3, line=0.4, adj=0, "MDS summary plot")
mtext(side=3, line=-0.275, adj=0.085, "A", outer=TRUE)
mtext(side=3, line=-0.275, adj=0.585, "B", outer=TRUE)</pre>
```

Cross-validation to determine the optimum number of features

Cross-validation for a range of choices of number of features

```
## Cross-validation to determine the optimum number of features
## 10-fold (x4). Warning messages are omitted.
## Accuracy measure will be: tissue.mfB.cv$acc.cv
tissue.mfB.cv <- cvdisc(GolubB, cl=tissue.mfB, nfeatures=1:23,
nfold=c(10,4), print.progress=FALSE)
## Defective measures will be in acc.resub (resubstitution)
## and acc.sel1 (select features prior to cross-validation)
tissue.mfB.badcv <- defectiveCVdisc(GolubB, cl=tissue.mfB,
foldids=tissue.mfB.cv$folds, nfeatures=1:23)
##
## Calculations for random normal data:
set.seed(43)
rGolubB <- matrix(rnorm(prod(dim(GolubB))), nrow=nrow(GolubB))
rtissue.mfB.cv <- cvdisc(rGolubB, cl=tissue.mfB, nfeatures=1:23,
nfold=c(10,4), print.progress=FALSE)
rtissue.mfB.badcv <- defectiveCVdisc(rGolubB, cl=tissue.mfB,
```

```
nfeatures=1:23,
foldids=rtissue.mfB.cv$folds)
```

Which features?

```
genelist <- matrix(tissue.mfB.cv$genelist[1:3, ,], nrow=3)
tab <- table(genelist, row(genelist))
ord <- order(tab[,1], tab[,2], tab[,3], decreasing=TRUE)
tab[ord,]</pre>
```

Subsection 9.6.4: Graphs derived from the cross-validation process

```
## Uses tissue.mfB.acc from above
tissue.mfB.scores <-
cvscores(cvlist = tissue.mfB.cv, nfeatures = 3, cl.other = NULL)
scoreplot(scorelist = tissue.mfB.scores, cl.circle=NULL,
prefix="B-cell subset -", fg='gray')</pre>
```

The key role of cross-validation

Subsection 9.6.5: Estimating contrasts, and calculating False Discovery Rates

```
fitG <- lmFit(vG, designG)
contrs <- c("b_f-b_m", "b_f-PBm", "b_m-PBm")
contr.matrix <- makeContrasts(contrasts=contrs,
levels=levels(tissue.mfB))
fit2 <- contrasts.fit(fitG, contr.matrix)
fit2 <- eBayes(fit2)</pre>
```

From p-values to false discovery rate (FDR)

```
print(topTable(fit2, number=5), digits=2)

summary(decideTests(fit2))
## Try also
## summary(decideTests(fit2, p.value=0.001))
```

Distributional extremes

Section 9.7: Causal inference from observational data — balance and matching

Subsection 9.7.1: Tools for the task

```
PGtheme <- DAAG::DAAGtheme(color=TRUE)
library(DAAG)
if(!require(grid))return("Package 'grid' is not installed -- cannot proceed")
dsetnames <- c("nsw-ctl", "nsw-trt", "psid1", "psid2", "psid3",</pre>
                "cps1", "cps2", "cps3")
colrs <- c("gray", "black", PGtheme$superpose.line$col[1:3])</pre>
lty \leftarrow c(1,2,1,1,1)
lwd \leftarrow c(1,0.75,0.75,0.75,0.75)
denplot <-
  function(sel=c(1:2,6:8), yvar="re75", offset=30, ylim=c(0,1.75),
    from=NULL, at=c(.5,1,1.5), labels=paste(at), bw="nrd0",
    ylab="Density", takelog=TRUE, col.axis="black"){
      nzre <- unlist(lapply(list(subset(nswdemo, trt==0),</pre>
                                   subset(nswdemo, trt==1),
                                   psid1, psid2, psid3, cps1, cps2, cps3)[sel],
                              function(x){z <- x[,yvar]; z[z>0]}))
num <- unlist(lapply(list(subset(nswdemo, trt==0), subset(nswdemo, trt==1),</pre>
                            psid1, psid2, psid3, cps1, cps2, cps3),
                      function(x){z \leftarrow x[,yvar]; sum(z>0)}))
xy <- data.frame(nzre=nzre, fac = factor(rep(dsetnames[sel], num[sel]),</pre>
                  levels=dsetnames[sel]))
if(takelog) {
```

```
y <- log(xy$nzre+offset)</pre>
xlab <- paste("log(", yvar, "+", offset, ")", sep="")} else</pre>
  {
  y <- xy$nzre
  xlab <- yvar</pre>
densityplot(~ y, groups=fac, data=xy, bw=bw, from=from,
  scales=list(y=list(at=at, labels=labels, col=col.axis), tck=0.25),
  plot.points=FALSE, col=colrs[1:5], lwd=lwd, lty=lty,
  key=list(x=0.01, y=0.99, text=list(dsetnames[sel[3:5]]), col=colrs[3:5],
           cex=0.75, lines=list(lwd=rep(1.5,3)), between=1),
  par.settings=list(col=colrs, lty=lty, cex=0.75, lwd=lwd,
                    fontsize=list(text=9, points=5)),
  fg="gray", ylim=ylim, ylab=ylab, xlab=xlab)
## Plot base graph; overlay with lattice graphs on same page
par(fig=c(0,1,0,1), mar=c(0,0,0,0))
plot(0:1,0:1, axes=FALSE, type="n", bty="n", xlab="", ylab="")
legend(x="top",legend=dsetnames[1:2], lty=1:2, lwd=c(1,0.75),
       col=colrs[1:2], bty="n", ncol=2, yjust=0.75)
print(denplot(), position=c(0, 0, 0.32, 0.505), newpage=FALSE)
print(denplot(1:5, ylab=" ", col.axis="white"),
      position=c(0.21, 0, .53, 0.505), newpage=FALSE)
print(denplot(ylab=" ", yvar="re78", col.axis="white"),
      position=c(0.47, 0, 0.79, 0.505), newpage=FALSE)
print(denplot(1:5, ylab=" ", yvar="re78", col.axis="white"),
      position=c(0.68, 0, 1, 0.505), newpage=FALSE)
## Age
print(denplot(yvar="age", takelog=FALSE, ylim=c(0,0.1), from=16,
      at=c(.02,.04,.06,.08), labels=c(".02",".04",".06",".08")),
      position=c(0, 0.475, 0.32, .98), newpage=FALSE)
print(denplot(1:5, yvar="age", takelog=FALSE, ylim=c(0,0.1), from=16,
      at=c(.02,.04,.06,.08), labels=c(".02",".04",".06",".08"),
      ylab=" ", col.axis="white"),
      position=c(0.21, 0.475, .53, .98), newpage=FALSE)
## educ
print(denplot(1:5, yvar="educ", takelog=FALSE, ylim=c(0,0.5), bw=0.5,
      at=c(.1,.2,.3,.4), ylab="""),
      position=c(0.47, 0.475, .79, .98), newpage=FALSE)
print(denplot(yvar="educ", takelog=FALSE, ylim=c(0,0.75), bw=0.5,
      at=c(.1,.2,.3,.4), ylab=" ", col.axis="white"),
      position=c(0.68, 0.475, 1, .98), newpage=FALSE)
```

```
addControl <-
function(control, offset=30){
  nam <- deparse(substitute(control))
  if(nam=="nswdemo")nsw0 <- nswdemo else
    nsw0 <- rbind(control, subset(DAAG::nswdemo, trt==1))
  nsw0$z75 <- factor(nsw0$re75==0, labels=c("0",">0"))
  nsw0$ethnicid <- factor(with(nsw0, ifelse(black==1, "black",
        ifelse(hisp==1, "hisp", "other"))), levels=c("other","black","hisp"))
  nsw0 <- nsw0[, -match(c("black","hisp"), names(nsw0))]
  nsw0
}</pre>
```

```
with(trtdat, table(pres74,z75))
```

Subsection 9.7.2: Regression comparisons

Regression calculations

```
nsw.gam <- gam(log(re78+30)~ trt + ethnicid + z75 + nodeg + s(age) + s(educ) + log(re75+30), data=nsw)
```

Subsection 9.7.3: The use of scores to replace covariates

Subsection 9.7.4: Two-dimensional representation using randomForest proximities

```
suppressPackageStartupMessages(library(randomForest))
form <- trt ~ age + educ + ethnicid + marr + nodeg + z75 + re75log
nsw.rf <- randomForest(form, data=nsw, sampsize=c(297,297))</pre>
```

```
p.rf <- predict(nsw.rf,type="prob")[,2]</pre>
sc.rf < log((p.rf+0.001)/(1-p.rf+0.001))
omitn <- match(c("PropScore","weights","subclass"), names(dat2RF), nomatch=0)</pre>
matchISO.rf <-matchit(trt ~ age + educ + ethnicid + marr + nodeg + z75 +
                      re75log, ratio=1, data=dat2RF[,-omitn], distance=isoScores[,1])
## summary(match.rf,un=F,improvement=F)
## summary(match.rf, un=F, interactions=T, improvement=F)$sum.matched[,1:4]
## In the first place, look only at the first 4 columns
dat1RF <- match.data(matchISO.rf, distance="PropScore")</pre>
dat1RF.lm <- lm(re78log ~ trt, data = dat1RF, weights = weights)</pre>
                    # Allows use of `vcovCL()` from the `sandwich` package
library(sandwich)
lmtest::coeftest(dat1RF.lm, vcov. = vcovCL, cluster = ~subclass)
## Check for increase in number with non-zero earnings
dat1RF.glm <- glm(I(re78>0) ~ trt, data = dat1RF, weights = weights,
                  family=binomial)
lmtest::coeftest(dat1RF.glm, vcov. = vcovCL, cluster = ~subclass)
```

Derivation and investigation of scores

```
library(mgcv)
formG <- trt ~ ethnicid + marr+ z75 + s(age) + s(educ) + s(re75log)
nsw.gam <- gam(formG, family=binomial(link="logit"), data=nsw)</pre>
pred <- predict(nsw.gam, type='response')</pre>
table(nsw$trt, round(pred))
## Alternative
library(splines) ## Fit normal cubic splines using splines::ns()
formNS <- trt ~ ethnicid + marr+ z75 + ns(age,2) +
ns(educ) + ns(re75log,3)
nsw.glm <- glm(formNS, family=binomial(link="logit"), data=nsw)</pre>
pred <- predict(nsw.glm, type='response')</pre>
table(nsw$trt, round(pred))
cbind(AIC(nsw.glm,nsw.gam), BIC(nsw.glm, nsw.gam))
## Include factor by factor and variable interactions with ethnicid
## and marr (Result not shown)
formGx <- trt ~ (ethnicid+marr+z75)^2 + s(age, by=ethnicid)+
                s(educ, by=ethnicid) + s(re75log,by=ethnicid)+
                s(age, by=marr) + s(educ, by=marr) + s(re75log,by=marr)
```

```
nswx.gam <- gam(formula = formGx, data = nsw, family=binomial(link = "logit"))</pre>
predx <- predict(nswx.gam, type='response')</pre>
table(nsw$trt, round(predx))
AIC(nsw.glm,nsw.gam,nswx.gam)
library(MatchIt)
## Use data frame that omits re74. Otherwise matchit() will generate NAs
## where they occur in re74, even though re74 is not in the model formula.
nswG <- nsw[, c("trt", "age", "educ", "ethnicid", "marr", "nodeg", "z75",</pre>
                "re75log", "re78log", "re78")]
formG <- trt ~ ethnicid + marr+ z75 + s(age) + s(educ) + s(re75log)
match.gam <- matchit(formula = formG, data = nswG, method = "nearest",</pre>
                     distance = "gam", link = "logit", reestimate=TRUE)
datG <- match.data(match.gam, distance="PropScore")</pre>
## Summary information
match.gam
## summary(match.gam,un=F,improvement=F)
## summary(match.gam, un=F, interactions=T, improvement=F)$sum.matched[,1:4]
## In the first place, look only at the first 4 columns
suppressPackageStartupMessages(library(gridExtra))
suppressPackageStartupMessages(library(ggplot2))
suppressPackageStartupMessages(library(cobalt))
gg1<- cobalt::love.plot(match.gam, position="bottom", grid=TRUE,
                         star.char="",stars='raw') +
  ggtitle("A: Differences from balance") +
  theme(plot.title = element_text(hjust=0, vjust=0.5, size=11),
        plot.margin=unit(c(9,15,0,9), 'pt'))
sub <- match(with(subset(datG, trt=="Control"), subclass),</pre>
             with(subset(datG, trt=="Treat"), subclass))
datGpaired <- cbind(subset(datG, trt=="Treat"),</pre>
                     with(subset(datG, trt=="Control")[sub,],
                     cbind("Cre78log"=re78log,"CPropScore"=PropScore)))
gg2 <- ggplot(datGpaired)+
  geom_point(aes(PropScore,I(re78log-Cre78log)), size=1)+
  geom_smooth(aes(PropScore,I(re78log-Cre78log)), method = "gam",
              formula = y \sim s(x, bs = "cs")) +
  xlab("Propensity score for treated")+
  ylab("Treatment vs control differences") +
  ggtitle("B: Treatment vs control differences") +
  theme(plot.title = element_text(hjust=0, vjust=0.5, size=11),
```

plot.margin=unit(c(9,9,0,15), 'pt'))

```
grid.arrange(gg1, gg2, ncol=2)
```

```
library(sandwich)
datG.lm <- lm(re78log ~ trt, data = datG, weights = weights)
## With 1:1 matching, the weights argument is not really needed
## Print first two coefficients only.
lmtest::coeftest(datG.lm, vcov. = vcovCL, cluster = ~subclass)[1:2,]
## Check number whose income was greater than 0
datG.glm <- glm(I(re78>0) ~ trt, data = datG, weights = weights, family=binomial)
lmtest::coeftest(datG.glm, vcov. = vcovCL, cluster = ~subclass)[1:2,]
```

Alternative matching approaches

Subsection 9.7.5: Coarsened exact matching

```
form <- trt ~ age + educ + ethnicid + marr + nodeg + z75 + re75log
match5.cem <- matchit(formula=form, data=nswG, method="cem", cutpoints=5)
datcem5 <- match.data(match5.cem)
match6.cem <- matchit(formula=form, data=nswG, method="cem", cutpoints=6)
datcem6 <- match.data(match6.cem)
## Show the effect of adding another cutpoint
match5.cem
match6.cem</pre>
```

```
library(sandwich)
datcem5.lm <- lm(re78log ~ trt, data = datcem5, weights = weights)
## The function vcovHC() provides cluster robust standard errors
lmtest::coeftest(datcem5.lm, vcov. = vcovHC)
## Estimate treatment effect on number with some earnings:
datcem6.glm <- glm(I(re78>0) ~ trt, data = datcem6, weights = weights,
family=binomial)
lmtest::coeftest(datcem6.glm, vcov. = vcovHC)
```

Section 9.8: Multiple imputation

```
suppressPackageStartupMessages(library(mice))
Boys <- with(subset(mice::boys, age>=9),
             data.frame(age=age, loghgt=log(hgt), logbmi=log(bmi), loghc=log(hc)))
(Pattern <- md.pattern(Boys, plot=F))
set.seed(31)
                    # Set to reproduce result shown
PatternB <- rbind(Pattern[-c(1,nrow(Pattern)), -ncol(Pattern)],</pre>
             c(0,1,1,1), c(0,1,0,0), c(0,0,1,0))
boys <- rbind(ic(Boys),</pre>
              ampute(cc(Boys), pattern=PatternB, freq=c(.3,.15,.15,.2,.1,.1),
                      prop=0.75) $amp)
md.pattern(boys, plot=FALSE)
set.seed(17)
                    # Set to reproduce result shown
out <- capture.output(</pre>
                               # Evaluate; send screen output to text string
  boys.mids <- mice(boys, method='pmm', m=8) )</pre>
impDFs <- complete(boys.mids, action='all') # Returns a list of m=8 dataframes</pre>
## Average over imputed dataframes (use for exploratory purposes only)
impArray <- sapply(impDFs, function(x)as.matrix(x), simplify='array')</pre>
boysAv <- as.data.frame(apply(impArray, 1:2, mean))</pre>
fits <- with(boys.mids, lm(logbmi~age+loghgt))</pre>
pool.coef <- summary(pool(fits)) # Include in table below</pre>
## 2) Regression that leaves out rows with NAs
omitNArows.coef <- coef(summary(lm(logbmi~age+loghgt, data=boys)))</pre>
## 3) Regression fit to average over data frames after imputation
boysAv.coef <- coef(summary(lm(logbmi~age+loghgt, data=boysAv)))</pre>
## 4) Fit to original data, with 36 rows had missing data
Orig.coef <- coef(summary(lm(logbmi ~ age+loghgt, data=Boys)))</pre>
ctab <- cbind(summary(pool(fits))[,2:3], omitNArows.coef[,1:2], boysAv.coef[,1:2],</pre>
      Orig.coef[,1:2])
tab <- setNames(cbind(ctab[,c(1,3,5,7)], ctab[,c(2,4,6,8)]),
                 paste0(rep(c('Est', 'SE'), c(4,4)), rep(1:4, 2)))
round(tab,3)
```

Time series cross-sectional data – an example

```
airquality <- datasets::airquality
airq <- cbind(airquality[, 1:4], day=1:nrow(airquality))
    # 'day' (starting May 1) replaces columns 'Month' & 'Day')
## Replace `Ozone` with `rt4ozone`:
airq <- cbind(rt4ozone=airq$Ozone^0.25, airq[,-1])</pre>
```

```
airq.imp <- mice(airq, m=20, print=FALSE)
## 20 imputations shows up issues of concern very clearly</pre>
```

```
## Code for figure
out <- micemd::overimpute(airq.imp)</pre>
```

Some further points

Section 9.9: Further reading

Data with more variables than observations

Causal inference

Multiple imputation

Section 9.10: Exercises

9.3

```
library(DAAG)
oz::oz(sections=c(3:5, 11:16))
names(possumsites)[1:2] <- c("long", "lat")
with(possumsites, {
points(long, lat);
text(long, lat, row.names(possumsites), pos=c(2,4,2,2,4,2,2))
})</pre>
```

```
data(wine, package='gclus')
mat <- with(wine,
   round(1-cor(cbind(Alcohol, Malic, Magnesium, Phenols, Flavanoids)),2))
colnames(mat) <- rownames(mat) <- 1:5
print(mat)</pre>
```

9.9a

```
confusion` <-
function(actual, predicted, digits=4){
  tab <- table(actual, predicted)
  confuse <- apply(tab, 1, function(x)x/sum(x))
  print(round(confuse, digits))
  acc <- sum(tab[row(tab)==col(tab)])/sum(tab)
  invisible(print(c("Overall accuracy" = round(acc,digits))))
}
data(Vehicle, package="mlbench")
lhat <- MASS::lda(Class ~ ., data=Vehicle, CV=TRUE)$class
qhat <- MASS::qda(Class ~ ., data=Vehicle, CV=TRUE)$class
DAAG::confusion(Vehicle$Class, lhat)
DAAG::confusion(Vehicle$Class, qhat)
randomForest::randomForest(Class ~ ., data=Vehicle, CV=TRUE)</pre>
```

9.9c

9.10

```
library(ape); library(MASS)
library(DAAGbio)
primates.dna <- as.DNAbin(primateDNA)
primates.dist <- dist.dna(primates.dna, model="K80")
primates.cmd <- cmdscale(primates.dist)
eqscplot(primates.cmd)
rtleft <- c(4,2,4,2)[unclass(cut(primates.cmd[,1], breaks=4))]
text(primates.cmd, labels=row.names(primates.cmd), pos=rtleft)</pre>
```

```
d <- dist(primates.cmd)
sum((d-primates.dist)^2)/sum(primates.dist^2)</pre>
```

9.11

9.15

```
wine$Class <- factor(Wine$Class)
wine.rf <- randomForest(x=wine[,-1], y=wine$Class)

if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
  code <- knitr::knit_code$get()
  txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))
  writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/ch9.R")
}</pre>
```

10 Appendix A: The R System – A Brief Overview

On options for working with the code see the vignettes Ch1-Learning and UsingCode.

* Packages required (plus any dependencies)

DAAG dplyr tidyr tibble MASS gplots plotrix latticeExtra RColorBrewer Additionally, knitr and Hmisc are required in order to process the qmd source file.

The R System – A Brief Overview

Section 10.1: Getting started with R

Subsection 10.1.1: Learn by typing code at the command line

```
> ## Arithmetic calculations. See the help page `?Arithmetic` {-}
> 2*3+10
                    # The symbol `*` denotes 'multiply'
> ## Use the `c()` function to join results into a numeric vector
> c(sqrt(10), 2*3+10, sqrt(10), 2^3) # 2^3 is 2 to the power of 3
> ## R knows about pi
> 2*pi*6378
                # Approximate circumference of earth at equator (km)
?help
                   # Get information on the use of `help()`
                  # Or, type help(sqrt)
?sqrt
?Arithmetic
                   # See, in similar vein ?Syntax
?!<!
                   # `?Comparison` finds the same help page
## Two commands on one line; Use ';' as separator
2*3*4+10; sqrt(10) ## Try also `cat(2*3*4+10, sqrt(10), sep='n')
## Convert CO2 carbon emissions from tonnes of carbon to tonnes of CO2
3.664*c(.53, 2.56, 9.62) ## Data are for 1900, 1960 & 2020
```

```
## Use `cat()` to print several items, with control of formatting cat(2*3*4+10, sqrt(10), '\n')
```

Assignment

```
## Convert from amounts of carbon to amounts of CO2 (billions of tonnes)
## and assign result to a named object
fossilCO2vals <- c(.53, 2.56, 9.62)*3.664 # Amounts in 1900, 1960, and 2020
# Equivalently `fossilCO2vals <- c(.53, 2.56, 9.62)*rep(3.664,3)`
## To assign and print, enclose in round brackets
(fossilCO2vals <- c(.53, 2.56, 9.62)*3.664)</pre>
```

```
3.664*c(.53,2.56, 9.62) -> fossilCO2vals
```

Entry of data at the command line, a graphics formula, and a graph

```
Year <- c(1900, 1920, 1940, 1960, 1980, 2000, 2020)

CO2 <- c(.53,.96,1.32,2.56,5.32,6.95,9.62)*3.664

## Now plot Carbon Dioxide emissions as a function of Year

plot(CO2 ~ Year, pch=16, fg="gray")
```

Grouping vectors togeher into data frames

```
CO2byYear <- data.frame(year=Year, co2gas=CO2)
CO2byYear  # Display the contents of the data frame.
rm(Year, CO2)  # Optionally, remove `Year` and `Carbon` from the workspace
plot(co2gas ~ year, data=CO2byYear, pch=16)

sqrt(10)  # Number of digits is determined by current seting
options(digits=2)  # Change until further notice,
sqrt(10)
```

Wide-ranging information access and searches

Section 10.2: R data structures

Subsection 10.2.1: Vectors, dates, and arrays

```
vehicles <- c("Compact", "Large", "Midsize", "Small", "Sporty", "Van")
c(T, F, F, T, T, F) # A logical vector, assuming F=FALSE and T=TRUE
## Character vector</pre>
```

```
## Character vector
mammals <- c("Rat", "Pig", "Rat", "Mouse", "Pig")
## Logical vector
rodent <- c("TRUE", "FALSE", "TRUE", "FALSE", "TRUE", "FALSE")
## From character vector `mammals`, create factor
mfac <- factor(mammals)
levels(mfac)
table(mfac)</pre>
```

Dates

```
day1 <- as.Date(c("2022-01-01", "2022-02-01", "2022-03-01"))
as.numeric(day1)  # Days since 1 January 1970
day1[3] - day1[2]</pre>
```

The use of square brackets to extract subsets of vectors

```
## Specify the indices of the elements that are to be extracted x <-c(3, 11, 8, 15, 12, 18) x[c(1,4:6)] # Elements in positions 1, 4, 5, and 6 ## Use negative indices to identify elements for omission x[-c(2,3)] # Positive and negative indices cannot be mixed ## Specify a vector of logical values. x > 10 # This generates a vector of logical values x[x > 10]
```

```
bodywt <- c(Cow=465, Goat=28, Donkey=187, Pig=192)
bodywt[c("Goat", "Pig")]</pre>
```

Matrices and arrays

```
arr123 <- array(1:24, dim=c(2,4,3))
## This prints as three 2 by 4 matrices. Print just the first of the three.
arr123[, 2, 1]  # Column 2 and index 1 of 3rd dimension
attributes(arr123)</pre>
```

Subsection 10.2.2: Factors

if(all(mf1==mf2))print(mf1)

```
gender <- c(rep("male",691), rep("female",692))
gender <- factor(gender) # From character vector, create factor
levels(gender) # Notice that `female` comes first

Gender <- factor(gender, levels=c("male", "female"))

mf1 <- factor(rep(c('male','female'),c(2,3)), labels=c("f", "m"))
## The following has the same result
mf2 <- factor(rep(c('male','female'), c(2,3)))</pre>
```

```
sum(gender=="male")
```

levels(mf2) <- c("f","m") # Assign new levels</pre>

```
table(chickwts$feed) # feed is a factor
source <- chickwts$feed
levels(source) <- c("milk","plant","plant","plant","plant","plant")
table(source)</pre>
```

Ordered factors

```
stress <- rep(c("low", "medium", "high"), 2)
ord.stress <- ordered(stress, levels=c("low", "medium", "high"))
ord.stress
ord.stress >= "medium"
```

Missing values in values of factors

Subsection 10.2.3: Operations with data frames

```
Cars93sum <- DAAG::Cars93.summary # Create copy in workspace Cars93sum
```

```
Cars93sum[4:6, 2:3]  # Extract rows 4 to 6 and columns 2 and 3

Cars93sum[6:4,]  # Extract rows in the order 6, 5, 4

Cars93sum[, 2:3]  # Extract columns 2 and 3

## Or, use negative integers to specify rows and/or columns to be omitted

Cars93sum[-(1:3), -c(1,4)]  # In each case, numbers must be all +ve or all -ve

## Specify row and/or column names

Cars93sum[c("Small", "Sporty", "Van"), c("Max.passengers", "No.of.cars")]
```

Data frames vs matrices

```
names(Cars93sum)[3] <- "numCars"
names(Cars9sum) <- c("minPass", "maxPass", "numCars", "code")</pre>
```

Using a data frame as a database - with() and within()

```
## trees (datasets) has data on Black Cherry Trees
with(trees, round(c(mean(Girth), median(Girth), sd(Girth)),1))

with(DAAG::pair65,  # stretch of rubber bands
    {lenchange = heated-ambient
        c(mean(lenchange), median(lenchange))
})

## Add variables `mph` and `gradient` to `DAAG::nihills`
nihr <- within(DAAG::nihills, {mph <- dist/time; gradient <- climb/dist})</pre>
```

Extracting rows from data frames

```
unlist(Cars93sum[1,])
```

Subsection 10.2.4: Data manipulation functions used in earlier chapters

Subsection 10.2.5: Writing data to a file, and reading data from a file

Data input from the RStudio menu — data frames vs tibbles

Subsection 10.2.6: Issues for working with data frames and tibbles

Extraction of columns from data frames and tibbles

```
sites <- DAAG::possumsites  # sites is then a data frame
sites[,3]  # returns a vector
sites[,3, drop=FALSE]  # returns a 1-column data frame

dplyr::as_tibble(sites)[,3]  # returns a 1-column tibble
dplyr::as_tibble(sites)[[3]]  # returns a vector
sites[[3]]  # returns a vector</pre>
```

Conversion between data frames and tibbles

```
attributes(DAAG::possumsites)[['row.names']]
possumSites <- tibble::as_tibble(DAAG::possumsites, rownames="Site")
possumSites</pre>
```

Subsection 10.2.7: Lists

```
## Summary statistics for 31 felled black cherry tree
## Median (middle value), range, number, units
htstats <- list(med=76, range=c(low=63,high=87), n=31, units="ft")
              # Show first two list elements only
htstats[1:2]
## The following are alternative ways to extract the second list element
                   # First list element (Can replace `2` by 'range')
htstats[2]
htstats[2][1] # A subset of a list is a list
htstats[[2]]; htstats$range; htstats[["range"]]
unlist(htstats[2]) # Contents of second list element, with composite names
unlist(htstats[2], use.names=F) # Elements have no names
tstats <- with(MASS::shoes, t.test(B, A, paired=TRUE))</pre>
names(tstats)
                   ## Names of list elements. See `?t.test` for details.
tstats[1]
                    ## Type tstats[1] to see the first list element
## Compact listing of contents list elements 1 to 5, which are all numeric
unlist(tstats[1:5]) ## With `unlist(tstats)` all elements become character
```

Section 10.3: Functions and operators

Subsection 10.3.1: Common useful built-in functions

```
## Data indices
length()  # number of elements in a vector or a list
order()  # x[order(x)] sorts x (by default, NAs are last)
which()  # which indices of a logical vector are `TRUE`
which.max()  # locates (first) maximum (NB, also: `which.min()`)
```

```
## Data manipulation
c() # join together (`concatenate`) elements or vectors or lists
diff()
          # vector of first differences
sort()
            # sort elements into order, by default omitting NAs
            # reverse the order of vector elements
rev()
t()
            # transpose matrix or data frame
             # (a data frame is first coerced to a matrixwith()
            # do computation using columns of specified data frame
with()
## Data summary
mean() # mean of the elements of a vector
median()
            # median of the elements of a vector
            # minimum and maximum value elements of vector
range()
unique()
            # form the vector of distinct values
## List function arguments
args()
             # information on the arguments to a function
## Obtain details
head() # display first few rows (by default 6) of object
ls()
            # list names of objects in the workspace
## Print multiple objects
            # prints multiple objects, one after the other
cat()
## Functions that return TRUE or FALSE?
all()
            # returns TRUE if all values are TRUE
any()
            # returns TRUE if any values are TRUE
is.factor() # returns TRUE if the argument is a factor
          # returns TRUE if the argument is an NA
is.na()
             # NB also is.logical(), etc.
seq(from =1, by=2, length.out=3) # Unabbeviated arguments
seq(from =1, by=2, length=3) # Abbreviate `length.out` to `length`
```

Subsection 10.3.2: User-written functions

```
distance <- c(148,182,173,166,109,141,166)
mean.and.sd(distance)
```

```
## Execute the function with the default argument:
mean.and.sd()

## Thus, to return the mean, SD and name of the input vector
## replace c(mean=av, SD=sdev) by
list(mean=av, SD=sdev, dataset = deparse(substitute(x)))
```

Subsection 10.3.3: Generic functions, and the class of an object

Subsection 10.3.4: Pipes — a "Do this, then this, . . ." syntax

```
mean(rnorm(20, sd=2))
20 |> rnorm(sd=2) |> mean()
```

```
logmammals <- MASS::mammals |> log() |> setNames(c("logbody","logbrain"))
## Alternatively, use the ability to reverse the assignment operator.
MASS::mammals |> log() |> setNames(c("logbody","logbrain")) -> logmammals
## This last is more in the spirit of pipes.
```

```
MASS::mammals |>
  log() |>
  setNames(c("logbody","logbrain")) |>
  (\(d)lm(logbrain ~ logbody, data=d))() |>
  coef()
```

Subsection 10.3.5: Operators

```
## Multiple of divisor that leaves smallest non-negative remainder
c("Multiple of divisor" = 24 %/% 9, "Remainder after division" = 6)
```

Section 10.4: Calculations with matrices, arrays, lists, and data frames

Calculations in parallel across all elements of a vector

```
x <- 1:6
log(x)  # Natural logarithm of 1, 2, ... 6
log(x, base=10)  # Common logarithm (base 10)
log(64, base=c(2,10))  # Apply different bases to one number
log(matrix(1:6, nrow=2), base=2)  # Take logarithms of all matrix elements</pre>
```

Patterned data

```
seq(from=5, to=22, by=3) # The first value is 5.
rep(c(2,3,5), 4) # Repeat the sequence (2, 3, 5) four times over
rep(c("female", "male"), c(2,3)) # Use syntax with a character vector
```

Subsection 10.4.1: Missing values

```
nbranch <- subset(DAAG::rainforest, species=="Acacia mabellae")$branch
nbranch  # Number of small branches (2cm or less)

mean(nbranch, na.rm=TRUE)

nbranch == NA  # This always equals `NA`

is.na(nbranch)  # Use to check for NAs

nbranch[is.na(nbranch)] <- -999
  # `mean(nbranch)` will then be a nonsense value</pre>
```

NAs in modeling functions

```
options()$na.action # Version 3.2.2, following startup
```

Counting and identifying NAs – the use of table()

```
with(DAAG::nswdemo, table(trt, re74>0, useNA="ifany"))
```

Infinities and NaNs

```
summary(DAAG::primates)
primates <- DAAG::primates</pre>
                    # `plotCI() function in package `gplots`
gplots::plotCI()
plotrix::plotCI()
                    # `plotCI() function in package `plotrix`
sessionInfo()[['basePkgs']]
## List just the workspace and the first eight packages on the search list:
search()[1:9]
data(package="datasets")
Section 10.5: Brief notes on R graphics packages and functions
Subsection 10.5.1: Lattice graphics — a step beyond base graphics
grog <- DAAG::grog</pre>
chr <- with(grog, match(Country, c('Australia', 'NewZealand')))</pre>
  # Australia: 1; matches 1st element of c('Australia', 'NewZealand')
  # NewZealand: 2; matches 2nd element
plot(Beer ~ Year, data=grog, ylim=c(0, max(Beer)*1.1), pch = chr)
with(grog, points(Wine ~ Year, pch=chr, col='red'))
legend("bottomright", legend=c("Australia", "New Zealand"), pch=1:2)
title(main="Beer consumption (1, pure alcohol)", line=1)
                          ## Loads both lattice and the add-on latticeExtra
library(latticeExtra)
gph <- xyplot(Beer+Wine ~ Year, groups=Country, data=grog)</pre>
update(gph, par.settings=simpleTheme(pch=19), auto.key=list(columns=2))
## Or, condition on `Country`
xyplot(Beer+Wine+Spirit ~ Year | Country, data=grog,
       par.settings=simpleTheme(pch=19), auto.key=list(columns=3))
```

xyplot(csoa~it | tint*target, groups=agegp, data=tinting, auto.key=list(columns=2))

tinting <- DAAG::tinting

```
cuckoos <- DAAG::cuckoos
av <- with(cuckoos, aggregate(length, list(species=species), FUN=mean))
gph <- dotplot(species ~ length, data=cuckoos, alpha=0.4) +</pre>
  as.layer(dotplot(species ~ x, pch=3, cex=1.4, col="black", data=av))
update(gph, xlab="Length of egg (mm)")
## Alternatives, using `layer()` or `as.layer()`
avg <- with(cuckoos, data.frame(nspec=1:nlevels(species),</pre>
                              av=sapply(split(length, species), mean)))
dotplot(species ~ length, data=cuckoos) +
  layer(lpoints(nspec~av, pch=3, cex=1.25, col="black"), data=avg)
dotplot(species ~ length, data=cuckoos) +
  as.layer(dotplot(nspec~av, data=avg, pch=3, cex=1.25, col="black"))
## Specify panel function
dotplot(species ~ length, data=cuckoos,
  panel=function(x,y,...){panel.dotplot(x, y, pch=1, col="gray40")
    avg <- data.frame(nspec=1:nlevels(y), av=sapply(split(x,y),mean))</pre>
    with(avg, lpoints(nspec~av, pch=3, cex=1.25, col="black")) })
```

Combining separately created graphics objects

Subsection 10.5.2: Dynamic graphics – the rgl package

```
vignette('plot3D', package='plot3D')
```

Section 10.6: Plotting characters, symbols, line types and colors

```
ycol < -2.1 - (0:9) * 2.1
ftype <- c("plain", "bold", "italic", "bold italic", "symbol")</pre>
yline <- 4.2
ypmax <- 20
farleft < -7.8
plot(c(-4, 31), c(4.25, ypmax), type = "n", xlab = "", ylab = "",
axes = F)
chh \leftarrow par()$cxy[2]
text(0:25, rep(ypmax + 0.8 * chh, 26), paste(0:25), srt = 90,
cex = 0.75, xpd = T)
text(-1.5, ypmax + 0.8 * chh, "pch = ", cex = 0.75, xpd = T)
points(0:25, rep(ypmax, 26), pch = 0:25, lwd=0.8)
letterfont <- function(ypos = ypmax, font = 2) {</pre>
par(font = font)
text(-1.35, ypos, "64-76", cex = 0.75, adj = 1, xpd = TRUE)
text(19 - 1.35, ypos, "96-108", cex = 0.75, adj = 1)
points(c(0:12), rep(ypos, 13), pch = 64:76)
points(19:31, rep(ypos, 13), pch = 96:108)
text(farleft, ypos, paste(font), xpd = T)
text(farleft, ypos - 0.5, ftype[font], cex = 0.75)
}
plotfont <- function(xpos = 0:31, ypos = ypmax, font = 1,
sel32 = 2:4, showfont = TRUE) {
par(font = font)
i <- 0
for (j in sel32) {
i < -i + 1
maxval < - j * 32 - 1
if(j==4)maxval <- maxval-1</pre>
text(-1.35, ypos - i + 1, paste((j - 1) * 32, "-",
maxval, sep = ""), cex = 0.75, adj = 1, xpd = TRUE)
if(j!=4)
points(xpos, rep(ypos - i + 1, 32), pch = (j - 1) *
32 + (0:31)
else
points(xpos[-32], rep(ypos - i + 1, 31), pch = (j - 1) *
32 + (0:30)
lines(rep(-1.05, 2), c(ypos - length(sel32) + 1, ypos) +
c(-0.4, 0.4), xpd = T, col = "grey40")
```

```
if (showfont) {
text(farleft, ypos, paste("font =", font, " "), xpd = T)
text(farleft, ypos - 0.5, ftype[font], cex = 0.75,
xpd = T
}
plotfont(ypos = ypmax - 1.5, font = 1, sel32 = 2:4)
for (j \text{ in } 2:4) letterfont(ypos = ypmax - 2.1 - 1.4 * j, font = j)
plotfont(ypos = ypmax - 9.1, font = 5, sel32 = 3)
plotfont(xpos = c(-0.5, 1:31), ypos = ypmax - 10.1, font = 5,
sel32 = 4, showfont = FALSE)
par(font = 1)
ltypes <- c("blank", "solid", "dashed", "dotted", "dotdash",</pre>
"longdash", "twodash")
lcode <- c("", "", "44", "13", "1343", "73", "2262")
for (i in 0:6) {
lines(c(4, 31), c(yline + 4.5 - 0.8 * i, yline + 4.5 - 0.8 * i
0.8 * i), lty = i, lwd = 2, xpd = T)
if (i == 0)
numchar <- paste("lty =", i, " ")</pre>
else numchar <- i
text(farleft, yline + 4.5 - 0.8 * i, numchar, xpd = TRUE)
text(farleft + 3.5, yline + 4.5 - 0.8 * i, ltypes[i +
1], cex = 0.85, xpd = TRUE)
text(farleft + 7.5, yline + 4.5 - 0.8 * i, lcode[i +
1], cex = 0.85, xpd = TRUE)
```

```
ycol <- -2.1 - (0:9) * 2.1
ftype <- c("plain", "bold", "italic", "bold italic", "symbol")
yline <- 4.2
ypmax <- 20
farleft <- -7.8
plot(c(-4, 31), c(4.25, ypmax), type = "n", xlab = "", ylab = "",
axes = F)
chh <- par()$cxy[2]
text(0:25, rep(ypmax + 0.8 * chh, 26), paste(0:25), srt = 90,
cex = 0.75, xpd = T)
text(-1.5, ypmax + 0.8 * chh, "pch = ", cex = 0.75, xpd = T)
points(0:25, rep(ypmax, 26), pch = 0:25, lwd=0.8)
letterfont <- function(ypos = ypmax, font = 2) {
par(font = font)</pre>
```

```
text(-1.35, ypos, "64-76", cex = 0.75, adj = 1, xpd = TRUE)
text(19 - 1.35, ypos, "96-108", cex = 0.75, adj = 1)
points(c(0:12), rep(ypos, 13), pch = 64:76)
points(19:31, rep(ypos, 13), pch = 96:108)
text(farleft, ypos, paste(font), xpd = T)
text(farleft, ypos - 0.5, ftype[font], cex = 0.75)
plotfont <- function(xpos = 0:31, ypos = ypmax, font = 1,
sel32 = 2:4, showfont = TRUE) {
par(font = font)
i <- 0
for (j in sel32) {
i <- i + 1
maxval <- j * 32 - 1
if(j==4) maxval <- maxval-1
text(-1.35, ypos - i + 1, paste((j - 1) * 32, "-",
maxval, sep = ""), cex = 0.75, adj = 1, xpd = TRUE)
if(j!=4)
points(xpos, rep(ypos - i + 1, 32), pch = (j - 1) *
32 + (0:31)
points(xpos[-32], rep(ypos - i + 1, 31), pch = (j - 1) *
32 + (0:30)
lines(rep(-1.05, 2), c(ypos - length(sel32) + 1, ypos) +
c(-0.4, 0.4), xpd = T, col = "grey40")
if (showfont) {
text(farleft, ypos, paste("font =", font, " "), xpd = T)
text(farleft, ypos - 0.5, ftype[font], cex = 0.75,
xpd = T)
}
plotfont(ypos = ypmax - 1.5, font = 1, sel32 = 2:4)
for (j \text{ in } 2:4) letterfont(ypos = ypmax - 2.1 - 1.4 * j, font = j)
plotfont(ypos = ypmax - 9.1, font = 5, sel32 = 3)
plotfont(xpos = c(-0.5, 1:31), ypos = ypmax - 10.1, font = 5,
sel32 = 4, showfont = FALSE)
par(font = 1)
ltypes <- c("blank", "solid", "dashed", "dotted", "dotdash",</pre>
"longdash", "twodash")
lcode <- c("", "", "44", "13", "1343", "73", "2262")
for (i in 0:6) {
```

```
lines(c(4, 31), c(yline + 4.5 - 0.8 * i, yline + 4.5 -
0.8 * i), lty = i, lwd = 2, xpd = T)
if (i == 0)
numchar <- paste("lty =", i, " ")
else numchar <- i
text(farleft, yline + 4.5 - 0.8 * i, numchar, xpd = TRUE)
text(farleft + 3.5, yline + 4.5 - 0.8 * i, ltypes[i +
1], cex = 0.85, xpd = TRUE)
text(farleft + 7.5, yline + 4.5 - 0.8 * i, lcode[i +
1], cex = 0.85, xpd = TRUE)
}</pre>
```

Font families

Colors

```
library(RColorBrewer)
palette(brewer.pal(12, "Set3"))
```

```
if(file.exists("/Users/johnm1/pkgs/PGRcode/inst/doc/")){
code <- knitr::knit_code$get()
txt <- paste0("\n## ", names(code),"\n", sapply(code, paste, collapse='\n'))
writeLines(txt, con="/Users/johnm1/pkgs/PGRcode/inst/doc/Appendix.R")
}</pre>
```